Surface Topography: Metrology and Properties

TOPICAL REVIEW

Quantitative characterization of surface topography using spectral analysis

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Abstract
Roughness determines many functional properties of surfaces, such as adhesion, friction, and (thermal and electrical) contact conductance. Recent analytical models and simulations enable quantitative prediction of these properties from knowledge of the power spectral density (PSD) of the surface topography. The utility of the PSD is that it contains statistical information that is unbiased by the particular scan size and pixel resolution chosen by the researcher. In this article, we first review the mathematical definition of the PSD, including the one- and two-dimensional cases, and common variations of each. We then discuss strategies for reconstructing an accurate PSD of a surface using topography measurements at different size scales. Finally, we discuss detecting and mitigating artifacts at the smallest scales, and computing upper/lower bounds on functional properties obtained from models. We accompany our discussion with virtual measurements on computer-generated surfaces. This discussion summarizes how to analyze topography measurements to reconstruct a reliable PSD. Analytical models demonstrate the potential for tuning functional properties by rationally tailoring surface topography—however, this potential can only be achieved through the accurate, quantitative reconstruction of the PSDs of real-world surfaces.

1. Introduction

1.1. The importance of surface roughness and of the power spectral density (PSD) as a description of topography
The surface roughness of a part has tremendous influence on its functionality. This has been reviewed extensively, e.g. in [1] and elsewhere. Roughness affects not only surface properties—such as hydrophobicity [2], optical and plasmonic behavior [3], adhesion [4–6], friction and Casimir forces [7]—but also ‘bulk’ properties, such as fracture toughness and fatigue resistance [8]. For this reason, surface treatments are commonly used to control surface finish, and dozens of reference standards exist to describe measurement techniques and desired characteristics (such as ASME B46 [9], ISO 4287 [10], ISO 25178 [11] and SEMI MF1811 [12]).

The power spectral density (PSD) of a surface is a mathematical tool that decomposes a surface into contributions from different spatial frequencies (wavevectors). Mathematically, the PSD is the Fourier transform of the autocorrelation function of the signal, which contains just the power (and not the phase) across a range of wavevectors [13–15]. This allows identification of the spatial frequencies that can be found in the signal. Figure 1 illustrates the PSD schematically for a regular dual-sinusoidal surface, a self-affine fractal surface, and the non-self-affine surface of a frozen capillary wave. Overall, the primary utility of the PSD is that it contains statistical information on the surface topography—which is largely unbiased by the choice of a particular scan size and pixel resolution picked by the researcher.

Many contact mechanics models (dstaking back to [16]) have been constructed to compute functional properties of surfaces as a function of roughness. Specifically, the PSD of a surface can be used to compute quantities such as contact stiffness, adhesion, and the true contact area (which is distinct from the apparent contact area and which has relevance to friction [17–19] and electrical [20] and thermal [21] conductance across the contact). Many of these models (such as Persson’s scaling theory for contact [22–24]) take as an input the entire PSD of the surface. Other models (such as [5, 6, 25–30]) use mathematical manipulations such that the only input parameters are three scalar quantities: the root-mean-
square (RMS) height \( h_{rms} \), the RMS slope \( h'_{rms} \), and the RMS curvature \( h''_{rms} \)—all of which are calculated most reliably from the PSD itself, as discussed in section 1.2. No matter which type of contact mechanics model is used, the central challenge for the practical application of these models is the measurement and calculation of accurate, reliable PSDs of real-world surfaces.

1.2. Using the power spectrum to compute accurate values for RMS height, slope, and curvature of a surface

In an ideal case of perfect knowledge of surface topography as a continuous map of heights \( h(x, y) \) at in-plane position \( x, y \) with zero mean, the values of \( h_{rms} \), \( h'_{rms} \), and \( h''_{rms} \) can be computed directly from the real-space topography. In this case, the RMS height is given by

\[
h_{rms} = \langle h^2 \rangle = \langle f \rangle = A_0^{-1} \int f(x, y)\,dx\,dy,
\]

where the angle brackets denote the average over the \( x\)-\( y \) plane of the enclosed function \( f \). The RMS slope is computed as

\[
h'_{rms} = \langle \nabla h^2 \rho \rangle
\]

and the RMS curvature is computed as

\[
h''_{rms} = \frac{1}{2}\langle \nabla^2 h^2 \rangle.
\]

For many contact properties, just one of these three quantities will dominate the behavior. Specifically, the contact stiffness of an interface between two rough surfaces is primarily dependent on roughness at the large scale—and can be quantitatively related to the RMS height \( h_{rms} \) [26–29]. Analytical models and atomistic simulations show that other important quantities—such as the large-scale adhesive properties of a contacting junction [5, 6, 31], and the true contact area [25, 30, 32]—depend only on the smallest scales of roughness and therefore can be related to \( h'_{rms} \) and \( h''_{rms} \). Note that \( h_{rms} \), \( h'_{rms} \), and \( h''_{rms} \) are single-valued and unique for a particular surface, assuming perfect knowledge of that surface.

However, the central problem [33] of applying these analytical contact models to real-world surfaces is that it is not possible to determine a perfect description of surface topography as a continuous map of heights. Rather, we can only experimentally measure heights at discrete points, with finite resolution and over some finite measurement region. It is therefore not possible to directly determine \( h_{rms}, h'_{rms} \), and \( h''_{rms} \) from real-space measurements.

From any real-space measurement, we can of course compute approximate values for RMS height (typically denoted \( R_s \) if obtained from a line scan and \( S_q \) if obtained from an area scan), RMS slope \( S_{2\phi} \), and RMS curvature \( S_{2\phi} \). However, only in ideal experimental measurement with infinite scan size and infinite resolution will these be representative of the surface (i.e. in the ideal case, \( S_q \equiv h_{rms}, S_{2\phi} \equiv h'_{rms} \), and \( S_{2\phi} \equiv h''_{rms} \)). In real experiments, the measured value of \( S_q \) will differ from the true \( h_{rms} \) in cases where the measured region is insufficiently large or the resolution is insufficiently fine to accurately sample the topography. In almost every case, the measured values of \( S_{2\phi} \) and \( S_{2\phi} \) will differ from \( h'_{rms} \) and \( h''_{rms} \), because the true local slope and curvature are computed from a discrete set of height information which is subject to experimental noise. Noise at small scales affects gradients and higher-order derivatives more than absolute height measurement. Additionally, there are multiple (infinite) ways to discretize the gradient operator, \( \nabla h \).

Two common standard slope measures \( S_{2q} \) (ISO 25178–2 [10]) and \( S_{2q} \) (ASME B46.1 [9]) use a
first- and a sixth-order finite difference stencil for $\nabla h$, respectively. Both can introduce either smoothing or artificially sharp corners, which will lead to inaccuracies. Smaller measurements with higher resolution will result in values of $S_{a\infty}$ and $S_{\Delta q}$ that are closer to $h_{\text{rms}}'$ and $h_{\text{rms}}''$ for the surface, but processing and instrumental artifacts tend to be most severe when pushing the resolution limits of an instrument.

The resolution to the problem of an incomplete measurement of a surface is to reconstruct the PSD of the surface, as completely and accurately as possible. Specifically, one can compute, report, and analyze the PSD across many length scales. In some cases, it is possible to reconstruct an entire, complete PSD using multiple measurements and multiple techniques. Even in cases where the complete surface PSD cannot be measured due to instrument limitations, the partial PSD is extremely useful. Specifically: the PSD enables the calculation of upper and lower bounds for $h_{\text{rms}}'$ and $h_{\text{rms}}''$ under certain assumptions; it can be used to identify and/or mitigate instrumental artifacts; and it can be extrapolated (in some cases) to learn about a surface beyond the limits of the measurement technique.

1.3. Challenges for the accurate experimental determination of the PSD

The PSD has been fruitfully applied to surface topography measurements for decades [34–37]. Many useful reference texts (for example [13]) describe in detail the calculation of the PSD for a surface. There is also an international reference standard specifically describing its calculation (SEMI MF1811 [12]). Nevertheless, the PSD is underutilized quantitatively (except for extracting the Hurst exponent, which is related to the fractal dimension of a surface (see appendix A.3)), and is used inconsistently. We believe that there are three critical challenges that must be addressed to enable the use of the PSD as a quantitative tool for prediction of properties. We discuss them in the following sections and strive to establish strategies for computing, analyzing, and reporting quantitative PSDs.

Challenge A: While the mathematical description of the PSD is well-defined, there are several variations in the way it can be computed from measured data—each of which will lead to different numerical results.

Challenge B: The theoretical PSD that is used in many mathematical descriptions of surfaces such as contact mechanical models, is complete, accurate, and describes an ensemble of surfaces or a surface of infinite spatial extent. By contrast, the experimental PSD that is computed from a single measurement is incomplete (bandwidth-limited), inaccurate (artifact-prone), and describes a single iteration of a surface, which is finite in extent.

Challenge C: Finally, recent contact theories predict that surface properties such as contact area and adhesion depend strongly on the smallest-scale of roughness. The current state-of-the-art experimental characterization techniques contain instrumental and analysis artifacts at these scales, which must be detected and mitigated.

The following sections serve to explain these challenges and present strategies on how to mitigate them. To demonstrate these, we have created virtual surfaces whose structure we know down to the smallest scale. By comparing the input PSD that was used to create the surface to the output PSD that was measured from the surface, the results demonstrate the influence of bandwidth and other instrumental limitations on the measured PSD. The purpose of the present paper is to discuss and review the application of spectral analysis to real, experimental surfaces. This topic has particular relevance to the application of self-affine scaling models; however, the concepts discussed apply equally well to the characterization of all surfaces—whether self-affine over a certain range or not.

2. Challenge A: Variations in the mathematical definition of the PSD

Many experimental investigations report the PSD as a useful measure of roughness for surfaces. However, a review of recent atomic force microscopy (AFM)-based PSD measurements on advanced-technology surfaces demonstrates significant inconsistency in the ways that PSDs are defined, computed, and analyzed. Even the units of the reported PSDs vary from $m^2$ [38, 39] to $m^3$ [40–42] to $m^4$ [43–45] to ‘arbitrary units’ [46, 47]. Furthermore, the PSD is frequently used qualitatively to distinguish surfaces [42, 45] and, when used quantitatively, is typically only used to determine fractal dimensions [39, 46]. There is also variation among these investigations in the calculation and interpretation of this value.

Here, we define and distinguish the various functions that are all referred to in common usage as the ‘PSD’. This section of the main text distinguishes conceptually between types of PSD; rigorous mathematical definitions are included in appendix A.1. Each of the various types of PSD is the Fourier transform of the height autocorrelation function. By virtue of the convolution theorem of Fourier analysis, this can be computed as the square of the Fourier transform of the surface height $h(x, y)$. However, this calculation can be applied in several different ways to 1D and 2D signals. To concretely distinguish the different varieties and demonstrate their mathematical differences, the calculation of the root mean square height $h_{\text{rms}}$ is shown for each.

2.1. Conceptual understanding of the different curves called ‘PSD’

For a 1D signal (e.g., a line-scan from a stylus profilometer, as shown in figure 2(a)), the PSD $C^{1D}$ is a symmetric, 1D function defined in frequency space (figure 2(b)). The units of $C^{1D}$ are $(m^3)$ and the units of $q_x$ are $(m^{-1})$, such that the area under the curve, which is equal to $h_{\text{rms}}$ has the correct units of $(m^2)$:
is typically shown. However, a pseudo-

d. This still has the correct units of (m²):

A 2D topographic scan h(x, y) shown in (d) can be represented by (e) a 2D PSD \(C_{q x q y}(q_x, q_y)\) that is a surface in reciprocal space. A 2D PSD is also commonly computed (according to equation (5) to enable easier comparison with \(C_{1D}^{\text{iso}}\)). Note that the units and absolute values are distinct between \(C_{2D}^{\text{iso}}\) and \(C_{\text{pseudo-1D}}\).

There are exact mathematical expressions relating these forms to each other (see appendix A.2 for more detail).

By contrast, for a 2D signal (e.g., a topography map, figure 2(f)), the area under the curve must be multiplied by 2 in order to account for the region where \(q_x < 0\):

\[
h_{\text{rms}}^2 = \frac{1}{\pi} \int_{0}^{\infty} C_{1D}^{\text{iso}}(q_x) dq_x.
\]

Note that equations (1) and (2) assume that the mean of the signal is zero, because it is common to subtract the mean value from the data.

By contrast, for a 2D signal (e.g., a topography map, as shown in figure 2(d)), the full 2D PSD \(C_{2D}^{\text{iso}}\) is a surface in frequency space, shown in figure 2(e). The units of \(C_{2D}^{\text{iso}}\) are (m⁴) and the units of \(q_x\) and \(q_y\) are each (m⁻¹). In this case, the volume under the PSD surface is equal to \(h_{\text{rms}}^2\) which still has units of (m⁴). Converting \((q_x, q_y)\) to polar coordinates \((q, \theta)\) and integrating out \(\theta\) yields a factor of 2\(\pi\) since \(C_{2D}^{\text{iso}}\) is invariant with \(\theta\). The result is:

\[
h_{\text{rms}}^2 = \frac{1}{2\pi} \int_{0}^{\infty} q C_{2D}^{\text{iso}}(q) dq.
\]

Many commercial software packages report a 1D PSD with units of (m³) even for 2D surfaces because 1D PSDs are in more common usage; however, there are different ways to compute this 1D PSD from 2D data—and there can be differences in the resulting values. For instance, one method is to take the 1D PSD of each line of data, and then to take an average over all lines; we refer to this as \(C_{1D}^{\text{iso}}\) for 2D data. Another method is to report a pseudo-1D PSD,

\[
C_{\text{pseudo-1D}}(q) = \frac{1}{\pi} C_{2D}^{\text{iso}}(q)
\]

which follows \(C_{1D}^{\text{iso}}\) at intermediate wavevectors for self-affine surfaces. However, \(C_{\text{pseudo-1D}}\) cannot be treated in the same way as \(C_{1D}^{\text{iso}}\) when computing scalar parameters. This is easily seen by inserting equation (5) into (4) to obtain

\[
h_{\text{rms}}^2 = \frac{1}{2\pi} \int_{0}^{\infty} \frac{q}{\pi} C_{2D}^{\text{iso}}(q) dq.
\]

which differs by a factor of 2\(\pi\) from equation (2).

To complicate the picture, all of the previously described functions—\(C_{1D}^{\text{iso}}, C_{1D}^{\text{iso}+}, C_{2D}^{\text{iso}}, C_{\text{pseudo-1D}}\) are imprecisely referred to as ‘the PSD’ [38–46]. However, it is clear that they must be treated differently when quantitative values are computed. There are exact mathematical expressions relating the various forms to each other (see appendix A.2 for more detail).

Most software packages are limited to reporting the 1D representations \(C_{1D}^{\text{iso}}\) or \(C_{\text{pseudo-1D}}\), as they are

\[
\text{Figure 2. Examples of the various curves that are generally referred to as ‘PSD’}.\ A 1D line scan \(h(x)\) shown in (a) can be represented by (b) a 1D PSD \(C_{1D}^{\text{iso}}(q_x)\), but is more commonly indicated by (c) a one-sided 1D PSD \(C_{\text{pseudo-1D}}(q_x)\) that omits wavevectors \(q_x < 0\). A 2D topographic scan \(h(x, y)\) shown in (d) can be represented by (e) a 2D PSD \(C_{2D}^{\text{iso}}(q_x, q_y)\) that is a surface in reciprocal space. A 2D PSD is also commonly computed (according to equation (5) to enable easier comparison with \(C_{1D}^{\text{iso}}\)). Note that the units and absolute values are distinct between \(C_{2D}^{\text{iso}}\) and \(C_{\text{pseudo-1D}}\).
more amenable to conventional PSD analysis (most of which was developed for 1D time-series data). Indeed, the international reference standard SEMI MF1811 [12] only discusses 1D profile measurements and some limited extension to 2D isotropic surfaces. These quantities are complete representations of the full C2D only in cases where the surface is isotropic. In the more general case, the mathematical integration over the full surface of C2D is required in order to compute accurate quantitative surface parameters. The mathematical use of C∞ instead of C2D for a non-isotropic surface will not lead to errors in the value of hrms as computed in equation (4), but will lead to errors in h′ rms and h″ rms—which are discussed in the next section. Yet many software packages do not report the full C2D and may obscure the fact that the surface is anisotropic—leading to quantitative values computed from C∞ that are unrepresentative of the surface.

2.2. Self-affine surfaces, and the resulting simplifications to the PSD

It has been shown that a wide range of natural and synthetic surfaces [48]—from coastlines [49] to mountain ranges [50, 51] to fracture surfaces [52] to machined surfaces [53]—show characteristics of self-affine (also called fractal) scaling [34, 36, 54–56]. The underlying picture is that roughness consists of asperities (bumps), which are covered with smaller asperities, which in turn are covered with smaller asperities, etc, as was described by Archard in 1957 [16]. The PSD of a perfectly self-affine surface has a power-law dependence on the spatial frequency of roughness, and its exponent is related to the fractal dimension [36] of the surface (figure 1(f)). The calculations in section 2.1 and appendices A.1 and A.2 apply to all surfaces, regardless of self-affinity. However, in cases where self-affine roughness is observed, some of the calculations can be simplified.

Power spectra of real surfaces often show \( C^\infty(q) \propto q^{-2-2H} \) over many (but not all) scales [32, 36]. This is the signature of a self-affine surface with Hurst exponent \( H \) (see appendix A.3 for more details). At small wavelength \( \lambda \) (large wavevector \( q \)), the power-law will be cut off by the atomic spacing. The details of this cutoff are still debated. For the sake of the discussion in this paper, we assume that a sharp cutoff happens at a wavelength \( \lambda_0 \) (wavevector \( q_0 = 2\pi/\lambda_0 \)). At large wavelength \( \lambda_0 \) the power law typically crosses over to constant power, \( C^\infty(q) = \text{const} \) for \( q < q_L = 2\pi/\lambda_0 \). We assume that this ‘roll-off’ can only extend up to a wavelength \( \lambda_0 \) (wavevector \( q_0 = 2\pi/\lambda_0 \)) where the power drops to zero. The idealized power-spectrum of a self-affine, randomly rough surface is therefore:

\[
C^\infty(q) = C_0 \begin{cases} 0 & \text{if } q < q_0 \\ q_L^{-2} q^{-2H} = \text{const.} & \text{if } q_0 \leq q < q_L \\ q^{-2H} & \text{if } q_L \leq q < q_0 \\ 0 & \text{if } q \geq q_0 \end{cases}
\]  

(7)

where \( q_0 < q_L < q_0 \) and \( C_0 \) is a constant. This power-spectrum is illustrated schematically in figure 1(f). The figure illustrates two Hurst exponents (solid and dashed line) with identical RMS slope \( h^\prime_{\text{rms}} = 0.1 \).

Self-affine scaling typically extends over many decades such that \( q_0 \gg q_L \). In this limit, we obtain simple analytical expressions for RMS slope and curvature (see also appendix A.1):

\[
(h^\prime_{\text{rms}})^2 = \frac{1}{2\pi} \int_0^\infty q^2 C^\infty(q)q dq = \frac{1}{4\pi} \frac{C_0}{1-H} q_L^{-2H}
\]  

(8)

\[
(h^\prime_{\text{rms}})^2 = \frac{1}{8\pi} \int_0^\infty q^2 C^\infty(q)q dq = \frac{1}{16\pi} \frac{C_0}{2-H} q_L^{-2H}
\]  

(9)

It is important to note that because \( q_0 \gg q_L \), those expressions do not depend on \( q_0 \) and \( q_L \). Slope and curvature are entirely determined by the structure at the smallest scales of the surface. Even if we did not have a sharp cutoff at \( q_L \), the integral expressions that give \( h^\prime_{\text{rms}} \) and \( h^\prime_{\text{rms}} \) would be dominated by what happens at the smallest scales because of the power-law scaling of the PSD. This is graphically illustrated in figure 3.

By contrast, the expression for RMS height depends on the power at the scale \( q_0 \) where the power-law region ends. With no roll-off \( q_0 \equiv q_L \) we get

\[
(h^\prime_{\text{rms}})^2 = \frac{C_0}{4\pi H} q_L^{-2H}
\]  

(10)

while for a very large roll-off region \( q_0 \ll q_L \) we obtain

\[
(h^\prime_{\text{rms}})^2 = \frac{C_0}{4\pi} \left(1 + \frac{1}{H}\right) q_L^{-2H}
\]  

(11)
Both expressions depend only on $C_0$ and $q_L$ and neither on $q_s$ nor on $q_r$.

We can analytically compute the PSD $C^{1D+}(q)$ of a line profile that corresponds to the 2D, isotropic PSD given by equation (7). The resulting expression is (see appendix A.3 for more detail):

$$ C^{1D}(q_s) \approx \frac{q}{\pi} C^{iso}(q_s) \sqrt{1 - \left(\frac{q}{q_s}\right)^2} $$

(12)

which forms the basis for the pseudo-1D PSD defined in equation (5). However, the 1D PSD will taper off approximately as $\sqrt{1 - (q/x)^2}$ rather than drop to zero sharply. This behavior is illustrated in figure 2(c).

### 2.3. Strategies for the calculation of PSD and scalar roughness quantities

We suggest the following guidelines for measuring and reporting a PSD. For quantitative numerical calculations such as in [25, 57], $C^{iso}$ is the appropriate form of the PSD to use. However, the surface should also be checked for anisotropy, as most of the contact models assume isotropic roughness. For a 2D image of topography, this can be assessed by computing the surface PSD $C^{iD}$, in which the presence or absence of radial symmetry should be obvious, or by comparing $C^{iD}$ against $C^{pseudo-iD}$, which should be identical for an isotropic surface.

If anisotropy is detected, it must be determined whether its origin is true anisotropy in the surface or artificial anisotropy introduced by the measurement technique, as is common for characterization techniques with a preferred direction. (For instance, in AFM the direction parallel to the scanning axis is sampled in less than one second, while topography in the perpendicular direction may take several minutes to measure: the latter is therefore much more prone to artifacts from drift.) This can be determined by repeating the measurement on the same surface in three different directions: horizontal, vertical and a third, oblique direction. If the $C^{iD}$ as calculated in these three directions is identical, then the surface is likely isotropic and, for self-affine surfaces, $C^{iso}$ can be approximated using $C^{iso}(q) = C^{iD}(q) \pi/q$. If, on the other hand, $C^{iD}$ is very different in the three directions, then the surface is likely to be truly anisotropic.

### 3. Challenge B: Reconstructing the theoretical PSD and scalar roughness parameters from incomplete data

In this section, we will distinguish among the PSD of a measurement, the PSD of a surface, and the PSD of a class or an ensemble of surfaces. One can always mathematically compute the PSD from any topography measurement carried out on a subsection of the full surface; however, the computed spectrum from any single measurement can only yield information for a limited range of wavelengths and will typically contain instrumental artifacts. Even ignoring instrumental artifacts (which are discussed in more detail in section 4), care is required to accurately reconstruct the full theoretical PSD to avoid artifacts due to: aperiodicity; artifacts of stitching; and small sample sizes. Thus, the calculation of upper and lower bounds of $h_{rms}, h_{rms}$ and $h_{rms}$ will be discussed.

To demonstrate each of these issues, we have analyzed computer-generated surfaces that were ‘measured’ at various locations and resolutions. These synthetic measurements are, of course, free of measurement artifacts and therefore are useful for demonstrating the errors that can arise due to the analysis procedure, rather than the experimental procedure. These synthetic, computer-generated surfaces were created under the assumptions of the random process model of surface topography [34, 54, 55]. We start with a well-defined self-affine PSD in the form of equation (7), which we refer to as $C^{iso}_{input}(q)$, which is the input PSD. We then construct a surface as a superposition of waves of the form $h_{ij}(\tau) = \chi(q) \exp[iq\cdot\tau + i\phi(q)]$. The amplitude $\chi(q)$ is chosen randomly from a Gaussian distribution with a standard deviation given by the square-root of the PSD, $[C^{iso}_{input}(q)]^{1/2}$, for each $q$. We additionally chose a random phase shift $\phi(q)$ within the interval $[0, 2\pi)$ from a uniform distribution; however, this phase-shift does not affect the measured PSD. This way of creating synthetic PSDs is often referred to as a Fourier-filtering algorithm. All synthetic surfaces discussed here have a Hurst exponent of $H = 0.8$.

### 3.1. PSD of a single measurement—aperiodicity and the need for windowing

Experimental surfaces and their measurements will never be fully periodic. This introduces problems with a straightforward application of the Fourier transform described in section 2 that treats the signal as periodic; thus, windowing [58] of the data is required [59]. The nonperiodic data is multiplied with a periodic windowing function that goes smoothly to zero at the edges of the topography image. In Fourier space, this window acts as a low-pass filter, removing the high-frequency components that are introduced by edges of the topography. While windowing is common practice in the signal-processing community, many conventional windows have a maximum that is set to unity. When using self-affine scaling laws to compute scalar roughness parameters, the sum rules that are used to compute $h_{rms}, h_{rms}$ and $h_{rms}$ must be conserved. Therefore, the area beneath the squared window $w$ needs to be normalized to return the length $L$ or area $L_x L_y$ of the window’s support:

$$ \int_0^L w(x)dx = L \text{ or } \int_0^{L_x} \int_0^{L_y} w(x, y)dxdy = L_x L_y. $$

(13)
Here we demonstrate the effect of aperiodicity, and artifacts that can arise from improper windowing using a realization of a well-defined random and statistically isotropic, periodic surface of \(2048 \times 2048\) pixels that is shown in figure 4(a). We assume the full surface is a \(2 \mu m \times 2 \mu m\) scan of a surface topography. The example surface used in this section has no power below a wavelength of 30 nm and has a Hurst exponent of 0.8.

In the present paper, we exclusively use the Hann window \([58]\), but results obtained with other windowing functions are virtually indistinguishable. The 1D Hann window is given by

\[
w_{\text{Hann}}^{1D}(x) = \left( \frac{2}{3} \right)^{1/2} \left( 1 - \cos 2\pi x / L_x \right),
\]

where \(L_x\) is the length of the signal in the relevant direction (here designated \(x\)). Note that equation (14) fulfills the normalization conditions discussed above. In computing the 1D PSD, we apply this 1D window to each line scan independently, \(h_{\text{windowed}}(x) = w_{\text{Hann}}^{1D}(x)h(x)\). Computing the 2D PSD requires a windowing function that is a function of \(x\) and \(y\) position within the plane. A common construction is to use the product of two 1D windows, \(w_{\text{Hann}}^{1D}(x, y) = w_{\text{Hann}}^{1D}(x)w_{\text{Hann}}^{1D}(y)\). However, the resulting window is not rotationally symmetric. Even if the surface is isotropic, the PSD of the windowed surface will become anisotropic, as shown in figure 4(g). For these reasons, we use the radially symmetric Hann window,

\[
w_{\text{Hann}}^{2D}(x, y) = \left( \frac{3\pi}{8} - \frac{2}{\pi} \right)^{-1/2} \times \left\{ \begin{array}{cl} 1 + \cos \frac{2\pi \sqrt{X^2 + Y^2}}{\min(L_x, L_y)} & \text{for } X^2 + Y^2 < \left[ \min(L_x, L_y)/2 \right]^2 \end{array} \right.
\]

where \(X = x - L_x/2\) and \(Y = y - L_y/2\) and the function is equal to zero everywhere the inequality is not satisfied.

Figure 4(b) shows the 1D PSD \(C_{q,1}^{1D}\) of the periodic surface shown in figure 4(a), alongside \(C_{q,2}^{2D}\). These results show that the 1D and 2D PSDs for self-affine surfaces are equivalent, except for the region near the short-wavelength cutoff where the 1D PSD tapers off smoothly and the 2D PSD is cut off sharply (see equation (12)). Both fall right on top of the input PSD shown by the black solid line. The PSD obtained with a Hann window, figure 4(c), follows the same power-law as the straight computation of the PSD and also recovers the same power in absolute terms. There are minor variations in the fluctuation of the PSD.

We now emulate a measurement of this surface by cutting out the central \(1 \mu m \times 1 \mu m\) section, which is no longer periodic. The resulting surface is shown in figure 4(d) and the non-windowed and windowed PSDs are shown in figures 4(e) and (f). It is clear that windowing is crucial to reconstructing the true PSD for non-periodic data. Critically, the non-windowed PSD makes the self-affine scaling appear to extend to the resolution limit of the measurement. In terms of windowing, the non-periodicity acts like a square window whose Fourier transform has \(1/q\) asymptotic behavior. The tail of the 1D PSD in figure 4(e) therefore scales as \(1/q^2\), as shown by the dashed line in that plot. Note that this looks like self-affine scaling with a Hurst exponent of \(H = 0.5\). For a real measurement of a real-world surface (where the true PSD is unknown), it could be erroneously concluded that self-affinity extends much further than it actually does. This artifact is particularly problematic when estimating the contact area using analytical models, as it would cause significant inaccuracy in the calculation of \(h_{\text{rms}}^\prime\) and \(h_{\text{rms}}^\prime\), which in turn causes errors in many predicted surface properties calculated from the PSD. Figure 4(f) shows that using the appropriate window resolves this issue.

Finally, figures 4(g) and (h) illustrate the effect of using a square and a radial window, respectively, on the 2D PSD. The window symmetry can clearly be seen in both images. The square window introduces apparent asymmetry into the 2D PSD: a vertical and a horizontal line of increased power. The use of a radial window does not bias the 2D PSD in any direction.

3.2. PSD of a set of measurements

Mathematically and in simulations, the PSD of a surface can be perfectly understood over the whole frequency range. By contrast, the range of spectral information for a single measurement is limited [15] the maximum wavelength is determined by the size \(L_x \times L_y\) of the domain over which the measurement was taken; the minimum wavelength is determined by the pixel size. The limit of accurate spectral information may be further reduced by instrumental artifacts [60], but this section is not considering instrumental artifacts. By combining multiple measurements, an individual technique is capable of providing spectral information over the range of wavelengths from the minimum instrument resolution to the maximum analysis size.

The most common measurement techniques employ scanning probes (i.e. stylus profilometry and AFM) or light and/or x-rays (i.e. optical profilometry and angle-resolved scattering) to probe topography [61]. These techniques are summarized in table 1. Stylus profilometry [62] has been used in some form for over 100 years and drags a sharp needle (typically 2–10 µm radius, but sometimes less than 1 µm) across a surface and records the deflection as a measure of surface topography. While the procedure is extremely robust and versatile, it is also relatively time-intensive, can cause sample damage, and can be limited by the relatively large tip. AFM (or, more generally, scanning probe microscopy) [63] is a class of techniques where a nanoscale tip is raster scanned over a surface. This tip can be in contact (contact mode), in intermittent contact (tapping mode), or out-of-contact (non-contact AFM or scanning tunneling microscopy). In all cases, the tip is used to sense the vertical position of the surface over an array of pixels...
Figure 4. Non-periodicity of measured data. A synthetic periodic surface (a) containing $2048 \times 2048$ pixels was created with self-affine scaling properties and Gaussian statistics. The power spectra are obtained using a straight FFT (b) and an FFT after applying a Hann window to the full topography (b). The black solid lines show the input PSD that was used to generate these surfaces, and match well with measured values (red, blue lines). By contrast, the $1 \mu m \times 1 \mu m$ central section of the topography (d) has been cropped from the surface shown in (a), and the resulting smaller surface is aperiodic. PSDs of the cropped region are then computed using a straight (e) and windowed (f) FFT. Without windowing, the PSD picks up an artificial contribution $\propto q^{-2}$ (dashed line in panel (e)) at large wavevectors (small wavelengths). The dashed line in panel (f) shows the PSD if a conventional normalization of the window is used, rather than normalizing to conserve the sum rules (section 3.1). Further, the use of the square of a simple 1D Hann window (g) can introduce artificial asymmetry into the PSD; while the radial Hann window (h) faithfully reproduces the PSD.

in a grid. Scanning probe microscopy can measure the smallest-scale features of any of the conventional surface characterization approaches and can achieve atomic resolution on ultra-flat samples. For rough samples, the lateral resolution is severely degraded by instrument noise and tip artifacts (see section 3).

Optical profilometry [64] refers to a family of techniques that use either phase shifts of monochromatic light or optical coherence of white light to determine the vertical position of each pixel in an analysis region. Using analysis algorithms and fitting routines (such as those discussed in [65, 66]), sub-nanometer resolution can be achieved in the vertical direction while the lateral resolution is diffraction limited to typical ranges of 500 nm–1 $\mu m$. The analysis size depends on the chosen microscope objective; while it is typically limited to approximately 5 mm for a single image, an encoded stage plus digital image registration can be readily used to stitch together multiple images to achieve 100 mm regions for analysis. Light scattering [13] and x-ray scattering [67] are a class of techniques that compare incident and scattered beams to measure changes in intensity as a function of wavelength. The details of this topic are beyond the scope of this paper, but unlike the other techniques discussed, scattering techniques do not measure the real-space topography of the sample, but rather use theories of wave-surface interactions [68] to relate the measured spectrum of the scattered beam to the spectrum of the surface. The scattered beam can be affected differently by different types of surfaces [69], and can be strongly affected by surface properties other than topography [67], and therefore various
assumptions about the surface are often required. X-ray scattering can, in theory, sample roughness down to the wavelength of the x-ray (less than 1 Å), but in practice instrumental and measurement difficulties limit this to several nanometers.

To recover the complete PSD of a surface, it is typically necessary to combine the results from various techniques and multiple measurements per technique that are all performed on the same surface. The resultant PSDs can be stitched together over many orders of magnitude [15, 51, 70]. However, the process of stitching can introduce artifacts into the final PSD which cause a lack of overlap between scales [15]. To demonstrate this, we created a very large (65 536 pixels per side) computer-generated surface that has a well-defined PSD. We used the central 50 000 pixels per side section of this surface as our realization of a 100-μm by 100-μm surface. We used the central 50 000 pixels per side (per side) computer-generated surface that has a well-defined PSD. We used the central 50 000 pixels per side section of this surface as our realization of a 100-μm by 100-μm surface. The surface was then ‘measured’ (see figures 3(a)–(c)) with scan sizes down to 1 μm on a side. Each ‘measurement’ had 500 by 500 pixels, therefore the pixel size scaled with scan size—as is common in measurement techniques such as AFM and optical profilometry. The measurement of the images was carried out by picking a random height value within each sub-pixel. Other ways to ‘sample’ each pixel were also tested (i.e. picking subpixels at equal distances and finding the highest point in each pixel), but these details did not affect the results.

Figure 5(d) shows the PSD stitched from three individual measurements at 100 μm, 10 μm and 1 μm image size taken in the middle of the surface. These results show first that sub-images must be tilt-compensated to eliminate errors at low q, as are evident in the inset image. The surfaces are untitled by subtracting an inclined plane \( h(x, y) = ax + by + c \) to the topography such that the average slope of the surface is zero. While such a tilt compensation is commonplace in AFM measurements, there are other measurements where it is not typically done. An example is the technique recommended in [59] where one very large surface measurement is taken using optical reflectometry—then this large measurement is divided into sub-images for individual analysis. Even in this case, the sub-images must be tilt compensated so that each one is flat. When the sub-images are correctly windowed and correctly tilt-compensated, the overlay of PSDs is nearly perfect.

Finally, experimental measurements will almost always be limited to a range of wavevectors, such that the ‘true’ surface PSD can never be fully determined. To account for this, any surfaces that show self-affine scaling over all or part of the spectrum can be fitted with a power-law. This enables the use of equations (8)–(11) to compute \( h_{rms} \) and \( h_{rms}' \) for known bounds on long- and short-wavelength cutoffs. For example, a lower bound on \( q_{\ell} \) can be obtained from the measurement at the lowest resolution (if the PSD levels off) and an upper bound to \( q_{s} \) is \( 2 \pi /L \) where \( L \) is the maximum length of the geometry of the surface under investigation. Similarly, a lower bound on \( q_{s} \) is \( 2 \pi /a_{0} \) where \( a_{0} \) is some characteristic interatomic spacing. The upper bound on \( q_{s} \) is given by the resolution of the instrument used for the highest resolution topography measurement; a possible measure for scanning probe techniques is described in section 4. These upper- and lower bounds for the cutoffs directly lead to upper and lower bounds on the scalar roughness parameters. Ideally, these bounds should be reported rather than a single value in all roughness investigations.

| Technique                  | Approx. maximum analysis size | Approx. lateral resolution limit | Frequency range | Advantages                                      | Limitations                                                                 |
|----------------------------|-------------------------------|---------------------------------|-----------------|------------------------------------------------|-----------------------------------------------------------------------------|
| Stylus profilometry        | 50 mm (200 mm with stitching) | 1 μm                            | \( 10^5 \left( 10^4 \right)^{-1} \text{ m}^{-1} \) | Sub-nanometer height resolution. Unaffected by sample’s optical properties | Contact method, can cause damage. Measurement is time-intensive. Tip shape can introduce artifacts |
| Optical profilometry       | 5 mm (100 mm with stitching)  | 1 μm                            | \( 10^5 \left( 10^2 \right)^{-1} \text{ m}^{-1} \) | Rapid, non-contact. Sub-nanometer height resolution | Diffraction-limited spatial information. Difficult to use on very rough surfaces. Artifacts from transparent thin films |
| Light and x-ray scattering | 5 mm                          | 10 nm                           | \( 10^5 \left( 10^9 \right)^{-1} \text{ m}^{-1} \) | Rapid, non-contact. Insensitive to vibration | Relies on models and assumptions to relate scattered beam to surface spectrum. Does not recover real-space topography |
| Scanning probe microscopy  | 100 μm                        | 1 Å                             | \( 10^{5} \left( 10^{11} \right)^{-1} \text{ m}^{-1} \) | High-resolution                          | Small areas. Sensitive to vibration. Tip shape can introduce artifacts |

Table 1. Bandwidth limitations of various techniques for measuring surface topography. The ranges indicated cannot be achieved in a single measurement, but rather represent the ultimate limits of the techniques if multiple measurements are performed from the highest scan size. Note that the frequency range for reliable data will be significantly reduced due to tip-size and noise artifacts, as discussed in section 4.
3.3. Strategies to reconstruct a multi-scale PSD

In light of the various factors discussed in this section, it is recommended that a tilt correction and a window (in this order) always be applied in every PSD calculation. We recommend that conclusions never be drawn about a surface from a single topography measurement. Surfaces may not be self-affine over all length scales and, even those that are, will have high and low-frequency cutoffs that affect the calculation of roughness parameters. Rather, it is recommended to compute a ‘master PSD’ of the surface by combining PSDs from many topography measurements over as large a frequency range as possible. Not only should this master PSD include multiple surfaces and multiple locations per surface, but it should also include multiple different analysis techniques and a variety of sampling sizes for each technique. Finally, once this master PSD is computed, it should be used to compute upper and lower bounds for $h_{\text{rms}}$, $h'_{\text{rms}}$, and $h''_{\text{rms}}$. These will serve as a guide as to whether more precise measurements are required for a given surface, and also will enable the calculation of uncertainty in surface properties predicted from contact models.

4. Challenge C: Accurately measuring topography at the smallest scales

Every measurement technique introduces artifacts in the measurement. Here we focus on artifacts introduced by scanning probe microscopy, because it provides the highest-resolution information of any conventional surface measurement technique and is commonly used to characterize surfaces at the nanoscale. AFM introduces many artifacts [71] into the measured data, including tip size effects, drift, acoustic and electronic noise, and image bow. In the context of prediction of surface properties, the most significant are the effect of tip size and instrument noise [72]. The purpose of this section is to introduce the particular significance of these problems with respect to scalar roughness parameters. We show the effects on our synthetic surfaces and discuss practical upper limits on the frequency range that can be measured reliably.
4.1. Limitations of scanning probe microscopy for rough surfaces at the small scale

The finite size of the tip presents two related, but distinct problems: tip convolution and feature deletion/creation. Tip convolution is the apparent blunting of sharp features because of the moving point-of-contact between the feature and the finite-size tip [73]. For example, the AFM measurement of an infinitely sharp spike will yield an inverse image of the tip itself. This effect has been extensively investigated [74–76] and algorithms have been developed to mathematically compensate, and reconstruct the unconvoluted surface topography [77–80]. However, feature deletion/creation occurs when the bluntness of the tip prevents the sampling of topography that is smaller than a certain spatial wavelength, and/or creates kinks in the data when there is a trench that is too narrow for the tip to reach the bottom [81]. While it is common in hard materials, it can also occur for soft samples when the surface fully conforms to the blunt tip, such that roughness features below a characteristic size cannot be sampled [82]. In these cases, there is no mathematical route to reconstruct the original surface from the measured data. The surface has not been accurately sampled and is indistinguishable from a range of other surfaces.

At present, there is no simple solution to solving the high-frequency problem for AFM. However, the investigator can determine the maximum frequency that can be reliably measured for a certain technique on a certain surface. This enables the determination of uncertainty on any quantities computed from the PSD.

The radius of the AFM probe determines the maximum spatial frequency that can be measured [72, 83, 84]. For a simple sine wave surface, the minimum wavelength \( \lambda_c \) that can be sampled by a spherical-ended tip of radius \( R_{\text{tip}} \) is given by [81, 85, 86]:

\[
\lambda_c = 2\pi R_{\text{tip}}
\]

(16)

where \( \chi \) is the amplitude of the sine wave.

Figure 6. Effect of tip radius. A 0.1 \( \mu \text{m} \) square synthetic surface containing 200 pixels per side has been created from a subsection of a periodic self-affine surface 0.5 \( \mu \text{m} \) in size. When the surface is scanned with an infinitely sharp tip (a), the original surface is recovered. When the surface is scanned with an ideal spherical tip of radius 40 nm (b), some blunting is readily apparent. The blunted surface is used (after the application of a Hann window) to compute a PSD (d). The PSD of the scanned surface shows a \( q^{-4} \) noise term at large wavevectors (small wavelengths). The asymptotic behavior is given by equation (17) and shown by the dashed line. The black solid line shows the input PSD that was used to generate these surfaces. Note that the transition from real data to artifact-prone data is not readily visible in the computed PSD. Rather it is up to the investigator to determine the reliability cutoff—using equation (19)—and to discard any PSD data for \( q \) above this cutoff.
4.2. Effect of instrumental white noise

Each measurement instrument introduces noise into the process. The source of noise is manifold and ranges from thermally-induced oscillations in cantilever-based measurements [87] to thermal and shot noise in the measurement electronics [88]. We here test the influence of noise on the measurements of PSDs by simply assuming uncorrelated white noise. This is an appropriate model for many sources of noise, such as thermal or shot noise.

White noise has a constant, wavelength-independent PSD. We introduce it into our synthetic surfaces by adding a random height, drawn from a Gaussian distribution with standard deviation $\sigma_{\text{noise}}$, to each topographic point. From equation (A.8) we immediately see that this leads to a constant contribution to the PSD with power $C_{\text{noise}}^i = l_i \sigma^2_{\text{noise}}$.

$$C_{\text{noise}}^{\text{i}} = l_i \sigma^2_{\text{noise}}. \quad (21)$$

It is interesting to note that the power depends not just on amplitude but also explicitly on measurement resolution. However, $\sigma_{\text{noise}}$ is a length which depends on the details of how the measured signal is converted into height information; this conversion may itself be affected by measurement resolution, care must therefore be taken in interpreting equations (21) and (22) with respect to their scaling with resolution.

To demonstrate the effect of white noise, we calculate the PSD of a synthetic surface with additional noise consisting of 2000 × 2000 pixels. For each pixel $i,j$ we draw a random height $\Delta h_{ij}$ from a Gaussian distribution with standard deviation $\sigma_{\text{noise}}$ and add $\Delta h_{ij}$ to the topography map. The corresponding PSDs are shown in figure 7; $\sigma_{\text{noise}}$ was set to 20% of the RMS height $h_{\text{rms}}$. The dashed lines in figure 7 show that white noise appears as a region of constant power for both 1D and 2D PSDs (panels (a) and (b), respectively), which is well-described by equations (21) and (22). When the resolution is decreased to 200 × 200 pixels, the power of
the noise increases as predicted. Note that for the low-resolution scan the noise intersects the power-law of the self-affine surface at a wavelength that is larger than the cutoff. The PSD then tapers off smoothly to constant power.

The fact that white noise leads to a region of constant power in 1D and 2D PSDs can be used to detect it. The ratio of equations (21) and (22) should be equal to the scan resolution in the y-direction, \( l_y \). Figure 7(c) shows this ratio as a function of wavevector \( q \) for our synthetic surface. The dashed line shows \( C^{iso} / C^{iso} + 3 \) intersects this line is the limit of reliability of the noisy data. Values at higher \( q \) are affected by instrumental noise.

4.3. Strategies to detect and mitigate small-scale artifacts

Care must be taken when measuring and analyzing spectral content at the smallest scales. Perhaps the most concerning aspect of this reliability issue is that PSDs can be mathematically computed for arbitrarily small scan sizes and correspondingly large wavevectors. In many cases, the data will appear normal and there will be no inherent indication of where this tip size ‘reliability cutoff’ lies. Further, the effect on the PSD after this cutoff is not predictable. Depending on the profile and the tip shape, it may smooth out or roughen the profile, and may change the shape of the PSD [81]. Therefore, when performing AFM, it is recommended to determine the tip radius (by direct imaging [44, 89] or by tip-reconstruction numerical algorithms [75, 79]) and then to use equations (19) and (20) to determine the maximum frequency that can be accurately measured using that combination of tip and surface. Then the noise must be analyzed, either qualitatively by looking for a level-off, or quantitatively by computing \( C^{iso} / C^{iso} + 3 \) to determine the limit of the measurement noise. Spectral content beyond these limits should neither be trusted, nor used for analysis. One can therefore specify only a lower bound for the high-frequency cutoff—and all properties that are computed from the PSD should reflect the uncertainty in this value. If higher-resolution information is required, then sharper tips and low-noise equipment must be used, but even the sharpest tips have a limit of approximately \( 3-5 \text{-nm radius} \). For most surfaces that are not atomically smooth, this limits the small-wavelength cutoff to \( >10 \text{ nm} \). For this reason, new approaches must be devised in the future for transcending the limits of conventional techniques and characterizing rough surfaces down to the smallest scales.

5. Conclusions

Recent analytical models and numerical simulations make predictions for functional properties, such as macroscopic contact properties (e.g., stiffness, contact area, and adhesion), on the basis of the power spectral density (PSD) of a surface. We have demonstrated three significant experimental challenges that hinder the application of these models to real-world surfaces—along with strategies to mitigate each one. First, there are several different well-accepted methods for the calculation of the PSD that result in different quantitative values. We reviewed these and showed how they relate to each other, and then discussed the considerations of using each. For analytical models of contact between randomly rough surfaces, the 2D PSD is the appropriate one to use—rather than the ‘pseudo-1D PSD’ that is computed by many software packages and recommended by international standards. Second, the analytical theories assume knowledge of the PSD of an infinite surface across a wide frequency range, while experimental measurements are plagued by edge effects and are necessarily limited to a narrow frequency band. We reviewed the use of windowing and stitching to reliably combine multiple measurements, and discussed the calculation of upper and lower bounds on the true values of the scalar roughness parameters for a surface. Third, common techniques for surface topography measurement provide inaccurate data in the highest-frequency regime. We showed the consequences of this issue for contact models and discussed the calculation of a ‘reliability cutoff’—beyond which measured PSD data should not be trusted, analyzed, or reported.

We have demonstrated these various considerations using computer-generated surfaces. This analysis has shown that various specific types of artifacts can be found in experimentally-measured PSDs:

- Not properly accounting for the aperiodicity of the data by using the appropriate windows introduces a component \( q^{-2} \) into the 1D PSD (\( q^{-3} \) in the 2D PSD). This artifact appears identical to self-affine scaling with a Hurst exponent \( H = 0.5 \), but can be avoided by computing the PSD with a properly normalized, radially symmetric window.
- Not properly accounting for surface tilt introduces an overestimation of the PSD at low wavevectors, causing a lack of overlap between PSD measurements from scans of different sizes. This artifact can be avoided through tilt-compensation when computing the PSD.
- Tip shape in AFM measurements introduces a spurious component \( q^{-4} \) into the 1D PSD (\( q^{-3} \) in the 2D PSD). This artifact can be partially mitigated by the use of sharper tips, but cannot be avoided entirely. We have proposed a parameter-free expression that allows AFM users to identify the region of the PSD that is unreliable and should not be reported or used for calculation.
- Instrumental noise leads to constant power at the highest \( q \). This artifact is easy to detect since it occurs in both 1D and 2D PSDs. We have proposed an approach for detecting the noise limit from the ratio of 2D and 1D PSDs.
Taken together, this article provides theoretical and practical guidance for the application of analytical roughness models to real-world surfaces.

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Appendix

A.1. Mathematical definition of the PSD

Many excellent references describe the calculation of the PSD, such as [32]. We will review the salient features here to establish the conventions used in this paper, and also to rigorously demonstrate the origins of some of the mathematical inconsistencies between the different types of PDSs which were mentioned in section 2.

We start by assuming that the surface topography is given by a continuous function \( h(x, y) \) with in-plane position \( x, y \) and periodicity \( L_x, L_y \) in both Cartesian directions. We note that the choice of a function \( h(x, y) \) excludes surfaces with overhangs that are sometimes observed experimentally. Due to the periodicity of \( h(x, y) \), the Fourier transform \( \hat{h}_{q_x, q_y} \) is nonzero at discrete lattice points only. This type of transform is commonly referred to as the Fourier series, but we will use the term transform throughout the rest of the document. (We denote continuous functions using a typical function designation, \( h(x, y) \), and we will represent discrete functions using subscripts, \( h_{x,y} \), here and elsewhere in this manuscript). The forward and inverse transforms in the plane are given by the expressions

\[
\hat{h}_{q_x, q_y} = \int_A h(x, y) e^{-i q_x x + q_y y} \, dx \, dy \quad (A.1)
\]

\[
h(x, y) = \frac{1}{L_x L_y} \sum_{q_x, q_y} \hat{h}_{q_x, q_y} e^{i q_x x + q_y y} \quad (A.2)
\]

where the integral is over the full area \( A = L_x L_y \) of the periodic topography profile and the sum runs between negative and positive infinity in steps of \( 2\pi/L_x \) and \( 2\pi/L_y \) in \( x \) and \( y \), respectively.

Some authors use a pre-factor of \( 1/\sqrt{A} \) in front of both transforms or a factor of \( 1/A \) in front of the forward transform. Using one of these other conventions will change some of the pre-factors in the equations throughout this paper. It does also influence the units of the Fourier transform and hence the values obtained for the PSD. It is therefore important to agree on a canonical formulation for the forward and inverse transforms.

Using equation (A.1) gives units of the Fourier transform \( \hat{h} \) as \((m^2)\). In most real-world situations \( h(x, y) \) is not described as a well-defined continuous function, but rather is measured only at discrete sites on a regular rectangular grid with ‘pixel’ size \( L_x \times L_y \). Here \( L_x = N_x/L_x \) and \( L_y = N_y/L_y \) where \( N_x \) and \( N_y \) is the number of grid points in the \( x \)- and \( y \)-direction, respectively. In this case, the integral becomes a sum over all lattice sites and the sum in the inverse transform runs (in steps of \( \Delta q_x = 2\pi/L_x \)) from \( q_x = 0 \) to \( 2\pi(N_x-1)/L_x \) rather than from \( q_x = -\infty \) to \( \infty \). Equations (A.1) and (A.2) then become the forward and inverse discrete Fourier transforms (DFT),

\[
\hat{h}_{q_x, q_y} = \frac{1}{L_x L_y} \sum_{x, y} h_{x,y} e^{-i q_x x + q_y y} \quad (A.3)
\]

\[
h_{x,y} = \frac{1}{L_x L_y} \sum_{q_x, q_y} \hat{h}_{q_x, q_y} e^{i q_x x + q_y y}. \quad (A.4)
\]

The most common numerical algorithm to compute the DFT is the fast Fourier transform (FFT) [90]. Because it is pervasively used, the terms FFT and DFT are often used synonymously. Note that in algorithmic implementations of equations (A.3) and (A.4), \( L_x = l_x = 1 \) and \( 1/L_x L_y = 1/N_x N_y \) where \( N_x \times N_y \) are the dimensions of the DFT grid.

We now define the 2D PSD as

\[
C_{q_x, q_y}^{2D} = |\hat{h}_{q_x, q_y}|^2 \quad (A.5)
\]

\( C_{q_x, q_y}^{2D} \) has units of \((m^4)\). Equation (A.5) removes all phase information from \( \hat{h}_{q_x, q_y} \) but retains its amplitude. Because the reciprocal space product \( g_{q_x, q_y} = \hat{h}^*_{q_x, q_y} \cdot \hat{h}_{q_x, q_y} \) (here \( * \) is the complex conjugate) becomes a convolution, \( g(x, y) = \int h^*(x-x', y-y') h(x', y') \, dx' \, dy' \) in real space, equation (A.4) is the Fourier transform of the height autocorrelation function.

We show below that this normalization in combination with equations (A.1) and (A.2) makes the PSD independent of sample size and allows comparison of PSDs measured over different areas. Equation (A.5) is compatible with the definition of the 2D PSD in SEMI MF1811[12]. Theoretical work by Persson uses a different normalization of the transform and the PSD, such that:

\[
C_{q_x, q_y}^{2D, P} = C_{q_x, q_y}^{2D} / 4\pi^2. \quad (A.6)
\]

An important equality that derives from the properties of the convolution is Parseval’s theorem, which relates the real-space power to the reciprocal-space power. Given the definition of the Fourier transform in equations (A.1) and (A.2), Parseval’s theorem has the form

\[
\int_A |h(x, y)|^2 \, dx \, dy = A^{-1} \sum_{q_x, q_y} |\hat{h}_{q_x, q_y}|^2 = \sum_{q_x, q_y} C_{q_x, q_y}^{2D} \quad (A.7)
\]

The PSD \( C_{q_x, q_y}^{2D} \) contains certain scalar roughness parameters as simple sum rules. These are in particular
the RMS roughness \( h_{\text{rms}} = \sqrt{\langle h^2 \rangle} \), RMS slope \( h'_{\text{rms}} = \sqrt{\langle (\nabla h)^2 \rangle} \) and RMS curvature \( h''_{\text{rms}} = \frac{1}{2} \sqrt{\langle (\nabla^2 h)^2 \rangle} \). By virtue of Parseval’s theorem, we find

\[
h'_{\text{rms}} = \frac{1}{A} \int h(x, y) \, dx \, dy = \frac{1}{A} \sum_{q_{x},q_{y}} q_{x} C_{q_{x}q_{y}}^{2D}(q_{x},q_{y}) \tag{A.8}
\]

\[
(h'_{\text{rms}})^2 = \frac{1}{A} \int (\nabla h(x, y))^2 \, dx \, dy = \frac{1}{A} \sum_{q_{x},q_{y}} q_{x}^2 + q_{y}^2 \left| \hat{h}_{q_{x}q_{y}} \right|^2 \tag{A.9}
\]

\[
h''_{\text{rms}} = \frac{1}{4A} \sum_{q_{x},q_{y}} q_{x} C_{q_{x}q_{y}}^{2D}(q_{x},q_{y}) \tag{A.10}
\]

Note that these expressions are independent of area \( A \) and hence well defined even in the limit \( A \to \infty \).

A.2. Relating the 2D PSD to that of a line profile

1D power spectra are much more common than the previously discussed 2D version due to their use in electrical engineering with time-varying electrical signals. For the same surface \( h(x, y) \) the PSD of a line profile can be obtained from the 1D Fourier series

\[
\hat{h}_{q_{x}}(y) = \int_{L_{x}} h(x, y) e^{-i2\pi q_{x}x} \, dx \tag{A.15}
\]

\[
h(x, y) = \int_{L_{y}} \hat{h}_{q_{x}}(y) e^{i2\pi q_{x}x} \, dq_{x} \tag{A.16}
\]

The 2D Fourier transform given in equations (A.1) and (A.2) is simply the consecutive application of this formula in the \( x- \) and \( y- \) direction. The 1D PSD is then given by

\[
C_{q_{x}}^{1D}(y) = L_{x}^{-1} \left| \hat{h}_{q_{x}}(y) \right|^2 \tag{A.17}
\]

which has units of \((\text{m}^3)\) and depends explicitly on \( y \), i.e. the line of the scan. PSDs for line scans are typically reported as averages over multiple scans, denoted with a line over \( C \),

\[
\overline{C}_{q_{x}}^{1D} = L_{y}^{-1} \int \hat{h}_{q_{x}}^{1D}(y) dy \tag{A.18}
\]

Note that in the main text, the line over the \( C \) is dropped for brevity. By virtue of Parseval’s theorem, we can express this averaged 1D PSD in terms of the 2D PSD,

\[
\overline{C}_{q_{x}}^{1D} = L_{y}^{-1} \sum_{q_{y}} C_{q_{x}q_{y}}^{2D} \tag{A.19}
\]

or for continuous \( q_{x}, q_{y} \),

\[
\overline{C}_{q_{x}}^{1D}(q_{y}) = \frac{1}{2\pi} \int_{-\infty}^{\infty} C_{q_{x}q_{y}}^{2D}(q_{x},q_{y}) \, dq_{x} \tag{A.20}
\]

Note that a 2D power-spectrum can be reconstructed from the 1D power spectrum if we assume that the surface is isotropic, i.e. if \( C_{q_{x}q_{y}}^{2D}(q_{x},q_{y}) = C_{\text{iso}}^{2D}(q_{x},q_{y}) \), with \( q = \sqrt{q_{x}^{2} + q_{y}^{2}} \) and \( dq_{x} dq_{y} = q_{x} dq_{x} dq_{y} \). Substituting \( q_{x} \) for \( q \) in equation (A.20) gives

\[
\overline{C}_{q_{x}}^{1D}(q_{y}) = \frac{1}{\pi} \int_{q}^{\infty} \frac{q C_{\text{iso}}^{2D}(q_{x},q_{y}) \, dq_{x}}{\sqrt{q_{x}^{2} - q^{2}}} \tag{A.21}
\]

Inversion of this expression yields

\[
C_{\text{iso}}^{2D}(q_{x},q_{y}) = -2 \int_{q}^{\infty} \frac{1}{\sqrt{q_{x}^{2} - q^{2}}} \overline{C}_{q_{x}}^{1D}(q_{y}) \, dq_{x} \tag{A.22}
\]

As discussed in the following section, for self-affine surfaces this is approximately equal to \( \pi \overline{C}_{q_{x}}^{1D}(q_{y}) \).

A.3. Power spectra for self-affine random surfaces

The 2D PSD of a self-affine, randomly rough surface with Hurst exponent \( H \) follows the power-law expression \( C_{\text{iso}}^{2D}(q) \propto q^{-2-2H} \) (see also equation (7)). Self-affinity implies a certain scaling behavior that is
encoded into this power-law. If all lengths are scaled by a factor of $C$, then $q' = q/C$ and

$$C_{iso}(q')d^3q' = C_{iso}(q/C)d^3q/C^2 = C^{2H}C_{iso}(q)d^3q,$$

i.e. heights need to be rescaled by $C^H$ to give a surface with the same (statistical) roughness. This is the essence of self-affinity [36]. The Hurst [56] exponent $H$ of the power-law is related to the fractal dimension $D = 3 - H$. The values of $H$ are typically characteristic for the process that formed the surfaces. Typical values for $H$ on scales from atoms to mountains are between 0.7 and 0.9 (e.g. [50,52]).

From equations (A.11)–(A.13) we can easily derive the prefactor $C_0$ in $C_{iso}(q)$ = $C_0q^{-2+2H}$ (see also equation (7)) given either $h_{rms}$ and $q_L$, or $q_{rms}$ and $q_L$. Under the assumption of scale separation $q_s \gg q_L$ we get:

$$C_{iso}(q) = 4\pi(1 - H)\left(h_{rms}/q^2_L\right)^{2-2H},$$

(A.24)

$$C_{iso}(q) = 4\pi\alpha H\left(h_{rms}/q^2_L\right)^{2-2H}$$

with

$$\alpha = \left\{ \begin{array}{ll}
1 & \text{for } q_s = q_L, \\
(1 + H)^{-1} & \text{for } q_s \ll q_L.
\end{array} \right.$$

(A.25)

It is instructive to compute the PSD of a line scan of such an ideal isotropic and self-affine surface. We apply equation (A.20) to convert the 2D PSD with $q_s = q_L$ into

$$C_{ID}(q_s) = \frac{C_0}{\pi} q_s^{-2+2H} \left( \frac{1}{2} H^2 - \frac{3}{2} - 1 - \left(\frac{q_s}{q_L}\right)^2 \right) \sqrt{\frac{q_s}{q_L}} - 1,$$

(A.26)

where $F_1$ is the Gauss hypergeometric function. The expression takes a particularly simple form for $H = 1/2$ where $F_1(1/2; 3/2; 3/2; z) = 1/\sqrt{1 - z}$. This yields

$$C_{ID}(q_s) \approx \frac{C_0}{\pi} q_s^{-2+2H} \sqrt{\frac{1 - \left(\frac{q_s}{q_L}\right)^2}{\left(\frac{q_s}{q_L}\right)^2}},$$

(A.27)

and turns out to be an excellent approximation to equation (A.25) even for $H = 1/2$. The overall scaling is, as expected, $q_s^{-1-2H}$ i.e. same as the 2D PSD but with a power increased by one. The additional factor $\sqrt{1 - (q_s/q_L)^2}$ tapers the function off to exactly zero at $q_s = q_L$. The deviation from $x^{-1-2H}$ starts to become significant at about $q_s/q_L > 0.1$.

We can therefore express

$$C_{ID}(q_s) \approx \frac{q_s}{\pi} C_{iso}(q_s) \sqrt{1 - \left(\frac{q_s}{q_L}\right)^2},$$

(A.28)

which forms the basis for the pseudo-1D PSD defined in equation (5).

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