# Using the Wigner-Ibach surmise to analyze terrace-width distributions: history, user's guide, and advances 

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ABSTRACT A history is given of the applications of the simple expression generalized from the surmise by Wigner and also by Ibach to extract the strength of the interaction between steps on a vicinal surface, via the terrace width distribution (TWD). A concise guide for use with experiments and a summary of some recent extensions is provided.

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## 1 Introduction

Measurement of the terrace width distribution (TWD) $\check{P}(\ell)$ of vicinal surfaces is now used routinely to find the dimensionless strength $\tilde{A}$ of the elastic repulsion between steps. The University of Maryland [1] and the Forschungszentrum Jülich [2] have been at the vanguard of this progress. Use of an extension of the Wigner surmise from random matrix theory has resolved ambiguities on how best to estimate $\tilde{A}$ from the variance. This paper discusses the history of this development and the crucial role played by Harald Ibach in divining from physical insight an analytic expression for the TWD associated with the special case $\tilde{A}=0$ that is essentially the same as the simple expression that Wigner surmised describes (albeit, ultimately, not exactly) the distribution of the energy differences of adjacent levels in a nucleus when the coupling has unitary symmetry. This insight spawned my long collaborative effort with FZ Jülich to develop generalizations appropriate to arbitrary vicinal surfaces and to corroborate the viability of the resulting formulation to account for extensive experimental data. In addition to a personal history of this progress (with a few new results), I collect (with enhancements) in one place several tables from publications and present a concise "User's Guide" for applying the formalism and a short discussion of some recent developments, e.g., for azimuthally misoriented vicinal surfaces and for nonequilibrium situations. Space limitations preclude fuller discussions or an authoritative update on experimental progress. Three of the four figures are heretofore unpublished. Also interwoven are comments stimulated by several penetrating

[^0]questions, many by Harald Ibach, that arose during talks I gave in Germany that forced me to sharpen my thinking. These are topics typically skirted in publications as being well-established, but my experience is that they are not generally well understood. The use of "we" below is not a polite affectation but instead reflects the crucial role played by my many collaborators in this extended series of studies.

To set the stage (cf. Fig. 1), the direction perpendicular to the terraces (which are densely-packed facets) is typically called $\hat{z}$. In "Maryland notation" the normal to the vicinal surface lies in the $x-z$ plane, and the distance $\ell$ between steps is measured along $\hat{x}$, while the steps run along the $\hat{y}$ direction. In the simplest and usual approximation, the repulsions between adjacent steps arise from two sources: an entropic or steric interaction due to the physical condition that the steps cannot cross, since overhangs cannot occur in nature. The second comes from elastic dipole moments due to local atomic relaxation around each step, leading to frustrated lateral relaxation of atoms on the terrace plane between two steps. Both interactions are proportional to $1 / \ell^{2}$.


FIGURE 1 Plots of generic lattice configurations and associated TWDs, to illustrate essential features, as discussed in text, of steps on a vicinal surface (with no energetic interactions). Top left: "perfect" cleaved crystal. Top right: straight steps placed randomly, corresponding to a strictly 1D model. Lower left: meandering steps in a TSK model. Lower right: TWDs $P(s)$ associated with these distributions. The downward arrow emphasizes the greatly decreased chance of finding steps at very small separation when they meander, as compared to straight steps, due to the entropic repulsion


FIGURE 2 Illustration of how entropic repulsion and energetic interactions combine, plotted vs. the dimensionless energetic interaction strength $\tilde{A} \equiv a \tilde{\beta} /\left(k_{\mathrm{B}} T\right)^{2}$. The dashed straight line is just $\tilde{A}$. The solid curve above it is the combined entropic and energetic interactions, labeled $\tilde{A}_{\text {eff }}$ for reasons explained below. The difference between the two curves at any value of the abscissa is the dimensionless entropic repulsion for that $\tilde{A}$. The decreasing curve, scaled on the right ordinate, is the ratio of this entropic repulsion to the total dimensionless repulsion $\tilde{A}_{\text {eff }}$. It falls monotonically with $\tilde{A}$, passing through unity at $\tilde{A}=0$. See the discussion accompanying (16) for more information and explicit expressions for the curves

To illustrate the essence of $\check{P}(\ell)$, we consider in Fig. 1 its shape for 3 idealized configurations. There is just one characteristic length in the $\hat{x}$ direction, namely the average step separation $\langle\ell\rangle$. (Contrary to widespread misconception, $\langle\ell\rangle$ need not be a multiple of, or even simply related to, the substrate lattice spacing. It is the step height times $\cot \varphi$, where $\varphi$ is the arbitrary misorientation, as shown in Fig. 1.) Therefore, we consider $P(s)=\langle\ell\rangle^{-1} \check{P}(\ell)$, where $s \equiv \ell /\langle\ell\rangle$, a dimensionless length. For a "perfect" cleaved crystal, $P(s)$ is just a spike $\delta(s-1)$. If, as intrinsic to 1 D models, the steps are imagined as uncooked spaghetti dropped at any position with probability $1 /\langle\ell\rangle, P(s)$ is a Poisson distribution $\exp (-s)$. Actual steps do meander, as one study most simply in a terrace-step-kink (TSK) model. In this model, the only excitations are kinks (with energy $\varepsilon$ ) along the step. (This is a good approximation at low temperature $T$ since adatoms or vacancies on the terrace cost several $\varepsilon$ [ $4 \varepsilon$ in the case of a simple cubic lattice]. The entropic repulsion due to step meandering dramatically decreases the probability of finding adjacent steps at $\ell \ll\langle\ell\rangle$. To preserve the mean of one, $P(s)$ must also be smaller than $\exp (-s)$ for large $s$.

If there is an additional energetic repulsion $A / \ell^{2}$, the magnitude of the step meandering will decrease, narrowing $P(s)$. As $A \rightarrow \infty$, the width approaches a delta function. Note that the energetic and entropic interactions do not simply add. In particular, there is no negative (attractive) value of $A$ at which the two cancel each other. Thus, for strong repulsions, steps rarely come close, so the entropic interaction plays a smaller role, while for $A<0$, the entropic contribution increases, as illustrated in Fig. 2 and worked out in detail below.

## 2 History

Investigation of the interaction between steps on vicinal surfaces is a core part of the flourishing field of exploring the properties of these technologically important and scientifically rich systems, as discussed in several excellent
reviews [1-3]. The earliest studies seeking to extract $A$ from TWDs used the mean-field-like Gruber-Mullins [4] approximation, in which a single active step fluctuates between two fixed straight steps $2\langle\ell\rangle$ apart. Then the energy associating with the fluctuations $x(y, t)$ is
$\Delta \mathcal{E}=-\beta(0) L_{y}+\int_{0}^{L_{y}} \beta(\theta(y)) \sqrt{1+\left(\frac{\partial x}{\partial y}\right)^{2}} \mathrm{~d} y$,
where $\beta$ is the step free energy per length (or line tension [5] ${ }^{1}$ ) for a step at orientation $\theta$ relative to the mean direction of the step (and the direction of the fixed, bounding steps), and $L_{y}$ is the size of the system along the mean step direction (i.e., the step length with no kinks). We expand $\beta(\theta)$ as the Taylor series $\beta(0)+\beta^{\prime}(0) \theta+\frac{1}{2} \beta^{\prime \prime}(0) \theta^{2}$ and recognize that the length of the line segment has increased from $\mathrm{d} y$ to $\mathrm{d} y / \cos \theta \approx \mathrm{d} y(1+$ $\left.\frac{1}{2} \theta^{2}\right)$. For close-packed steps, for which $\beta^{\prime}(0)=0$, it is well known that (using $\theta \approx \tan \theta=\partial x / \partial y$ ),
$\Delta \varepsilon \approx \frac{\tilde{\beta}(0)}{2} \int_{0}^{L_{y}}\left(\frac{\partial x}{\partial y}\right)^{2} \mathrm{~d} y, \quad \tilde{\beta}(0) \equiv \beta(0)+\beta^{\prime \prime}(0)$,
where $\tilde{\beta}$ is the step stiffness [6]. Most treatments gloss over the fact that the stiffness has the same definition for steps with arbitrary in-plane orientation. The key point is that to create such steps, one must apply a "torque" [7] which exactly cancels the linear term $\beta^{\prime}(0) \theta$. Stasevich [8] provides a more formal proof.

Since $x(y)$ is taken to be a single-valued function that is defined over the whole domain of $y$, the 2 D configuration of the step can be viewed as the worldline of a particle in 1D by recognizing $y$ as a time-like variable. Since the steps cannot cross, these particles can be described as 1D fermions. We also see from (2) that in the $(1+1) \mathrm{D}$ formulation, $\partial x / \partial y \rightarrow \partial x / \partial t$, with the form of a velocity, so that the stiffness plays the role of a mass; indeed, it serves as the inertial parameter of steps in this fermion perspective (though not with regard to actual dynamics in response to external forces [9]). Moreover, stiffness is one of the three ingredients of the very-successful step-continuum model [1].

Pursuing this analogy for polymers in 2D, de Gennes [10] showed nearly four decades ago that this problem could be mapped into the Schrödinger equation in 1D, with the thermal energy $k_{\mathrm{B}} T$ replacing $\hbar$. Then in the Gruber-Mullins approximation [4], the step with no energetic interactions becomes a particle in a 1 D infinite well of width $2\langle\ell\rangle$, with well-known groundstate properties
$\psi_{0}(\ell) \propto \sin \left(\frac{\pi \ell}{2\langle\ell\rangle}\right) ; P(s)=\sin ^{2}\left(\frac{\pi s}{2}\right) ; E_{0}=\frac{\left(\pi k_{\mathrm{B}} T\right)^{2}}{8 \tilde{\beta}\langle\ell\rangle^{2}}$.

Thus, it is the kinetic energy of the ground state in the quantum model that corresponds to the entropic repulsion (per length)

[^1]of the step. In the exact solution for the free energy expansion of the equilibrium crystal shape [11], the numerical coefficient in the corresponding term is $\frac{1}{6}$ rather than $\frac{1}{8}$. Note that $P(s)$ peaks at $s=1$ and vanishes for $s \geq 2$.

Suppose, next, that there is an energetic repulsion $U(\ell)=$ $A / \ell^{2}$ between steps. In the 1 D Schrödinger equation, the prefactor of $-\partial^{2} \psi(\ell) / \partial \ell^{2}$ is $\left(k_{\mathrm{B}} T\right)^{2} / 2 \tilde{\beta}$; hence, $A$ only enters the problem in the dimensionless combination $\tilde{A} \equiv$ $A \tilde{\beta} /\left(k_{\mathrm{B}} T\right)^{2}[12]^{2}$. In the Gruber-Mullins picture, the potential (per length) experienced by the single active particle is (with $\ell \equiv \ell-\langle\ell\rangle)$ :

$$
\begin{align*}
\tilde{U}(\check{\ell}) & =\frac{\tilde{A}}{(\check{\ell}-\langle\ell\rangle)^{2}}+\frac{\tilde{A}}{(\check{\ell}+\langle\ell\rangle)^{2}} \\
& =\frac{2 \tilde{A}}{\langle\ell\rangle^{2}}+\frac{6 \tilde{A} \check{\ell}^{2}}{\langle\ell\rangle^{4}}+\frac{10 \tilde{A} \check{\ell}^{4}}{\langle\ell\rangle^{6}}+\ldots \tag{4}
\end{align*}
$$

The first term is just a constant shift in the energy. For $\tilde{A}$ sufficiently large, the particle is confined to a region $|\check{\ell}| \ll\langle\ell\rangle$, so that we can neglect the fixed walls and the quartic term, reducing the problem to the familiar simple harmonic oscillator, with the solution:
$\psi_{0}(\ell) \propto \mathrm{e}^{-\check{\ell}^{2} / 4 w^{2}} ; P_{\mathrm{G}}(s) \equiv \frac{1}{\sigma_{\mathrm{G}} \sqrt{2 \pi}} \exp \left[-\frac{(s-1)^{2}}{2 \sigma_{\mathrm{G}}^{2}}\right]$,
where $\sigma_{\mathrm{G}}=(48 \tilde{A})^{-1 / 4}$ and $w=\sigma_{\mathrm{G}}\langle\ell\rangle$.
For $\tilde{A}$ of 0 or 2, it turns out, as we shall see, that the TWD can be computed exactly. For these cases, (3) and (5), respectively, provide serviceable approximations. It is (5) that is prescribed for analyzing TWDs in the most-cited resource on vicinal surfaces [1]. Indeed, it formed the basis of initial successful analyses of experimental STM data [13]. However, it has some notable shortcomings. Perhaps most obviously, it is useless for small but not vanishing $\tilde{A}$, for which the TWD is highly skewed, not resembling a Gaussian, and the peak, correspondingly, is significantly below the mean spacing. For large values of $\tilde{A}$, it significantly underestimates the variance or, equivalently, the value of $\tilde{A}$ one extracts from the experimental TWD width: Ihle et al. [14] point out that in the Gruber-Mullins approximation the TWD variance is the same as that of the active step, since the neighboring step is straight. For large $\tilde{A}$ the fluctuations of the individual steps on an actual vicinal surface become relatively independent, so the variance of the TWD is the sum of the variance of each, i.e., twice the step variance. Given the great (quartic) sensitivity of $\tilde{A}$ to the TWD width, this is problematic. As experimentalists acquired more high-quality TWD data, other approximation schemes were proposed, all producing Gaussian distributions with widths $\propto \tilde{A}^{-1 / 4}$, but with proportionality constants notably larger than $48^{-1 / 4}=0.38$.

For the "free-fermion" $(\tilde{A}=0)$ case, Joós et al. [15] developed at the beginning of the 1990's a sequence of analytic approximants to the exact but formidable expression [16,17] for $P(s)$. The procedure is based on a discrete version of the problem, representing the fermion-like steps in terms of second-quantized operators and taking note that the TWD is

[^2]not just a pair-wise step-step correlation function but actually a many-particle correlation function in which we demand that no step lie between the steps at 0 and $\ell$. The $n$th approximant behaves well at smaller $s$ but eventually passes through 0 around $s=n$ and then behaved non-physically. Specifically, the leading behavior of each approximant is $1-$ $[\sin (\pi s) /(\pi s)]^{2} \approx \frac{1}{3}(\pi s)^{2}$. In the asymptotic region, a different approach shows leading behavior for large $s$ is $P(s) \propto$ $s^{7 / 4} \exp \left[-\frac{1}{8}(\pi s)^{2}\right]$, where the proportionality constant has the numerical value $\sim 9.46$. (The analytic form is in [15].) This expression works to surprisingly small $s$, reproducing the peak semi-quantitatively. (It would have gone to even smaller $s$ than indicated in Fig. 4 of [15] had the two leading asymptotic corrections not been included!) Finally, [15] as well as a slightly earlier paper [18] draw the analogy between the TWD of vicinal surfaces and the distribution of spacings between interacting (spinless) fermions on a ring, the CalogeroSutherland model [19-22], which in turn for three particular values of the interaction - in one case repulsive $(\tilde{A}=2)$, in another attractive $\left(\tilde{A}=-\frac{1}{4}\right)$, and lastly the free-fermion case ( $\tilde{A}=0$ ) - could, astonishingly, be solved exactly by connecting to random matrix theory [17, 23, 24]; Fig. 5 of [15] depicts the three resulting TWDs.

This was the state of affairs when H. Ibach presented me (seemingly sometime during his sabbatical stay in College Park during 1996-1997) with Fig. 3, in which he achieved outstanding agreement with [numerical approximation visually indistinguishable from] the exact solution by plotting what essentially amounts to
$P_{2}(s)=\frac{32}{\pi^{2}} s^{2} \exp \left(-\frac{4}{\pi} s^{2}\right)$,
where the subscript of $P$ refers to the exponent of $s$, and the pair of numerical factors ensure that the distribution is normalized and has a mean of one. He challenged me to explain


FIGURE 3 The graph for the expression deduced by H. Ibach for the TWD of steps with no energetic repulsion (solid curve), essentially (6) surmised by Wigner, while the points are generated using the formalism for the approximants, given in [15]
his insight, and Fig. 3 graced my office wall, greeting me each morning, for well over a year. Not hearing back from me, he put the remarkable expression (albeit with some typos that obscured its potency) into [25]. On Feb. 2, 1998, Safi Bahcall gave an intriguing condensed-matter seminar at University of Maryland on "Superconductivity and Random Matrices" (cf. [26]). In subsequent discussions he rifled through a preprint of Guhr et al.'s seminal review of random matrix theory [27], and I spotted Fig. 12 therein, noting the similarity to Fig. 5 in [15], as well as the accompanying discussion of the Wigner surmise, their equation (3.50) being the first I had seen of what will be (9) below.

The other two curves correspond to corresponding Wigner surmises for those cases:
$P_{4}(s)=\left(\frac{64}{9 \pi}\right)^{3} s^{4} \exp \left(-\frac{64}{9 \pi} s^{2}\right) ;$
$P_{1}(s)=\frac{\pi}{2} s^{1} \exp \left(-\frac{\pi}{4} s^{2}\right)$.
In random matrix literature the exponent of $s$, viz. 1,2 , or 4 , is called $\beta$, due to an analogy with inverse temperature in one justification. To avoid possible confusion with the step free energy per length $\beta$ or the stiffness $\tilde{\beta}$ for vicinal surfaces, we have called it instead by the Greek symbol that looked most similar, $\varrho$. The Appendix offers transparent arguments on how the three kinds of symmetry lead to the associate exponents 1 , 2, and 4. As seen most clearly by explicit plots, e.g., Fig. 4.2a of Haake's text [28], $P_{1}(s), P_{2}(s)$, and $P_{4}(s)$ are excellent approximations of the exact results for orthogonal, unitary, and symplectic ensembles, respectively, and these simple expressions are routinely used when confronting experimental data in a broad range of physical problems [27,28]. (The agreement is particularly outstanding for $P_{2}(s)$ and $P_{4}(s)$, which are the germane cases for vicinal surfaces: the variance is $1 \%$ below and $0.4 \%$ above the exact values, respectively; for $P_{1}(s)$ it is $4-1 / 2 \%$ below. This agreement is significantly better than any other approximation (cf. Table 2 of [29]) and far better than the Gruber-Mullins approximation, as depicted in Fig. 1 of [30], Fig. 1 of [31], and Fig. 2 of [32].)

With the values $\varrho=1,2$, and 4 , the three specific expressions (6)-(8) comprising the so-called Wigner surmise [27, 28] can be written as a single formula

$$
\begin{equation*}
P_{\varrho}(s)=a_{\varrho} s^{\varrho} \exp \left(-b_{\varrho} s^{2}\right) \tag{9}
\end{equation*}
$$

where the constants $b_{\varrho}$, which fixes its mean at unity, and $a_{\varrho}$, which normalizes $P(s)$, are
$b_{\varrho}=\left[\frac{\Gamma\left(\frac{\varrho+2}{2}\right)}{\Gamma\left(\frac{\varrho+1}{2}\right)}\right]^{2}, \quad a_{\varrho}=\frac{2\left[\Gamma\left(\frac{\varrho+2}{2}\right)\right]^{\varrho+1}}{\left[\Gamma\left(\frac{\varrho+1}{2}\right)\right]^{\varrho+2}}=\frac{2 b_{\varrho}^{(\varrho+1) / 2}}{\Gamma\left(\frac{\varrho+1}{2}\right)}$.

As noted above the Calogero-Sutherland model provides a connection between random matrix results, notably the Wigner surmise, and the distribution of spacings between fermions in 1D interacting with dimensionless strength $\tilde{A}$. Specifically,
$\tilde{A}=\frac{\varrho}{2}\left(\frac{\varrho}{2}-1\right) \Leftrightarrow \varrho=1+\sqrt{1+4 \tilde{A}}$.

For an arbitrary system, there is no reason that $\tilde{A}$ should take on one of the three special values. However, it seems impossible to generalize the arguments in the Appendix to general values. For our purposes, the obvious solution is to use (11) for arbitrary $\varrho$ or $\hat{A}$. Curiously, this form has not been applied in conventional investigations involving RMT, but instead other analytic phenomenological expressions, e.g., those proposed by Brody [37] ${ }^{3}$ and by Izrailev [38] ${ }^{4}$ are used (cf. [27]). Such applications typically involves mixtures of systems of two of the symmetries; neither they nor other models involving crossover between well-defined symmetries [39] are analogous to systems based on the Calogero-Sutherland model for arbitrary $\varrho$.

For arbitrary $\varrho$ there is no symmetry-based justification of distribution based on the Wigner surmise of (9). Nonetheless, we have argued that it provides a viable, arguably optimal interpolation scheme between the two special values of $\varrho$ and also out to the Grenoble expression for nearly-infinite repulsion [30,31]; we have also used it successfully to analyze experimental data [31, 40, 41]. For brevity, we refer hereafter to this set of formulas, (9) and (10) as the GWD (generalized Wigner surmise); elsewhere [29, 31, 41] we have called it CGWD (continuum generalized Wigner distribution).

The moments of the GWD can be expressed simply in terms of $b_{\varrho}$ :
$\mu_{n}^{\prime} \equiv \int_{0}^{\infty} a_{\varrho} s^{\varrho+n} \exp \left(-b_{\varrho} s^{2}\right) \mathrm{d} s=\frac{\Gamma\left(\frac{\varrho+1+n}{2}\right)}{b_{\varrho}^{n / 2} \Gamma\left(\frac{\varrho+1}{2}\right)}$.
The variance $\sigma_{\mathrm{W}}^{2}=\mu_{2}=\mu_{2}^{\prime}-1$ of the GWD is then
$\sigma_{\mathrm{W}}^{2}=\frac{\varrho+1}{2 b_{\varrho}}-1 \approx \frac{1}{2 \varrho}-\frac{3}{8 \varrho^{2}}+\frac{3}{16 \varrho^{3}}-\frac{7}{384 \varrho^{4}}+\mathcal{O}\left(\varrho^{-5}\right)$,
for large values of $\varrho$, as given in equation (A8) of [41]. Note that $b_{\varrho}$ cancels in the ratio:
$\frac{\mu_{n}^{\prime}}{\left(\mu_{2}^{\prime}\right)^{n / 2}}=\frac{\Gamma\left(\frac{\varrho+1+n}{2}\right)}{\left(\frac{\varrho+1}{2}\right)^{n / 2} \Gamma\left(\frac{\varrho+1}{2}\right)}$.
Similarly, the peak of $p_{\varrho}(s)$ (i.e., its mode) occurs at
$s_{\max }=\left(\frac{\varrho}{2 b_{\varrho}}\right)^{1 / 2} \approx 1-\frac{\mathrm{e}^{-\varrho}}{4}-\frac{1}{4 \varrho}$.
For the special values of $\varrho$, the peak positions are tabulated in Table 1. We further note that $s_{\max }=0.96,0.97$, and 0.98 occur for $\varrho=6.1,8.2$, and 12.4 , respectively. When dealing with experimental data, such shifts from $s_{\max }=1$ would be difficult to distinguish.

While there are some formal justifications for the GWD as an optimal description of TWDs for arbitrary $\tilde{A}$, arguably the most convincing argument is a comparison of the predicted

[^3]
${ }^{\mathrm{a}}$ Covariance of adjacent terrace widths: $\operatorname{cov}\left(s_{1}, s_{2}\right) \equiv \frac{\left\langle\left(s_{1}-\left\langle s_{1}\right\rangle\right)\left(s_{2}-\left\langle s_{2}\right\rangle\right)\right\rangle}{\left.\left[\left\langle\left(s_{1}-\left\langle s_{1}\right\rangle\right)^{2}\right\rangle\left(s_{2}-\left\langle s_{2}\right\rangle\right)^{2}\right\rangle\right]^{1 / 2}}=\sigma^{-2}\left[\left\langle s_{1} s_{2}\right\rangle-1\right]=-1+\left\langle\left(s_{1}+s_{2}-2\right)^{2}\right\rangle / 2 \sigma^{2}$
TABLE 1 Tabulation of various measurable properties of terrace-width distributions $P(s)$ [where $s$ is the terrace width normalized by its average value] based on exact results at the three soluble values of the dimensionless interaction strength $\tilde{A}$, the corresponding generalized Wigner distribution (GWD) expression, and the various Gaussian approximations: Gruber-Mullins (GM), modified Grenoble [14, 30, 33] (G), and Saclay [34](S). (In the original Grenoble approximation $[14,33], \sigma^{2} \propto \tilde{A}^{-1 / 2}$ rather than $\tilde{A}_{\text {eff }}^{-1 / 2} \equiv 2 / \varrho$, but with the same prefactor as indicated in this table.) $\mathrm{SHO} \Rightarrow$ simple harmonic oscillator, i.e. uniformly spaced steps (energies). The extreme case $\varrho=0$, for which exact results are trivial, is included to dramatize trends in $\varrho$ [35] ${ }^{6}$. As $\varrho$ increases, the TWD becomes narrower, more symmetric, and more nearly Gaussian. Anticorrelations of neighboring terrace-width fluctuations increase. For the three exactly-solvable (non-trivial) cases, the GWD provides an excellent approximation, far better than any alternative. [Adapted from [30], with most results for the "Exact" case from [17, 27], with new evaluations for symplectic case from [36]]
variance with numerical data generated from Monte Carlo simulations. We include in the comparisons some Gaussian approximations (viz. approximation schemes which lead to TWDs that are Gaussians), alluded to earlier, e.g., in Table 1. For these the dimensional variance of the TWD must scale like $\langle\ell\rangle^{2}$. The approximations differ in how the dimensionless part depends on $\tilde{A}$. The approximation developed in Grenoble by Ihle, Pierre-Louis, and Misbah [14, 33] focuses on the limit of large $\varrho$, neglecting the entropic interaction in that limit. While the variance $\sigma^{2} \propto \tilde{A}^{-1 / 2}$, the proportionality constant is 1.8 times that in the Gruber-Mullins case (cf. Table 2). One improve this approximation, especially for repulsions that are not extremely strong, by including the entropic interaction in an average way. This is done by replacing $\tilde{A}$ by
$\tilde{A}_{\text {eff }}=\left(\frac{\varrho}{2}\right)^{2}=\tilde{A}+\frac{\varrho}{2}$.
(cf. (11).) In this modified scheme, $\sigma^{2} \propto \varrho^{-1}$.
The physical meaning of $\tilde{A}_{\text {eff }}$ has not been adequately discussed heretofore. It corresponds to the full strength of the inverse-square repulsion between steps, i.e., the modification due to the inclusion of entropic interactions. From (16) it is obvious that the contribution of the entropic interaction, viz. the difference between the total and the energetic interaction,
as discussed in conjunction with Fig. 2, is $\varrho / 2$. This quantity, as noted then, depends sensitively on $\tilde{A}$. Note also that, remarkably, the ratio of the entropic interaction to the total interaction is $(\varrho / 2) /(\varrho / 2)^{2}=2 / \varrho$; this is the fractional contribution that is plotted in Fig. 2.

Barbier et al. at Saclay [34, 42-44] consider instead the $m$-terrace width relative to $m\langle\ell\rangle$, comparing the variance for large $m$ to the coefficient of $\ln m$ expected from roughening theory, they conclude that $\sigma^{2}=4 /\left(\pi^{2} \varrho\right)$, i.e., the same form as the modified Grenoble approximation but with a proportionality constant $K$ that is $82 \%$ as large. This information is summarized in Table 2. Note that for the Saclay and Wigner approaches, one must assume all steps interact with an inverse square repulsion; for the others, one can treat either that assumption or allow only nearest neighbors to interact energetically.

In Fig. 4 we compare how well all these theoretical approximations account for the behavior extracted from a wellcontrolled numerical experiment, Metropolis Monte Carlo simulations [45] of a TSK model. For this model (with $A=0$ ), the characteristic distance between close approaches of neighboring steps is [46]
$y_{\text {coll }}=\langle\ell\rangle^{2} \tilde{\beta} / 4 k_{\mathrm{B}} T=(\langle\ell\rangle / 2)^{2} \sinh ^{2}\left(\varepsilon / 2 k_{\mathrm{B}} T\right)$,

| Vicinal | $T(\mathrm{~K})$ | $\sigma^{2}$ | $\varrho$ | $\tilde{A}$ | $W / G$ |
| :--- | ---: | :--- | :--- | :--- | :--- |
| $\mathrm{Pt}(110)-(1 \times 2)[48]$ | 298 |  | 2.2 | 0.13 | - |
| $\mathrm{Cu}(19,17,17)[41,49]$ | 353 | 0.122 | 4.1 | 2.2 | 0.77 |
| $\mathrm{Si}(111)[50]$ | 1173 | 0.11 | 3.8 | 1.7 | 0.96 |
| $\mathrm{Cu}(1,1,13)[25,41]$ | 348 | 0.091 | 4.8 | 3.0 | 1.27 |
| $\mathrm{Cu}(11,7,7)[41,49]$ | 306 | 0.085 | 5.1 | 4 | 1.37 |
| $\mathrm{Cu}(111)[41,49]$ | 313 | 0.084 | 5.0 | 3.6 | 1.39 |
| $\mathrm{Cu}(111)[41,49]$ | 301 | 0.073 | 6.0 | 6.0 | 1.58 |
| $\mathrm{Ag}(100)$ | 300 | 0.073 | 6.4 | 6.9 | 1.58 |
| $\mathrm{Cu}(1,1,19)[25,41]$ | 320 | 0.070 | 6.7 | 7.9 | 1.64 |
| $\mathrm{Si}(111)-(7 \times 7)[51]$ | 1100 | 0.068 | 6.4 | 7.0 | 1.67 |
| $\mathrm{Si}(111)-(1 \times 1) \mathrm{Br}[13]$ | 853 | 0.068 | 6.4 | 7.0 | 1.67 |
| $\mathrm{Si}(111)-\mathrm{Ga}[52]$ | 823 | 0.068 | 6.6 | 7.6 | 1.67 |
| $\mathrm{Si}(111)-\mathrm{Al} \sqrt{3}[53]$ | 1040 | 0.058 | 7.6 | 10.5 | 1.85 |
| $\mathrm{Cu}(1,1,11)[34]$ | 300 | 0.053 | 8.7 | 15 | 1.95 |
| $\mathrm{Cu}(1,1,13)[25,41]$ | 285 | 0.044 | 10 | 20 | 2.12 |
| $\mathrm{Pt}(111)[54]$ | 900 | 0.020 | 24 | 135 | 2.59 |

TABLE 2 Compendium of experiments measuring the variances of terrace width distributions of vicinal systems, as of the end of the last decade. The estimate of $\tilde{A}$ is obtained from the (normalized) variance using (18), except for the first-row entry, which is based on a direct fit using the 2-parameter GWD (cf. 6th item of "User's Guide"). W/G stands for the ratio of the estimates of the interaction strength based on Wigner and Gruber-Mullins perspectives, $\tilde{A}_{\mathrm{W}} / \tilde{A}_{\mathrm{G}}$, as given in (19). (Condensed from [31])
which is about 35 at $k_{\mathrm{B}} T / \varepsilon=1 / 2$ and $\langle\ell\rangle=10$, used in our most extensive calculations. For $A>0$, meandering is suppressed, making this distance is larger. In most cases, we used $L_{y}=2000 \gg y_{\text {coll }}$, and $N=100$ steps $^{7}$. We used a standard high-quality random-number generator (Ran3 [47]) and averaged over 100 runs using different initial seeds. In these runs the variance reached its steady-state value after about 3000 MCS (Monte Carlo steps per site); we started "taking data" after $10^{4} \mathrm{MCS}$, recording results every 10 MCS until reaching $3 \times 10^{4} \mathrm{MCS}$ [29].

The excellent agreement between the GWD expression and numerical data generated with Monte Carlo simulations is displayed in Fig. 4. The various predictions of the variance are plotted vs. $\tilde{A}$. A logarithmic scale is used for the horizontal axis so as not to give undue visual emphasis to larger values of $\tilde{A}$ nor to blur the region of rapid variation for small (but nonvanishing) $\tilde{A}$, for which an exact calibration point exists. The Wigner result is essentially given by (18). Table 2 shows that the physical values of $\tilde{A}$ range from near 0 up to the mid teens. Some pathology is presumably involved in the rare cases in which larger values are observed. There are relatively few reports of small but non-zero values of $\tilde{A}$. A reason might well be that any of the Gaussian approximations manifestly fail in this regime, so that before the recognition of the utility of the Wigner distribution, one could not deal quantitatively with small $\tilde{A}$ [48].

To heighten the contrast, the data in Fig. 4 can be replotted (in Fig. 2b of [29]) using as the ordinate $\varrho(\tilde{A}) \times \sigma^{2}(\tilde{A})$, so that approximations for which $\sigma^{2} \propto \tilde{A}_{\text {eff }}^{-1 / 2}$ become horizontal lines. With such rescaling, then, the Saclay and the modified Grenoble (all steps interacting) predictions appear as lines at ordinates 0.405 and 0.495 , respectively (cf. Table 1). The solid curve representing the GWD rises slowly, from 0.4 at $\tilde{A}=1$ toward $1 / 2$, capturing a similar rise of the MC data.

[^4]

FIGURE 4 The variance $\sigma^{2}(\tilde{A})$ vs. $\tilde{A}$ on a logarithmic scale, plotted for the GWD (light solid curve) and for the various Gaussian approximations: the modified Grenoble (short-short-long dashed for all steps interacting, shortlong dashed for NN step interactions only), Saclay (short-short-long-long dashed line), and Gruber-Mullins (long dashed for all steps interacting, short dashed for NN steps only). Monte Carlo data are shown as •'s, with statistical errors less than the size of the symbols. See text for discussion. (Fig. 2a of [29])

## 3 User's guide: Synopsis of findings

References [2] and [40] contain several figures showing applications, to experimental TWDs, of the perspective discussed above. In the following we summarize several specific ideas that should be of use in confronting data.

### 3.1 Interaction strength from variance

From (13) and (11), one can estimate the variance from $\tilde{A}$, but experimentalists usually seek the reverse, measuring the variance of the TWD and seeking to extract $A$. An excellent estimate [41] of the GWD-predicted $\tilde{A}$ from the variance, based on series expansion of (13), is:
$\tilde{A} \approx \frac{1}{16}\left[\left(\sigma^{2}\right)^{-2}-7\left(\sigma^{2}\right)^{-1}+\frac{27}{4}+\frac{35}{6} \sigma^{2}\right]$,
with all four terms needed to provide a good approximation over the full physical range of $\tilde{A}$. The Gaussian methods described earlier essentially use just the first term of this expression and adjust the prefactor.

### 3.2 Gaussian fits of the GWD

Since TWDs for strong repulsions are well described by Gaussians, the GWD is well approximated by a Gaussian in this limit. In [41] a quantitative assessment is given of how closely the two distributions correspond as a function of $\varrho$. At the calibration point (for which an exact solution exists) for repulsive interactions ( $\varrho=4$ ), the relative difference of the standard deviation of a Gaussian fitted to $P_{\varrho}(s)$ from the actual standard deviation of this GWD (viz. the square root of the second moment of $P_{\varrho}(s)$ about its mean of unity) is $\sim 1 \%$, and decreases monotonically with increasing $\varrho$. (The $\chi^{2}$, however, is poorer with a Gaussian than with the GWD.) For this range ( $\varrho \geq 4$ ) differences between estimates of $\tilde{\tilde{A}}$ obtained from GWD and the various Gaussian-fit
methods come primarily from different philosophies of extracting $\tilde{A}$ from $\sigma$ rather than from differences in the fitting schemes.

Gaussian approximations assume the peak (mode) is at $s=1(\ell=\langle\ell\rangle)$; in fact, since peak of the GWD must lie below one to achieve a mean of one; the tabulations in Table 1 bear this out. Only to the extent that the mode is close to unity is a Gaussian approximation reasonable. Formulas have been derived [41] indicating the errors in fitting $\varrho$ due to errors in the first or zeroth moment of the distribution. Another criterion for the validity of the Gaussian fitting function is that the TWD not be noticeably asymmetric about its peak, i.e., that it be negligible for $\ell>2\langle\ell\rangle$.

### 3.3 Analyzing TWD skewness: unfulfilled hopes

When $\varrho$ is too small for the TWD to meet the criteria for adequate fitting by a Gaussian, we hoped that one could obtain reliable estimates of $\tilde{A}$ by analyzing the skewness. Although we tried a variety of formulas and schemes [30, 40], it turned out that in the end a fit to the GWD was needed, so that skewness did not offer a shortcut to $\varrho$.

### 3.4 Correcting estimates based on Gruber-Mullins

When dealing with tabulations of data analyzed in the traditional way [1] based on the Gruber-Mullins perspective, it is useful to recast (18) in a form that indicates the factor by which the estimate $\tilde{A}_{\text {W }}$ based on GWD exceeds the traditional estimate $\tilde{A}_{\mathrm{G}}$ :
$\tilde{A}_{\mathrm{W}} / \tilde{A}_{\mathrm{G}}\left[\equiv A_{\mathrm{W}} / A_{\mathrm{G}}\right] \approx 3-21 \sigma^{2}+\frac{81}{4} \sigma^{4}+\frac{35}{2} \sigma^{6}$.
Since $\tilde{A}=A \tilde{\beta} /\left(k_{\mathrm{B}} T\right)^{2}$, the ratio of the physical interaction strengths is the same as that of the dimensionless strengths.

### 3.5 Alternative: fitting with gamma distribution

Although the GWD is a simple, single-parameter function, it is not a "canned" distribution and that one needs to input gamma functions. For the smaller values of $\varrho$ when a Gaussian is inappropriate, there is a preprogrammed function that is available: the gamma distribution
$P_{\Gamma}(x)=\frac{x^{\alpha-1} \mathrm{e}^{-x / \theta}}{\Gamma(\alpha) \theta^{\alpha}}$,
is serviceable if one recasts the data in terms of $x$ as $s^{2}$; one the can identify $\theta$ as $b_{\varrho}^{-1}$ and, more importantly, $\alpha$ as $(\varrho+1) / 2$. This approach has not yet been tested with actual data.

### 3.6 Wigner distribution as a 2-parameter fit

The GWDs giving the best fits of experimental TWDs sometimes have first moments that differ somewhat from the first moments of the data, especially in cases termed "poor data" $[40,41]$ which exhibit a small "hump" at large values of $s$, beyond the mode. Moreover, it may be desirable to determine the scaling length (the "effective mean", which equals the first moment for ideal GWDs) and the variance in a single fitting procedure rather than to predetermine
this length from the first moment. This "refined" scaling implies that the argument of $P_{\varrho}$ should be $\ell / \bar{\ell}$, where $\bar{\ell}$ denotes the characteristic length determined along with $\varrho$ in a twoparameter least-squares fit of the data to a GWD, leading to the replacement ${ }^{8}$ :

$$
\begin{equation*}
P_{\varrho}(s) \rightarrow(\langle\ell\rangle / \bar{\ell}) P_{\varrho}(s\langle\ell\rangle / \bar{\ell}), \quad \text { i.e. } \quad(\langle\ell\rangle / \bar{\ell}) P_{\varrho}(\ell / \bar{\ell}) . \tag{21}
\end{equation*}
$$

In the specific applications to data for copper [31], $\langle\ell\rangle / \bar{\ell}$ tends to be greater than unity, typically by several percent, but it is unclear whether this is true for semiconductors (or even for other metals). In our Monte Carlo simulations [29], where we have greater control of purity and uniformity than in experiments, the optimal $\bar{\ell}$ is essentially identical to $\langle\ell\rangle$ : a twoparameter fit is unnecessary.

### 3.7 Effects of lattice discreteness

For large values of $\tilde{A}$, the continuum picture breaks down and one must confront the discreteness of the lattice both in actual physical systems and in Monte Carlo simulations. This problem is discussed extensively in [31] and [29] but are not repeated here since the relevant values of $\tilde{A}$ are larger than found in many physical systems. By adapting the celebrated VGL model [55], we have found [29] that the roughening criterion translates to $\tilde{A}_{\mathrm{R}}=\langle\ell\rangle^{4} / 6$. Note that vicinal surfaces are rough, in contrast to the flat terrace orientation. Thus, for $\langle\ell\rangle=3$, e.g., when $\tilde{A}_{\mathrm{R}}$ exceeds $\sim 14$ (so above the physical range), the vicinal orientation becomes a facet. In some cases such as $\mathrm{Si}(113)$, the steps are exceedingly straight and uniformly spaced, making them excellent templates for growth of nanowires [56]. It is, therefore, likely that this surface is a facet rather than a rough vicinal.

### 3.8 Estimate of number of independent measurements

In order to estimate uncertainties in the determination of the TWD and, ultimately, $\tilde{A}$, it is important to have a realistic value of the number of independent measurements, a number generally much smaller than the total number of measurements. This problem is discussed in [41]. The upshot is that the number of "independent" terrace widths is reduced from $L_{y} N$ (the number of atomic spacings across the sampled area along the mean step direction times the number of steps) by up to nearly two orders of magnitude, due to slow decay of correlations perpendicular to the steps. Thus, several STM images are needed to obtain decent statistics.

## 4 New developments and directions <br> 4.1 Pair correlation function

The TWD amounts to a many-particle correlation function since one insists that there be no step between 0 and $\ell$ (or $s$ ). Instead one can study the step-step correlation function $h_{\varrho}(s)$, essentially the probability of finding a pair of steps

[^5]separated by $\ell$ regardless of how many steps lie in between. Until $\ell$ reaches $\sim\langle\ell\rangle$ there is little difference between the two, but then the pair correlation begins to rise and peak near $2\langle\ell\rangle$ and at subsequent multiples of $\langle\ell\rangle$, with a decreasing envelope around the oscillations, so that eventually $h \rightarrow\langle\ell\rangle^{-1}$. For fitting experimental data (in this case, $\mathrm{Si}(111)$ at $1100^{\circ} \mathrm{C}$ ), we [57] used an asymptotic expression [58]
$\langle\ell\rangle h_{\varrho}(s) \sim-\frac{1}{\pi^{2} \varrho s^{2}}+2 \sum_{j=1}^{\infty} \frac{d_{j}^{2}(\varrho)}{(2 \pi s)^{4 j^{2} / \varrho}} \cos (2 \pi j s)$,
$d_{j}(\varrho)=\Gamma\left(1+\frac{2 j}{\varrho}\right) \prod_{m=1}^{j-1}\left(\frac{2 m}{\pi \varrho}\right) \Gamma^{2}\left(\frac{2 \pi m}{\varrho}\right)\left(\frac{2 m}{\varrho}\right)$,
that works well for $s>1 / 2$, better than the conventionally used harmonic lattice approximation, since the conventional "harmonic" approximation [59] is insufficiently accurate. For smaller $s$ we patched onto $a_{\varrho} s^{\varrho}$. While Ha [60] derived a general exact solution for $h_{\varrho}(s)$, it is computationally intractable.

### 4.2 Fokker-Planck

Recently, Pimpinelli et al. [9], starting from Dyson's 1D Coulomb gas model and making plausible assumptions, derived a Fokker-Planck equation
$\frac{\partial P(s, \tilde{t})}{\partial \tilde{t}}=\frac{\partial}{\partial s}\left[\left(2 b_{\varrho} s-\frac{\varrho}{s}\right) P(s, \tilde{t})\right]+\frac{\partial^{2}}{\partial s^{2}}[P(s, \tilde{t})]$,
where $\tilde{t} \equiv t / \tau$, and the characteristic time $\tau$ is $\langle\ell\rangle^{2}$ over the squared strength of the noise in the underlying Langevin equation. The steady-state solution of (23) is the GWD. With it we can describe with compact analytic expressions the evolution toward equilibrium of steps from several experimentally relevant initial distributions: perfect cleaved crystals $(P(s)=\delta(s-1)$ as in Fig. 1), step bunches, and prequench equilibrated distributions at different temperatures $\left(P_{\varrho_{0}}(s)\right)$. The decay time $\tau$ of the difference of variance of $P(s, t)$ from its saturation value, we have also found [61], can be related to underlying atomistic processes in kinetic Monte Carlo simulations of the evolution of an initially constrained vicinal surface.

These ideas have broad ramifications. In econophysics the same formalism arises for the distribution of the means of stock prices in the Heston model [62]. Similar behavior may occur in precipation patterns at geothermal hot springs [63].

The argument [9] for (23) bolsters the formal justification for the GWD, as does work by Richards et al. [64] that uses a two-particle Calogero model [19, 20] (harmonically bound interacting spinless fermions on a line).

### 4.3 Azimuthal misorientation

Most of the above has tacitly assumed that the steps are close-packed or at least oriented along high symmetry directions. For vicinal surfaces that are misoriented in the azimuthal in addition to the polar orientation, there are complications in applying the GWD formalism, particularly in determining the dependence of $\tilde{A}-$ ultimately of $\tilde{\beta}$ and $A-$ on in-plane misorientation $\theta$. For $\tilde{\beta}$ we have made substantial progress recently, again prompted by collaboration with
the Jülich group [65] and by persistent probing questions by Ibach. For the $\{100\}$ and $\{111\}$ faces of fcc cubic crystals, we have derived surprisingly simple formulas for the $\beta(\theta)$ and $\tilde{\beta}(\theta)$ for low $T$ (compared to the terrace roughening temperature) $[66,67]$; in the case of Cu and other noble metals, this criterion is easily satisfied for room $T$. For arbitrary $T$, we have generated a more complicated analytic expression that is straightforward incorporate in continuum simulations, in particular finite-element codes [68]. We have also carried out ab initio calculations of the characteristic energies of the latticegas models used to understand $\tilde{\beta}(\theta)$ for Cu , to insure that these numbers are consistent with those estimated from experiments using statistical-mechanics reasoning [69]. Fuller discussion is beyond the limits of the article.

Much less is known about $A(\theta)$, but again the Jülich group is at the forefront of these investigations [70]. Another important open is the relation between $A$ and surface stress [71]. There has been no successful extension of the seminal theory for isotropic substrates [3,72] to account quantitatively for $A$ of a realistic surface [73].

## 5 Conclusion

The GWD has proved a powerful tool to link the study of the step spacings on vicinal surfaces to very general ideas of fluctuations. It is remarkable that H . Ibach could deduce its form, for the $A=0$ case, from physical insight. His perspicacious graph (Fig. 3, spawned great progress in understanding and exploiting the fluctuations of the spacings between steps. Even if one chooses to fit with a Gaussian, the theory of GWD provides the most reliable way to extract the strength of the step repulsions from the TWD. In addition to clarifying the equilibrium properties of steps, these ideas are now helping us to understand non-equilibrium aspects, notably the relaxation of steps toward equilibrium. Pimpinelli and I are also actively investigating applications to various aspects of growth in surface and interface problems.

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## Appendix

## How the Hamiltonian symmetries lead to the Wigner surmise exponents

This appendix, based on [74] (see also §4.3 of [28]), describes how the exponents of $s$ in (6)-(7) come about. For physicists, random matrix theory is rooted in the study of energy eigenvalues in nuclear physics. As recounted beautifully in [27], conventional statistical mechanics treats ensembles of identical physical systems with the same Hamiltonian but different
initial conditions. In contrast, Wigner considered ensembles of dynamical systems governed by different Hamiltonians $\mathscr{H}$ having some common symmetry property and sought generic properties of such ensembles associated with the symmetry. Dyson extended this work to show that there are three symmetries of the matrices of the Hamiltonian: orthogonal (real symmetric) matrices correspond to time-reversal invariance with rotational symmetry, unitary (complex) matrices to violated time-reversal invariance (as for electrons in a magnetic field), symplectic (see (2.3) of [27], §2.4 of [17]) to time-reversal invariance with broken rotational symmetry and $1 / 2$-integer spin. Wigner then, for convenience, considered Gaussian weightings,

$$
\begin{equation*}
p(\mathscr{H}) \sim \exp \left[-b N \operatorname{tr}\left(\mathscr{H}^{2}\right)\right], \tag{A.1}
\end{equation*}
$$

with the idea that for large matrices the fluctuations of the eigenvalues should be independent of the weight factors as well as of the specific form of the level spectrum. (There is no information about the average values.)

As described in [74], we start with the simplest case, an orthogonal ensemble ( $\beta=1$ ) with $N=2$ (just 2 particles), with $\mathscr{H}$ having diagonal elements $h_{11}$ and $h_{22}$, and off-diagonal element $h_{12}\left(=h_{21}\right)$. We focus on the Jacobean associated with a change of variables for the Gaussianensemble probability distribution function in going from the joint probability distribution function $p\left(E_{1}, E_{2}\right)$ for adjacent eigenenergies $E_{1}, E_{2}$ to $P(s) \mathrm{d} s$; we define variables $\vec{h} \equiv$ $\left(h_{11}+h_{22}\right) / 2, u \equiv h_{11}-h_{22}$ and $s \equiv\left(u^{2}+4 h_{12}^{2}\right)^{1 / 2}=\mid E_{2}-$ $E_{1} \mid$. Thus, $E_{1,2}=\bar{h} \pm s / 2$. We must now take into account all possible matrix elements $h_{11}, h_{22}, h_{12}$. From (A.1) and with $\mathrm{d} h_{11} \mathrm{~d} h_{22}=\mathrm{d} \bar{h} \mathrm{~d} u$,

$$
\begin{align*}
& \iiint p \mathrm{~d} h_{11} \mathrm{~d} h_{22} \mathrm{~d} h_{12} \\
& \quad=\int \mathrm{d} s \iint \exp \left[-2 b\left(E_{1}^{2}+E_{2}^{2}\right)\right] \mathrm{d} \bar{h} \mathrm{~d} u\left|\frac{\mathrm{~d} h_{12}}{\mathrm{~d} s}\right| . \tag{A.2}
\end{align*}
$$

Hence we can identify the inner double integral of (A.2) as $p(s)$. Since $h_{12}= \pm(1 / 2)\left(s^{2}-u^{2}\right)^{1 / 2},\left|\mathrm{~d} h_{12} / \mathrm{d} s\right|=$ $(s / 2)\left(s^{2}-u^{2}\right)^{-1 / 2}$. Since $E_{1,2}$ do not depend on $u$, we can pull the exponential out of the integration over $u$, leaving the elementary integral
$(s / 2) \int_{-s}^{s}\left(s^{2}-u^{2}\right)^{-1 / 2} \mathrm{~d} u=\pi s / 2$.
Next, since $2\left(E_{1}^{2}+E_{2}^{2}\right)=s^{2}+4 \bar{h}^{2}$, the integration over $\bar{h}$ is also elementary, and we are left with $P(s) \sim s \exp \left(-b s^{2}\right)$. This exact result for $N=2$ provides an excellent approximation for large $N$ as well. Indeed, near the level crossing (corresponding to $s \rightarrow 0$ ) the problem tends to reduce to a $2 \times 2$ problem. The integral for $P(s)$ over variables $u$ and $h_{12}$ includes in its integrand a Dirac delta function $\delta\left(s-\left[u^{2}+4 h_{12}^{2}\right]^{1 / 2}\right)$, which vanishes for $s=0$ only when the two [squared] independent variables do. Hence, $P(s) \propto s$, corresponding to a circular shell in parameter space, and we again have $\beta=1$.

For unitary ensembles there is an additional independent parameter since $h_{12}^{2}$ becomes $\left(\operatorname{Re} h_{12}\right)^{2}+\left(\operatorname{Im} h_{12}\right)^{2}$. Hence,
$P(s) \propto s^{2}$, corresponding to a spherical shell in parameter space, i.e., $\beta=2$. The argument for the symplectic ensemble leads similarly - but via quaternions or Pauli spin matrices to $\beta=4$.

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[^1]:    ${ }^{1}$ Line tension and step free energy per length are equivalent here since the surface is maintained at constant (zero) charge, but not in an electrochemical system, where the electrode potential $\varphi$ is fixed rather than the surface charge density conjugate to $\varphi$.

[^2]:    ${ }^{2}$ This expression assumes that $\tilde{\beta}$ does not depend on $\tilde{A}$.

[^3]:    ${ }^{3} p_{q}^{\mathrm{Bro}}(s)=c_{q} s^{q} \exp \left(-c_{q} \hat{q} s^{q+1}\right), c=\hat{q} \Gamma^{q+1}(\hat{q}), \hat{q} \equiv 1 /(q+1)$.
    ${ }^{4} p_{\varrho}^{\mathrm{Izr}}(s)=A_{\varrho} s^{\varrho} \exp \left[-\left(\pi^{2} \varrho / 16\right) s^{2}-\left(B_{\varrho}-\pi \varrho / 4\right) s\right]$.

[^4]:    ${ }^{7}$ For $\tilde{A}=50$, we found it safest to increase to $L_{y}=3000, N=200$.

[^5]:    ${ }^{8}$ Since $s$ is still determined from the raw data as $\ell /\langle\ell\rangle$, the refined scaling translates into replacing $s$ by $s\langle\ell\rangle / \bar{\ell}$ in the argument of the distribution. If the integration variable $s$ were similarly replaced, then the refined scaling would amount to a redefinition of a dummy variable, normalization would still be realized; however, the independent variable is kept as $s$ [31].

