LETTER TO THE EDITOR

Supporting random wave models: a quantum mechanical approach

Juan Diego Urbina and Klaus Richter

Institut für Theoretische Physik, Universität Regensburg, D-93040 Regensburg, Germany

E-mail: juan-diego.urbina@physik.uni-regensburg.de

Received 23 April 2003, in final form 17 July 2003
Published 10 September 2003
Online at stacks.iop.org/JPhysA/36/L495

Abstract
We show how two-point correlation functions recently derived within non-isotropic random wave models can be obtained in the appropriate limit in terms of the exact Green function of the quantum system. Since no statistical model is required for this derivation, this shows that taking the wavefunctions as Gaussian processes is the only assumption of those models. We also show how for clean systems the two-point correlation function based on an energy average defines a Gaussian theory which substantially reduces the spurious contributions coming from the normalization problem.

PACS numbers: 05.45.Mt, 05.40.—a

Since Berry’s seminal paper in 1977 [1], the so-called random wave model (RWM) has become by far the most popular and successful tool to describe the statistical properties of wavefunctions of classically chaotic systems. In this approach wavefunctions are modelled by a random superposition of plane waves. Its applications range from the realm of optics [2], passing by the general problem of wave mechanics in disordered media [3, 4] to important issues in mesoscopic systems [5]. Owing to this robustness this approach has been regarded as the indicator of wave signatures of classical chaotic dynamics [6].

The reasons for this success can be traced back to two fundamental points. First, it can be formally shown that such a random wavefunction is a stationary random process [7] (roughly speaking a function taking random values at each point); second, such a random process is Gaussian, namely, it is uniquely characterized by a two-point correlation function which expresses fundamental symmetries, like the isotropy of free space. The fact that the process is Gaussian represents a considerable advantage in an operational sense since it provides us with a set of rules to cope with averages over complicated expressions in the way Wick’s theorem and its variants do. At the same time the generality of the random wave two-point correlation makes the theory a remarkably good approximation when the effect of the boundaries can be neglected, as for bulk properties.
When applied to real quantum systems, however, there still remain limitations related to the above-mentioned ingredients. Such limitations have recently received attention in the context of nodal lines and nodal domains statistics [6]. Concerning the Gaussian assumption, a formal proof showing that a chaotic wavefunction is indeed a Gaussian process is still lacking. Even more, as noted in [8] the Gaussian distribution explicitly contradicts the normalization condition for the wavefunction. In practical terms this means that, when dealing with statistics beyond the two-point correlation function, the Gaussian distribution produces spurious non-physical contributions, and attempts to construct a RWM respecting the normalization constraint lead to severe mathematical difficulties [9]. Still this assumption is supported by many arguments based on random matrix theory [5], quantum ergodicity [6], information theory [9] and Berry’s original semiclassical picture [1]. Impressive numerical results also support the conjecture at the level of one-point statistics [10], and evidence for higher order statistics is given in [5, 6]. Hence, it is appealing to look for a RWM which minimizes the spurious contributions due to the normalization problem while keeping the wavefunction distribution still Gaussian.

Concerning the isotropic character of the theory, constructing a random superposition of waves satisfying both the Schrödinger equation and boundary conditions turns out to be at least as difficult as solving the full quantum mechanical problem by means of standard techniques. To our knowledge the attempts in the direction of a non-isotropic RWM can only deal with highly idealized boundaries such as an infinite straight wall [11], a linear potential barrier [12] and the edge between two infinite lines enclosing an angle of a rational multiple of \( \pi \) [13]. Also in [14] a modification of the RWM to include finite size effects is presented. The fact that these approximations already produce non-trivial deviations from the isotropic case indicates the relevance of including arbitrary boundaries.

Our aim in this letter is twofold: first, we shall show that the mentioned results for the two-point correlation function defining the non-isotropic and finite-size RWM can be derived from quantum mechanical expressions, namely, they are independent of any statistical assumption about the wavefunction. Second, we shall show how for a statistical description of wavefunctions using an energy ensemble average, the spurious contributions coming from the normalization problem are of order \( O(1/N) \) with \( N \) being the number of members of the ensemble, making their effect negligible for high energies.

1. Isotropic and non-isotropic random wave models

We consider solutions of the Schrödinger equation \( \left(-\frac{k^2}{2m} \nabla^2 + V(\vec{r})\right)\psi_n(\vec{r}) = E_n\psi_n(\vec{r}) \) for a closed system where the corresponding classical dynamics is chaotic (in the following we take \( 2m = 1 \)). The RWM assumes the statistical description of an ensemble of wavefunctions mimicked by a random superposition of \( M \) plane waves, \( \psi^r(\vec{r}) = \sum_{j=1}^{M} a_j \exp(ik \cdot \vec{r}) \). Here, \( a_1, \ldots, a_M \) is a set of independent random variables with a variance such that the wavefunction \( \psi^r(\vec{r}) \) is normalized on average, and \( k(\vec{r}) = \sqrt{e - V(\vec{r})/\hbar} \) is the local wave number with \( e \) the mean energy of the states under study.

For the sake of comparison we shall focus on the following averages, used for the nodal counting statistics (we follow the notation of Berry [11]):

\[
\begin{align*}
B(\vec{r}) & := \langle (\psi(\vec{r})^2) \rangle \\
D_x(\vec{r}) & := \left\langle \left( \frac{\partial \psi(\vec{r})}{\partial x} \right)^2 \right\rangle \\
D_y(\vec{r}) & := \left\langle \left( \frac{\partial \psi(\vec{r})}{\partial y} \right)^2 \right\rangle \\
K_x(\vec{r}) & := \left\langle \psi(\vec{r}) \frac{\partial \psi(\vec{r})}{\partial x} \right\rangle \\
K_y(\vec{r}) & := \left\langle \psi(\vec{r}) \frac{\partial \psi(\vec{r})}{\partial y} \right\rangle.
\end{align*}
\]
We wish to stress, however, that the RWM, as it is Gaussian, can deal with far more general averages. For billiard systems the isotropic RWM (denoted by a superscript $i$) is given by the ensemble

$$\psi^i(\vec{r}) = \sqrt{\frac{2}{J}} \sum_{j=1}^{J} \cos(kx \cos \theta_j + ky \sin \theta_j + \phi_j)$$

and the average $\langle \cdots \rangle$ is defined by integration over a set of independent random phases $\phi_j \in (0, 2\pi)$. We also choose $\theta_j = \frac{2\pi j}{J}$, where the limit $J \to \infty$ is always taken after averaging over the phases $\phi$. Explicit calculation then gives the following results [2]:

$$B^i(\vec{r}) = 1 \quad D_i^x(\vec{r}) = \frac{k^2}{2} \quad D_i^y(\vec{r}) = \frac{k^2}{2} \quad K^i_\parallel(\vec{r}) = 0.$$  

(3)

These results represent bulk approximations to the real situation, since boundary effects are completely neglected. In order to improve this limitation, the following ensemble of non-isotropic (ni) superpositions of random waves was introduced in [11] to take into account the effect of a straight infinite boundary at $y = y_0$ on which we demand the wavefunction to satisfy Dirichlet (D) or Neumann (N) boundary conditions:

$$\psi^D(\vec{r}) = \sqrt{\frac{4}{J}} \sum_{j=1}^{J} \sin k(y - y_0) \cos(kx \cos \theta_j + \phi_j)$$

$$\psi^N(\vec{r}) = \sqrt{\frac{4}{J}} \sum_{j=1}^{J} \cos k(y - y_0) \cos(kx \cos \theta_j + \phi_j).$$

(4)

(5)

With the averaging procedure as in the isotropic case, one obtains for the Dirichlet (upper sign) and Neumann (lower sign) cases [11]:

$$B^{ni}(\vec{r}) = 1 \mp J_0(2k(y - y_0))$$

$$D_i^{ni}(\vec{r}) = \frac{k^2}{2} (1 \mp J_0(2k(y - y_0)) \mp J_2(2k(y - y_0)))$$

$$D_i^{ni}(\vec{r}) = \frac{k^2}{2} (1 \pm J_0(2k(y - y_0)) \mp J_2(2k(y - y_0)))$$

$$K_y^{ni}(\vec{r}) = \pm k J_1(2k(y - y_0)).$$

(6)

For more general situations where the confining potential is smooth (S), Bies and Heller [12] introduced the following ensemble of random Airy functions $Ai(x)$ to satisfy locally the Schrödinger equation for a linear ramp potential $V(x, y) = Vy$:

$$\psi^S(\vec{r}) = \frac{1}{\sqrt{J}} \sum_{j=1}^{J} \tilde{\Psi}_i[\Psi(y, Q_j)] \exp[i(Q_j x + \phi_j)].$$

(7)

Here

$$\Psi(y, Q) = \left(\frac{V}{\hbar^2}\right)^\frac{i}{4} (y - y_0) + \left(\frac{\hbar^2}{V}\right)^\frac{i}{4} Q^2$$
and \( y_0 = e/V \) is the turning point in the direction of the linear ramp, fixed by the mean energy \( e \) of the eigenstates under study. The phases \( \phi_j \) are defined as usual and provide the averaging, while \( Q_j \in [-\infty, \infty] \). Explicit calculation then gives [12, 15]:

\[
B^S(\vec{r}) = \int_0^\infty \text{Ai}^2 [\Psi(y, Q)] \, dQ \\
D^S(\vec{r}) = \int_0^\infty \text{Ai} \, dQ \\
D_y^S(\vec{r}) = \int_0^\infty \text{Ai} \, dQ \\
K_y^S(\vec{r}) = \int_0^\infty \text{Ai} \, dQ
\]

where \( \text{Ai}^\prime(x) \) is the derivative of the Airy function.

2. The quantum description

We consider a set of normalized solutions \( \psi_n(\vec{r}) \) of the Schrödinger equation with non-degenerate eigenvalues \( E_n \) lying inside the interval \( W = [e - \frac{\delta e}{2}, e + \frac{\delta e}{2}] \). Considering \( W \) as a range of energies with almost constant mean level spacing \( \Delta(e) \), the number of states within the interval is 

\[
N = \frac{\delta e}{\Delta(e)} \quad \text{(in general } N = \int_{e-\frac{\delta e}{2}}^{e+\frac{\delta e}{2}} \rho(E) \, dE \text{ with } \rho(E) \text{ being the level density).}
\]

In the high-energy limit we are interested in, \( N \gg 1 \) with \( \frac{\delta e}{2} \ll 1 \) are well-defined limits which we shall always assume implicitly. The two-point correlation function

\[
F(\vec{r}_1, \vec{r}_2) := \langle \psi(\vec{r}_1) \psi^*(\vec{r}_2) \rangle := \frac{1}{N} \sum_{E_n \in W} \psi_n(\vec{r}_1) \psi^*_n(\vec{r}_2)
\]

(9)

can be used to calculate the averages in equation (1) by differentiation:

\[
B(\vec{r}) := \left[ F(\vec{r}_1, \vec{r}_2) \right]_{\vec{r}_1=\vec{r}_2=\vec{r}} \\
D_y(\vec{r}) := \left[ \frac{\partial^2}{\partial y_1 \partial y_2} F(\vec{r}_1, \vec{r}_2) \right]_{\vec{r}_1=\vec{r}_2=\vec{r}} \\
D_y(\vec{r}) := \left[ \frac{\partial^2}{\partial y_1 \partial y_2} F(\vec{r}_1, \vec{r}_2) \right]_{\vec{r}_1=\vec{r}_2=\vec{r}} \\
K_y(\vec{r}) := \frac{1}{2} \left( \frac{\partial}{\partial y_1} \frac{\partial}{\partial y_2} F(\vec{r}_1, \vec{r}_2) \right)_{\vec{r}_1=\vec{r}_2=\vec{r}}
\]

(10)

It is convenient to use the Green function of the system,

\[
G(\vec{r}_1, \vec{r}_2, E + i0^+) = \sum_{n=1}^{\infty} \frac{\psi_n(\vec{r}_1) \psi^*_n(\vec{r}_2)}{E - E_n + i0^+}
\]

(11)

to obtain the expression

\[
F(\vec{r}_1, \vec{r}_2) = \frac{\Delta(e)}{2\pi i} \frac{1}{\delta e} \int_{-\infty}^{\infty} (G^*(\vec{r}_1, \vec{r}_2, E + i0^+) - G(\vec{r}_2, \vec{r}_1, E + i0^+)) \, dE.
\]

(12)

Note that this is an exact result and the common approximation \( F(\vec{r}_1, \vec{r}_2) \sim (G^*(\vec{r}_1, \vec{r}_2, E + i0^+) - G(\vec{r}_2, \vec{r}_1, E + i0^+)) \) [16] is not valid in general and requires further assumptions. Even more, the additional energy integration in equation (12) will turn out to be essential.

Different approximations to the Green function valid under different situations can now be used to study the corresponding wavefunction statistics.
3. The bulk contribution and finite size effects

For billiard systems the bulk \((b)\) results are obtained by using the free propagator given in two dimensions by the Hankel function \(G^0(\vec{r}_2, \vec{r}_1, E + i0^+) = \frac{i}{\pi} H^{(1)}_0(\sqrt{E} |\vec{r}_1 - \vec{r}_2|)\) instead of the exact Green function. The corresponding contribution to the two-point correlation is

\[
F^b(\vec{r}_1, \vec{r}_2) = \frac{1}{A} \delta_\varepsilon \int_{e_1 - \frac{\varepsilon}{2}}^{e_2 + \frac{\varepsilon}{2}} J_0 \left( \frac{\sqrt{E}}{\hbar} |\vec{r}_1 - \vec{r}_2| \right) dE
\]

where \(A\) is the billiard area. Using equations (10) and (13) we easily recover the results in equation (3) correctly normalized. Further analysis of expression (13) shows that it reduces to Berry’s result \([1]\)

\[
\langle \psi(\vec{r}_1) \psi(\vec{r}_2) \rangle^b = \frac{1}{A} J_0 \left( \frac{\sqrt{E}}{\hbar} |\vec{r}_1 - \vec{r}_2| \right)
\]

when \(|\vec{r}_1 - \vec{r}_2| \ll \sqrt{4A/\pi}\), while it decays much faster for \(|\vec{r}_1 - \vec{r}_2| \geq \sqrt{4A/\pi}\) as long as \(\delta_\varepsilon \geq \sqrt{\pi e/4A}\). Noting that \(\sqrt{4A/\pi}\) is just the average system linear size \(L\) we see that equation (13) actually defines a RWM which incorporates finite size effects when the average is taken on scales larger than the ballistic Thouless energy, \(\epsilon_{\text{Th}} = \hbar \sqrt{\varepsilon}/L\). Equation (13) then provides an analytical expression for the correlation function defined in \([14]\).

4. The case of an infinite straight barrier

For this situation we construct Green functions with the correct parity under the reflection symmetry with respect to the line \(y = y_0\) by means of the method of images. The symmetric and antisymmetric combinations give the corresponding two-point correlation for Dirichlet (upper sign) and Neumann (lower sign) boundary conditions as

\[
F^D,N(\vec{r}_1, \vec{r}_2) = \frac{1}{A} \delta_\varepsilon \int_{e_1 - \frac{\varepsilon}{2}}^{e_2 + \frac{\varepsilon}{2}} \left[ J_0 \left( \frac{\sqrt{E}}{\hbar} \sqrt{(x_1 - x_2)^2 + (y_1 - y_2)^2} \right) \pm J_0 \left( \frac{\sqrt{E}}{\hbar} \sqrt{(x_1 - x_2)^2 + (2y_0 - y_1 - y_2)^2} \right) \right] dE.
\]

Using this correlation function and equation (10) we obtain the averages defined in equation (1). Berry’s results equation (6) are again obtained in the limit of very short distances to the boundary \(|y - y_0| \ll \sqrt{\frac{A}{\pi}}\).

5. The infinite, smooth barrier

For a particle in the potential \(V(x, y) = Vy\), the Schrödinger equation is separable. The solutions along the \(x\) direction are plane waves and in the \(y\) direction Airy functions. Hence we have

\[
\psi_{k,e}(x, y) = \left( \frac{8\pi^3}{4\hbar^3 V} \right)^{\frac{1}{2}} \exp(-ikx) \text{Ai} \left( \frac{V}{\hbar^2} \right)^{\frac{1}{2}} \left( y - \frac{eV}{V} \right)
\]

and for the Green function

\[
G(\vec{r}_1, \vec{r}_2, E + i0^+) = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} \psi_{k,e}(\vec{r}_1) \psi_{k,e}^*(\vec{r}_2) \frac{1}{E - e - \hbar^2 k^2 + i0^+} de dk.
\]
Including this result into equation (12) we find for the correlation function:

\[
F^2(\vec{r}_1, \vec{r}_2) = \left( \frac{2\pi^3}{\hbar^4 V} \right)^{\frac{1}{3}} \frac{1}{N} \int_{-\infty}^{\infty} \int_{0}^{\infty} \cos(k(x_1 - x_2)) \left[ \frac{V}{\hbar^2} \right]^\frac{1}{3} \left( y_1 - \frac{E - \hbar^2 k^2}{V} \right) \right] \left[ \frac{V}{\hbar^2} \right]^\frac{1}{3} \left( y_2 - \frac{E - \hbar^2 k^2}{V} \right) \] \text{d}k \text{d}E.
\]

(18)

Together with the relations (10) again, one obtains equation (8) in the limit \(\Delta(\varepsilon) \to 0\).

To summarize so far, we obtained the one-point averages, equation (1), for a closed system from pure quantum mechanical considerations without appealing to any statistical assumption about the wavefunction. The known RWM results (3), (6), (8) are then derived in the appropriate limits (either short distances or infinite system size). The generalization to any other average bilinear in the wavefunction is straightforward.

6. The normalization problem

There is a prominent counter argument against the Gaussian assumption first presented to our knowledge in [4] and further explored in [9], which deserves special attention. It is the apparent contradiction between the normalization of the members of the ensemble and the Gaussian distribution of the wavefunction amplitudes. Mathematically this can be stated in the following way. Consider the functional

\[
\eta[\psi] = \int |\psi(\vec{r})|^2 \text{d}\vec{r}
\]

where \(\psi(\vec{r})\) is a member of the ensemble we use to describe the statistical properties of the wavefunction. One constraint we must impose is the normalization of all \(\psi(\vec{r})\), expressed by the vanishing of the ensemble variance \(\text{Var}(\eta) = \langle |\eta[\psi]|^2 \rangle - \langle |\eta[\psi]| \rangle^2\). If the wavefunction’s distribution is Gaussian, we obtain

\[
\text{Var}(\eta) = 2 \int \int |\langle \psi(\vec{r}_1) \psi^*(\vec{r}_2) \rangle|^2 \text{d}\vec{r}_1 \text{d}\vec{r}_2.
\]

(20)

This is in clear contradiction to the normalization condition \(\text{Var}(\eta) = 0\). Here we show that this variance is of order \(\text{Var}(\eta) = O(1/N)\). To this end we recall the definition of the two-point correlation, equation (9), and note that the \(\psi_n(\vec{r})\) are eigenfunctions of the same Hamilton operator, i.e., they form an orthonormal set:

\[
\int \psi_i(\vec{r}) \psi^*_j(\vec{r}) \text{d}\vec{r} = \delta_{i,j}.
\]

(21)

Then it is easy to obtain the following composition rule for the two-point correlation:

\[
\int \langle \psi(\vec{r}_1) \psi^*(\vec{r}_2) \rangle \langle \psi(\vec{r}_1) \psi^*(\vec{r}_2) \rangle \text{d}\vec{r} = \frac{1}{N} \langle \psi(\vec{r}_1) \psi^*(\vec{r}_2) \rangle.
\]

(22)

Since \(\langle \psi(\vec{r}_1) \psi^*(\vec{r}_2) \rangle\) converges in the limit \(\frac{\varepsilon}{\Delta} \sim \text{const}, N \to \infty\), we see that indeed \(\text{Var}(\eta)\) converges to zero as \(O(1/N)\).

This behaviour of \(\text{Var}(\eta)\), relying on the fact that an energy ensemble average is taken, differs from the case of disordered systems where the lack of orthogonality between the different members of the ensemble (since they are eigenfunctions of different Hamiltonians corresponding to different disorder realizations) leads to an expression for \(\text{Var}(\eta)\) of order \(O(1)\).
It is important to note that the extra energy average is essential to satisfy equation (22). For example, a simple calculation shows that Berry’s result, equation (14), does not satisfy the composition rule, while our result (13) does it as long as $\delta e \gtrsim e_{Th}$. This is a particular case of a more general statement saying that under certain conditions approximate Green functions will make the correlation function satisfy the composition rule. The proof of this result requires the use of semiclassical techniques and will be presented elsewhere [17].

7. Concluding remarks

We have shown that all the two-point correlation functions used to fix the different random wave models, (isotropic, non-isotropic, for a smooth boundary and including finite size effects) can be derived in the appropriate limit of the exact quantum mechanical expressions. To this end we do not use any statistical assumption about the wavefunctions; in fact, these results are independent of the character of the classical system, i.e., regular, mixed or chaotic\(^1\). Also, we showed that for clean chaotic systems the use of an energy ensemble reduces the spurious contributions coming from the normalization problem without affecting the Gaussian assumption. This result only requires the consistent use of the quantum mechanical definition of the correlation function.

A Gaussian field with a correlation given in terms of the exact Green function of the system and the energy average carefully taken into account represents a generalization which includes all known RWMs as limiting cases and successfully takes into account boundary and normalization effects for any closed, clean chaotic system. For general shapes (such as those studied in [13]) and boundary conditions (such as the mixed case presented in [15]) the exact quantum approach presented here cannot be analytically performed. An approach using the semiclassical Green function to derive the asymptotic expressions for the correlations presented here and in [13, 15] is then an adequate method and will be the subject of a separate communication [17].

Acknowledgments

We thank Peter Schlagheck, Martin Sieber and Marko Turek for helpful conversations. JDU is indebted to Sven Gnutzmann, Georg Foltin and Uzy Smilansky for encouraging and important remarks and acknowledges the kind hospitality at the Weizmann centre of complex systems in Rehovot, Israel, where this work was finished. This work was supported by the Deutsche Forschungsgemeinschaft through the Graduiertenkolleg ‘Nonlinearity and Nonequilibrium in Condensed Matter’.

References


\(^1\) The reason is that for integrable systems the energy average in equation (9) mixes wavefunctions with different symmetries. Even when this can be corrected working with a Green function belonging to the particular subspace under study, the Gaussian assumption is known to fail in this case.

See also: Bogomolny E and Schmidt C 2002 *Phys. Rev. Lett.* **88** 114101


