Particles, waves and trajectories:
210 years after Young’s experiment

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Abstract. Mermin’s “shut up and calculate!” somehow summarizes the most widely accepted view on quantum mechanics. This conception has led to a rather constraining way to think and understand the quantum world. Nonetheless, a closer look at the principles and formal body of this theory shows that, beyond longstanding prejudices, there is still room enough for alternative tools. This is the case, for example, of Bohmian mechanics. As it is discussed here, there is nothing contradictory or wrong with this hydrodynamical representation, which enhances the dynamical role of the quantum phase to the detriment (to some extent) of the probability density. The possibility to describe the evolution of quantum systems in terms of trajectories or streamlines is just a direct consequence of the fact that Bohmian mechanics (quantum hydrodynamics) is just a way to recast quantum mechanics in the more general language of the theory of characteristics. Misconceptions concerning Bohmian mechanics typically come from the fact that many times it is taken out of context and considered as an alternative theory to quantum mechanics, which is not the case. On the contrary, an appropriate contextualization shows that Bohmian mechanics constitutes a serious and useful representation of quantum mechanics, at the same level as any other quantum picture, such as Schrödinger’s, Heisenberg’s, Dirac’s, or Feynman’s, for instance. To illustrate its versatility, two phenomena will be briefly considered, namely dissipation and light interference.

1. Introduction
In 2013 we have celebrated the 100th anniversary of Bohr’s atomic model [1, 2], which led to the development of quantum mechanics in the 1920s. However, we have also celebrated the 210th anniversary of Young’s famous two-slit experiment. In 1803 Thomas Young presented to the Royal Society his experimental “proof of the general law of the interference of two portions of light” [3, 4]; more than one hundred years later this experiment has become one of the most influential ones in physics [5], particularly due to its tight connection with quantum mechanics. The fact is vividly expressed at the beginning of the third volume of the Feynman’s Lectures on Physics [6], where we read:

In this chapter we shall tackle immediately the basic element of the mysterious behavior in its most strange form. We choose to examine a phenomenon which is impossible, absolutely impossible, to explain in any classical way, and which has in it the heart of quantum mechanics. In reality, it contains the only mystery. We cannot make the mystery go away by “explaining” how it works. We will just tell you how it works. In telling you how it works we will have told you about the basic peculiarities of all quantum mechanics.
The phenomenon referred to here, namely quantum interference, is precisely the same phenomenon observed by Young—although for light instead of massive particles.

Nowadays the enormous success of quantum mechanics is indisputable. Not only this theory explains the most fundamental aspects of the physical world, but it has also given rise to technological applications with a direct impact on our daily life. Now, 210 years after Young’s experiment, what do we really know about quantum systems? Unfortunately, not much (if anything at all). That is, although we have a very accurate theory, our understanding of this theory still relies on the ideas prevailing in the late 1920s and 1930s, strongly linked to the experimental capabilities at that moment. At present fine experiments can be performed in the time domain, reproducing Young’s experiment particle by particle. Even though the general conception of quantum systems is still anchored in somewhat old-fashioned, self-imposed constraints, which have more to do with positivist prejudices than with limitations of the theoretical framework of quantum mechanics [7, 8]. Mermin’s quotation “shut up and calculate!” [9, 10] accurately summarizes this position. Obviously, this has constituted (and still constitutes) an important obstacle to the development and advance of new ways to understand the quantum world, particularly those relying on the concept of trajectory. This is the case, for example, of Bohmian mechanics.

The purpose of this communication is to show that there is nothing contradictory or wrong with Bohmian mechanics. Rather than a matter of taste, the discussion will show that this approach is just another representation (a hydrodynamical one) of quantum mechanics, at the same level as other more standard representations, e.g., Schrödinger’s, Heisenberg’s, Dirac’s, Feynman’s, etc. Actually, although it goes beyond the scopes of this work, leaving aside aspects commonly associated with Bohmian mechanics, like the possibility of hidden variables or the ontology of the wave function, it can readily be seen that it is just a direct translation of quantum mechanics into the more general language of the theory of characteristics [11]. Now, why Bohmian mechanics? Because it focuses on the quantum phase, which is not a quantum observable, although it plays a decisive role on quantum system dynamics. Bearing this in mind, this work is organized as follows. The essential elements of this approach are introduced in Sec. 2. In Sec. 3 a brief overview on how this approach and similar ones have been applied to different physical problems is presented. This makes readily apparent the nature of Bohmian mechanics as a theory of characteristics, even if the latter’s formalism is not explicitly discussed. In Sec. 4 Bohmian mechanics is applied to two different phenomena, namely dissipation and light interference, in order to show its versatility. To conclude, in Sec. 5 a series of final remarks are summarized.

2. Waves, trajectories and quantum mechanics
In appearance, the formulation of quantum mechanics in the Schrödinger representation keeps a close analogy with classical wave theory. Within this representation, quantum systems are described by a probability amplitude or wave function, \( \Psi \), which evolves according to the partial differential equation

\[ i\hbar \frac{\partial \Psi}{\partial t} = -\frac{\hbar^2}{2m} \nabla^2 \Psi + V\Psi. \]  

(1)

This equation describes the transport of probability instead of energy, as it happens with classical waves. Nonetheless, as Eq. (1) is formulated, this is not evident, since it rather displays the form of a typical diffusion equation (with a complex diffusion coefficient). After Max Born proposed the statistical interpretation, we say that the probability density, \( \rho \equiv |\Psi|^2 \), gives the probability that the quantum system has (or is in) a particular configuration. Therefore, a representation of \( \rho \) in time gives us information on how the configurational probability of the system evolves (is transported) in time, i.e., which are the configurations (e.g., positions) where we have more or less chance to find the system. If we look at a typical outcome obtained from an interference
experiment, even in the case of very large and complex molecular systems [12], we find that these particles distribute according to $\rho$, in agreement with Born’s statistical interpretation. However, particles are detected one by one at different places. The individual positions (evolutions) of these particles, however, are no quantum observables.

Motivated by that fact, in 1952 David Bohm proposed [13] a model that could explain such individual arrivals at the same time that could also account for the collective particle behavior, all without appealing to von Neumann’s reduction postulate. He regarded the individual systems as “hidden variables”. To develop this interpretative model, Bohm considered Schrödinger’s equation (1) plus the nonlinear (polar) transformation

$$\Psi(r, t) = \rho^{1/2}(r, t)e^{iS(r, t)/\hbar}. \tag{2}$$

This transformation relates the complex-valued fields ($\Psi, \Psi^*$) with the real-valued fields ($\rho, S$), where $S$ describes the local variation of the quantum phase. Substituting (2) into the time-dependent Schrödinger equation and then separating the real and imaginary parts of the resulting equation, one finds

$$\frac{\partial \rho}{\partial t} + \nabla \cdot \left( \rho \frac{\nabla S}{m} \right) = 0, \tag{3}$$

$$\frac{\partial S}{\partial t} + \frac{(\nabla S)^2}{2m} + V - \frac{\hbar^2}{2m} \nabla^2 \rho^{1/2} = 0. \tag{4}$$

The continuity (or conservation) equation (3) rules the ensemble dynamics, i.e., the number of particles described by the probability density has to remain constant; the quantum Hamilton-Jacobi equation (4) describes the time-evolution of the phase field, where the last term on the right-hand side is the so-called quantum potential. These two equations make more apparent the physical meaning of Eq. (1) as a transport equation. In analogy to the classical Hamilton-Jacobi equation, if $S$ is identified with the classical action, then a momentum $p = \nabla S$ can be postulated (Bohm’s momentum). This momentum can be expressed as $p = m \hat{r}$, which gives rise to the guidance equation of motion

$$\hat{r} = \frac{\nabla S}{m}. \tag{5}$$

The local velocity field $\mathbf{v} = \hat{r}$, which depends directly on the quantum phase, thus governs the individual system dynamics and, in virtue of Eq. (3), gives rise to averages that are in agreement with the results obtained directly from Schrödinger’s equation. It is interesting to note at this point that Eq. (5) does not need to be postulated, but it arises automatically within the theory of characteristics [11]. The integration of this equation in whichever evolution parameter (e.g., time) generates the corresponding characteristics perpendicular to $S$-surfaces of constant phase.

Previous to Bohm’s model, de Broglie formulated a similar one [14] where particles were assumed to be singularities guided by the wave. Both models pursued essentially the same idea, namely to explain the ensemble behavior of quantum systems at the same time that their individual motion was also properly described. The meaning of individual system, however, is rather uncomfortable, since its (individual) evolution is not accessible. To understand this idea with a simple example, think of the stream of a river: its precise characterization does not provide any information at all on the individual motion of its molecular constituents. In this regard, it is probably more precise the approach developed by Erwin Madelung in 1926. Shortly after Schrödinger proposed his equation, Madelung provided a clear prescription [15] to formulate such an equation in hydrodynamic form. Accordingly, quantum systems could be visualized in terms of a series of streamlines, which would follow the flow associated with the system probability density. That is, we know nothing about the precise motion of individual systems.
Figure 1. The different representations of quantum mechanics provide us with a different description of quantum systems. Typically quantum systems are thought from a complementary viewpoint (left). Bohmian mechanics, however, gives preference to the configuration space and emphasizes transport-related properties (right).

(they could be evolving in a Brownian-like fashion, for example), but still we can determine how the ensemble probability flows throughout the corresponding configuration space.

Following Madelung’s view, Eq. (5) needs not be postulated, but it is a consequence of the fact that Eq. (3) is a (real-valued) transport equation. This allows us to establish a direct analogy with classical fluid dynamics. Hence, making use of the local probability current density,

\[ J = \frac{\hbar}{m} \text{Im} (\Psi^* \nabla \Psi) = v \rho, \]  

(6)

where \( v \) is a local (hydrodynamic) velocity field, one finds the expression

\[ v = \frac{J}{\rho} = \frac{\nabla S}{m}, \]  

(7)

which is equivalent to the above Bohmian velocity. The integration in time (or whichever parameter) of Eq. (7) generates a family of streamlines or paths (for each wave function considered) along which the quantum fluid propagates, just as in the case of a classical fluid. As it can be inferred, the first equality goes beyond Bohmian mechanics and allows to define streamlines in any system characterized by a certain density and a vector that transports it through the corresponding configuration space, regardless whether such a density describes a quantum system or not. Notice that in this case, instead of the set of equations (3) and (4), we have a set of hydrodynamic equations, as shown by Takabayasi [16–18].

Taking into account the fact that Bohmian mechanics is a different theory, but only a reformulation of quantum mechanics, one may wonder why it is worth using. The answer is very simple, it is just a matter of which aspect of the physical system one wishes to stress. This is nicely illustrated in Fig. 1. Typically, we understand quantum systems in a somewhat dichotomic way, e.g., either wave or particle, position or momentum, energy or time, etc, emphasizing the so-called complementarity (see left panel). However, we may also be interested in transport issues, for which Bohmian mechanics constitutes a more convenient picture (right panel), since it describes the probability flow in configuration space without caring about any other complementary aspect. In this regard, we find, for example, that probably one of the most relevant and distinctive properties revealed by this quantum hydrodynamic formulation, is the non-crossing [19, 20], i.e., the fact that quantum fluxes (described in terms of bunches of Bohmian trajectories) cannot cross in configuration space at the same time. This is in compliance with the outcomes obtained from a recent photon-by-photon realization of Young’s experiment [21]. As a direct consequence of the non-crossing, one is able to establish well-defined quantum probability
tubes [22, 23], i.e., tubes in configuration space along which the integral of the probability density at a given time remains constant.

3. Hydrodynamic approaches in the literature

As seen above, the hydrodynamic language of Bohmian mechanics enables a visualization of quantum systems in terms of streamlines that follow the flow of the probability density. This simple pictorial representation attracted the attention of the chemical physics community immediately after the first wave-packet propagation schemes were developed and used in the field by the end of the 1960s. Pioneers in this field were McCullough and Wyatt, who studied the quantum dynamics of collinear atom-diatom reactions in hydrodynamic terms [24–26], just in a period when classical trajectories were in fashion. The following quotation, extracted from [25], gives a very clear idea of what this community was looking for:

Classical mechanics gives an amazingly good description of the probability density and flux patterns during most of the reaction; however, the classical and quantal descriptions begin to diverge near the end of the reaction. Essentially, the classical reaction terminates before the quantal reaction. The dynamic behavior of the reaction is hydrodynamically turbulent, as shown by transient whirlpool formation on the inside of the reaction path.

This ended up with the development of the so-called quantum-trajectory methods [27] at the end of the 1990s, just something that David Bohm himself thought to be the path to follow to describe the physical systems.

In above case, a hydrodynamical viewpoint was adopted, without making an explicit calculation of streamlines or trajectories (hydrodynamical vector fields were used instead). Nevertheless, a few years later, in the middle of the 1970s, Hirschfelder and coworkers materialized Madelung’s ideas when trying to recast solutions of Schrödinger’s equation in a pictorial way for tunneling and vortical dynamics. As before, the underlying motivation was seeking for a better understanding of the phenomena studied [28]:

This paper has resulted from an effort to get a better understanding of quantum mechanics by making a thorough study of a very simple problem, […] . The mathematics is simple, but the analysis is far-reaching.

Again, these studies came from the field of chemical physics. In this sense, it is interesting to note that the first studies of barrier tunneling came from this community [28], thus predating those that later on appeared following Bohm’s ideas [29].

In the 1980s we find a convergence to Bohmian mechanics from the field of molecular magnetism [30, 31], where the quantum hydrodynamics was also in fashion. In particular, in [30] we read:

A representation of the electron flow induced by the external field can be extremely useful to understand molecular magnetism. To this end, maps reporting modulus and trajectory of quantum-mechanical current density revealed a fundamental tool, whose importance could be hardly overestimated.

By means of these representations one can observe very convolved trajectories describing the electron current densities induced by external magnetic fields acting on different types of molecular systems [31]. At an applied level, “these tools provide fundamental help for rationalization of magnetic response properties, such as magnetizability and nuclear magnetic shielding” [31].

1 The references here provided are rather scarce, but I think that they will be enough for the reader interested in further enquiring about the use of streamlines/trajectories in the literature.
The same motivation can be found even earlier in electromagnetism, where the tradition of explaining phenomena in terms of rays was stronger and therefore there was not an urgent need to appeal to the Madelung-Bohm scheme. Thus, in 1952, the same year that Bohm published his work on hidden variables, Braunbek and Laukien published [32] a work where they studied the diffraction by an edge (a perfectly conducting half-plane) by means of lines of average electromagnetic energy flow, obtained from the analytical solution provided to this problem by Sommerfeld in 1896 [33]. About 20 years later (again in the 1970s, when the computational tools were already more sophisticated), Prosser produced [34] the first trajectories for Young’s two slit experiment and provided an explanation in terms of “photon” trajectories [35]. These trajectories preceded those obtained by Dewdney et al. for matter waves using Bohm’s model [36]. Later on, different authors have treated the problem of electromagnetism in terms of trajectories (an account can be found in [37]), until in 2011 Steinberg and coworkers performed an experiment from which the first trajectories were inferred experimentally [21]. At a more applied level, and in consonance with Madelung’s viewpoint, we also find works dealing with streamlines in wave optics [38, 39] or transport through billiards [40], for example.

Of course, one could regard the trajectories for electromagnetic fields (radiation) as similar to those for matter waves (massive particles). In the end, Maxwell’s equations can be seen as equivalent to Schrödinger’s one [41]. However, we can move apart from these scenarios, and still we find analogous streamline-based descriptions with similar purposes. This is the case with sound waves, for example, some of which started appearing in the middle 1980s [42–46]. In particular, in Ref. [42] we read:

A method is presented for computing the energy streamlines of a sound source. This enables charts to be plotted showing, as continuous lines, the flow paths of the sound energy from the vibrating surface to the nearfield and beyond. Energy streamlines appear to be a new construct; they have some similarities to the velocity streamlines used in fluid dynamics. Examples of the energy streamlines are given for the point-driven plate in water. […] These streamlines make it easier for the eye to follow the energy flow from the source into the nearfield and beyond. These paths are complicated, in some cases, but are of considerable interest from several points of view.

This is precisely the role played by Bohmian mechanics in the study of quantum mechanical systems! This role is clearer and clearer as one goes through Ref. [42] in more detail. While searching through the literature, one finds remarkable the fact that in other areas of physics dealing with waves (other than quantum mechanics or, by extension, quantum optics) the visualization of flows is particularly relevant. Physicists have tried to develop methods based on characteristics, like those described above, or just stroboscopic interferometric ones [47], in order to explore and understand the behavior of the fluid qualitative and quantitatively. This is in contrast with the reluctance found in quantum mechanics to treat systems on equal footing.

According to the above discussion, the interest generated by the calculation of streamlines associated with the transport of some quantity —e.g., probability, electromagnetic energy, pressure energy— has led to a series of approaches that all converge to the very same need: dealing with a tool that allows one to objectively monitor the transport of such a quantity. In this regard, if quantum mechanics is separated from any other wave theory, one may end up concluding that Bohmian trajectories represent real trajectories pursued by real particles (or, in general, degrees of freedom), i.e., they are hidden variables. However, if this theory is properly contextualized, we find that there is nothing that allows us to establish a direct connection between the possible motion of real particles and this type of trajectories [20, 48]. Comparing with other theories, and as it will be seen in the next Section, as much we can say that a Bohmian particle is a particle that obeys a Bohmian dynamics, i.e., according to a (local) average drift momentum, which provides the particle with nonlocal (global) hydrodynamic-like information. Such a particle allows us to infer dynamical properties of the quantum fluid, which are usually
“hidden” when studied by means of the wave function formulation. That is, Bohmian particles are the quantum equivalent of classical tracer particles (or just tracers) that can be found in other areas of physics and chemistry.

4. Some applications

4.1. Dissipative Bohmian mechanics

The first context where we are going to apply Bohmian mechanics in the sense described above, namely as a hydrodynamic picture of quantum mechanics, is that of dissipation. Thus, consider the well-known classical dissipative equation

$$m\dddot{x} + m\gamma \dot{x} + \frac{\partial V(x)}{\partial x} = 0,$$

where $m$ is the system mass and $\gamma$ is the friction coefficient. According to this equation, the dissipation undergone by the system is proportional to its speed at a given time, $\dot{x}$. In order to determine the quantum analog, i.e., to specify the Schrödinger equation corresponding to Eq. (8), we need to find a suitable Hamiltonian. One way to proceed is by considering [49] the change of variables

$$X = x, \quad P = me^{\gamma t} \dot{x} = pe^{\gamma t},$$

where $(x, p)$ denote the physical variables, with $p = m\dot{x}$, and $(X, P)$ the canonical ones. The latter allow us to define the conservative Hamiltonian

$$H(X, P) = \dot{X}P - L = \frac{P^2}{2m} e^{-\gamma t} + V(X)e^{\gamma t},$$

and satisfy the usual canonical relations

$$\dot{X} = \frac{\partial H(X, P)}{\partial P}, \quad \dot{P} = -\frac{\partial H(X, P)}{\partial X},$$

which enable the conservation of the total energy. If the initial energy is $E_0$ and therefore $H(X, P) = E_0$, the inverse change to the physical coordinates gives

$$H(x, p) = \frac{p^2}{2m} + V(x) = E_0 e^{-\gamma t},$$

i.e., the energy is lost exponentially at a constant rate ($\gamma$), as expected from the physical dissipative system described by Eq. (8). The dissipative model given by Eq. (10) is known as the Caldirola-Kanai model [50, 51] and constitutes one of the former attempts to express (8) in a Hamiltonian form.

In order to find the quantum analog of the Caldirola-Kanai model, we now make use of the standard quantization procedure, and associate the operators $\hat{X}$ and $\hat{P} = -i\hbar \partial / \partial \hat{X}$ with the canonical variables $X$ and $P$, respectively. Because of their canonicity, these operators satisfy the usual commutation relation $[\hat{X}, \hat{P}] = i\hbar$ —which does not hold for the “physical” operators, $[\hat{x}, \hat{p}] = i\hbar e^{-\gamma t}$. Accordingly, the quantum Caldirola-Kanai Hamiltonian arises by replacing $X$ and $P$ in Eq. (10) by the corresponding operators,

$$\hat{H}(\hat{X}, \hat{P}) = -\frac{\hbar^2}{2m} e^{-\gamma t} \frac{\partial^2}{\partial X^2} + e^{\gamma t} \hat{V}(\hat{X}),$$

and the corresponding Schrödinger equation, in the physical variable $x$, reads as

$$i\hbar \frac{\partial \Psi}{\partial t} = -\frac{\hbar^2}{2m} e^{-\gamma t} \frac{\partial^2 \Psi}{\partial x^2} + e^{\gamma t} V(x) \Psi.$$
Figure 2. Dissipative Bohmian trajectories for: (a) $\gamma = 0.3\omega_0$, (b) $\gamma = 2\omega_0$, and (c) $\gamma = 4\omega_0$, with $\omega_0 = 2\pi/\tau_0 \approx 0.628$ ($\tau_0 = 10$). To compare with, the frictionless Bohmian trajectories have also been included in each panel (gray dashed lines). The initial positions have been distributed according to the initial Gaussian probability density.

It is easy to show that the associated (dissipative) Bohmian trajectories obey the modified guidance equation

$$\dot{x} = \frac{J}{\rho} = \frac{1}{m} \frac{\partial S}{\partial x} e^{-\gamma t}. \quad (15)$$

Note the similarity between this equation and the classical analog, $\dot{x} = pe^{-\gamma t}/m$.

For potential functions $V(x)$ which are polynomials with a degree equal or smaller than two, one can find analytical solutions for initial wave functions with the shape of a Gaussian wave packet [49, 52]. Without entering details, just to illustrate the dissipative dynamics described by Eq. (14), we are going to consider a harmonic oscillator. The dissipative Bohmian trajectories for different values of the friction coefficient are displayed in Fig. 2. As it can be noticed, in the physical coordinates not only there is a clear violation of the usual commutation relation, which manifests in a vanishing dispersion of the wave packet, but the system approaches the bottom part of the potential well, thus going down the zero-point energy. This is a pathological behavior (in the physical variables) typical of the Caldirola-Kanai model, which is based on a continuous dissipation of the system energy due to the lack of a proper quantization. The latter only applies to the canonical variables $X$ and $P$. A similar behavior can be found, for example, when dealing with beables [53, 54]. The same correspondence can also be found if one considers an initial wave-packet superposition inside the harmonic potential, as shown in Fig. 3.

In order to avoid such an inconvenience, other models have been proposed in the literature as suitable quantum candidates of Eq. (8) [55–59]. In analogy to Brownian-type wave-function models [60], these models use to include a nonlinear term, thus also making the corresponding Schrödinger equation nonlinear. Of course, this nonlinear term is usually accompanied by a stochastic term that accounts for the random fluctuations of the medium that give rise to such a nonlinearity (just as in the Langevin equation for Brownian motion, for example). For example, in Kostin’s model [61] a nonlinear term depending on the phase of the wave function is considered, with the Schrödinger equation reading as

$$i\hbar \frac{\partial \Psi}{\partial t} = -\frac{\hbar^2}{2m} \frac{\partial^2 \Psi}{\partial x^2} + V\Psi + V_R\Psi + \gamma \left( S - \int \rho S dx \right) \Psi, \quad (16)$$

with $S = (\hbar/2i) \ln(\Psi/\Psi^*)$ and where $V_R$ is a random potential. Here we have focused on this model in particular because of the link that can be established between this model and Bohmian mechanics precisely through $S$, as it will be seen below.

In the full dissipative case, where $V_R = 0$, if we substitute the usual polar ansatz into Eq. (16) and then proceed as in standard Bohmian mechanics, we reach the modified quantum Hamilton-
Figure 3. Dissipative Bohmian trajectories for: (a) $\gamma = 0.3\omega_0$, (b) $\gamma = 2\omega_0$, and (c) $\gamma = 4\omega_0$, with $\omega_0 = 2\pi/\tau_0 \approx 0.628$ ($\tau_0 = 10$). To compare with, the frictionless Bohmian trajectories have also been included in each panel (gray dashed lines). The initial positions have been distributed according to the initial Gaussian probability density.

Jacobi equation
\[
\frac{\partial S}{\partial t} + \frac{1}{2m} \left( \frac{\partial S}{\partial x} \right)^2 + V + Q + \gamma S = 0, \tag{17}
\]
where $Q$ denotes the quantum potential (see Sec. 2). Assuming that the Bohmian trajectories are obtained from the usual equation of motion, $p = m\dot{x} = \partial S/\partial x$, and differentiating Eq. (17) with respect to $x$, we reach
\[
m\ddot{x} + m\gamma\dot{x} + \frac{\partial (V + Q)}{\partial x} = 0, \tag{18}
\]
where we have made use of the Lagrangian derivative
\[
\frac{d}{dt} = \frac{\partial}{\partial t} + \dot{x} \frac{\partial}{\partial x} = \frac{\partial S}{\partial t} + \frac{1}{m} \left( \frac{\partial S}{\partial x} \right) \frac{\partial}{\partial x}. \tag{19}
\]
Note that Eq. (18) has the same functional form as Eq. (8), except for the fact that it also includes a purely quantum force (given in terms of the space derivative of the quantum potential). That is, Bohmian mechanics shows a direct path to find the quantum analog of Eq. (8), consisting of just including the quantum potential. Kostin’s model has been proven very useful to determine energy bound states due to its convergence by dissipation [62].

4.2. “Photon” trajectories

Another field of interest concerning the applications of Bohmian mechanics is electromagnetism, as mentioned above, where a time-independent trajectory approach can be readily developed starting from Maxwell’s equations [37, 63]. Within this approach, where the time-dependence is removed by averaging over the oscillating electromagnetic field, the trajectories describe the flow of electromagnetic energy (analogous to the probability accounted for by the Bohmian trajectories). As shown by Prosser [34], this is possible by directly considering Maxwell’s equations defined by an electric field $\mathbf{E}(\mathbf{r})$ and a magnetic field $\mathbf{H}(\mathbf{r})$, the two key elements necessary to define the corresponding streamlines are the time-averaged electromagnetic energy density and Poynting vector (electromagnetic current density),
\[
U(\mathbf{r}) = \frac{1}{4} [\epsilon_0 \mathbf{E}(\mathbf{r}) \cdot \mathbf{E}^*(\mathbf{r}) + \mu_0 \mathbf{H}(\mathbf{r}) \cdot \mathbf{H}^*(\mathbf{r})], \tag{20}
\]
\[
\mathbf{S}(\mathbf{r}) = \frac{1}{2} \text{Re} [\mathbf{E}(\mathbf{r}) \times \mathbf{H}^*(\mathbf{r})], \tag{21}
\]
respectively; \( \mathbf{E}(r) \) and \( \mathbf{H}(r) \) denote respectively the spatial part of the electric and magnetic field vectors, which satisfy Maxwell’s equations and have been assumed to be harmonic, i.e.,

\[
\tilde{\mathbf{E}}(r) = \mathbf{E}(r)e^{-i\omega t}, \\
\tilde{\mathbf{H}}(r) = \mathbf{H}(r)e^{-i\omega t}.
\]

(22)

Since the electromagnetic energy density is transported through space in the form of the Poynting vector, a local velocity field can be defined [64] in analogy to (7), which reads as

\[
\mathbf{S}(r) = U(r)\mathbf{v}.
\]

(23)

The electromagnetic energy flow lines or “photon” paths are then obtained by integrating the equation

\[
\frac{dr}{ds} = \frac{1}{c} \frac{\mathbf{S}(r)}{U(r)}
\]

along the arc-length coordinate \( s \) (which can be referred to a proper time \( \tau = s/c \), with \( c \) being the speed of light). The analogy between light and massive particles becomes more apparent if the spatial parts of the electric and magnetic fields are expressed in terms of a scalar function \( \Psi \) that satisfies the Helmholtz equation and the corresponding boundary conditions [37, 63].

In order to show such an analogy, consider a monochromatic electromagnetic wave in vacuum incident onto a plate with two slits. The plate is on the \( XY \) plane, at \( z = 0 \). For simplicity, it is assumed that the fields are independent of the \( y \) coordinate. This assumption is well justified if the slits are parallel to the \( y \) axis and their width along this axis is much larger than along \( x \). In such a case from Maxwell’s equations one obtains two independent sets of equations. One involves the components \( H_x, H_z, \) and \( E_y \) of the electromagnetic field and is commonly regarded as \( E \)-polarized. The other involves the components \( E_x, E_z, \) and \( H_y \), and is known as \( H \)-polarization. The electric and magnetic fields behind the slits can be expressed as [63]

\[
\mathbf{E}(r) = -\frac{i\beta}{k} \frac{\partial \Psi}{\partial z} \mathbf{e}_x + \frac{i\beta}{k} \frac{\partial \Psi}{\partial x} \mathbf{e}_z + \alpha \Psi \mathbf{e}_y,
\]

(25)

\[
\mathbf{H}(r) = \frac{i\alpha}{\omega \mu_0} \frac{\partial \Psi}{\partial z} \mathbf{e}_x - \frac{i\alpha}{\omega \mu_0} \frac{\partial \Psi}{\partial x} \mathbf{e}_z + \frac{k\beta e^{i\phi}}{\omega \mu_0} \Psi \mathbf{e}_y.
\]

(26)

The scalar field \( \Psi \) at any \( z \) can be expressed as a Fresnel-Kirchhoff integral [65], which in general has to be numerically integrated once the initial condition is established. In particular, if one considers a grating with two Gaussian slits [66], the initial condition can be assumed to be a coherent superposition of the two diffracted waves,

\[
\Psi(x, 0) = \psi_1(x, 0) + \psi_2(x, 0),
\]

(27)

where

\[
\psi_i(x, 0) = \left( \frac{1}{2\pi \sigma_i^2} \right)^{1/4} e^{-(x-x_{0,i})^2/4\sigma_i^2} W(x - x_{0,i}, w_i),
\]

(28)

with \( i = 1, 2 \). A window function, \( W(x, w) \), has been added to each wave packet in order to provoke a truncation and, therefore, to analyze the eventual effects on the final interference pattern. In this case, this function is such that it is one within the extension covered by the corresponding slit (i.e., between \(-w_i\) and \( w_i\)), and zero everywhere else.

A series of electromagnetic energy flow lines or averaged “photon” trajectories are displayed in Fig. 4 (upper panel). As in the previous example, the initial positions have also been distributed according to the Gaussian weight. To compare with, the numerical values considered have been
Figure 4. Lower panel: Electromagnetic energy streamlines or average “photon” trajectories behind two Gaussian slits [66]. The numerical parameters used are [65]: $\sigma_1 = \sigma_2 = 0.3$ mm, $x_{0,1} = -x_{0,2} = 2.35$ mm, and $\lambda = 943$ nm. The trajectories have been distributed according to the corresponding initial Gaussian probability densities behind each slit. Upper panels: Transverse momentum, $k_x/k$, evaluated at the four distances reported in Ref. [21] (see red dashed lines in the lower panel) for full Gaussian slits (red solid lines) and slits truncated at $w_i = 1.9\sigma_i$ (blue dotted lines) and $w_i = 1.5\sigma_i$ (green dashed lines). To compare with, the experimental data (black circles) are also displayed. In these four calculations, asymmetric Gaussians were used, with parameters: $\sigma_1 = 0.307$ mm, $\sigma_2 = 0.301$ mm, $x_{0,1} = 2.335$ mm, $x_{0,2} = -2.355$ mm, and $\lambda = 943$ nm. Although the model is rather simple and makes no explicit use of the concept of “weak measurement”, there is a good agreement between the simulation and the trajectories inferred from the experimental data, which shows the suitability of Maxwell’s equations to describe this type of experiments. This adequacy is also seen when the transversal momentum obtained from the Poynting vector is compared to the corresponding experimental data, as shown in the upper panels for four different distances from the two slits. Notice that these two quantities are connected through the relation [67]

$$\frac{k_x}{k} = \frac{S_x}{S}.$$
where $S_x$ and $S$ refer to the $x$-component and modulus of the Poynting vector, respectively. In particular, in the four upper panels displayed in Fig. 4 the Gaussians have been considered to be asymmetric in order to find a better fitting with the experimental data (black circles). It can be seen that as the Gaussians are more severely truncated, the oscillations of $k_x/k$ undergo a remarkable increase. This means a stronger action of the quantum potential between adjacent interference fringes according to the usual Bohmian interpretation [36, 68].

5. Final remarks

In general, the position maintained with respect to Bohmian mechanics is that it constitutes an alternative interpretation to quantum mechanics (or even an alternative theory). From the above discussion, it is difficult to find arguments sustaining such claims —often used in a pejorative sense to refute any work in this area. We have seen that Bohmian mechanics should be rather regarded as an alternative and complementary representation of quantum mechanics, particularly when we note that it is the direct translation of the latter into the language of the theory of characteristics, which has nothing to do with hidden variables or ontological views.

In that sense, it is worth thinking for a while about the different representations that we have in classical and quantum mechanics. For the former, we admit different formulations, each one emphasizing a different physical aspect of the systems described. For example, Newton’s formulation relies on the relationship between the motion displayed by objects and the external forces that act on them. On the other hand, we also have Hamilton’s formulation, which is based on the concept of energy conservation. This allows us to tackle physical problems in a rather flexible way, choosing the formulation that better fits our needs, and at the same time to understand the same phenomenon from different perspectives.

Similarly in quantum mechanics there are also different formulations or ways to tackle the same problem, which are chosen according to their suitability —either analytical or numerical. For example, Schrödinger’s formulation stresses the time-evolution of the system (through its wave function) under the influence of a given Hamiltonian; any property of the system is synthesized from its wave function either at a given time (probability densities) or at two different times (correlation functions). With Heisenberg’s formulation, on the contrary, one focuses on the evolution in time of the operators associated with observables and how they act on a given (time-independent) system state. Half-way between both, the Dirac or interaction representation is more convenient to analyze the dynamics of systems interacting with other systems. Feynman’s path-integral representation, relying on the concept of classical action and trajectory, is suitable in the treatment of large systems. Other representations, such as the Wigner-Moyal or the Husimi ones, stress the role of the density matrix in phase space. All these well-known examples are equivalent formulations of quantum mechanics, which provide a different description of the same system. In the same way, Bohmian mechanics stresses the role of the quantum phase, which has determining consequences for quantum systems even though it is not an observable. Figure 1 summarizes this descriptive complementarity.

The trajectories that we obtain through Bohmian mechanics help us to visualize and understand the physics underlying quantum systems and phenomena by monitoring the flow of the probability. This appealing feature has been used in different contexts, not necessarily connected to Bohmian mechanics or, in general, to quantum mechanics. Here, for example, the dynamics of a dissipative system has been analyzed, making evident the pathologies of a well-known model and how they can be avoided. In this sense, notice that at a practical level the calculation of Bohmian trajectories is more convenient even if the Schrödinger equation has to be solved in order to compute them. First, the analysis of the system dynamics in terms of trajectories is simpler than in terms of probability densities, specially for two or more dimensions. Second, if instead of probabilities, one considers expectation values, the trajectories are still very appropriate, because the former only provide us with averaged information at every time, while
the latter allow us to visualize the dynamics of each particular part of the quantum state.

Furthermore, the same concepts and tools can be extended to other areas of physics, as we have seen in the example of the “photon” trajectories. Although the differential equations are different and the fields propagated have a different physical meaning, the underlying ideas are exactly the same. This is actually the way how the recent experimental realization of Young’s two slit experiment has been explained [21]. Among the many different ways that one could devise to join the transversal momentum function at different distances from the two slits, Bohmian mechanics provides a very clear prescription of how to do it, without incurring any kind of approximation or including any external element to quantum mechanics.

In that latter regard, I shall finish here with the following consideration. In order to explain his outcomes from the two-slit experiment, Young used the so-called Huygens’ construction. Accordingly, the position of a wavefront at a given time can be obtained by considering a wavefront at a previous time. Each point on this second wavefront is assumed to be a source of secondary waves, whose interference gives rise to the wavefront at the later time. The direction of travel of these wavefronts is what we call a ray. In the case of plane waves, the wavefronts are perpendicular to rays; in the case of interference, the shape of the wavefronts varies from point to point, that making intractable a ray description . . . from an analytical viewpoint! Should Young have had a computer, he would have been able to evaluate the rays locally, at each point . . . and would have discovered Bohmian mechanics more than 100 years earlier, for Bohmian trajectories constitute the convoluted generalization of the concept of ray. In other words, they are the characteristics that correspond to surfaces describing waves.

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