Can quantum mechanics be an emergent phenomenon?

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Abstract. We raise the issue whether conventional quantum mechanics, which is not a hidden variable theory in the usual Jauch-Piron’s sense, might nevertheless be a hidden variable theory in the sense recently conjectured by G. ’t Hooft in his pre-quantization scheme. We find that quantum mechanics might indeed have a fully deterministic underpinning by showing that Born’s rule naturally emerges (i.e., it is not postulated) when ’t Hooft’s Hamiltonian for be-ables is combined with Koopmann–von Neumann operatorial formulation of classical physics.

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1. Introduction

Although quantum mechanics (QM) has been the undisputed basis for most of progress in fundamental physics during the last 90 years or so, the extension of the current theoretical frontier to Planck’s scale physics, and recent enlargements of our experimental capabilities, may make the 21st century the period in which possible limits of quantum theory will be subjected to a thorough scrutiny. The basic premise of this paper is that QM is actually not a complete ontological system, but in fact it represents a very accurate low-energy approximation to a deeper level of dynamics. But what exactly the “deeper level dynamics” should be? There is a growing interest in this bone of contention which is partially fueled by the belief that in order to make a convincing synthesis of QM and general relativity (GR) a new conceptual paradigm is needed to tackle physics at very small space-time scales. There is, however, a fundamental discord in how this should be achieved. One way of thinking maintains that at high energies the rules of GR should be changed/modified while the rules of quantum physics should be kept untouched. Such a view is typically justified by arguing that QM is the most precise theory ever, being accurate to about one part in $10^{11}$ (for the latest precision tests of QED see, e.g. Ref. \cite{1}). As pointed by R. Penrose this view is deceptive \cite{2}: in fact GR has now been experimentally tested by measuring the orbit-period slowdown in the Hulse-Taylor binary pulsar PSR 1913 + 16 with accuracy to one part in $10^{14}$ — and this precision is apparently limited only by the
accuracy of clocks on Earth [3]. Another way of thinking holds that it is rather QM that should be replaced at high energy scales with a more fundamental paradigm, while GR may, or may not, be modified at such high energies. Along this line, motivated by the black-hole thermodynamics, G. ’t Hooft has hinted [4] that the fundamental rules should be, at very high energy scale — perhaps at the Planck scale — deterministic.

There is a long standing suspicion towards determinism that might underlie QM which mainly steams from Bell’s inequality (BI) [5] and its various generalizations [6]. While deterministic theories always satisfy Bell’s inequality, QM systems evidently experimentally violate BI [7, 8, 9]. This is then interpreted as a proof of the non-existence of a deterministic underpinning for QM. At this stage it should be stressed that BI’s are not a result about QM. They simply state that usual deterministic theories obeying the usual (Kolmogorovian) probability theory inevitably satisfy certain inequalities between the mean values of certain (suitably chosen) observed quantities. By “usual deterministic theories” one means theories that are realistic, i.e. the system has an intrinsic existence independent of observation, and local, i.e. sufficiently separated measurements should not influence each other. Though these assumptions about the underlying deterministic nature of QM are certainly intuitively plausible, they are too much rooted in our everyday laboratory-scale experience. Planck’s scale (∼ 10^{19} \text{ GeV}, i.e. ∼ 10^{-35} \text{ m}) deterministic dynamics (whatever this means) may, after all, look non-local to a human observer whose observational scale is at best few TeV (∼ 10^{-19} \text{ m}). In fact, according to ’t Hooft [10], the enormous amount of information that would be lost in the process of “coarse graining”, from 10^{19} \text{ GeV} to 10^{3} \text{ GeV} would lead to formation of equivalence classes, in the sense that many distinct states of the Planckian-scale dynamics would go into a single state of the observational-scale dynamics. For example, in Ref. [10] is proposed that, when the dynamical equations at be-ables level describe the configuration-space of a chaotic system, the equivalence classes could be related to the stable orbits of such system (e.g., limit cycles). The mechanism responsible for clustering of trajectories to equivalence classes is identified by ’t Hooft as information loss: after a while one cannot retrace back the initial conditions of a given trajectory, one can only say at what attractive trajectory it will end up. Explicit examples of be-able systems that give rise, at a macroscopic level, to a genuine quantum behavior can be found in Refs. [4, 10, 11, 12]. Applications of the outlined scenario have been given, e.g., in Refs. [11, 12, 13, 14, 15]. For other approaches see also Refs. [16, 17, 18].

The passage to quantum theory without conventional quantization procedure is called by ’t Hooft pre-quantization. In what follows we wish to use the pre-quantization concept to tackle the issue of Born’s rule from first principles, i.e. without need to postulate it. The paper is organized as follows: In Section 2 we present some fundamentals of ’t Hooft pre-quantization method. An extension of this is presented in the second half of Section 2. There we put forward some ideas concerning the dependance of ’t Hooft’s loss-of-information constraint on observer’s energy scale. A connection with Koopman–von Neumann (KvN) operatorial formulation of classical physics is discussed in Section 3. There we demonstrate in some detail that Born’s rule is, at the primordial energy scale, merely a convenient mathematical instrument for doing classical statistical physics, while other, equivalent means are also available. It is only at low-energy (observational) scales, where the operatorial and deterministic modes of description substantially depart, that Born’s rule starts to have its own independent existence. Various remarks and speculations are postponed till the concluding Section 4.

2. Essentials of ’t Hooft’s pre-quantization method

In this section we briefly outline ’t Hooft’s continuous-time pre-quantization method [10]. A discrete-time version, which employs cellular automata, is discussed, e.g. in Refs. [4, 12, 10], and will not be followed here. We start with the assumption that the dynamics at the primordial
deterministic level is described by the Hamiltonian

\[ H = \sum_i p_i f_i(q) + g(q). \]  

(1)

Here \( f_i \) and \( g \) are functions of \( q = \{q_1, \ldots, q_n\} \). One should note that the equation of motion for \( q_i \), i.e.

\[ \dot{q}_i = f_i(q), \]

(2)
is autonomous, since the \( p_i \) variables are decoupled. The above system is obviously deterministic, however its Hamiltonian is not bounded from below. It is also worthy of noting that

\[ H_q \]

for \( q \) observe that because \( \hat{\cdot} \)

see that the generator of time translations (i.e., the “Hamiltonian” operator), of the form

\[ \sum_i \frac{\partial^2 H}{\partial p_i \partial q_k}(\Delta t)^2 + \cdots = F_i(q(t), \Delta t), \]

(3)

where \( F_i \) is some function of \( q(t) \) and \( \Delta t \) but not \( p \). Since \( (3) \) holds for any \( \Delta t \) we get the Poisson bracket

\[ \{q_i(t'), q_k(t)\} = 0 \quad \text{for} \quad \forall \ t, t'. \]

(4)

Because of the autonomous character of Eq.(2) one can define a formal Hilbert space \( \mathcal{H} \) spanned by the states \( \{\{q\}\} \), and associate with \( p_i \) the operator \( \hat{p}_i = -i \partial / \partial q_i \). It is not difficult to see that the generator of time translations (i.e., the “Hamiltonian” operator), of the form

\[ \hat{H} = \sum_i \hat{p}_i f_i(q) + g(\hat{q}) \]

generates precisely the deterministic evolution \( \hat{q} \). Indeed, we first observe that because \( \hat{H} \) is generator of time translations then in Heisenberg’s picture

\[ \hat{q}_i(t + \Delta t) = e^{i\Delta t \hat{H}} \hat{q}_i(t) e^{-i\Delta t \hat{H}}, \]

(5)

which for infinitesimal \( \Delta t \) implies

\[ \hat{q}_i(t + \Delta t) - \hat{q}_i(t) = i\Delta t [\hat{H}, \hat{q}_i(t)] \quad \Rightarrow \quad \dot{\hat{q}}_i = f_i(\hat{q}). \]

(6)

On the other hand for arbitrary finite \( \Delta t \) we have from \( (5) \)

\[ \hat{q}_i(t + \Delta t) = \sum_{n=0}^{\infty} \frac{1}{n!}[\hat{H}, [\cdots [\hat{H}, \hat{q}_i(t)]] \cdots] = \hat{F}_i(\hat{q}(t), \Delta t). \]

(7)

In Eq.(7) \( \hat{H} \) appears in the generic term of the sum \( n \) times. On the other hand \( \hat{F}_i \) is some function of \( \hat{q}(t) \) and \( \Delta t \) but not \( \hat{p} \), which immediately implies that

\[ \{\hat{q}_i(t), \hat{q}_j(t')\} = 0, \]

(8)

for any \( t \) and \( t' \) (this in turn gives \( F_i = \hat{F}_i \)). Result \( (8) \) shows that the Heisenberg equation of motion for \( \hat{q}_i(t) \) in the \( q \)-representation is identical with the \( c \)-number dynamical Eq.(2). This is because \( \hat{q}_i(t + \Delta t) \) and \( \hat{q}_i(t) \) commute, and hence \( \hat{q}_i(t + \Delta t), \hat{q}_i(t), f_i(\hat{q}) \) and also \( d\hat{q}_i(t)/dt \) can be simultaneously diagonalized. In this diagonal basis we get back the \( c \)-numbered autonomous Eq.(2). In other words, operators \( \hat{q}_i \) evolve deterministically even after “quantization”. This evolution is only between base vectors. From the Schrödinger-picture point of view this means that the state vector evolves smoothly from one base vector to another (in Schrödinger picture base vectors are time independent and fixed). So at each instant the state vector coincides with some specific base vector. Because of this, there is no non-trivial linear superposition
of the state vector in terms of base vectors and hence no interference phenomenon shows up when measurement of \(q\)-variable is performed. Dynamical variables fulfilling Eq. (8) were first considered by Bell [5] who called them be-ables as opposed to observed dynamical variables which are in QM called observables.

Let us now come back to Eq. (1). One may immediately notice that \(H\) is unbounded from below. This fact should not disturb us too much since actual dynamics of be-ables is described by Eq. (2). The \(H\) was merely introduced to set up the parallel operatorial formulation of the be-able dynamics. In the following section we will see that this Hamiltonian has yet another useful role, namely it helps to formulate a classical statistical mechanics for be-ables. On the observational (i.e. emergent) level the Hamiltonians are key objects providing both the equations of motions and the energy of the system. The emergent Hamiltonians must be bounded from below. The concept of lower bound is, in 't Hooft’s proposal, just an emergent property formed during the coarse graining of the be-able degrees of freedom down to the observational ones. We can devise a simple toy model mechanism showing how a lower bound for \(H\) may develop. To this end consider \(\rho(\hat{q})\) to be some positive function of \(\hat{q}\) (but not \(\hat{p}\)) with \([\hat{\rho}, \hat{H}] = 0\). One then defines the splitting

\[
\hat{H} = \hat{H}_+ - \hat{H}_-,
\]

\[
\hat{H}_+ = \left(\hat{\rho} + \hat{H}\right)^2 \hat{\rho}^{-1} / 4, \quad \hat{H}_- = \left(\hat{\rho} - \hat{H}\right)^2 \hat{\rho}^{-1} / 4,
\]

where \(\hat{H}_+\) and \(\hat{H}_-\) are positive-definite operators satisfying

\[
[\hat{H}_+, \hat{H}_-] = [\hat{\rho}, \hat{H}_+] = 0.
\]

At this stage we introduce the “coarse-graining” operator \(\hat{\Phi}\) that describes the loss of information occurring during the passage from the be-able to observational scale. One possible choice is [1]

\[
\hat{\Phi}_E = \left(1 - e^{-(E_p - E)/E}\right)\hat{H}_-,
\]

where \(E\) refers to the observer’s energy scale, while \(E_p\) is the be-able energy scale, which we take to be Planck’s scale. The operator \(\hat{\Phi}_E\) is then implemented as a constraint on the Hilbert space \(\mathcal{H}\). So at each observational energy scale \(E\), the observed physical states \(|\psi\rangle_{\text{phys}}\) are given by the condition

\[
\hat{\Phi}_E |\psi\rangle_{\text{phys}} = 0.
\]

This equation identifies the states that are not affected by the coarse graining, i.e., states that are still distinguishable at the observational scale \(E\). The \(c\)-number factor \((1 - e^{-(E_p - E)/E})\) is not irrelevant and one cannot use directly \(\hat{H}_-\) instead of \(\hat{\Phi}_E\). In fact the constraint (12) is, according to Dirac’s classification of constraints [20], a first class primary constraint because \([\hat{\Phi}_E, \hat{\Phi}_E] = 0\) and \([\hat{\Phi}_E, \hat{H}] = 0\). First-class constraints generate gauge transformation and hence not only restrict the full Hilbert space (from \(\mathcal{H}\) to \(\mathcal{H}_c\)) but also produce equivalence classes of states, which are in general non local (e.g., points with a space-like separation are identified). Two states belong to the same class if they can be transformed into each other by a gauge transformation with the generator \(\hat{\Phi}_E\). Let \(\mathcal{G}_E\) be the one parameter group of these gauge transformations generated by \(\hat{\Phi}_E\). The equivalence classes built out of such gauge transformations represent at each scale \(E\) the physical states (i.e., observables). If \(\mathcal{O}_E\) denotes the space spanned by the observables, we can identify \(\mathcal{O}_E\) with a quotient space

\[
\mathcal{O}_E = \mathcal{H}_c / \mathcal{G}_E.
\]

1 This form might be inspired by the entropy considerations reported in [11]. We shall discuss this issue elsewhere.
The quotient space $\mathcal{O}_E$ (its structure and dimensionality) depends on the energy scale $E$. In particular at the level of be-ables where $E = E_p$ the constraint $\Phi_E$ is identically zero and the space of observables is directly the Hilbert space $\mathcal{H}$. On the other hand, when $E \ll E_p$, e.g. at scales available to a human observer, one has $\Phi_E = \hat{H}_L$. The latter is the constraint originally considered by ’t Hooft [4, 10]. In such a case the resulting physical state space, i.e. the space of observables, has the energy eigenvalues that are trivially bounded from below owing to

$$\hat{H}|\psi\rangle_{phys} = \hat{H}_+|\psi\rangle_{phys} = \hat{\rho}|\psi\rangle_{phys}. \quad (14)$$

Thus, in the Schrödinger picture the equation of motion

$$\frac{d}{dt}|\psi_t\rangle_{phys} = -i\hat{H}_+|\psi_t\rangle_{phys}, \quad (15)$$

has only positive frequencies on physical states. Above coarse-graining procedure was so far implemented in an operatorial mode of description of the be-able dynamics. As a result we have obtained on observational energy scales a Schrödinger equation with a “well behaved” (i.e., bounded from below) emergent Hamiltonian. Constraining procedure can be, in parallel, also applied to a deterministic mode of description of the be-able dynamics. This can be done through the Dirac–Bergmann algorithm [21]. Here one implements the constraint $\Phi_E$ directly on the phase space $\Gamma$ spanned by $p$ and $q$. This results in the constrained hyper surface $\Gamma_c$.

Because $\Phi_E$ Poisson-commutes with itself and with $H$, it is a first class primary constraint [21]. Similarly as in the operatorial case, this means that $\Phi_E$ not only restricts the phase space but also provides classes of equivalence for possible evolutions. This is because $\Phi_E$ is unable to determine the Lagrange multiplier in the “extended” Hamiltonian $H_e(q,p) = H(q,p) + \lambda(t)\Phi_E(q)$. In particular, the self-consistency equation

$$0 = \frac{d\Phi_E}{dt} = \{\Phi_E, H_e\} = \{\Phi_E, H\} + \lambda(t)\{\Phi_E, \Phi_E\}, \quad (16)$$

is trivially fulfilled without giving any information on $\lambda(t)$. This leads to the class of equivalent dynamics described by the equation $dq_i/dt = \{q_i, H_e\}$. Each of the possible trajectories in the class is parameterized by different $\lambda(t)$, see Fig.1. Two points belong to the same class if they can be transformed into each other by gauge transformation generated by $\Phi_E$. For infinitesimal transformations this means that for two equivalent dynamics with slightly different $\lambda(t)$

$$\delta q(t) = q_{\lambda(t)} - q_{\lambda'(t)} = \varepsilon\{q(t), \Phi_E\}, \quad (17)$$
with $\varepsilon = t(\lambda - \lambda')$. Gauge classes can be identified by choosing a single representative element from each class. This is done by a gauge-fixing condition $\chi$ such that $\{\Phi_E, \chi\} \neq 0$. The emergent physical space (space of classically observed degrees of freedom at scale $E$) is a quotient space $\Gamma^*_E = \Gamma_c / \mathcal{G}_E$. It should be noted that the identification of the equivalent classes is a non-local procedure, since at a given instant one generally identifies also points with a space-like separation. This is, in spirit, deeply different from the quotienting procedure used in gauge field theories. There the quotient space is obtained from the configuration space that in itself is non-physical (it contains huge amount of physically equivalent configurations). The physical space emerges only after one identifies each equivalence class with a distinct physical configuration.

In contrast, in our case we start with the configuration space (space of be-able dynamics) that emerges only after one identifies each equivalence class with a distinct physical configuration. A priori physical (it contains huge amount of physically equivalent configurations). The physical space is a quotient space obtained from the configuration space that in itself is non-physical. This is, in spirit, deeply different from the quotienting procedure used in gauge field theories.

3. Koopman–von Neumann formulation and ’t Hooft’s be-able dynamics

Let us come back to ’t Hooft’s Hamiltonian $H$ and try to understand its role a bit more. There are three immediate issues that should be addressed. These are: the issue of hermiticity, the issue of the canonical momenta $p_i$, and finally the issue of the ensuing state-vectors and related Born’s rule. The hermiticity is not a real problem at the energy scale $E_P$. There the dynamics is driven by Eq. (2) and the Hamiltonian is only a formal tool that allows to generate the corresponding evolution equations. Should one wish to have hermitian Hamiltonian from the very scratch, one may compensate for the non-hermitian ordering between $\hat{p}_i$ and $f_i(\hat{q})$ by adding the (non dynamical) function $g(\hat{q})$ to $\hat{H}$. This should be done in such a way that $g(\hat{q}) - g(\hat{q}) = \sum_i [\hat{p}_i, f_i(\hat{q})]$, so that $\hat{H}^\dagger = \hat{H}$. In the particular case when $g(\hat{q})$ is anti-hermitian the Hamiltonian is not only hermitian but also Weyl ordered. Indeed, in this case we have

$$\hat{H} = \sum_i \hat{p}_i f_i(\hat{q}) + g(\hat{q}) = \frac{i}{2} \sum_i (\hat{p}_i f_i(\hat{q}) + f_i(\hat{q})\hat{p}_i) + \frac{i}{2} \sum_i [\hat{p}_i, f_i(\hat{q})] + g(\hat{q})$$

$$= \frac{i}{2} \sum_i (\hat{p}_i f_i(\hat{q}) + f_i(\hat{q})\hat{p}_i) = W\left(\sum_i \hat{p}_i f_i(\hat{q})\right).$$

(18)

In this connection it is interesting to notice that the Weyl form also enters in the Pontryagin’s approach to quantization of non-Hamiltonian systems described by equations. The hermiticity itself is, however, an important concept at low energies, where the emergent Hamiltonian is directly interpreted as the energy of the system. However, in the existent models of ’t Hooft’s constraining conditions the emergent Hamiltonian is not only hermitian but also Weyl ordered. In the previous section we have seen that it also allows to formulate an operatorial version of the be-able dynamics. It should be noticed that, while at the Planck energy $E_P$ the Poisson bracket (or alternatively the commutator) of the be-able position with its kinetic momentum vanishes, i.e. $\{q_i(t), p_j^{kin}(t)\} \propto \{q_i(t), \dot{q}_j(t)\} = 0$, on the contrary the corresponding Poisson bracket with canonical momentum is the canonical one, $\{q_i(t), p_j(t)\} = 1$.

Since the be-able dynamics is formulated on the tangent rather than cotangent bundle, it is the kinetic rather than canonical momenta that is physically a more relevant object at $E_P$. The third and more pressing issue is the Born rule. We now show that the Born rule is closely related to Koopman–von Neumann’s (KvN) operatorial formulation of classical physics. To this end we can ask ourselves how an observer “living” in the be-able world would do statistical physics on systems described by Eq. (2). In fact, the simplest way is to go through the Hamiltonian $H$. In this case Koopman and von Neumann found a simple recipe for defining the
probability density function. Following KvN we define a “wave function” \( \psi(p,q,t) \) that evolves in time with the Liouville operator, i.e., the Hamilton field (summation over \( i \) is understood)

\[
\hat{H} = -i \partial_{p_i} H(p,q) \partial_{q_i} + i \partial_{q_i} H(p,q) \partial_{p_i},
\]

according to the equation

\[
i \frac{\partial}{\partial t} \psi = \hat{H} \psi.
\]  

(19)

The vectors \( \psi \) are complex wave functions on the phase space \( \Gamma = (p,q) \), with the normalization

\[
\int dp dq |\psi(p,q)|^2 = 1.
\]

The corresponding complex conjugation turns Eq.(20) to

\[
i \frac{\partial}{\partial t} \psi^* = \hat{H} \psi^*.
\]

(21)

By multiplying (20) with \( \psi^* \) and (21) with \( \psi \), and adding them together we obtain

\[
i \frac{\partial}{\partial t} \varrho = \hat{H} \varrho \iff \frac{\partial}{\partial t} \varrho = (-\partial_{p_i} H \partial_{q_i} + \partial_{q_i} H \partial_{p_i}) \varrho,
\]

(22)

where we have defined \( \varrho(p,q) = \psi^*(p,q) \psi(p,q) \). The second equation is nothing but the well-known Liouville equation for the probability density function of a (classical) statistical system whose dynamics is driven by a (classical) Hamiltonian \( H \). If one defines the scalar product between two wave functions \( \psi_1 \) and \( \psi_2 \) as \( \langle \psi_1 | \psi_2 \rangle = \int dp dq \psi_1^* \psi_2 \), then it is easy to show that \( \langle \psi_1 | \hat{H} \psi_2 \rangle = \langle \hat{H} \psi_1 | \psi_2 \rangle \), i.e., the Liouvillean \( \hat{H} \) is a self-adjoint operator, which implies that the norm of any state is conserved during the evolution. The latter is consistent with the interpretation of \( \varrho = \psi^* \psi \) as a probability density function on the phase space \( \Gamma \). Now, since the \( p \) variables are in ’t Hooft’s proposal only dummy variables (true degrees of freedom are be-ables, i.e., variables \( q \) the relevant density matrix is the reduced density matrix \( \tilde{\varrho}(q) = \int dp \varrho(p,q) \). By using ’t Hooft’s Hamiltonian (1) and integrating (22) over \( p \) we obtain

\[
\frac{\partial}{\partial t} \tilde{\varrho}(q) = -f_i(q) \partial_{q_i} \tilde{\varrho}(q) - [\partial_{q_i} f_i(q)] \tilde{\varrho}(q) = -\partial_{q_i} (f_i(q) \tilde{\varrho}(q))
\]

\[
= -i \hat{p}_i f_i(q) \tilde{\varrho}(q) = -i \hat{H}(p,q) \tilde{\varrho}(q),
\]

(23)

where \( \hat{H} \) is ’t Hooft’s Hamiltonian without the \( g \) term. If now we define

\[
\tilde{\psi}(q) = \int_{-\infty}^{\infty} dp \psi(p,q),
\]

(24)

we can use Eq.(20) (respectively Eq.(21)) to compute the evolution of \( \tilde{\psi}(q) \) (respectively \( \tilde{\psi}^*(q) \)) under the ’t Hooft’s Hamiltonian \( \hat{H} \), so that we obtain

\[
i \frac{\partial}{\partial t} \tilde{\psi}(q) = \hat{H}(p,q) \tilde{\psi}(q) \quad \text{and} \quad i \frac{\partial}{\partial t} \tilde{\psi}^*(q) = \hat{H}(p,q) \tilde{\psi}^*(q).
\]

(25)

Multiplying the first by \( \tilde{\psi}^*(q) \) and the second by \( \tilde{\psi}(q) \), summing together, and taking advantage of the particular form of ’t Hooft’s Hamiltonian, we get

\[
\frac{\partial}{\partial t} (\tilde{\psi}^*(q) \tilde{\psi}(q)) = -i \hat{H}(p,q) (\tilde{\psi}^*(q) \tilde{\psi}(q))
\]

(26)

which, confronted with (23), bring us to the identification \( \varrho(q) = \tilde{\psi}^*(q) \tilde{\psi}(q) \).
This is precisely the (configuration-space) Born rule. While in the previous section we have seen that the be-able dynamics can be equivalently described both by classical or (quantum-like) operatorial means, this section reinforces the picture by showing that this parallelism can be extended also to statistical physics. The parallel mode of description of course breaks when constraint \((12)\) is imposed. We thus see that ‘t Hooft’s Hamiltonian has a privileged role among classical Hamiltonians. In fact, as shown in Ref. [25], there are no other Hamiltonian systems with the peculiar property that their full quantum evolution coincides with the classical one.

We should finally note that the above canonical scenario has also a path-integral counterpart. In this case the KvN operatorial version of the deterministic system \([11]\) is taken over by the classical path integral of Gozzi et al \([25, 26]\). In fact, because both \(\tilde{\psi}(q)\) and \(\tilde{\varrho}(q) = |\tilde{\psi}(q)|^2\) satisfy the same equation of the motion – Schrödinger’s equation, the solutions can be written through the same evolution kernel, i.e.

\[
\tilde{\varrho}(q, t) = \int dq' K(q, t; q', t') \tilde{\varrho}(q', t'), \quad \text{and} \quad \tilde{\psi}(q, t) = \int dq' K(q, t; q', t') \tilde{\psi}(q', t').
\]  

(27)

Although both above kernels represent formally identical expressions, they are qualitatively different. The first kernel in (27) corresponds to a transition probability, while the second corresponds to a transition amplitude. The fact that the same kernel propagates both \(\tilde{\psi}\) and \(|\tilde{\varrho}|^2\) was, in the KvN framework, firstly recognized and discussed in Refs. [26]. The corresponding discussion for ‘t Hooft Hamiltonian system was done in Ref. [25]. The evolution kernel itself has the path integral representation

\[
K(q, t; q', t') = \mathcal{N} \int_{\xi(t) = \xi_1}^{\xi(t') = \xi_2} D\xi \exp \left[ i \int_t^{t'} dt \left( L(\xi, \dot{\xi}) \right) \right] = \mathcal{N} \int_{q(t) = q_1}^{q(t') = q_2} Dq \prod_i \delta[\dot{q}_i - f_i(q)].
\]

(28)

Here we have defined \(2N\) configuration-space coordinates as

\[
\xi_i = p_i, \quad i = 1, \ldots, N \quad \text{and} \quad \xi_i = q_i, \quad i = N + 1, \ldots, 2N,
\]

and the Lagrangian

\[
L(\xi, \dot{\xi}) = \frac{1}{2} \xi^i \omega_{ij} \dot{\xi}^j - H(\xi),
\]

(30)

(\(\omega\) is the \(2N \times 2N\) symplectic matrix). The factor \(\mathcal{N}\) is a normalization factor, and the measure can be explicitly written as

\[
\mathcal{N} \int_{\xi(t) = \xi_1}^{\xi(t') = \xi_2} D\xi = \mathcal{N} \int_{q(t) = q_1}^{q(t') = q_2} Dq Dp.
\]

(31)

The kernel representation (28) corresponds to the classical path integral of Gozzi et al \([26]\). The constraining procedure can be now applied directly to the path integral (28) either through Dirac–Bergmann \([21]\) or Faddeev–Jackiw procedure \([25, 27, 28]\). The actual constraining mechanism is however different when applied to path integrals representing transition amplitudes or to path integrals representing transition probabilities. In particular, for transition probabilities one should rewrite the path integral in the Feynman–Vernon closed-time-path form \([29]\), and then apply the constraint to each time branch independently. This lead to a different emergent kernel than in the case of transition amplitudes. So, similarly as in the operatorial framework, in the path-integral formulation the constraint \((12)\) removes the dichotomy of classical-quantal formulation of the be-able dynamics. It should be stressed that the symplectic structure in both kernels will be, however, still preserved, yet now with an effective Hamiltonian and with different degrees of freedom \([28]\).
Figure 2. Schematic representation of the emergent quantum description. At the primordial level $E_P$ both classical and quantal descriptions are equivalent. Also, both the state $\psi$ and the deterministic density matrix $\rho$ are described via the same evolution kernel $K$. As the energy scale lowers, the information is continuously lost in the process of coarse graining and the observer sees only effective dynamics, i.e. equivalence classes of primordial dynamics. The dashed lines remind that constraint (12) must be applied at all energy scales. Operatorial and deterministic description are coarse-grained together and at low energies are perceived as QM and SAD, respectively. On the statistical level SAD is described via a density matrix with a symplectic evolution kernel. The latter is what the observer perceives as CP.

4. Conclusions and outlooks

In this paper we have demonstrated that when KvN operatorial formulation of classical physics is combined with ’t Hooft’s Hamiltonian for be-ables, one naturally arrives at the Born’s rule without postulating it. This result substantiates ’t Hooft’s pre-quantization scheme and affirmatively answer to the question whether QM can be a hidden variable theory. It should be stressed that the be-able dynamics is fully local (see Eq.(1)). The non-locality that provides the loophole in Bell inequalities is realized in ’t Hooft’s pre-quantization by identifying the physical space of observables $\Gamma^*_E$ with a quotient space $\Gamma_c/G_E$ that has a non-local structure with respect to the physical space of be-ables $\Gamma$.

The picture one can draw from our discussion is the following (see Fig.2): at the primordial scale $E_P$ the dynamics is purely deterministic and it can be equivalently tackled with classical (description on the level of single trajectories) and operatorial (description on the level of evolution in the Hilbert space of states) means. As one lowers the energies, the typical length and time scales are getting larger, and the corresponding description is getting coarse-grained. This leads to a huge information loss and brings one to an effective level of description. A typical observer at $E \ll E_P$ has then at his/her disposal the operatorial mode of description — the Schrödinger equation (15) with the effective Hamiltonian $\hat{H}_+$, which he/she calls Quantum Mechanics. On a parallel footing, there is also available to him/her a non-operatorial, non-local, coarse-grained description of the deterministic system that propagates at energy $E_P$. This non-local imprint of the be-able dynamics$^2$ (SAD) has, however, the statistical density matrix that is described with the standard classical Liouville equation, or equivalently with a classical kernel for transition probabilities. The kernel itself can be represented via path integral with the Lagrangian that exhibits an explicit symplectic structure. It is thus natural for our observer to define as Classical Physics (CP) the physics experienced at the macroscopic statistical level.

$^2$ We denote the non-local imprint of the be-able dynamics as SAD to stress its formal connection with Einstein’s “spooky action at distance”.
This emergent macroscopic description exhibits a symplectic structure which is then identified as a typical signature of CP. In this view, one should not be surprised that BI's are violated at an observational scale. This is because our understanding of CP (on which Bell's inequalities are ultimately based) is not derived from SAD dynamics alone (where BI's could be easily violated) but only from its statistical behavior.

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