QUANTUM MECHANICS AS A SPACE-TIME THEORY

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ABSTRACT. We show how quantum mechanics can be understood as a space-time theory provided that its spatial continuum is modelled by a variable real number (qnumber) continuum. Such a continuum can be constructed using only standard Hilbert space entities. The geometry of atoms and subatomic objects differs from that of classical objects. The systems that are non-local when measured in the classical space-time continuum may be localized in the quantum continuum. We compare this new description of space-time with the Bohmian picture of quantum mechanics.

1. WHAT IS QUANTUM SPACE-TIME?

Both modern mathematics and modern physics underwent serious foundational crises during the 20th century.

The crisis in mathematics occurred at the beginning of the century and the main problem was to deal with certain infinities that are directly related to the concept of real number. Poincaré [31] explained this crisis in terms of different attitudes to infinity, related to Aristotle’s actual infinity and the potential infinity (the first attitude believes that the actual infinity exists, we begin with the collection in which we find the pre-existing objects, the second holds that a collection is formed by successively adding new members, it is infinite because we can see no reason why this process should stop). It led finally to the emergence of new, non-standard definitions of real numbers.

The crisis in physics concerns the interpretation of the quantum theory, the measurement problem and the question of non-locality.

In previous works we showed how in principle certain paradoxes of the quantum theory can be explained provided we enlarge our conception of number [10]. Our goal was to show how the basic axioms of quantum mechanics can be reformulated in terms of non-standard real numbers that we call qnumbers. It is our goal in the present paper to analyze non-locality and the concept of space-time at the light of the new conceptual tools that we developed in the past.

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Our main motivation is that most discussions of quantum mechanics use a background space-time that is the same as classical space-time, usually without any supportive arguments and even sometimes denying that quantum mechanics is a space-time theory. And yet many of the difficulties in understanding quantum phenomena derive from the use of classical space-time. We claim in the present paper that the space-time of quantum phenomena differs from that of classical phenomena in the nature of its continuum. According to our theory [10], the description of quantum phenomena requires a real number continuum that is not the classical continuum. It is not even a fixed element of the theory but varies with the quantum system in a way similar to the way the metric geometry of Einstein’s general relativity varies with the physical system [2]. This is not part of the usual paradigm of quantum theory but adopting it enables us to reformulate the paradoxes of the standard interpretation when each quantum system has its own real number continuum.

The points of quantum space are identified with triplets of quantum real numbers, which we call qrumbers to help distinguish them from quantum numbers of standard quantum theory. Qrumbers are real numbers that are taken as numerical values by quantum systems [2]. One important way in which they differ from standard real numbers is that each qrumber has a non-trivial extent to which it is valid. This extent depends upon the condition of the system. Moreover a copy of the standard reals is embedded in the qrumbers, in fact, to every extent, the standard rationals are dense in the qrumbers.

In this approach, a non-standard qrumber is never obtained as the output from the measurement of a quantity because a measurement is a process in which a standard rational number is obtained as an approximation to the qrumber value of the quantity whose extent is conditioned by the measurement [10]. The empirical fact behind this is that measurements can only produce standard rational numbers.

If we accept the identification of classical points in standard Euclidean space with triplets of standard real numbers, then a single point of quantum space may be approximated by different classical points depending on how the measurements are made. For example, it is possible that a quantum particle localized in the vicinity of a point in quantum space is not localized in the neighbourhood of a single classical point. Furthermore a quantum particle with qrumber values for its position and momentum has a trajectory in qrumber space [2]. This does not contradict the Heisenberg uncertainty relations which only restrict the product of the ranges of standard real number values of positions and momenta that can be prepared or measured [10]. The trajectories in qrumber space are obtained as solutions to equations of motion that are discussed in Ref. [2, 10]. They have the form reminiscent of the
classical Hamiltonian equations of motion but instead they are formulated in terms of the qrumber values of position and momentum. In a sense this result evokes a picture reminiscent of the Bohmian picture in which trajectories in a generalised configuration space are associated with a quantum system. However the ontologies of the two pictures are different at the level of kinematics and dynamics.

In this paper the standard Hilbert space formulation of quantum mechanics is interpreted as a space-time theory by using an approach to the coordination problem similar to that suggested by Dieks \[15\]. In his approach the numerical values of the position of a particle are taken to be attributes of the particle in the same way as its mass or charge. Then the assignment of numbers to the physical properties of a particle is made in such a way that the standard form of the physical laws governing the motion of the particle is maintained. We use a real number continuum given by the sheaf of Dedekind reals $\mathbb{R}_D(\mathcal{E}_S)$ in the topos of sheaves on the quantum state space $\mathcal{E}_S$. A qrumber is a local section of the sheaf $\mathbb{R}_D(\mathcal{E}_S)$ defined in section \[72\] \[21\].

Accordingly, the ontology of quantum particles is that of classical particles except that the numerical values of their attributes are given by qrumbers.

This definition mixes standard concepts of the Hilbert space but also sophisticated concepts that were developed in the framework of non-standard analysis. Nevertheless it is sufficient for the comprehension of the nextcoming results to visualise qrumbers associated to a set of quantum observable as a cloud of standard real numbers that we obtain by computing the average values of these observables while the density matrix associated to the system varies in a certain open set or extent. In this view, sharp values for the observables that are characteristic of the classical picture (in terms of standard real numbers) gets replaced by ”fuzzy” values.

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In this paper we focus on the changes in the concept of localisation. A particle that is localised in terms of the qrumber space may be non-localised when viewed from the classical real number space. For example, we consider Bell’s experiment for two massive identical spin-1/2 particles \[4\]. When the standard description is expressed in terms of qrumbers we can show that the qrumber distance between the identical particles is zero at the time that the spin measurements in the classically separated Stern-Gerlach apparatuses are carried out. We compare this with the Bohmian picture \[7\] in which the separation is zero only in average \[18\].
2. What is a space-time theory?

In this paper, we restrict our attention to space-time theories of particles the prototype of which was given by Newton in *Principia*.[30] In it the motion of a material point particle is described by giving the values of its spatial coordinates as functions of time. The space consists of the set of values of the spatial coordinates available to the material particle. The space of a Newtonian particle is three-dimensional Euclidean space, that is, the spatial coordinates of a classical particle at a particular instant are given by a triplet of three standard real numbers. Each triplet is identified with a point in a three dimensional Euclidean geometry. The properties of the points are defined abstractly through Euclid’s postulates which impose restrictions on the classes of real number continua that can be used to label the points, but the restrictions are not sufficient to uniquely define the real number continuum. Once a real number continuum is determined for the theory, then the numerical values of the spatial coordinates at an instant depend upon a frame of reference. The set of permissible frames of reference is invariant under the relativity group of the theory.

We define a space-time theory of a Galilean relativistic quantum system to be a theory of that system in which the particles have spatial coordinates that take numerical values at each instant of time. The set of values of the spatial coordinates available to the particles gives the spatial continuum, the set of instants available to the particles gives the time continuum. Mathematically these identifications are only valid up to an isomorphism.

Newton in Ref.[30] introduced the concepts of absolute and relative space,

“Absolute space, in its own nature, without relation to anything external, remains always similar and immovable. Relative space is some movable dimension or measure of the absolute space, which our senses determine by its position to bodies; and which is commonly taken for immovable space.”

We understand this distinction in the following way. When we have a physical theory which is expressed in mathematical terms we can understand absolute space as being just the abstract mathematical structure of space. For example, Newton’s absolute space is the purely axiomatic Euclidean geometry used in *Principia*. Then the distinction between absolute and relative space merely denotes the difference between an axiomatic geometry for space, which is a purely abstract mathematical construction, and what Einstein called the "practical" geometry of space that is obtained when the "empty conceptual schemata" of axiomatic geometry are coordinated with "real objects of experience". The recognition and identification of the "real objects of experience"
depends upon the structures that the theorist has imagined previously. In 1930 Einstein [20] noted,

"It seems that the human mind has first to construct forms independently before we can find them in things. Kepler’s marvelous achievement is a particularly fine example of the truth that knowledge cannot spring from experience alone but only from the comparison of the inventions of the intellect with observed fact."

The purely abstract mathematical construction that lies behind the present approach is a special case of the idea of a topos first developed by Grothendieck, then by Lawvere and Tierney to replace set theory as the proper framework for mathematics [29]. We will use a spatial topos, $\text{Shv}(X)$, the category of sheaves on a topological space $X$. The standard real number continuum is the prototype of a class of real number continua, which we call the Dedekind reals $\mathbb{R}_D(X)$, that are given by the sheaf of germs of continuous real valued functions on the topological space $X$. The open subsets of $X$ are the extents to which the numbers exist, they give the truth values. Hence the internal logic is intuitionistic in general but is Boolean when $X$ is the one point space or when the topology on $X$ is trivial.

In the following paragraphs we assume that a physical attribute of a physical system is a quantity that would yield a single-valued, standard or classical real number if it were measured. Thus each of the three components of a position vector is a physical attribute.

In the standard formulations of Galilean relativistic quantum mechanics, in a given reference frame, at each instant of time, each physical attribute of a quantum system is represented by a self-adjoint operator $\hat{A} \in \mathcal{A}$, an $O^*$-algebra [23]. The states of the quantum system are normalized positive linear functionals on $\mathcal{A}$. When restricted to the real algebra $\mathcal{A}_{sa}$ of self-adjoint operators the functionals map $\mathcal{A}_{sa}$ to $\mathbb{R}$. The set of functionals is called the quantum state space, $\mathcal{E}_S$, of the system. The qnumbers of a system are the Dedekind real numbers $\mathbb{R}_D(\mathcal{E}_S)$ on its quantum state space.

The classical mechanical analogue of quantum state space, $\mathcal{E}_S$, is isomorphic to the one point space $\{*\}$, so that the Dedekind real numbers for a classical system are $\mathbb{R}_D(\{*\}) = \mathbb{R}$, the standard real numbers. To see this, consider a system of massive particles in the Hamiltonian formalism of classical mechanics. In a given reference frame, at each instant of time, each physical attribute of the classical particles is represented by a real number, its value at that instant of time in that reference frame. Therefore the algebra $\mathcal{A}_{cl}$ of the representatives of all physical attributes at one time in a given reference frame is isomorphic to the algebra of real numbers $\mathbb{R}$. Then it follows that the set of normalized positive linear functionals on $\mathcal{A}_{cl}$ contains only the identity map. Therefore the space of “states” for a classical mechanical system is the one point space and hence $\mathbb{R}_D(\{*\}) = \mathbb{R}$.
The concept 'state' employed in the preceding paragraph is different from that normally used in classical mechanics. Here, as in standard quantum mechanics, a state is a normalized positive linear functional on the algebra of physical quantities in a given reference frame at a given time. In classical mechanics, a state is given by a point \((q, p)\) in phase space. The latter is both determinative, because \((q, p)\) is the initial condition for Hamilton’s equations of motion that uniquely determines the future states of the system, and generative, because every physical quantity is given by a real-valued function of \((q, p)\). The quantum states do not have the generative property and therefore do not play the same role as the classical mechanical states. If the classical mechanical states are taken to be ontic, the quantum states can be taken to be epistemic.

2.1. Arguments that QM is not a space-time theory. The following is a summary of the argument that quantum mechanics is not a space-time theory [15].

1. The Hilbert space formalism is self-sufficient and does not need a space-time manifold as a background. There is no special role for position, all physical quantities have the same status; they are all represented by 'observables', i.e. by self-adjoint operators.

2. Furthermore, the position and momentum quantities are represented by operators that do not commute. It then follows from the standard interpretation that they cannot both have well-defined values at the same time. If the eigenvalue-eigenstate link is accepted (this is the rule which says that a quantity has a definite value only if its system is in an eigenstate of the corresponding observable) a particle cannot have both a well-defined position and a well-defined momentum at any instant and hence cannot have a trajectory because non-commuting operators don’t have common eigenstates.

3. The nub of the argument is that "the standard mathematical formalism of quantum theory strongly deviates from its classical counterpart in that physical magnitudes are not represented by functions on space (or on phase space)". The operators used for the representation of physical magnitudes in quantum theory are not automatically associated with real number values.

We think that these arguments do not prove that quantum mechanics is not a space-time theory, rather they show that a particular interpretation of the Hilbert space formalism of quantum mechanics does not have a space-time picture.

1. The first point is dubious because many of the problems of the standard interpretation of QM, such as the problems of locality, can be related to the absence of quantum space. It has been asserted of this interpretation that "all the problems centre around the relation between - on the one hand - the values of physical quantities, and - on
the other hand - the results of measurements” [24]. These values and results are usually taken to be the same type of entities, by the precedent of classical physics, but the ancient Greek “measurement problem” - that the results of measurements should be rational numbers, but the value of the ratio of the diagonal to the side of a square is not - should caution us. The values of physical quantities can be a type of real number more general than the numbers obtained as the results of measurements. Moreover the standard interpretation does not adequately distinguish the representation of a physical quantity from the values that the physical quantity may take. This appears to be a prejudice based on their confluence in classical mechanics.

2. The second point re-emphasizes the problems of the standard interpretation. The eigenvalue - eigenstate link hypothesis is ambiguous; it is not clear whether it refers only to measured values. Dirac states that “a measurement always causes the system to jump into an eigenstate of the dynamical variable that is being measured” [16] but does not say from where the state has jumped, c.f., the discussion in Bell [4].

We will re-interpret the standard mathematical formalism of QM to reveal an underlying space that generalises the space of classical mechanics. From this point of view, the main error of the standard interpretation was to not examine the mathematical structure of quantum mechanics with sufficient care to discover the spatial structure existing within it. It is worth noting that in the Bohmian approach [7] where the privileged observable (beable) is the position, the eigenvalue-eigenstate link hypothesis is not fulfilled in general, even when the observable that we consider is the position. Indeed if this hypothesis was fulfilled, the fact that trajectories are supposed to exist would imply that wave functions are not spread in space. The eigenvalue-eigenstate link is also not true when we consider velocities. This does not lead to any inconsistency in the Bohmian picture where instantaneous velocities are not directly measurable [7].

3. Again this comment is true only if by ”space” we mean the space of standard real numbers. It shows that quantum mechanics cannot be represented as a space-time theory with standard real numbers. In classical mechanics, the algebra of the physical quantities is represented by real numbers that are identified with the entities that represent their values. This leads to the identification of physical quantities, such as the energy $H(p,q)$, that are given by functions on phase space with the values that their functions take at points on phase space. It is this identification that is not maintained in quantum mechanics, but

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3 This hypothesis is often not listed in the axioms for QM, e.g. it is not in Ref. [9], because operators don’t always have eigenstates. In particular the position and momentum operators have no eigenvalues, although we can construct states that yield approximate eigenvalues [34].
we claim that classical space (or phase space) does not provide the underlying continuum for quantum mechanics. The qr-number values of physical quantities in quantum mechanics are given by continuous functions on the standard quantum state space (defined in section 7.2).

A brief review of the development of the real number concept is relevant, especially in view of the identification of real numbers with points on a straight line, initiated by Descartes, that opened the way to the use of functions to describe the relations between geometrical objects and points. The ancient Greeks, who developed what we call Euclidean geometry, had not made that identification [22]; magnitudes such as lengths, angles, areas and volumes were measured using geometrical motions of translations and rotations of line segments, triangles etc. aided by concepts of similarity and congruence that depended upon the Archimedean principle that we discuss in section 4. It is not surprising that they only used ratios of natural numbers for if we only admit numbers that arise as the outputs from measurements we only admit rational numbers. It is not until we have a mathematical theory that describes what happens between the outcomes and inputs that we need numbers more general than rational numbers to ensure that the equations of the theory have numerical solutions. The properties of the equations and their solutions are used to deepen our understanding of the physical world. It is in this sense that Dedekind’s statement, “Numbers are free creations of the human intellect, they serve as a means of grasping more easily and more sharply the diversity of things.” [14], may be understood.

3. QNUMBERS; REAL NUMBERS FOR QUANTUM MECHANICS

The numerical values taken by physical quantities in quantum mechanics differ from standard real numbers in much the same way as variable functions differ from constants. Like functions they have domains of definition over which they may vary. Their definition comes from topos theory.

**Definition 1.** The qr-number continuum for a quantum particle is the continuum of Dedekind reals $\mathbb{R}_D(\mathcal{E}_S)$ given by the sheaf $\mathcal{C}(\mathcal{E}_S)$ of continuous real-valued functions in the topos $\mathcal{Sh}(\mathcal{E}_S)$ of sheaves on $\mathcal{E}_S$.

To apply this definition to quantum systems, we start from the standard Hilbert space formulation of quantum mechanics as given by von Neumann [34]. In it the physical quantities are represented by the self-adjoint operators defined on dense subsets of Hilbert space or, in the case of bounded operators, on the whole Hilbert space. These self-adjoint operators belong to an $O^*$ algebra $\mathcal{A}$, called the algebra of observables, if the symmetric product is used they form a real algebra $\mathcal{A}_{sa}$ [23]. The states are the other main ingredient of the standard
Hilbert space formulation of quantum mechanics. In the initial formulation the pure states are given by elements (vectors) belonging to the Hilbert space. The concept of state was later generalised to include impure states which are given by positive bounded self-adjoint trace class operators of trace 1. As noted above, the general concept of state is that of positive continuous linear functionals on \( \mathcal{A} \) (see also section 7.2). The pure states are given by projection operators onto one dimensional subspaces of the Hilbert space, which connects them to the original definition in terms of Hilbert space vectors, at least up to a complex multiplicative factor of modulus 1. We denote by \( \mathcal{E}_S \) the Schwartz subspace of the state space described in the appendix.

In the standard theory \( \operatorname{Tr} \hat{\rho} \hat{A} \) is the average or expectation value of the quantity represented by \( \hat{A} \) in the state given by \( \hat{\rho} \). In our theory for each operator \( \hat{A} \in \mathcal{A}_{sa} \) and each open subset \( W \in \mathcal{E}_S \), \( \operatorname{Tr} \hat{\rho} \hat{A} \) defines a real valued function with domain \( W \). In the topos of sheaves on the topological space \( \mathcal{E}_S \), the sheaf of Dedekind reals \( \mathbb{R}_D(\mathcal{E}_S) \) is isomorphic to the sheaf of continuous real-valued functions on \( \mathcal{E}_S \), so that a qrumber defined to extent \( W \) is a continuous function defined on \( W \). With the appropriate topology on \( \mathcal{E}_S \) the functions defined using \( \operatorname{Tr} \hat{\rho} \hat{A} \) are continuous and thus define qrumbers.

It is worth noting that, beside the fact that it shows how non-standard numbers can be used in order to reformulate the quantum theory, our approach also makes it possible to reformulate the postulates of quantum mechanics in terms of average (expectation) values only. This must be put in parallel with the results of S. Weigert who showed that one obtains, using the expectation values of a qurum of quantities, a closed system of linear differential equations that describes the quantum evolution of a system of spin \( s \).

For a Galilean relativistic quantum mechanical system, \( \mathcal{E}_S \) is the set of all normalised, strongly positive linear functionals on the enveloping algebra of the irreducible representation of the Lie algebra of the extended Galilean group. The irreducible representation of the enveloping Lie algebra of the extended Galilean group, labelled by \( (m, U, s) \) with central element \( \hbar I \), is unitarily equivalent to the tensor product \( M \otimes M_s \) of the \textit{Schrödinger} representation \( M \) of the algebra of the Canonical Commutation Relations (CCR-algebra) generated by the operators \( \{ \vec{P}, \vec{X}, \hbar I \} \) with \( M_s \) the irreducible representation of dimension \( 2s+1 \) of the Lie algebra of the rotation group \( SO(3) \). We choose the topology on \( \mathcal{E}_S \) to be the weakest that makes continuous all the functions of the form \( a_Q(\hat{\rho}) = \operatorname{Tr}(\hat{\rho} \hat{A}) \) for self-adjoint operators \( \hat{A} \in M \otimes M_s \). Then the functions \( a_Q \) form a subobject \( A \) of \( \mathbb{R}_D(\mathcal{E}_S) \) on \( \mathcal{E}_S \). Each qr-number defined to extent \( W \) is either a continuous function of the \( a_Q(W) \) or a constant real valued function on \( W \). The sheaf \( \mathcal{C}(W) \) of continuous real-valued functions over the open set \( W \subset \mathcal{E}_S \) can extended to a sheaf over \( \mathcal{E}_S \) by prolongation by zero.
A physical interpretation of these non-standard real numbers may be linked, via the interpretation of a quantum state \( \hat{\rho} \) as representing an ideal preparation process, to the requirement that the whole experimental arrangement must be included in the determination of physical quantities \( [8] \). However our model differs from this requirement in two important ways: firstly, the physical processes that constitute the preparation process may occur naturally without the intervention of experimentalists and, secondly, the preparation procedures are represented by open sets of states.

If a physical attribute of a particle is represented by the self-adjoint operator \( \hat{A} \) then the quumber values that the quantity can take are given by functions \( a_Q(W) \) defined on open sets \( W \) by \( a_Q(\hat{\rho}) = \text{Tr} \hat{\rho} \hat{A} \) for all \( \hat{\rho} \in W \). The open set \( W \) is the extent to which the quantum particle exists; we say that \( W \) is the ontic state of the particle. If the ontic state of the particle is \( W \) then the quumber value of the quantity represented by \( \hat{B} \) is given by a function \( b_Q(W) \) defined on \( W \). In this sense the quumber values of any quantity are determined by the ontic state \( W \). An ontic state \( W \) is specified by the set of all non-empty open subsets of \( W \). Since all conditions are defined pointwise, if \( U \neq \emptyset \) and \( U \subseteq W \) then a quumber \( a_Q(U) \) will satisfy any condition satisfied by \( a_Q(W) \). For example, \( \epsilon \) sharp collimation, see \([10]\), is a property of the ontic state \( W \) because if it holds on \( W \) it holds on all \( V \subset W \). If \( \hat{X} \) is the triplet of self-adjoint operators that represent the position of a particle then its quumber positions are given by the continuous functions \( \vec{x}_Q(W) \) on open sets \( W \). The set of all its quumber positions constitute the quantum particle’s three dimensional space, whose geometric properties depend upon the structure of the underlying continuum of \( \mathbb{R}_D(\mathbb{E}_S) \) \([32]\). \( \mathbb{R}_D(\mathbb{E}_S) \) is a field that is partially ordered but not totally ordered, in particular, trichotomy does not always hold. Also it is not Archimedean in the sense that there are qrumbers \( a \in \mathbb{R}_D(\mathbb{E}_S) \) such that \( a > 0 \) but there is no natural number \( n \in \mathbb{N} \) such that \( n > a \). \( \mathbb{R}_D(\mathbb{E}_S) \) is a complete metric space with respect to a distance function derived from the norm function \( | \cdot | : \mathbb{R}_D(\mathbb{E}_S) \rightarrow \mathbb{R}_D(\mathbb{E}_S) \) that takes a to \( \max(a, -a) \) \([32]\).

With this we can define a distance function between particle positions in \( \mathbb{R}_D(\mathbb{E}_S)^3 \) that is used to define localisation. Quantum localisation is different from classical localisation. For example, suppose that the z-coordinate of a particle has the quumber value \( z_Q(W) \), where \( W \) is the union of disjoint open subsets, \( W = U \cup V \) with \( U \cap V = \emptyset \). \( z_Q(W) \) is a single quumber value and hence represents a single point on the z-axis in quantum space. But if there are standard real numbers \( z_1 < z_2 < z_3 < z_4 \), with \( z_1 < Z_Q(U) < z_2 \), \( z_3 < Z_Q(V) < z_4 \) and \( z_2 \ll z_3 \) then the single quumber value may be
viewed classically as a pair of separated intervals of standard real numbers and hence non-localised. This example of the difference between quantum and classical localisation can be used to understand the 2-slit experiments. A particle may be localised in terms of the qnumber values of its position but not localised in terms of the classical standard real number values. The qnumber distance between a pair of particles may be small even though the classical distance between them is large.

In the Bohmian approach, non-locality [19; 7; 4] is related to the fact that the configuration space to which particles belong is not 3-dimensional but 3N dimensional. In our approach we generalise the concept of real number on which the concept of spatial point relies.

4.1. What is a quantum particle? A Galilean relativistic quantum particle is an entity localised in the qnumber space with the following characteristics which generalize those of a classical particle [5]:

(i) it has permanent properties which always possess qnumber values that may be approximated by standard real number values which are revealed in observations expressed by counterfactual empirical propositions (such as 'if I had performed such and such experiments, I would have obtained such and such standard real number outcomes'),

(ii) it has individuality,

(iii) it can be re-identified through time.

We must recognize the distinction between intrinsic properties that are independent of the condition of the particle, how it was prepared or what interactions it is undergoing and the extrinsic properties that may depend upon the condition of the particle. Particles with the same intrinsic properties are normally said to be identical [26]. For Galilean relativistic particles in quantum mechanics, the intrinsic properties are the mass $m$, a positive standard real, the internal energy $U$, a standard real, and the spin $s$, a natural number or half a natural number. $(m, U, s)$ label the irreducible projective unitary representations of the Galilean group $G$ [28]. The basic properties of the irreducible representations of $G$ are given an appendix.

It turns out that the concepts of indistinguishability (:= two entities are indistinguishable if they agree with respect to all their attributes) and identity (:= two entities are identical if they are the same object) are not equivalent in general [13].

Nevertheless, in quantum mechanics it is true that for all fundamental particles identity implies indistinguishability and vice versa.

We will now study two particle systems in our approach.

5. Two particle systems

5.1. Two different particles. In the standard mathematical framework the Hilbert space for the quantum mechanics of two different particles of unequal masses $m_1$ and $m_2$, internal energies $U_1$ and $U_2$ and
arbitrary spins $s_1$ and $s_2$ is given by the tensor product $\mathcal{H}(1) \otimes \mathcal{H}(2)$ of the Hilbert spaces $\mathcal{H}(1)$ and $\mathcal{H}(2)$ that are respectively the carrier spaces for the irreducible projective unitary representations of the Galilean group $\mathcal{G}$ labelled by $(m_1, U_1, s_1)$ and $(m_2, U_2, s_2)$. For each $j = 1, 2$, $\mathcal{H}(j) = \mathcal{H}_j(ccr) \otimes \mathbb{C}^{2s_j+1}$ where $\mathcal{H}_j(ccr) = L^2(\mathbb{R}^3)$ for both $j = 1, 2$.

We will write

$$\mathcal{H}(1, 2) = \mathcal{H}(1) \otimes \mathcal{H}(2).$$

The physical quantities associated with particle 1 include operators, representing elements of the enveloping algebra of the Lie algebra of $\mathcal{G}$, that are essentially self-adjoint on the Schwartz subspace $S$ of the spatial Hilbert space $L^2(\mathbb{R}^3)$ acting on $\mathbb{C}^{2s_1+1}$. To simplify the notation we will usually denote a tensor product of operators associated with particle 1 by a single symbol $A(1)$. Then the vector operators for particle 1 are position $\vec{X}(1) = \vec{X} \otimes I_{s_1}$, momentum $\vec{P}(1) = \vec{P} \otimes I_{s_1}$, angular momentum $\vec{L}(1) = \vec{L} \otimes I_{s_1}$ and spin $\vec{s}(1) = I_1 \otimes \vec{s}$, where $I_{s_1}$ is the identity matrix on $\mathbb{C}^{2s_1+1}$ and $I_1$ is the identity operator on $L^2(\mathbb{R}^3)$. We take the Schwartz space $S(1)$ to be the tensor product $S \otimes \mathbb{C}^{2s_1+1}$. The physical quantities associated with particle 2 have a similar tensor product composition and will be simply denoted by operators $B(2)$ acting on $S(2) = S \otimes \mathbb{C}^{2s_2+1} \subset L^2(\mathbb{R}^3) \otimes \mathbb{C}^{2s_2+1}$. They include the vector operators for position $\vec{X}(2) = \vec{X} \otimes I_{s_2}$, momentum $\vec{P}(2) = \vec{P} \otimes I_{s_2}$, angular momentum $\vec{L}(2) = \vec{L} \otimes I_{s_2}$ and spin $\vec{s}(2) = I_2 \otimes \vec{s}$, where $I_{s_2}$ is the identity matrix on $\mathbb{C}^{2s_2+1}$ and $I_2$ is the identity operator on $L^2(\mathbb{R}^3)$.

Because the tensor product $S \otimes \mathbb{C}^{2s_2+1}$ is denoted by $S(j)$ for $j = 1, 2$, then $S(1, 2) = S(1) \otimes S(2)$ represents the two particle Schwartz space.

The algebra $\mathcal{A}_{sa}(1, 2)$ of operators representing physical quantities associated with the compound system contains elements of the form $A(1) \otimes B(2)$ as well as other operators, such the total energy $H(1, 2) = \Sigma_{j=1}^3 P_j(1)^2/(2m_1) \otimes I(2) + I(1) \otimes \Sigma_{j=1}^3 P_j(2)^2/(2m_2) + V(1, 2)$ where the interaction operator $V(1, 2)$ is generally not of the form $A(1) \otimes B(2)$. When considered as part of the compound system the properties of particle 1 are represented by operators of the form $A(1) \otimes I(2)$ and the properties of particle 2 are represented by operators of the form $I(1) \otimes B(2)$, with $I(j) = I_j \otimes I_{s_j}, j = 1, 2$.

Schwartzian state space $\mathcal{E}_S(1, 2)$ is the state space of the compound system. $\mathcal{E}_S(1, 2)$ is contained in the convex hull of projections $\mathcal{P}(1, 2)$ onto the one-dimensional subspaces of $\mathcal{H}(1, 2)$ spanned by unit vectors $\psi(1, 2)$ belonging to the two particle Schwartz space $S(1, 2)$. Each state $\hat{\rho} \in \mathcal{E}_S(1, 2)$ is a trace class positive bounded operator on $\mathcal{H}(1, 2)$ with trace 1 which can be written as $\hat{\rho} = \sum \lambda_n P_n(1, 2)$. For all $n$, $\lambda_n \geq 0$, $\sum \lambda_n = 1$ and the $P_n(1, 2)$ are orthogonal projections onto one-dimensional subspaces in $S(1, 2)$. For each $A(1, 2) \in \mathcal{A}_{sa}(1, 2)$ we
define $a(1, 2)_Q(\rho) = \text{Tr} \hat{\rho} A(1, 2)$ for $\rho \in \mathcal{E}_S(1, 2)$. $\mathcal{E}_S(1, 2)$ is given the weakest topology that makes all the functions $a(1, 2)_Q$ continuous.

Physical quantities that are represented by operators of the form $A(1) \otimes B(2)$ have qrumber values $(a(1) \otimes b(2))_Q(W)$ defined on open subsets $W \in \mathcal{E}_S(1, 2)$. Even though as operators $A(1) \otimes B(2) = (A(1) \otimes I(2))(I(1) \otimes B(2))$ it is generally not true that $(a(1) \otimes b(2))_Q(W) = (a(1) \otimes 1(2))_Q(W))(1(1) \otimes b(2))_Q(W))$ because any open set $W$ will contain non-product states from $\mathcal{E}_S(1, 2)$. Of course we can rewrite $(a(1) \otimes 1(2))_Q(W) = b(2)_Q(W(2))$ where $W(2)$ is the open subset of $\mathcal{E}_S(2)$ that is obtained by partial tracing $W \in O(\mathcal{E}_S(1, 2))$ over an orthonormal basis in $\mathcal{H}(1)$. In a similar fashion $(a(1) \otimes 1(2))_Q(W) = a(1)_Q(W(1))$ where $W(1)$ is the open subset of $\mathcal{E}_S(1)$ that is obtained by partial tracing $W \in O(\mathcal{E}_S(1, 2))$ over an orthonormal basis in $\mathcal{H}(2)$.

Therefore $(a(1) \otimes b(2))_Q(W) \neq a(1)_Q(W(1)) \times b(2)_Q(W(2))$ unless for every $\rho \in W, \hat{\rho} = \hat{\rho}(1) \otimes \hat{\rho}(2)$ where $\hat{\rho}(j) \in W(j), j = 1, 2$. Because quantum systems can be entangled, the latter condition is not satisfied in general and qrumerons do not always factorize.

Of particular interest are the qrumber values of the kinematical variables of either of the two particles. For example, on an open set $W \in \mathcal{E}_S(1, 2)$; the position vector of particle 1 will have qrumber value $\vec{x}(1)_Q(\hat{W}(1))$, the momentum vector of particle 1 will have qrumber value $\vec{p}(1)_Q(\hat{W}(1))$, while the position vector of particle 2 will have qrumber value $\vec{x}(2)_Q(\hat{W}(2))$ and the momentum vector of particle 2 will have qrumber value $\vec{p}(2)_Q(\hat{W}(2))$, where for $j = 1, 2, \hat{W}(j)$ is an open subset of $\mathcal{E}_S(j)$ that is obtained by partial tracing $W(j)$ over an orthonormal basis in $\mathcal{C}^{2s_j+1}$, where $W(j)$ was obtained by tracing over an orthonormal basis in $\mathcal{H}(k), k \neq j$. The qrumber distance between the two particles when the compound system is in the open set $W \in \mathcal{E}_S(1, 2)$ is $d(1, 2)_Q(W)$ given by, where $\vec{x}(1) = \vec{X}(1) \otimes I_{s_j}$ etc.,

\begin{equation}
\|(\vec{x}(1) \otimes I(2) - (I(1) \otimes \vec{x}(2))_Q(W))\|_3 = \|\vec{x}(1)_Q(\hat{W}(1)) - \vec{x}(2)_Q(\hat{W}(2))\|_3.
\end{equation}

That is, for each $\hat{\rho}(1, 2) \in W$,

\begin{equation}
d(1, 2)_Q(\hat{\rho}(1, 2)) = \| \text{Tr} \hat{\rho}(1) \vec{X}(1) - \text{Tr} \hat{\rho}(2) \vec{X}(2) \|_3.
\end{equation}

where $\hat{\rho}(j)$ is obtained by partially tracing $\hat{\rho}(1, 2)$ over an orthonormal basis in $\mathcal{C}^{2s_j+1}$ after having partially traced over an orthonormal basis in $\mathcal{H}(k), k \neq j$. $\|\vec{V}\|_3 = (\vec{V} \cdot \vec{V})^{1/2}$ is the standard Euclidean norm of the vector $\vec{V} \in \mathbb{R}^3$.

Consider the following situation; at $t = 0$ particles 1 and 2 have the same qr-number position and their qr-number momenta are of the same magnitude but in opposite directions, that is, $(\vec{x}(1) \otimes I(2))_Q(W) = (I(1) \otimes \vec{x}(2))_Q(W)$, and $(\vec{p}(1) \otimes I(2))_Q(W) = -(I(1) \otimes \vec{p}(2))_Q(W)$. If the particles move freely then at any time $t > 0$, [10]
\[(\vec{x}(1) \otimes I(2))_Q(W)(t) = (\vec{x}(1) \otimes I(2))_Q(W) + \frac{i}{\hbar}m_1(\vec{p}(1) \otimes I(2))_Q(W)\]

and \[(I(1) \otimes \vec{x}(2))_Q(W)(t) = (I(1) \otimes \vec{x}(2))_Q(W) + \frac{i}{\hbar}m_2(I(1) \otimes \vec{p}(2))_Q(W).\]

Therefore at \(t > 0\) the qrumber distance between the particles is

\[
d(1, 2)_Q(W)(t) = \frac{(m_1 + m_2)}{(m_1m_2)}t||\vec{p}(1) \otimes I(2))_Q(W)||_3.\]

where \(||(\vec{p}(1) \otimes I(2))_Q(W)||_3 = ||(I(1) \otimes \vec{p}(2))_Q(W)||_3.\)

Each particle has its own continuous qr-number trajectory, viz., the two particles move in opposite directions along a straight line in qr-number space. But they do not have trajectories in standard real number space. However, at any instant of time, we can show that one of the pair of particles is in a region of classical space while the other is in a second classical region, well-separated from the first. This is done by finding standard real numbers that approximate the qrnumers \((\vec{x}(1) \otimes I(2))_Q(W)(t)\) and \((I(1) \otimes \vec{x}(2))_Q(W)(t)\). That is, we can, in principle, determine an approximate classical trajectory for each particle from its qrumber trajectory. This contrasts with the standard interpretation in which Heisenberg’s uncertainty relations prevents us from determining a continuous trajectory in classical space by which a particular particle could be labelled and its path followed over time.

If the particles move freely then the total momentum is conserved so that for all times \(t\),

\[(\vec{p}(1) \otimes I(2))_Q(W)(t) = -(I(1) \otimes \vec{p}(2))_Q(W)(t).\]

Thus each particle has both a definite qrumber value for its position and momentum for all times, these values remain correlated until some interaction with the particles breaks the symmetry. Of course measurement of the position and momentum of either particle will yield approximate standard real number values which are in agreement with Heisenberg’s uncertainty relations [10].

5.2. Two identical particles. When the two particles are identical then the standard quantum theory does not distinguish between them.

In the standard mathematical framework the Hilbert spaces for the quantum mechanics of two identical particles of mass \(m \neq 0\), internal energy \(V\) and spin \(s\) is given by the symmetric subspace (bosons), or the anti-symmetric subspace (fermions), of the tensor product \(\mathcal{H}(1, 2) = \mathcal{H}(1) \otimes \mathcal{H}(2)\) of the Hilbert spaces \(\mathcal{H}(1)\) and \(\mathcal{H}(2)\) that are identical copies of the carrier space for the irreducible projective unitary representation \(U\) of the Galilean group. \(U\) has labels \((m, V, s)\). For each \(j = 1, 2\), \(\mathcal{H}(j) = \mathcal{H}(ccr) \otimes \mathbb{C}^{2s+1}\) where \(\mathcal{H}(ccr) = L^2(\mathbb{R}^3)\). The symmetric tensor product is written \((\mathcal{H}(1) \otimes \mathcal{H}(2))_+\) and the antisymmetric tensor product is written \((\mathcal{H}(1) \otimes \mathcal{H}(2))_-\). We will write

\[\mathcal{H}(1, 2)_\pm = (\mathcal{H}(1) \otimes \mathcal{H}(2))_\pm.\]
If $\Pi_\pm$ represent, respectively, the orthogonal projection operators from $\mathcal{H}(1,2)$ to $\mathcal{H}(1,2)_+$ or to $\mathcal{H}(1,2)_-$ then for any vector $u \otimes v \in \mathcal{H}(1,2)$, $\Pi_\pm (u \otimes v) = \frac{1}{2} (u \otimes v \pm v \otimes u)$. It is easy to show that the operators that represent physical quantities associated with systems of identical particles must be invariant under permutation of the particles \[26\]. For if we let $\mathcal{P}$ denote the unitary operator on $\mathcal{H}(1,2)$ that interchanges the vector states of the two particles, $\mathcal{P}(u \otimes v) = v \otimes u$, then $\mathcal{P} = 2\Pi_+ - I$ where $I$ is the identity operator on $\mathcal{H}(1,2)$. We then observe that $\mathcal{H}(1,2)_\pm$ are the eigenspaces of $\mathcal{P}$ corresponding to the eigenvalues $\pm 1$. An operator respects the bosonic or fermionic identity of the particles if it sends vectors in one of these eigenspaces to the same eigenspace, therefore it must commute with $\mathcal{P}$. Hence the operators that represent bosonic or fermionic physical quantities must be invariant under permutations.

That is the physical quantities associated with a system of two identical particles, bosons or fermions, are represented by operators $A(1,2)$ that are symmetric under interchange of the labels of the particles. The set of such operators form an algebra $(\mathcal{A}(1,2))_+$. Examples of operators $A(1,2) \in (\mathcal{A}(1,2))_+$ are $A(1) \otimes A(2)$ and $A(1) \otimes I(2) + I(1) \otimes A(2)$ where the $A(j)$ are operators, built from products of operators representing elements of the enveloping algebra of the Lie algebra of $G$, that are essentially self-adjoint on the Schwartz subspace $S$ of the configuration Hilbert space $L^2(\mathbb{R}^3)$ tensored with symmetric spin matrices acting on $\mathbb{C}^{2s+1}$. The $I(j) = I_j \otimes I_s$, are the identity operators on $\mathcal{H}(j) = \mathcal{H}(ccr) \otimes \mathbb{C}^{2s+1}$, $j = 1, 2$.

It follows that all states $\hat{\rho}(1,2)$ of a system of two identical particles must also be symmetric under the interchange of the particles’ labels. This holds for both bosons and fermions. Therefore state space of a system of two identical particles is contained in the symmetric Schwarzian state space $(\mathcal{E}_S(1,2))_+$. The open subsets of $(\mathcal{E}_S(1,2))_+$ are the restrictions of the open subsets of $\mathcal{E}_S(1,2)$ to $(\mathcal{E}_S(1,2))_+$. They give the weakest topology that makes continuous all the functions $a(1,2)_Q$ from $(\mathcal{E}_S(1,2))_+$ to $\mathbb{R}$;

\[
(7) \quad a(1,2)_Q(\hat{\rho}(1,2)) = \text{Tr} \hat{\rho}(1,2) A(1,2)
\]

for $A(1,2) \in (\mathcal{A}(1,2))_+$, when $\hat{\rho}(1,2) \in (\mathcal{E}_S(1,2))_+$. In particular, these functions are continuous with respect to the topology on $(\mathcal{E}_S(1,2))_+$ given by the restriction of the trace norm topology $\nu(\hat{\rho}(1,2)) = \text{Tr} |\hat{\rho}(1,2)|$ to $(\mathcal{E}_S(1,2))_+$.

The kinematical quantities associated with a system of two identical Galilean particles of mass $m \neq 0$ and spin $s$ have qumber values defined to extents $W \in O((\mathcal{E}_S(1,2))_+)$ by continuous functions; $\bar{x}(1,2)_Q, \bar{p}(1,2)_Q, \bar{L}(1,2)_Q$, defined for $\hat{\rho}(1,2) \in W$ by $\bar{x}(1,2)_Q(\hat{\rho}(1,2)) = \text{Tr} \hat{\rho}(1,2) \bar{X}(1,2)$, where $\bar{X}(1,2) = \bar{X}(1) \otimes I(2) + I(1) \otimes \bar{X}(2)$, etc. The two particle system has a trajectory in qumber space that is defined
by the time evolution of these values \[10\]. To obtain trajectories for the individual particles inside this system we define their kinematical quantities following the method used when the particles were not identical.

**Proposition 2.** In a system of two identical particles, the qumber value of any property of particle 1 is the same as the qumber value of the corresponding property of particle 2 when both qumbers are defined on the same open subset \(W(1, 2) \subset (\mathcal{E}_S(1, 2))_+\).

**Proof.** Suppose a physical property of a single particle of type \((m, V, s)\) is represented by the operator \(A\), so that when considered as part of the compound system the property of particle 1 is represented by an operator of the form \(A(1) \otimes I(2)\) and the property of particle 2 is represented by an operator of the form \(I(1) \otimes A(2)\). Either \(A(1) = A_1 \otimes I_{s_1}, A(2) = A_2 \otimes I_{s_2}\) when \(A\) represents a kinematical quantity or \(A(1) = I_1 \otimes A_{s_1}, A(2) = I_2 \otimes A_{s_2}\) when \(A\) represents a spin variable.

Consider the operator \(B(1, 2) = A(1) \otimes I(2) - I(1) \otimes A(2)\). \(B(1, 2) = -B(2, 1)\) is skew-symmetric with respect to the interchange of particle labels. Therefore for any state \(\hat{\rho}(1, 2) \in (\mathcal{E}_S(1, 2))_+\),
\[
\text{Tr} \hat{\rho}(1, 2)B(1, 2) = -\text{Tr} \hat{\rho}(1, 2)B(1, 2)
\]
because \(\mathcal{P}^{-1}\hat{\rho}(1, 2)\mathcal{P}^{-1}B(1, 2)\mathcal{P} = \text{Tr} \hat{\rho}(2, 1)B(2, 1)\). Therefore
\[
\text{Tr} \hat{\rho}(1, 2)(A(1) \otimes I(2)) = \text{Tr} \hat{\rho}(1, 2)(I(1) \otimes A(2))
\]

It is important to be clear that this result says only that when particles 1 and 2 are identical then the qumber values of the physical quantities of particle 1 are the same as those of particle 2. This does not imply that the measured values of quantities associated with particle 1 must be the same as the measured values of the quantities associated with particle 2. Nevertheless this result has considerable consequences for the concept of locality for identical particles, because it says that for each \(\hat{\rho}(1, 2) \in W(1, 2)\),
\[
\text{Tr} \hat{\rho}(1, 2)(\vec{X}(1) \otimes I_{s_1}) \otimes I(2) = \text{Tr} \hat{\rho}(1, 2)(I(1) \otimes \vec{X}(2) \otimes I_{s_2}).
\]
Therefore the qumber distance, \(d(1, 2)_Q(W(1, 2)) = \|((\vec{X}(1))_Q(W(1, 2)) - (\vec{X}(2))_Q(W(1, 2)))\|_3\), between identical particles is zero.

5.2.1. **EPR-type experiment.** Let us now consider an experiment which uses two identical massive particles that are produced in such a way that the sum of their momenta is zero. The experimenter prepares the momenta to be along the line of a standard real number vector \(\vec{a}\). The two particle system is defined to an extent that is given by an \(\epsilon\)-neighbourhood, \(W(1, 2)\), of the symmetrised pure state \(\Pi_{\epsilon}\) that projects onto the vector \(\frac{1}{2}(u_L(1) \otimes v_R(2) \pm v_R(1) \otimes u_L(2))\) where \(L\) and
$R$ refer to wave packets propagating along opposite directions following the line parallel to $\vec{a}$.

If $\epsilon$ is small enough, the reduced states of the single particles belong to open sets that are a fifty-fifty convex combination of an $\epsilon$-neighbourhood of the projector onto $u_L$ and an $\epsilon$-neighbourhood of the projector onto $v_R$ (see section 7.2). The qumber value of the position and velocity of an individual particle are then close to zero for all times for each particle (both 1 and 2). We physically interpret this property as follows: as a consequence of indistinguishability, the individual particle trajectories reduce to those of the centre of mass of the system. If now we consider two particles trajectories in the qumber approach, this is no longer true. Indeed, in our approach, we can describe the position of the pair by the qumbers $\frac{1}{2}(\hat{x}^i(1) \otimes \hat{x}^j(2))_{Q}(W(1, 2)), (i, j = 1, 2, 3)$. If we compute their value, up to negligible corrections that are proportional to $\epsilon$, using the appropriate system of reference axes we obtain the value zero for all except $(\hat{\vec{a}} \cdot \hat{\vec{x}})(1) \otimes (\hat{\vec{a}} \cdot \hat{\vec{x}})(2))_{Q}(W(1, 2))$ which has a qumber value on $W(1, 2)$ which is close to the standard real value $-(v \cdot t)^2$. $v$ is the absolute value of the velocity of each component along the $\vec{a}$ direction and $t$ the time that has elapsed since their emission from the source.

Combining this information about the trajectories with our previous observation of the fact that the center of mass does not depart from the origin, we can infer the correct picture for the pair trajectory, i.e. that one particle moves to the left and the other moves to the right, with equal speed $v$.

It is interesting to note that, similar to what happens in quantum optics, the pictures that we get about one and two particle trajectories are, in a sense, complementary and, to some extent, independent. This is not true in the Bohmian approach where individual trajectories are supposed to contain all the relevant information about the physical reality of quantum systems. Actually, in the example that we considered above, the two particles are entangled which explains why the knowledge of the pair does not reduce to the knowledge of its single components. The explanation of non-local features of the system is also different in the Bohmian approach and in ours because in the former it is due to the non-locality of the quantum potential and undirectly to the existence of an absolute spatio-temporal reference frame while in the latter non-locality is a property of conventional space-time, not of the quantum space. Moreover the measurement process itself is non-local as we shall now discuss with the help of the example provided by the EPR-Bohm-Bell experiment that we shall discuss now.

5.2.2. EPR-Bohm-Bell experiment for identical particles. Consider an EPR-Bohm-Bell experiment which uses two identical spin one-half massive particles. We can obtain the usual quantum mechanical results for
the experiment while maintaining that the two identical particles always have a qrumber position and hence are localised in qrumber space.

The two particle system is now defined to an extent that consists of an $\epsilon$-neighbourhood, $W(1, 2; s_1, s_2)$ of the symmetrised pure state $\Pi_\pm \otimes \pi(s_1, s_2)$ that projects onto the vector $\frac{1}{2}(u_L(1) \otimes v_R(2) \pm v_R(1) \otimes u_L(2)) \otimes (|+s_1 > \otimes |+s_2 > + |-s_1 > \otimes |+s_2 >)$ where $L$ and $R$ label the opposite directions along the line parallel to $\vec{a}$ and $|\pm s_j >$ represents a spin up (down) polarisation state along the directions $\vec{b}$ orthogonal to $\vec{a}$ for particle $j$. If we trace over the spin degrees of freedom the pure state $\Pi_\pm \otimes \pi(s_1, s_2)$ reduces to the pure spatial state $\Pi_\pm$ which is of the same form as that used for spinless particles. Therefore before the spins interact with the magnetic fields, the discussion of the previous section applies; the individual particle trajectories reduce to those of the centre of mass of the system and the qrumber values of position and velocity of the individual particles are then close to zero for all times for each particle. It is only when the two particle trajectories are calculated that we see that one particle travels to the left and the other to the right, but we cannot tell which.

In order to carry out an EPR-Bohm-Bell experiment we must check the spin correlations of the particles. This can be done by letting them pass through Stern-Gerlach devices with magnetic fields orthogonal to the direction $\vec{a}$. The positions of the particles are then measured after their passage through the Stern-Gerlach devices.

As we discuss with greater detail in Ref. [10] there exist several possible ways to describe the evolution of the system in the absence of measurement. For instance one could let it evolve according to the standard unitary (Schroedinger) evolution, or one could define an Hamilton-like evolution at the level of the qrumbers. In the present case, taking account of the gyromagnetic coupling of the spins, the two evolutions would be equivalent up to $\epsilon$ when the system is defined to an extent that consists of an $\epsilon$ neighbourhood $W(1, 2; s_1, s_2)$ around the symmetrised pure state $\Pi_\pm \otimes \pi(s_1, s_2)$.

By choosing $\epsilon$ small enough the trajectories will be enclosed in regions of volume comparable to that due to the spread of the wave function. In well conceived experiments we measure the positions of the particles with detectors larger than this spread [17], so that it does not really matter which type of evolution we adopt in our description of the evolution of the system, provided that the time of passage through the magnetic region of the Stern-Gerlach devices is short enough.

After the passage through these regions and before reaching the detectors, the two particle system is defined to an extent given by an $\epsilon$-neighbourhood of the projection $\Psi_{s_1,s_2}^{+,-}(1, 2)$ onto the vector $\Psi_{s_1,s_2}^{+,-}(1, 2) = \frac{1}{2}(u_L(1) \otimes v_R(2) \otimes |+s_1 > \otimes |+s_2 > + |-s_1 > \otimes |+s_2 > (\pm) v_R^-(1) \otimes u_L^+ (2)) \otimes |-s_R(1) > \otimes +s_L(2) > + u_L^-(1) \otimes v_R^+(2) \otimes |+s_L(1) > \otimes |+s_R(2) > (\pm) v_R^+(1) \otimes u_L^-(2)) \otimes |+s_R(1) > \otimes |-s_L(2) >)$ where $L^\pm$ and $R^\pm$
refer to wave packets propagating to the upper (lower) left and right parts of the plane spanned by $\vec{a}$ and $\vec{b}_K$, $K = L, R$, the direction of the magnetic field on the left or right. $\pm s_K(j)$ denotes spin up/down in the direction of $\vec{b}_K$, $K = L, R$ for the $j$th particle.

Tracing over the spin variables we obtain a reduced spatial state $P_\pm(1, 2)$ given by
\[
\frac{1}{4}(P_{u_L(1)}^+ \otimes P_{v_R(2)}^- + P_{u_L(2)}^- \otimes P_{v_R(1)}^+ + P_{v_R(1)}^+ \otimes P_{u_L(2)}^- + P_{v_R(2)}^- \otimes P_{u_L(1)}^+).
\]
If the wave packets are assumed to be non-overlapping narrow Gaussians then the single particle reduced state for the $j$th particle is
\[
\rho_\pm(j) = \frac{1}{4}(P_{u_L(j)}^+ + P_{u_L(j)}^- + P_{v_R(j)}^+ + P_{v_R(j)}^-),
\]
which is the same state for both bosons and fermions so we’ll suppress the $\pm$. The extent that the $j$th particle exists is given by an $\epsilon$-neighbourhood of $\rho(j)$.

If $\epsilon$ is small enough, the $\epsilon$-neighbourhood of $\rho(j)$ is a convex combination with equal weights of $\epsilon$-neighbourhoods of the pure state projectors onto the four vectors $u^\pm_L, v^\pm_R$.

In each reduced spatial state $P_\pm(1, 2)$, the pure states for particles 1 and 2 are paired. If we associate a segment of a qrumber trajectory with a single particle pure state via its $\epsilon$-neighbourhood then the qrumber trajectory for the identical particles is composed of a pair of distinct trajectories in the crossing diagram ($> -- <$). In the first of the pair, one of the particles, we cannot tell which, is at the top of the descending branch of the cross on the left and the other particle is at its bottom on the right; in the second of the pair, one of the particles, we cannot tell which, is at the top of the ascending branch of the cross on the right and the other particle is at its bottom on the left.

This situation clearly differs from the Bohmian picture in which either the ascending branch is selected or the descending one, but not both at the same time [18].

As we discuss in Ref.[10], it is at the moment of the measurement, when detectors placed beyond the region of the magnetic field click, one of the two branches of the qrumber trajectory is selected in a process equivalent to the collapse of a wave-function. We show in that paper that the probability associated to the respective branches necessarily obeys the Born rule in order to derive a self-consistent formulation of the qrumber approach. Clearly the interaction with the measuring apparatus is non-local in the usual sense when the branches of the cross on the left and right are separated by spacelike distances (where we refer here to distances measured in standard Euclidean space).

In a sense the problem of non-locality for identical particles can be seen to disappear when qrumbers are used because the qrumber distance between the particles 1 and 2 is zero, provided that we accept that space-time of quantum systems is described by its qrumber continuum rather than the classical real number continuum.
6. Conclusions

We have shown that if we are willing to accept that the spatial continuum is not given a priori but is an artefact of the theory then Galilean relativistic quantum mechanics has a space-time that is as real as the standard space-time of classical physics. Galilean relativistic atoms and sub-atomic particles exist in a space whose points are labelled by qrumbers rather than standard real numbers, they move along trajectories in the space of qrumbers. Their properties have qrumber values always. In this setting the ontology of Galilean relativistic atoms and sub-atomic particles is similar to that of classical particles except that the values taken by their attributes are qrumbers not standard real numbers. Accepting that we can have a non-standard spatial continuum for a physical theory allows us to better understand the theory, for example the problems of non-locality in the standard theory of quantum mechanics disappear when the qrumber spatial distance between quantum particles is used. For example, we showed in an explicit example of a Bell-type experiment how identical particles have zero qrumber spatial separation and hence their interactions are in a sense not non-local. The mystery of the two slit experiment is removed because the qrumber position of a quantum particle can be such that the one qrumber position can cover more than one standard spatial position of a classical particle.

Nevertheless, the quantum wholeness remains present because, due to the presence of entanglement, 2 particle behavior does not reduce to 1 particle behavior.

That the quantum space of a quantum system is different from standard Euclidean space does not reduce the efficacy of the space-time view for understanding physics. The standard real number continuum of classical physics presents much the same class of difficulties as the qrumber continuum: we never can observe all the points on the trajectory of a moving particle, there are many more points in the continuum than we can ever observe. There is one important difference, the points of quantum space are faithfully represented by qrumbers which exist to varying extents reflecting the conditions on the physical system under which the quantities have a value. The dependence of the condition of the physical system raises new possibilities. The universality of the numerical values taken by quantities is an idealisation that has real limitations; even in classical physics the length of a metal rod depends upon the ambient temperature. This paper only discusses Galilean relativistic particle theories. we think that will be possible to extend this approach to different relativistic quantum field theories when their symmetry groups are used in place of the Galilean group.
7. Appendices.

7.1. The standard mathematical formalism. We review the mathematical formalism of standard quantum mechanics.

A Galilean relativistic particle of mass $m$, internal energy $U$ and spin $s$ is associated with an irreducible projective unitary representation of the Galilean group $G$. $G$ is parameterised as follows $g = (b, \bar{a}, \bar{v}, R)$: $b \in \mathbb{R}$ for time translations, $\bar{a} \in \mathbb{R}^3$ for space translations, $\bar{v} \in \mathbb{R}^3$ for pure Galilean transformations or velocity translations, $R \in SO(3)$ for rotations about a point. Classically $g \in G$ acts upon the classical space-time coordinates sending $(\vec{x}, t)$ to $(\vec{x}', t') = (R\vec{x} + \bar{a} + t\bar{v}, t + b)$ then

\[(b', \bar{a}', \bar{v}', R')(b, \bar{a}, \bar{v}, R) = (b' + b, \bar{a}' + R'\bar{a} + b\bar{v}', \bar{v}' + R'\bar{v}, R'R)\]

The irreducible projective unitary representation $g \in G \mapsto U(g)$, labelled by $(m, U, s)$ where $m$ is a positive real number, $U$ is a standard real number and $s$ is a natural number or half a natural number, acts on the Hilbert space $\mathcal{H} := \mathcal{L}^2(\mathbb{R}^3) \otimes \mathbb{C}^{2s+1}$. The elements of $\mathcal{H}$ are $(2s + 1)$-component vectors of square integrable functions $\{\psi_i(\vec{x}) : \vec{x} \in \mathbb{R}^3\}_{i=1}^{2s+1}$. The corresponding space-time functions $\psi_i(\vec{x}, t)$ are defined using the generator $H = \frac{1}{2m}\vec{\beta} \cdot \vec{P} + U$ of the time translations, $\psi_i(\vec{x}, t) := (exp(-iHt/\hbar)\psi_i)(\vec{x})$. Then

\[U(g)\psi_i(\vec{x}, t) = exp(i\alpha_m) \Sigma_j D^i_j(R) \psi_j(R^{-1}(\vec{x} - \vec{v}(t - b) - \bar{a}), t - b)\]

where $\alpha_m = \alpha_m(b, \bar{a}, \bar{v}, R; \vec{x}, t) := \left[-\frac{1}{2}m\vec{v} \cdot \vec{v}(t - b) + \bar{m}\vec{v} \cdot (\vec{x} - \bar{a})\right]/\hbar$ and $D^i_j(R)$ are the matrix elements of the irreducible projective representation of $SO(3)$ on $\mathbb{C}^{2s+1}$.

If $H$ is the infinitesimal generator of the subgroup of time translations, $P_1, P_2, P_3$ are the infinitesimal generators of the subgroup of spatial translations along the three orthogonal axes of the standard basis, $K_1, K_2, K_3$ are the infinitesimal generators of the subgroup of velocity translations along those axes and $J_1, J_2, J_3$ are the infinitesimal generators of the subgroup of rotations around those axes then the Lie algebra of the extended Galilean group, with elements $(\theta, g); \theta \in \mathbb{R}, g \in G$, is generated by $H, \vec{P}, \vec{K}, \vec{J}$ and the central element $hI$ whose Lie brackets with all the other 10 elements vanish. The other Lie brackets are

\[\left[J_i, A_j\right] = \epsilon_{ijk} A_k, \vec{A} = \vec{P}, \vec{K}, \vec{J}; \left[H, B_i\right] = 0, \vec{B} = \vec{P}, \vec{J}; \left[H, K_i\right] = -P_i\]

and

\[\left[K_i, K_j\right] = 0, \left[P_i, P_j\right] = 0, \left[K_i, P_j\right] = m\hbar I_{ij}\]

As well as $\hbar I$, the element $U := H - \frac{1}{2m}\vec{\beta} \cdot \vec{P}$ of the enveloping algebra commutes with all the infinitesimal generators. $U$ may be identified as the internal energy of the particle. We usually take $U = 0$. The vector
position operator for the particle is $\vec{X} = \frac{1}{m}\vec{K}$, the vector operator $\vec{L} = \vec{X} \times \vec{P}$ is the orbital angular momentum. The spin vector operator is $\vec{S} = \vec{J} - \vec{L}$. The operator $\vec{S} \cdot \vec{S}$ commutes with all the elements of the Lie algebra and in any irreducible unitary representations $\vec{S} \cdot \vec{S} = s(s + 1)$ with $s$ an integer or half-integer.

The irreducible representation of the enveloping Lie algebra of the extended Galilean group labelled by $(m, U, s)$ with central element $\hbar I$ is unitarily equivalent to the tensor product $\tilde{M} \otimes M_s$ of the Schrödinger representation $M$ of the algebra of the Canonical Commutation Relations (CCR-algebra) generated by the operators $\{\vec{P}, \vec{X}, \hbar I\}$ with the irreducible matrix representation $M_s$ of dimension $2s + 1$, of the Lie algebra of the rotation group $SO(3)$.

The $Schrödinger$ representation of the CCR-algebra $M$ is the representation in which the Hilbert space is $\mathcal{H}(ccr) = \mathcal{L}^2(\mathbb{R}^3)$. $X_j$ is represented by multiplication by the real variable $x_j$ and $P_j$ by $(1/i)$ times the operator of differentiation with respect to $x_j$. Let $\mathcal{S}(\mathbb{R}^3)$ denote the Schwartz space of infinitely differentiable functions of rapid decrease on $\mathbb{R}^3$. Then the physical quantities are represented by self-adjoint elements in the closure $\tilde{M}$ of the CCR-algebra $M$, where $\tilde{M}$ is the smallest closed extension of $M$. $\tilde{M} = \{X \mid X \in M$ and $X$ is the restriction to $\mathcal{S}(\mathbb{R}^3)$ of the Hilbert space closure of $X\}$.

7.2. Qrumber. We use a real number continuum given by the sheaf of Dedekind reals $\mathbb{R}_D(\mathcal{E}_S)$ in the topos of sheaves on the quantum state space $\mathcal{E}_S$. A qrumber is a local section of the sheaf $\mathbb{R}_D(\mathcal{E}_S)$ $\mathbb{E}_S$ is the Schwartz subspace of the state space, consisting of those states $\hat{\rho}$ that are rapidly decreasing convex combinations of projection operators $\hat{P}_j$ onto one dimensional subspaces spanned by vectors in Schwartz space $\mathcal{S}$, that is, $\hat{\rho} = \sum_{j=1}^{\infty} \lambda_j \hat{P}_j$ with $\lambda_j \geq 0$ and $\lim_{j \to \infty} \lambda_j j^n = 0$ for all positive integers $n$.

The topology on $\mathcal{E}_S$ is the weakest that makes continuous all the functions of the form $a_Q(\hat{\rho}) = \text{Tr}(\hat{\rho} \cdot \hat{A})$ for self-adjoint operators $\hat{A} \in \mathcal{A}$. 
$M \otimes M_\nu$. Then the functions $a_Q$ form a subobject $A$ of $\mathbb{R}_D(\mathcal{E}_S)$ which is the sheaf of locally linear functions on $\mathcal{E}_S$. Each $q$-number defined to extent $W$ is either a continuous function of the $a_Q(W)$ or a constant real valued function on $W$. The sheaf $\mathcal{C}(W)$ of continuous real-valued functions over the open set $W \subset \mathcal{S}$ can extended to a sheaf over $\mathcal{E}_S$ by prolongation by zero.

One useful type of open set is the $\epsilon$-neighbourhood of a state $\rho_0$.

**Definition 3.** The $\epsilon$-neighbourhood of a state $\rho_0$ is $\nu(\rho_0; \epsilon) = \{\rho : Tr|\rho - \rho_0| < \epsilon\}$. That is, it is an open ball, in the trace norm topology, of radius $\epsilon$ centred on $\rho_0$.

When $\epsilon$ is small, we can get a good idea of the properties of $a_Q(\nu(\rho_0; \epsilon))$ from those of $Tr(\rho_0 \cdot A)$.

The following result is used in the discussions of identical particles. Let $\Psi(1, 2) = \frac{1}{\sqrt{2}}(\psi_+(1)\psi_-(2) \pm \psi_-(1)\psi_+(2))$ be an entangled wave function of particles 1 and 2, the normalised single particle wave functions being orthogonal, $\langle \psi_+(j), \psi_-(j) \rangle = 0, j = 1, 2$. The corresponding pure state is $P_{\Psi(1, 2)} = \frac{1}{2}(P_{\psi_+(1)} \otimes P_{\psi_-(2)} + P_{\psi_-(1)} \otimes P_{\psi_+(2)} + V_{-\psi_+(-1)} \otimes V_{-\psi_+(-2)} - V_{-\psi_+(-1)} \otimes V_{-\psi_+(-2)})$, where the partial isometries $V_{-\psi_+(-j)} = |\psi_+(j)\rangle < \psi_-(j)\rangle = V_{-\psi_+(-j)^*_j}, j = 1, 2$. The reduced state of particle $j$ is the mixed state, $\rho_0(j) = \frac{1}{2}(P_{\psi_+(j)} + P_{\psi_-(j)})$, for both bosons and fermions.

**Lemma 4.** If $\epsilon < 1$ then the $\epsilon$-neighbourhood of $\rho_0(j)$ does not contain any pure state. If $\epsilon$ is small enough it essentially only contains fifty convex combination of the $\epsilon$-neighbourhoods of $P_{\psi_+(j)}$ and $P_{\psi_-(j)}$, that is, $\nu(\rho_0(j); \epsilon) = \nu(P_{\psi_+(j)}; \epsilon) + \nu(P_{\psi_-(j)}; \epsilon), j = 1, 2$.

**Proof.** We will prove the result for particle 1.

If $\chi(1)$ is a unit vector with $\langle \chi(1) | \psi_{\pm}(1) \rangle = \gamma_{\pm}$ then the vectors $\{\psi_{+}(1), \psi_{-}(1), \chi(1)\}$ span a subspace of dimension at most 3. Then it is easy to calculate $Tr|P_{\chi(1)} - \rho_0(1)| = 1 + (1 - |\gamma_+|^2 - |\gamma_-|^2)^{\frac{1}{2}}$ which is less than $\epsilon$ only if $\epsilon > 1$. Therefore, when $\epsilon < 1$ every state $\rho \in \nu(\rho_0(1); \epsilon)$ is impure.

Now $\frac{1}{2}\nu(P_{\psi_+(1)}; \epsilon) + \frac{1}{2}\nu(P_{\psi_-(1)}; \epsilon)$ is contained in $\nu(\rho_0(1); \epsilon)$. Because if $\rho = \frac{1}{2}\rho_+ + \frac{1}{2}\rho_-$ with $\rho_j \in \nu(P_{\psi_j(1)}; \epsilon)$ for $j = +, -$, then since $\nu$ determines a norm on the space of trace class operators,

$$Tr|\rho - \rho_0(1)| \leq \frac{1}{2}(Tr|\rho_+ - P_{\psi_+(1)}| + Tr|\rho_- - P_{\psi_-(1)}|) < \epsilon.$$  

There are other states in the $\epsilon$-neighbourhood of $\rho_0(1)$. They are of the form, $\rho = \lambda P_{\psi_+(1)} + (1 - \lambda)P_{\psi_-(1)}$ where $\frac{1}{2} - \frac{\epsilon}{2} < \lambda < \frac{1}{2} + \frac{\epsilon}{2}$. They are impure states which are convex combinations of the same two states as $\rho_0(1)$ is. When $\epsilon$ is small they behave like $\rho_0(1)$ itself.

Therefore if a system of two identical particles exists to the extent $\nu(P_{\psi(1, 2)}; \epsilon)$ then the jth particle exists to the extent $\nu(\rho_0(j); \epsilon) = \nu(P_{\psi(j)\psi(j)}; \epsilon)$.
\[ \frac{1}{2} \nu(P_{\psi_+(j)}; \epsilon) + \frac{1}{2} \nu(P_{\psi_-(j)}; \epsilon) \]. Therefore, the \( i \)th coordinate quantum value of the \( j \)th particle's position is

\[ \frac{1}{2} x_i^Q( \nu(P_{\psi_+(j)}; \epsilon)) + \frac{1}{2} x_i^Q( \nu(P_{\psi_-(j)}; \epsilon)). \]

In the EPR-type experiment, we used \( \psi_+ = u_L \) and \( \psi_- = v_R \), which are assumed to be non-overlapping narrow Gaussians, \( u_L \) moving with speed \( v \) to the left along the axis \( \vec{a} \), \( v_R \) moving with speed \( v \) to the right along \( \vec{a} \). Then the quantum value of \( \vec{a} \cdot \vec{x}(j) \) for the \( j \)th particle is approximately zero.

On the other hand, when we measure \( \vec{a} \cdot \vec{x}(j) \) for the \( j \)th particle in the vicinity of either of the states \( P_{\psi_+(j)} \) or \( P_{\psi_-(j)} \), we will get a value, which leads us to deduce that the \( j \)th particle has a \( \vec{a} \cdot \vec{x}(j) \) quantum value to both the extents \( \nu(P_{\psi_+(j)}; \epsilon) \) and \( \nu(P_{\psi_-(j)}; \epsilon) \). It follows from the orthogonality of the wave functions and \( \epsilon < 1 \) that the extents are disjoint.

The calculation for the two particles trajectories is based on the assumption that the particles are moving freely. Since we have assumed that for each particle the single particle wave functions are non-overlapping, it is easy to show that the terms

\[ Tr_P \Psi(1,2) (\hat{\mathcal{P}}_i(1) \otimes \hat{X}_j(2) + \hat{X}_i(1) \otimes \frac{\hat{P}^j(0)}{m_2}) \]

\[ Tr_P \Psi(1,2) (\hat{X}_i(1) \otimes \hat{X}_j(2)) \] and

\[ Tr_P \Psi(1,2) (\hat{P}_i(1) \otimes \hat{P}_j(2)) \]

vanish for all \( i, j \) except for the component \( (a,a) \). For instance,

\[ Tr_P \Psi(1,2) (\hat{X}_a(1) \otimes \hat{X}_a(2)) = -(v,t)^2, \]

where \( X_a = \vec{a} \cdot \vec{X}/(\vec{a} \cdot \vec{a}) \). These results extend to the open set \( W = \nu(P_{\psi_+(1,2)}; \epsilon) \) by continuity.

The extension of these results to an EPR-Bohm-Bell experiment with two identical spin one-half massive particles is straightforward: the quantum trajectories obtained for spinless particles in the EPR-experiment are split by the interaction between the spin and the magnetic field. The pair of quantum trajectories become four quantum trajectories.

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References

[1] M. Adelman and J. V. Corbett, Applied Categorical Structures 3, 79 (1995).
[2] M. Adelman and J. V. Corbett: "Quantum Mechanics as an Intuitionistic form of Classical Mechanics" Proceedings of the Centre Mathematics and its Applications, pp 15-29, ANU, Canberra (2001).

[4] J. S. Bell, Phys. 1, 195 (1964).

[4] J. S. Bell, Speakable and Unspeakable in Quantum Mechanics”, pp 117-118 (Cambridge University Press, 1993).

[5] M. Bitbol, “Are there particles and quantum jumps?”, in R. Nair (ed), Mind, Matter and Mystery, pp 52-53, Scientia, New Delhi (2001).

[6] D. Bohm: Quantum theory, Prentice Hall, Englewood Cliffs, N.J. (1951).

[7] D. Bohm, Phys. Rev. 85, 166 (1952).

[8] N. Bohr, “Über die Antwendung der Quantentheorie auf den Atombau. I. Die Grundpostulate der Quantentheorie”, Z. Phys., 13 (1923) p 118, “one is obliged ... to always keep in mind the domain of application” and Atomic Theory and the Description of Nature, Cambridge (1932), p 73, “constantly keep the possibilities of definition as well as of observation before the mind”

[9] C. Cohen-Tannoudji, B. Diu and F. Laloe, Mecanique quantique, (Hermann, Paris, 1977).

[10] J. V. Corbett and T. Durt, Quantum Mechanics re-interpreted in qr-numbers, (2004) preprint.

[11] J. V. Corbett, The classical limit of trajectories in qr-number space, (2005) preprint.

[12] J. V. Corbett The Pauli Problem, State Reconstruction and Quantum-Real Numbers, ROMP, (2005)

[13] N. C. A. da Costa, D. Krause, S. French, "The Schroedinger Problem" in M. Bitbol, O. Darrigol (ed) Erwin Schroedinger, Philosophy and the Birth of Quantum Mechanics, pp 450-453, Editions Frontiers, Gif-sur-Yvette, France.

[14] R. Dedekind, "Was sind und was sollen die Zahlen"., Braunschweig, 1887, SIII.

[15] Dieks, D. "Space and Time in Particle and Field Physics", Stud. Hist.Phil.Mod.Phys., 32, 217-241 (2001)

[16] P. A. M. Dirac, "The Principles of Quantum Mechanics", 4th ed., Oxford University Press (1958) p 36.

[17] T. Durt: PhD thesis: "From quantum to classical, a toy model", Vrije Universiteit Brussel, February 1996.

[18] T. Durt and Y. Pierseaux: "Bohm’s interpretation and maximally entangled states" , Phys. Rev. A 66 (2002) 052109-052120.
[19] A. Einstein, B. Podolsky, and N. Rosen, Phys. Rev. 47, 777 (1935).

[20] Einstein, A. "Johannes Kepler", Frankfurter Zeitung, 9 November, 1930, on the 300th anniversary of Kepler’s death, republished in "Ideas and Opinions", Crown Publishers, Inc., (1954), pp 256-259.

[21] Einstein, A. "Geometry and Experience", Lecture before the Prussian Academy of Sciences, 27 January, 1921, republished in "Ideas and Opinions", Crown Publishers, Inc., (1954) pp 227-232.

[22] Hartshorne, R. Geometry: Euclid and Beyond, Springer, New York, Berlin, Heidelberg (2000)

[23] A. Inoue, Tomita-Takesaki Theory in Algebras of Unbounded Operators Lecture Notes in Mathematics; 1699, Springer (1999), K. Schmudgen, Unbounded Operator Algebras and Representation Theory Operator Theory: Vol. 37, Birkhauser Verlag (1990).

[24] Isham C.J. and Butterfield, J. "Some possible Roles for Topos Theory in Quantum Theory and Quantum Gravity", Fndns of Physics (2000)

[25] M. Jammer, The Conceptual Development of Quantum Mechanics, p175 New York (1966)

[26] J. M. Jauch, Foundations of Quantum Mechanics, p275-280, Addison-Wesley, Reading Massachusetts, (1968)

[27] W. Lawvere, “Quantifiers as Sheaves”, Actes Congres Intern. Math. 1, (1970).

[28] J.-L. Lévy-Leblond, “Galilei group and Galilean invariance”, in E.M. Loebl (ed), Group Theory and Its Applications, Academic Press, New York, pp 221-299, (1971).

[29] S. MacLane and I. Moerdijk, Sheaves in Geometry and Logic (Springer–Verlag, New York, 1994).

[30] Newton, I. Philosophiae Naturalis Principia Mathematica English translation, University of California Press, 1962).

[31] Poincare, H., “La logique de l’infini”, Dernieres Pensees, Flammarion, (1913), pp. 84 - 89. English translation, Mathematics and Sciences: Last Essays , Translator J W Bolduc, Dover, (1963) Chapter V.

[32] L. N. Stout, Cahiers Top. et Geom. Diff. XVII, 295 (1976); C. Mulvey, “Intuitionistic Algebra and Representation of Rings” in Memoirs of the AMS 148 (1974).

[33] R.G. Swan, The Theory of Sheaves (University of Chicago Press, Chicago and London, 1964) §3.2, pp 29-31.

[34] J. V. von Neumann: Mathematische grundlagen der Quanten-Mechanik , Springer-Verlag, Berlin (1932). English translation:
Mathematical Foundations of Quantum Mechanics, Princeton University Press, Princeton, 1955.

[35] S. Weigert: “Quantum Time Evolution in Terms of Nonredundant Expectation Values”, arXiv:quant-ph/9903103 v1 31 Mar 1999

[36] H. Weyl: Philosophy of Mathematics and Natural Science Appendix C, pp255-257, Princeton University Press, Princeton (1949).