MEASUREMENT IN QUANTUM PHYSICS

Michael Danos
Enrico Fermi Institute,
University of Chicago, Chicago, Illinois 60637, USA

and

Tien D. Kieu
School of Physics,
University of Melbourne, Parkville, Victoria 3052, Australia

Abstract

The conceptual problems in quantum mechanics – related to the collapse of the wave function, the particle-wave duality, the meaning of measurement – arise from the need to ascribe particle character to the wave function. As will be shown, all these problems dissolve when working instead with quantum fields, which have both wave and particle character. Otherwise the predictions of quantum physics, including Bell’s inequalities, coincide with those of the conventional treatments. The transfer of the results of the quantum measurement to the classical realm is also discussed.

1 Generalities

A vast literature exists on the interpretation of quantum mechanics in general, and on the meaning of measurement in quantum mechanics in particular. The discussion still takes place today; from small, semi-popular papers to highly technical large-scale programmatic treatments. To quote Bell, Ref. , who in discussing some of his articles in that book writes in the preface: “these [articles] show my conviction that, despite numerous solutions of the [measurement] problem ..., a problem of principle remains.” (See also Feynman, Ref. .) The most fundamental of the problems interfering with the understanding of quantum mechanics is indeed the problem of measurement, the “genesis of information,” and all the effects surrounding what has been termed “the Copenhagen collapse of the wave function,” which is not described by the Schrödinger equation . This process not only lies outside of the framework of quantum mechanics, but, being instantaneous, also violates relativistic causality. As we will see, all of this is closely related

*Visiting Scholar.
†Present address: CSRIO, Australia.
to the so-called “particle-wave duality” and is the source of Bell’s above mentioned “problem of principle.” Several different proposals to overcome Bell’s above “problem of principle” have been made. They all in some way break the framework of quantum theory. We shall not discuss these proposals, but refer the reader to the above mentioned literature citations [1, 2, 3, 4, 6].

Our point is that all problems associated with the subject known as “the problem of measurement in quantum mechanics” can be resolved without abandoning or supplementing quantum theory. That means that in this context invoking of “new physics” is not needed; even more strongly, it is counter-indicated.

A separate problem in the measurement process is the need to describe how a result of an interaction between the measured object and the apparatus on the quantum level is transferred to the classical level, e.g., to a pointer position. We shall address both these problems in our paper.

Quite generally, a theory is supposed to be able to make “predictions” in the sense that given a “state of the system” at some time, say, $t_0$, the theory must be capable of providing information on the state of the system at time $t$, where $t$ may be later or earlier than $t_0$. The “predictions” here can apply before the relevant experiment has been performed, or after. Also, the other terms, e.g., “the state of the system”, are defined within the theory. In quantum physics, which is our framework, this means (see Sections 7, 8) that given the state of the system as $\rho(t_0)$ the probability of an outcome, $O_k(t)$, is given by $Tr\{U(t,t_0) \rho(t_0) U(t,t_0)^\dagger O_k(t)\}$ where $U(t,t_0)$ is the evolution operator of the system, and where $k$ specifies the outcome. For example, the outcome could be a particle with spin-up emitted along a particular direction. In general there are many, usually infinitely many, possible outcomes. However, otherwise quantum physics is complete in that no hypotheses lying outside of the quantum physics framework are needed, for any and all circumstances.

Quantum physics does not make predictions on which of the possible outcomes actually will occur. In particular, it does not predict “when the event (e.g., the decay of the nucleus) actually will take place.” Such a prediction, being outside of quantum theory, would be in conflict with the Heisenberg uncertainty relations. One of the attempts to overcome this “limitation of the theory” is the deBroglie-Bohm pilot wave hypothesis [7]. This attempt supplements the wave
function by a “particle function” $X(t)$, called by the authors a “hidden variable.” Even though outside of quantum mechanics, this work in fact provided the stimulus for Bell to derive the famous Bell inequalities [8] which allow for the distinction by experiment between at least a large class of hidden-variable theories and quantum physics. The experiments by now have come out in favor of the quantum physics predictions [9].

On the other hand, the theory is silent on the choice of the initial conditions. In particular, one will have to chose $t_0$ sufficiently far back so that the “switching-on” transient is contained within ones treatment. This is for example the case when investigating the unperturbed [10] or perturbed [11] non-exponential part of the decay of unstable or metastable states. Another well-known case is that of the correlations and of the statistics in high-intensity particle beams. The theory will provide answers for any $\rho(t_0)$ and $O_k(t)$ even if the initial conditions actually are not, and can not be, realized in Nature. The reasons for this impossibility may be obvious, or may be extremely subtle. A discussion of this is given in Ref. [12]; it lies outside of the scope of the present paper.

We have used the term “outcome” rather than the usually employed term “measurement”. The reason for this is that “measurement” seems to imply the participation of an observer. By using the term “outcome” we want to indicate that one is free to define the system as broad or as narrow as one likes. Thus one may want to include the observer in the system, which the theory and the formalism allows; but this choice is irrelevant for our discussion.

In the present essay we want to show that Bell’s conceptual gap [4] disappears when recognizing that the particle-wave duality is only an artifact of quantum mechanics, related to the absence there of the particle aspects which are amputated when going from quantum field theory to quantum mechanics. The wave function contains only the wave aspects. Quantum fields (relativistic or non-relativistic) contain simultaneously both the particle and the wave aspects. The concepts required to construct the needed framework are very few, and very simple. In the present context the full interacting quantum field theory is not required; one does not have to go beyond the lowest order, i.e., no Feynman graphs containing loops need be considered. This limited theory is known to pose no mathematical difficulties [13]. This then is the subject we are going to address first (in Section 2). Next, in Section 3, we shall show in which way quantum mechanics (QM) is a sub-field of quantum physics (QP), and then we will be in the position to
describe the process of measurement. We do that in the next Sections by analyzing a series of experiments, and show that both the wave and the particle aspects are needed in the description of the measurement process. As we will see, no conceptual gaps remain when using both these aspects. In particular, no split between the quantum system and the conscious experimentalist is needed; the experimental apparatus, including the experimentalist, can be considered to be part of the quantum system.

Essentially all of the concepts needed in the description of the measurement in quantum physics are present already in the case of a two-slit experiment, Section 4, where the measuring arrangement consists of an array of detectors. The interference pattern which arises in response to the wave aspect emerges as the result of a large number of experiments, i.e., as a probability distribution. The particle aspect manifests itself in forbidding coincidences: in a weak beam situation only one counter at a time can register an event. Here already the “collapse of the wave function” and “communication at faster than the speed of light” between the counters of the array has to be operative if one wants to describe the situation in the frame of quantum mechanics. The last of the concepts, the sensitivity of the interference between different reaction channels to an intervening measurement, here also can be fully elucidated: a measurement which determines the slit through which the particle travelled destroys the interference pattern. Such a measurement changes the two-slit into a single-slit pattern.

The Einstein-Podolski-Rosen (EPR) experiment [14], discussed in Section 5, is the generalization of a two-slit experiment to a two-particle system; if one wants to describe it in quantum mechanics one must replace the 3-dimensional space of the two-slit experiment by a 6-dimensional configuration space. It requires a somewhat more complex experimental arrangement and also a more complex theoretical analysis. It is the simplest setup allowing for two-particle coincidences. When viewed within the frame of quantum mechanics in 3-dimensional position space it has indeed the well-known dramatic conceptual problems. Not so in quantum physics: the description of all these experiments is fully contained within its framework.

In the next two Sections we discuss in detail the completion of the quantum measurement, i.e., the reaction of a classical system (a pointer, Schrödinger’s cat) to the result in the quantum system. This involves the description of a classical system in terms of quantum physics; the mathematics needed for this description is sketched in Appendix A.
No split between the “quantum system” and the “classical apparatus” is needed; all has to be, and can be, considered from the quantum point of view. Another frequently ignored aspect is that measurement inescapably is an irreversible process, i.e., is associated with dissipation [12] taking place already on the quantum level. This point is important and must be taken into account whenever a complete description, including the measuring apparatus, is attempted.

The present paper does not address the question of the logical superstructure, denoted “the interpretation of quantum mechanics” in Refs. [2, 3]. The arguments and descriptions of these references in fact involve assumptions concerning the measurement process which are consistent with, actually follow from, the results of the present paper.

In summary, all aspects required in the description of the evolution of physical systems, including the act of measurement, are contained within quantum physics. Of course, the 19-th century dream of a fully deterministic description, which would transcend the frame of quantum physics, remains unfulfilled.

2 Quantum Physics

In order to make the paper self-contained we now collect the rudimentary aspects of quantum physics required for the present purpose [13].

In quantum physics the particular state under consideration is described by a state vector, \( |S\rangle \). Thus, for example, \( |S(x_1, x_2, t)\rangle \) represents a state such that at time \( t \) the system had two particles, one located at \( x_1 \), the other at \( x_2 \). The state vector for the system which has no particles is given the special notation \( |V\rangle \), and is denoted as “the vacuum.” The field operator, denoted by \( \Psi(x,t) \), *interrogates* the state vector for the presence of a particle at the point \( x,t \) in the form

\[
\Psi(x,t) |S(y,t)\rangle = \delta(x - y) |V\rangle \tag{1}
\]

with

\[
\Psi(x,t) | V \rangle = 0 \tag{2}
\]

(in these equations, and throughout in this paper, we shall use units such that \( \hbar \) and \( c = 1 \)). Hence one calls \( \Psi \) a “particle annihilation operator.” \( \Psi(x,t) \) and \( \bar{\Psi}(x,t) \) are defined to obey the
anti-commutation relations (commutation relations for Bosons)

\[ [\Psi(y,t), \bar{\Psi}(x,t)]_+ = \delta(x - y) \] (3)

where the \( \delta \)-function implies the structure of a point particle. Comparing (1) and (3) one sees that

\[ |S(x,t)\rangle = \bar{\Psi}(x,t) |V\rangle . \] (4)

In view of (4) one calls \( \bar{\Psi}(x,t) \) the “creation operator;” at the same time one sees from (3) that this operator creates a point particle at the position \( (x,t) \). The field operator \( \Psi(x,t) \) is defined to obey the appropriate equations of motion, e.g., in the nonrelativistic case the Schrödinger wave equation

\[(i \partial_t - H) \Psi(x,t) = 0 \] . (5)

If relativistic effects are important, the Dirac equation or other appropriate equations apply, in which case one has to consider the negative-energy solutions and so on but our arguments below will not be affected.

Equations (4) and (5) mean that \( \Psi(x,t) \) has particle character which propagates as a wave. In other words, the quantity \( \Psi(x,t) \) of quantum physics does not suffer from particle-wave duality; it has simultaneously both particle and wave characteristics. This, of course, is not possible in classical physics – nor in quantum mechanics.

Both for computational purposes and for visualization, it is useful to factorize \( \Psi \) into the particle and the wave aspects. This can be done by computing a complete set of \( c \)-number functions, say \( \psi_n(x,t) \) which obey the wave equation (5) \( \text{together with the boundary conditions appropriate to the system.} \) Then one can write the expansions

\[ \Psi(x,t) = \sum_n b_n \psi_n(x,t) \] (6)

for the field and

\[ \bar{\Psi}(x,t) = \sum_n b_n^\dagger \bar{\psi}_n(x,t) \] (7)

for the hermitian conjugate field. Inserting these definitions in (4) one sees that this equation is fulfilled if the quantities \( b_n, b_n^\dagger \) obey the anti-commutation relations

\[ [b_n, b_n^\dagger]_+ = \delta_{n,n'}, \] (8)

\[ [b_n^\dagger, b_n^\dagger]_+ = 0 = [b_n, b_n^\dagger]_+ \] (9)
which then leads to the completeness relation in the form

$$\sum_n \bar{\psi}_n(x,t) \psi_n(y,t) = \delta(x - y)$$

(10)

for the complete set of the solutions. It is useful to introduce the abbreviation

$$\Psi_n(x,t) = b_n \psi_n(x,t) .$$

(11)

Herewith

$$\Psi(x,t) = \sum_n \Psi_n(x,t) .$$

(12)

The content of the Eq. (11) can be expressed as: acting on the vacuum the operator $\bar{\Psi}_n(x,t)$ through the operator $b_n$ creates a particle in the state $\psi_n(x,t)$. Or, said differently, $b_n$ is the particle aspect, and $\psi_n(x,t)$ is the wave aspect of the quantum physics function $\Psi_n(x,t)$. For example, the anticommutation relations (8) ensure that at most one particle can occupy the state $\psi_n(x,t)$. In the next Section we will argue that $\psi_n(x,t)$ is linked to the probability interpretation of quantum mechanics.

We complete this description by giving the expression for the above-mentioned two-particle state vector

$$|S(x_1, x_2, t)\rangle = \bar{\Psi}(x_1, t) \bar{\Psi}(x_2, t) |V\rangle .$$

(13)

On the other hand the state vector of a system having a particle in the state $\psi_n(x,t)$ is

$$|S_n\rangle = b_n^\dagger |V\rangle .$$

(14)

Below we will deal mostly with systems of that kind.

3 Quantum Mechanics

The basic concept of quantum mechanics is “the wave function,” also called “the probability amplitude.” The wave function for the mode $n$, e.g., the state $n$ of the hydrogen atom, is denoted as $w_n(x,t)$. The meaning of this notation is defined as: given any position $x$, and any time $t$, the value of the wave function is the number $w_n$. The probability of finding the particle there is then $|w_n(x,t)|^2$. To compute the wave function itself one must solve the Schrödinger equation, or, if relativistic effects are important, the Dirac equation, imposing the appropriate boundary conditions on the solutions. Both these equations are wave equations.
How does this wave function emerge from quantum physics of the last Section? We make the ansatz

\[ w_n(x,t) = \langle V | \Psi(x,t) | S_n \rangle, \tag{15} \]

where

\[ |S_n\rangle = b^\dagger_n |V\rangle \tag{16} \]

is the state vector for the system in state \( n \). Equations (15) together with (11) yield

\[ w_n(x,t) = \psi_n(x,t) \tag{17} \]

which turns out to be consistent since both \( w_n(x,t) \) and \( \psi_n(x,t) \) fulfill the same equation and the same boundary conditions. This shows that the quantum mechanics wave function is the wave part of the quantum physics function. The function \( w_n(x,t) \) describes only “the wave aspects” of quantum physics; it lacks “the particle aspects” which have been lost in the interrogation (15). Thus, in contrast to the quantum physics function \( \Psi_n(x,t) \), which describes a particle propagating, i.e., moving through space and time, the quantum mechanics wave function \( w_n(x,t) \) describes a nothing propagating. The latter is a rather abstract entity, having led to many a fruitless search for the meaning of the deBroglie-Schrödinger wave function, and, with it, the meaning of quantum mechanics. In order to have a description which contains both the particle and the wave aspects one needs to work in quantum physics.

## 4 Preliminary Conclusions and Consolidation

From our discussion above one sees that in quantum physics complementarity, or, as it is also called, the particle-wave duality, is necessarily absent since the state function (11), \( \Psi_n(x,t) \), contains simultaneously both aspects. It describes the motion, i.e. the propagation, of a point particle through space and time. This propagation is that of a wave, which precludes the possibility of assigning a trajectory to that motion. In contrast, quantum mechanics simply lacks the particle concept, which is expressed by the interrogation formulae (1), (2), (3). The particle aspect has been eliminated from quantum mechanics at the point where the wave function was extracted from the quantum physics function in the interrogation (13). Thus, the quantity which has been left intact, the wave function \( w_n(x,t) \), describes the propagation of nothing in particular, as exemplified by the Cheshire cat, which had left, leaving only the grin behind.
In short, only the wave function $w_n(x,t)$ exists in quantum mechanics. Of course, the wave function is an exceedingly rich object, as can be seen from the scope of quantum mechanics. However, the particle concept is inescapably needed for the understanding, the interpretation, of the physical content of the results obtained upon computation of the wave function. It therefore must be re-inserted artificially “by hand.” (In the limited domain “quantum mechanics”, which does not include, for example, the radiative corrections, in the calculations themselves the particle aspect is not needed.) This then leads to the particle-wave duality, a logical gap, with the concomitant difficulty in reaching full understanding. This gap, which in fact is Bell’s above quoted “problem of principle”, is one of the factors, most likely the principal factor, which generated the aura of mystery and fog surrounding the subject “modern physics.”

5 The Two-Slit Experiment

Any measurement requires an interaction between “the system” and the “measuring device”, and in quantum physics every interaction involves the emission or absorption of a particle (recall the interaction term $\bar{\psi} \gamma^\mu A_\mu \psi$ of quantum electrodynamics: a particle is absorbed in the initial state and is re-emitted in a different, the final state; a photon is absorbed or emitted). Thus the measurement process lies outside of the framework of quantum mechanics. Let us discuss the process of measurement in terms of specific examples. This present discussion will require a somewhat more technical language than that of the previous Sections.

Consider the determination of a diffraction pattern in a photon two-slit experiment, and take the low-intensity case to avoid the complication of chance coincidences. The experimental arrangement thus consists of a photon source, an intervening screen with two (or one) slits, and an array of detectors behind the slit-screen to register the photons and make the data available to the experimentalist. The photon field, denoted here by $\varphi(x)$, can be expanded in any set of solutions of Maxwells equations. Here it will be convenient to employ for the two- and the one-slit cases two alternative such expansions, namely those which obey the boundary conditions required to account for the source, screen, two or one slits, etc. In the factorized form of Eqs. (3), (4) the field then is given by (we change of notation from the previous Sections and suppress the vector
character of the photon):

\[ \varphi(x) = \sum_n a_n^{(2)} f_n^{(2)}(x) \equiv \sum_n \varphi_n^{(2)}(x) \]  \hspace{2cm} (18)

if both slits are open, and

\[ \varphi(x) = \sum_n a_n^{(1)} f_n^{(1)}(x) \equiv \sum_n \varphi_n^{(1)}(x) \]  \hspace{2cm} (19)

if only one slit is open. (Of course, the right-hand sides of Eqs. (18) and (19) can be expanded in terms of either of \( f_n^{(k)}(x) \). Expansion in the \( \text{“wrong”} \) function converges uniformly except in the vicinity of the slits.) The individual terms of these two sets of solutions, \( f_n^{(2)}(x) \), \( f_n^{(1)}(x) \), which actually are the wave functions of quantum mechanics, are different since the boundary conditions for the two cases are different; in particular, the interference patterns described by these two solutions are different. In these fields, \( f_n^{(k)}(x) \) concerns the wave aspects, while \( a_n^{(k)} \) concerns the particle aspects: \( a_n^{(k)} \) creates, while \( a_n^{(k)} \) annihilates, a particle in state \( f_n^{(k)}(x) \). The two parts of the action of the detector, i.e., (i) the interaction with the photons, and (ii) the registration of a \( \text{“count”} \) and the transmission of the data to the user, and so on, factorize. The action (i) of the detector \( m \) tests for the presence of a particle by interrogating the state vector at the space-time point \( x_m \) (within the resolution of the detector); it is described by the absorption operator \( \varphi(x_m) \), both for the single-slit \( (k = 1) \) or two-slit \( (k = 2) \) case; see. Eqs. (18),(19).

Thus, for instance, the probability amplitude for a particle in state \( n \), being described by the state vector \( |S_n^{(k)}\rangle = a_n^{(k)\dagger}|V\rangle \), to be at the point \( x_m \), is

\[ \langle V|\varphi^{(k)}(x_m)|S_n^{(k)}\rangle = f_n^{(k)}(x_m) \]  \hspace{2cm} (20)

(The also correct form \( \langle V|\varphi^{(1)}(x_m)|S_n^{(2)}\rangle \) is inconvenient in that it results in a linear combination of the functions \( f_n^{(1)}(x_m) \).) We collect the description of the action (ii) of the detector in an appropriate operator \( \eta \), which also includes the detector efficiency. In this way, the action of the detector at the point \( x_m \) can be described by the detector function

\[ D_m = \sum_n \eta_m^n \varphi_n^{(k)}(x_m) \]  \hspace{2cm} (21)

The extent of the sum over \( n \) depends on the characteristics of the detector; \( \eta \) also contains the reaction of the measuring apparatus and hence is an appropriate quantum operator. The actual construction of the detector is of no importance here; as an example, the absorption of the photon
may result in the ionization of an atom, and the emitted electron may initiate a discharge as in a proportional counter.

For a system having an “incoming” photon in either the two-slit \((k = 2)\) or the one-slit \((k = 1)\) situation, the probability amplitude for the response of the detector \(m\) is

\[
A^{(k)}_m = \langle df| \otimes \langle V|D_m|S^{(k)}_n \rangle \otimes |di \rangle \sim \langle df|\eta_n^m|di \rangle \sim \eta_n^m \cdot f^{(k)}_n(x_m) \, ;
\] (22)
as to be expected the detector responds to the interference pattern of the photon field, described by \(f^{(k)}_n(x_m)\). Here \(\langle df|\eta|di \rangle\) denotes the expectation value describing the detector response, from state \(|di \rangle\) before the interaction to \(|df \rangle\) the state after detection. As always, the probability for counter \(m\) to respond is \(\left|A^{(k)}_m\right|^2\).

Both the particle and the wave aspect contributed to the result, Eq. (22). The particle aspect, the factor \(\varphi(x_m)\) contained in \(D_m\), absorbed/annihilated the photon, and this took place in a local manner, precisely at the (four-)point \(x_m\) in the detector \(m\); it also provided the factor \(f^{(k)}_n(x)\), i.e., the wave aspect, which is the appropriate solution to Maxwell’s equations.

Furthermore, once one detector has registered a photon, then no other detector can respond since the particle already has been absorbed. This follows from the expression for the probability amplitude, say \(A_c\), of a coincidence in detectors \(m\) and \(m'\) (dropping the irrelevant factors),

\[
A_c \sim \langle V|D_{m'}D_m|S^{(k)}_n \rangle \sim \langle V|D_{m'}|V \rangle = 0 \, ;
\] (23)
the probability for a coincidence thus vanishes in view of Eq. (2). Expressed in spoken language, the meaning of Eq. (23) is: “The answer to the question ‘can two detectors, \(D_m\) and \(D_{m'}\), absorb a single particle, \(S^{(k)}_n\)?’ is NO.” The absence of a coincidence is enforced by the particle aspect.

The requirement for a “decision” of hitting this one, or that one, but then no other detector, is the prototype of the need for the “collapse of the wave function” of quantum mechanics, which, of course, is not described by the equations of motion, e.g., the Schrödinger equation. Namely, in quantum mechanics the counter \(m'\) must somehow be made to “know” that the counter \(m\) has been hit. The wave function originally in general is non-zero at both places. Thus there is no reason for counter \(m'\) not to respond at the same time. To avoid such a coincidence one therefore in quantum mechanics must mimic the uniquely local character of the absorption process. This is accomplished by “collapsing the wave function”; essentially from \(f^{(k)}_n(x_m)\) to a delta-function
at \( x_m \) – or at \( x_{m'} \) if it was detector \( m' \) which had responded. This “collapse” is even more spectacular in the case of a more complicated reaction where the wave function extends not only over a small region of space but over a perhaps large number of reaction channels; to accomplish this feat “the needed signal: ‘collapse the wave function!’ may have to propagate faster than light”. (Of course, no such signal is available in quantum theory.) We will return to this point below in the discussion of the Einstein-Podolski-Rosen experiment.

In case one tries to check “which slit the photon passed through” one has to place a detector in the slit, say at \( x_s \). To know that the photon passed through this slit this detector would have to record a Compton scattering event. In this detection process the original photon is absorbed and a new photon is emitted, having a new energy and a new radiation pattern appropriate to the new geometry (as required by the dependence of the solution on the boundary conditions), i.e., radiation from within the slit. The Compton detector function then would have the form

\[
D_s \sim \eta_s \varphi^{(-)}(x_s) \varphi^{(k)}(x_s)
\]  

(24)

where \( \varphi^{(-)}(x_s) \) is the emission part of \( \varphi(x_s) \) for a photon with the new radiation pattern (that of only one slit open) constructed in analogy to Eqs. \( \text{[18]} \) or \( \text{[19]} \). \( \eta_s \) is as previously the operator describing the reaction of the detector, here the recoiling atom. The new radiation pattern is different from the old radiation pattern. In particular, it does not exhibit the two-slit interference pattern.

The two-slit interference pattern will disappear upon the Compton scattering of the photon even if nobody actually observes the counter, or even if the counter is broken. Such processes, of course, take place all the time; they are called “collimator scattering” and contribute to the experimental background. That means that the photon needs not “to know that it has been observed” to change the interference pattern. This way Nature in quantum physics “has an objective existence; it exists by itself” independent of observation. Hence the experimental apparatus, and by extension the experimenter, can be considered to be “part of the system”, without any difficulties, conceptual or otherwise.

Instead of putting a detector in one of the slits to find out which slit the particle passed through, one may place a screen over one of the slits after the particle has been emitted. This case requires the description by localized wave-packet states which will be introduced in the next
Section. Then, independently of whatever happened before or after, the state of the system is given by the solution of Maxwells equations appropriate to the boundary conditions valid at the time when the wave packet arrives at and passes through the slits. The state of the system at the time when the photon just has been emitted is given as always by

\[ \varphi(x,t) = \sum_n' C_n^{(2)} f_n^{(2)}(x) \left( a_n^{(2)} e^{-iEt} \right) + \sum_n' C_n^{(1)} f_n^{(1)}(x) \left( a_n^{(1)} e^{-iEt} \right) \]  

(25)

where the coefficients \( C_n^{(2)} \) and \( C_n^{(1)} \) are determined by the emission process. The primes at the summation signs indicate that the sums are to be taken over the functions appropriate to the case to avoid double counting. (If the screen with the slits is “very far” from the source, then \( C_n^{(1)} \) and \( C_n^{(2)} \) may be equal.) The probability for registering a count therefore will contain the factor \(|C_n^{(k)}|^2\), multiplying the square of the right-hand side of Eq. (22). Thus, in all of these experiments, the existence or non-existence of a diffraction pattern or a coincidence is determined by the particle aspects while the probabilities are given by the wave aspects; and no collapse takes place or is needed. In other words, the “yes or no” is determined by the particle aspects, the “how much” by the wave aspects of quantum physics.

6 The E-P-R Experiment

Particularly notorious as a “paradox” is the Einstein-Podolski-Rosen example [8, 14] which in quantum mechanics for its “explanation” combines the need for wave function collapse and action at a distance. With respect to the above two-slit example, the EPR setup has two added features (which lead to complications of detail, but not of principle): (i) the system contains two particles; and (ii) the time-dependence must be accounted for.

Consider the point (ii) first. An eigenstate of the Hamiltonian by definition has a precise energy. Hence the wave function of an eigenstate factorizes as

\[ \phi(x,t) = e^{-iEt} \phi(x) \]  

(26)

The probability to find the particle at the point \( x \), \(|\phi(x,t)|^2 = |\phi(x)|^2\), hence is independent of the time. With this solution thus it is impossible to specify a time as being before, during, or after the experiment. To achieve this possibility one must chose a suitable superposition of these
states. We follow the Weyl prescription. Thus for a free particle we chose the form
\[ w(E_k; x, t) = \mathcal{N} \int_{E_k - \Delta}^{E_k + \Delta} e^{i[px - E'(t-t_0)]} \, dE' \]  
(27)
where \( \mathcal{N} \) is the normalization constant. This function is localized: it peaks at \( x = 0 \) at the time \( t = t_0 \); as \( t \) increases the peak propagates towards larger \( x \) with a velocity the corresponding classical particle would have. As is well known, one must distinguish between the phase velocity, \( a = E/p \) and the group velocity, \( b = dE/dp \). It is the latter which corresponds to the classical particle velocity; in view of
\[ p = v \sqrt{m^2 + p^2} \]  
(28)
and
\[ E = \sqrt{m^2 + p^2}. \]  
(29)
we find \( b = v \). On the other hand \( a = 1/v \). The width \( \Delta \) of the superposition determines the width of the peak in \( x \); also, \( w(E_k; x, t_0) \) is a minimum-uncertainty wave function.

With functions of this kind one thus can describe, for example, the following: the particle was emitted from the source at time \( t_0 \), arrived at the scattering center at time \( t_1 \), and was absorbed in the detector at time \( t_2 \). From now on we will work exclusively with Weyl functions.

We explain the point (i) above directly in terms of the EPR setup. There at time \( t = t_0 \) two particles of spin \( s = 1/2 \), coupled to total spin \( S = 0 \), are emitted in opposite directions, go through a series of polarizers and analyzers to be finally absorbed in two widely separated detectors. The point of the experiment consists in changing the setup at random after the particles have been emitted and have become separated by such a distance that “they cannot communicate” without violating relativistic causality.

We need the following definitions. As the functions \( w \), Eq. (27), form a complete set the field operator can be expanded as
\[ \varphi(x, t) = \sum a_n \, w_n(x, t) \]  
(30)
Since we will only deal with the system when the particles are far from the source it is convenient to split the field into separate field operators for the two directions. Denoting spin “up” and “down” by the indices + and − respectively, taking the quantization to be along the z-direction,
we have for the split field operators (suppressing the time)

\[ \varphi_w(x) = \sum_n (a_n^+ w_n^+(x) + a_n^- w_n^-(x)), \quad x > 0 ; \]  

(31)

\[ \varphi_v(x) = \sum_n (b_n^+ v_n^+(x) - b_n^- v_n^-(x)), \quad x < 0 . \]  

(32)

Thus here \( w \) denotes a particle emitted “to the right,” and \( v \) “to the left”. Since the particles are described by Weyl functions they indeed fly apart. The state vector of the system then is

\[ |S_2\rangle = \mathcal{N} (a_n^\dagger b_{n-}^\dagger - a_{n-}^\dagger b_{n+}^\dagger) |V\rangle , \]  

(33)

with \( \mathcal{N} \) the normalization factor; note the absence of the summation over \( n \). In the following we will suppress the quantum number \( n \) for brevity.

The detectors are endowed with polarization analyzers, and are placed at \(-X\) and \(+X\) respectively with “very large” separation \( 2|X| \). Denoting the polarization of the analyzer by the subscript \( p = + \) or \(-\) the detector response is given as

\[ D_1 = a_+^\dagger w_+(x) \eta_+ + a_-^\dagger w_-(x) \eta_- \]  

(34)

and

\[ D_2 = b_+^\dagger v_+(x) \eta_+^\prime + b_-^\dagger v_-(x) \eta_-^\prime \]  

(35)

where detector 1 is at \( x = X \) and detector 2 at \( x = -X \). The probability for obtaining a coincidence, i.e., a count in both detectors, irrespective of whether the spin state is “up” or “down” then is given by (we suppress the detector states)

\[ A \sim \langle V | D_1 D_2 |S_2\rangle . \]  

(36)

The quantum probability for detection thus will have an interference term for spins at \( X \) opposite to that at \(-X\), as can be readily deduced from (31).

Now insert a polarization-sensitive filter in arm 1, such that only the “up” state is transmitted. This filter is represented by the projection operator

\[ F = a_+^\dagger a_+ \]  

(37)

which has the effect given by

\[ F a_+^\dagger |V\rangle = a_+^\dagger a_+ a_+^\dagger |V\rangle = a_+^\dagger |V\rangle \]  

(38)
\[ F a_+ \dagger |V\rangle = a_+ \dagger a_+ a_- \dagger |V\rangle = 0 . \]  (39)

Thus Eq. (36) is replaced by
\[ A \sim \langle V | D_1 D_2 F | S_2 \rangle . \]  (40)

Now only the term with spin “up” at \( X \) and “down” at \( -X \) survives. That means, that in a coincidence the detector in arm 1 “determines” the polarization of the particle in arm 2. And it does not matter at what time the filter was inserted in the beam path, as long as that took place before the arrival of the Weyl wave packet. All these results emerge directly in terms of the quantum physics functions; no collapse of a wave function, or transmission of a signal, is required.

To guarantee that, where appropriate, coincidences indeed do occur requires that the detector efficiencies be 100%.

A similar analysis can be carried out for the case of a spin-flip filter,
\[ T = a_+ \dagger + a_- \dagger a_+ \dagger , \]  (41)

the action of which is given by
\[ T a_+ \dagger |V\rangle = a_+ \dagger |V\rangle \]  (42)
or
\[ T a_- \dagger |V\rangle = a_+ \dagger |V\rangle . \]  (43)

Replacing in (40) \( F \) by \( T \) of (41), one finds that coincidence is achieved when both detected spins are of the same “orientation”. As above, it does not matter when the filter was inserted. Again, the counter in one of the arms “determines” the polarization of the particle in the other arm.

The importance of coupling to spin \( S = 0 \) is manifested by the minus sign in Eq. (33). As there is no privileged direction, one can choose the polarization detectors to be sensitive along the y-direction. Then the detector response function (34) is represented by \( a_\pm ^{\dagger} \) and \( b_\pm ^{\dagger} \), the annihilation operators quantized along y-orientation. The state vector of the system still can be represented by (33). However, it now is more convenient to expand the field operator (31) in the basis of y-oriented wave functions, and the above analysis will go through with “up” and “down” now referring to the y-axis. Whatever the representation of the field operator, we obtain the same result.

An interesting case arises when the analyzer of the detector in arm 1 measures the spin along the z-axis and that of arm 2 along the y-axis, say. The state then would appear to the detector
in arm 2 as having terms not only of the form $a^\dagger b^\dagger$ but also of the form $a^\dagger b^\dagger$. Therefore no strict yes-no coincidence rules would exist and only probability predictions would be possible. It is precisely these probabilities which are different in quantum physics and in classical probability. The analysis of this situation forms the basis for the Bell inequalities. All of the Bell predictions, Ref. [8], made in the framework of quantum mechanics are correct, i.e., are in full agreement with those derivable from the above quantum physics description – except for requiring an acausal propagation of the hypothetical signal “inducing the collapse of the wave function”.

7 Dissipation and Decoherence in Measurement

Above we demonstrated that the measurement process in quantum physics poses no conceptual problems when describing it in terms of quantum fields. In that discussion we paid no attention to the full measurement process, which actually can be split into two interdependent stages: the interaction with the quantum object, and the chain of amplification to the macroscopic “pointer position”. The essential point in the second stage is the transfer of the results of the measurement from the quantum physics (QP) part to the Classical Physics (CP) part of the apparatus.

So far we have concentrated only on the first of these stages, i.e., the very first interaction between the “system” and the “apparatus”. In the present Section we shall develop a more complete description, which will allow us to trace the process through to the completion of the second stage, i.e., from the beginning of the amplification chain to the display of the result of the measurement by a “pointer position”. Our aim is to derive the expression for the probability, say $P_p$, of the pointer position $p$ after the conclusion of the measurement process. In essence this amounts to the full treatment of the matrix element $\langle d_f | \eta | d_i \rangle$ of Eq. (22) postponed in Section 5.

Since the “yes or no” question has been answered in the first of the two stages of the overall measurement process, the second part which concerns the “how much” aspect could be discussed simply in the quantum mechanics (QM) framework. For completeness we will, however, also discuss it in the full QP frame.

The essential new aspects to be treated now will be: (i) the inclusion of dissipation which is inherent and inevitable in any actual measurement since any elementary quantum act of the measurement process itself is irreversible, i.e., dissipative [12]; and (ii) the process of decoherence,
which, as we will see, is the physical basis for the conversion from a quantum to a classical process [15, 17]. Superficially, these points seem similar but we will see that actually they differ in an essential manner. Of these two aspects, in particular (ii) requires that the description be done in terms of density matrices (see Appendix A), needed when dealing with non-interfering, “classical” objects. To give a full description we will have to assemble some needed QP quantities, in particular in the context of decoherence.

Concerning point (i), the very interaction underlying the measurement involves a time-reversal non-invariant evolution of the system object-plus-device, no matter whether the objects are quantum or classical. The dissipation in the macroscopic, “classical”, down-part of the chain of the overall process poses no problem. Our subject here thus is the analysis of the dissipation associated with the elementary, the quantum measurement process at the very beginning of the chain.

The point (ii) deals with the transition from the quantum to the classical part of the chain. This involves the description of classical objects in the QP frame. It is essential in resolving the Schrödinger cat paradox.

Begin with point (i), the initial interaction. For definitiveness we take as the example of the initial “quantum detector” a Compton proportional counter as used to determine “the slit the photon went through” in the two-slit experiment of Section 5. Such a counter is filled with a low-pressure gas and contains the needed electrodes. The photon may suffer Compton scattering on one of the electrons belonging to one of the gas molecules, the gas being a classical object of given temperature and pressure. We thus need the quantum description of that classical molecule, both of its motion and of its internal state, for example its rotation-vibration quantum numbers, and so on.

Consider the translational motion. Being confined to the counter volume, it can not be described by a plane wave. Instead, the basis states can be taken as Weyl packets, Eq. (27). Since it is a classical non-interfering state, it must be described by a density matrix (see Appendix A). That means, the density matrix must be made up of Weyl basis states, each of which is confined to the counter volume. Denoting the energy expectation of a Weyl basis state, \( \phi_i \), by \( E_i \), i.e.,

\[
E_i = \langle \phi_i | H | \phi_i \rangle ,
\]  

(44)
the elements of the density matrix will be

$$\omega_{ij} = Z \delta_{i,j} e^{-E_i/T},$$ (45)

with $Z$ the normalization constant, and $T$ the temperature. In (44) the index $i$ may encompass the internal quantum numbers of the molecule. (For room temperature and taking the gas of the counter to be argon we obtain the localization of the atoms, i.e., their position uncertainty, to be less than atomic dimensions, thus negligibly small.) The form of the density matrix for an individual molecule thus is of block-diagonal form: each Weyl basis state, being a wave function as in Eq. (27), is given by a block of fully interfering components, while no interferences exist between different blocks. The density matrix of the complete system, of the proportional counter, is given by the direct product of the individual-molecule density matrices.

To continue we need the QP expressions (see Appendix A) corresponding to the QM expressions used above in Eqs. (27), (44), (45). To that end we write the expansion of the Weyl states in the usual plane wave basis $\chi_i$:

$$\phi_i = \int dp g_i(p) \chi_i(p),$$ (46)

where the functions $g_i(p)$ form a suitable complete orthonormal set chosen in particular to ensure the fulfillment of the boundary conditions, i.e., vanishing outside of the counter. (The form of Eq. (46) is more general than that of Eq. (27).) Then the QP field operator is written

$$\psi_i = \sum_i B_i \phi_i,$$ (47)

where $B_i$ is the annihilation operator for the Weyl mode $i$,

$$B_i = \int dp g_i(p) b_i(p).$$ (48)

Herewith we can write the (bilinear) state vector for the initial-state density matrix

$$\bar{S}^{(0)} = \sum_{ij} B_i^\dagger |V\rangle \omega_{ij}^{(0)} \langle V|B_j \equiv \sum_{ij} \bar{S}_{ij}^{(0)}.$$ (49)

We shall call these quantities “density state vectors.” Similarly we have mutatis mutandis for the final state

$$\bar{S}^{(f)} = \sum_{ij} B_i^\dagger |V\rangle \omega_{ij}^{(f)} \langle V|B_j \equiv \sum_{ij} \bar{S}_{ij}^{(f)}.$$ (50)
which we will have to compute by considering the evolution of the system.

Denoting the photon Fock-space operators by \( a^{(0)} \), \( a^{(f)} \) for the initial and final photon state, respectively, we have for the complete density state vector

\[
S^{(k)} = \sum_{mn,ij} \bar{S}^{(k)}_{ij} \sigma_{mn}^\dagger a^{(k)}_m \sigma_{mn} a^{(k)}_n \equiv \sum_{mn,ij} S_{mn,ij}^{(k)}
\]

with \( (k) = (0), (f) \), and \( \sigma_{mn} \) the photon density matrix.

The interaction with the incoming photon is, as always,

\[
H(x) = ie \bar{\psi}(x) \gamma_{\mu} \psi(x) A^{\mu}(x) \ ;
\]

here \( \psi(x) \) can be expanded in terms of the functions \( \psi_i \), Eq. (47). As we are interested in a Compton process, the full interaction, including the rest of the apparatus, indicated as previously by \( \eta \), is

\[
I \eta = H(y) G(y, x) H(x) \eta.
\]

Here \( G \) is the relevant Green function, describing the propagation of the electron from \( x \) to \( y \).

Concerning dissipation in the measurement interaction induced by Eq. (53), every initial state, i.e., every incoming reaction channel, leads to several, actually to a very large number, of reaction channels. This is the signature of dissipation, as shown in Ref. \[12\], and as is immediately visible from the quantum Boltzmann formula for the entropy \( \mathcal{E} \) \[12\]:

\[
\mathcal{E}^{(k)} = - \sum_j \omega_{jj}^{(k)} \log \omega_{jj}^{(k)}
\]

where the superscript \( (k) \) indicates the initial or final state. There are very many more states, \( j \), in the final state than in the initial: the entropy has increased in the measurement interaction. It does not matter whether the final-state degrees of freedom are observed or not; they participate in the entropy Eq. (54). (See Ref. \[12\] concerning irreversibility.)

Now to the point (ii) above, the question of decoherence. That process arises when the full density matrix is replaced by the “effective” density matrix, i.e., that part of the density matrix associated with the relevant degrees of freedom, and when in that replacement the density matrix loses some or all of its off-diagonal elements by tracing over the unobserved degrees of freedom – which, in fact, is the only possible mechanism for decoherence \[15\]. (This tracing has the
same origin as the integration over the unobserved degrees of freedom when contracting a fully
differential cross section to a partially differential cross section.) This way we expect that for
the final state, being a classical system, the density matrix \( \omega_{ij}^{(f)} \) also will be diagonal, i.e., fully
decohered. However, since the incoming photon may be in a pure state, that expectation may
not be fulfilled; this has to be investigated.

The complete response of the system, the probability \( P_p \), then is given by the general expression
\[
P_p = \text{Tr}' \langle I^\dagger S^{(0)} I R \rangle.
\]
(55)
The structure of (55) is: the state after the initial interaction, \( I^\dagger S^{(0)} I \), undergoes evolution by
the action of the system evolution operator \( R \), associated with the operator \( \eta \). We now must
decide how far down the chain of evolution we want to follow. Thus one may include the observer
in the description; this would require knowledge of the QP mechanism of the brain action, which
we do not possess. We shall, however follow the chain in principle – even though not in detail –
down to the “pointer position”. As we will see, it is fully sufficient to break off the QP chain
much earlier, essentially immediately after the elementary act of the quantum measurement, and
continue its description as a CP object.

In Eq. (55) \( R \) is written in the density matrix form. Hence, in terms or the Tomonaga –
Schwinger evolution operator, i.e., the \( U \)-matrix [13], \( R \) is of the form
\[
R = U(t, t_0) U^\dagger(t, t_0)
\]
(56)
which is the full QP expression. It gives the state of the system at time \( t \), i.e., the fully differential
transition probability into all channels. To obtain the probability for a particular final state, say,
a given pointer position, one sums over all unobserved degrees of freedom, i.e., one computes the
full trace leaving out the observed variables – hence the prime at the trace symbol. Eq. (55) is a
QP expression.

We recognize that, in the notation of Section 5, the counter \( m \), Eq. (21) will respond for all
(photon) states \( \varphi_n(x_m) \) which are non-zero at \( x_m \). The different states \( n \) are here associated with
different recoils of the atom having undergone the Compton scattering. They provide the initial
states for the evolution operator \( R \), Eq. (56), i.e., the state of the system at time \( t_0 \). The operator \( R \)
contains the downstream dynamics, in particular the (hypothetical!) recoil detector. Even though
in our example of the Compton counter such a detector can not be built, in simpler cases it might
be possible to achieve the appropriate measurement [10]. Then in this step no decoherence would
occur; the interference terms between the reaction channels would be retained. Thus, in contrast
to dissipation, which is inevitable, *decoherence is not a basic law of nature*.

We now discuss the decoherence in some more detail, considering a system of adequate com-
plexity to qualify as a (potentially) classical apparatus. As the system evolves down the amplifi-
cation chain it not necessarily loses coherence immediately. That can be seen as follows.

Taking the system at the beginning of the step $c$ of the chain to be in a pure state, i.e., to be
given by a wave function (for brevity we write here in analogy to Eq. (15) only the wave function
part remaining after the evaluation of the Fock operator matrix elements)

$$\psi = \sum C_i \varphi_i$$  \hspace{1cm} (57)

then, at the next step, $c' = c + 1$,

$$\psi' = \sum C'_j \varphi_j$$  \hspace{1cm} (58)

and similarly with $c'' = c + 2$,

$$\psi'' = \sum C''_k \varphi_k$$  \hspace{1cm} (59)

The relation between the consecutive density matrices in terms of evolution operator $U$ is

$$\rho''_{kk'} = C''_k C'^*_{k'}$$

$$= \sum_{jj'} U_{kj} U^*_{{k'}j'} C'_j C'^*_{j'} = \sum_{jj'} U_{kj} U^*_{{k'}j'} \rho'_{jj'}$$

$$= \sum_{jj',ii'} U_{kj} U^*_{{k'}j'} U_{ji} U^*_{{i'}j'} C_i C'^*_{i'}$$

$$= \sum_{jj',ii'} U_{kj} U^*_{{k'}j'} U_{ji} U^*_{{i'}j'} \rho_{ii'}$$  \hspace{1cm} (60)

Here we have used the group character if the evolution operator

$$U(t_n, t_0) = \prod_{k=0}^{n-1} U(t_{k+1}, t_k)$$  \hspace{1cm} (61)

and have introduced the compact notation $U_{j,k} = U(t_j, t_k)$. The product of the U-matrices, upon
continuation to the end of the process, and evaluating the Fock operators, becomes the matrix of
the above operator $R$.  

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In a complex system the reaction channels tend to decouple. That means that such a decoupled state, say $k$, is connected with the initial state $i$ by a single chain of intermediate states, i.e., by a chain without loops $[17]$. If that state is not one of the actually observed states the density matrix at that point loses its relevant off-diagonal matrix elements, as discussed in detail in the next Section after Eq. (67). Hence the up-stream density matrices of the side chain in effect also lose their off-diagonal elements, and one may break off the product of the U-matrices at the point where this chain branches off, say at the state $j = j_0$, and replace the follow-up chain by the Kronecker $\delta_{j,j'}$, which means short-circuiting the tracing over the states at time $t$ and following all the branches back to the state $j_0$. In fact, in this case the intermediate density matrix, $\rho'$, in the row and the column $j_0$ then anyway in effect has vanishing elements, except the diagonal, which has the value $|C'_{j_0}|^2$. This thus will lead to reduction of the off-diagonal elements of the density matrix $\rho''$, without eliminating all of them. (Of course, the form $C''_k C''_{k'}^*$ for $\rho''_{kk'}$ then is not valid.) In this case the de-coherence is incomplete. Complete decoherence would result if all branches lead to unobservable, or, at least, unobserved output channels.

This formulation is fully general. Thus, for example, in the description of the quantum Zeno effect, Ref. [11], the “probe pulse” checking for the state of the system, could be described in the evolution from $\rho$ to $\rho'$, i.e., incorporated in the evolution operator $U'$. Similarly, an event in the history of the evolution, discussed in Refs. [2] and [3], is based on the factoring Eq. (61) and would properly be described in terms of the appropriate operators $U$.

Returning to our discussion, in the further evolution of the system in the next steps similar partial decoherence effects can take place. As decoherence is irreversible it is accumulative, and after a time it will be essentially complete. In the case of our proportional counter this full decoherence will need only very few steps, most probably only one. Further details concerning the mathematics of decoherence are given in Appendix B.

In summary, in any realistic situation there exist many degrees of freedom which participate in the measurement chain, and which are not being observed. In our example of the Compton process such is the recoiling ion, and the ionizations and delta rays arising from interactions of the recoiling Compton electron with the gas. No measurement of their characteristics is contemplated or possible. All they do is trigger the discharge in the counter. Hence here $R_{pq}$ is diagonal, i.e., it vanishes for $p \neq q$. That then eliminates any possibility of interference in the final state $[13]$.
the decoherence is complete.

Once the discharge in the counter has taken place, quantum effects become irrelevant, since in a decohered state coherence can not be restored. Furthermore, the amplification chain and what not is made of classical apparatus. From then on in the quantum description only mixed-state density matrices participate, exhibiting no interferences between the different possible outcomes.

8 The Schrödinger Cat as a Pointer

Even though the results of the previous Section implicitly provide the complete answer to the problem of the pointer, known as the Schrödinger cat, in view of the extensive discussions in the literature we shall give a rather detailed discussion of this case.

The experimental setup consists of a quantum measurement device, leading through a classical amplification chain to a two-position pointer indicating whether the quantum event has or has not taken place. Specifically, the pointer is taken to be a cat, and the two pointer positions are represented by the cat being alive or dead: the quantum event is amplified into killing of the cat. This setup thus is a particular realization of the generic quantum measuring apparatus. The question posed in this context is: Can one “know” whether the cat is alive or dead before looking in the box? More specifically: Is the cat dead or alive before (one of us!) looking? These questions have the same meaning, and permit the same answers, in both the classical and the quantum contexts.

The only question which is possible in the quantum and not in the classical context is: Before we look, is the cat in the linear superposition

$$\Psi_{\text{cat}} = a(t) \psi_{\text{live}} + b(t) \psi_{\text{dead}}$$

Can one observe interference between $\psi_{\text{live}}$ and $\psi_{\text{dead}}$? Does this wave function collapse into either “live” or “dead” upon our taking a look?

Recall the way interferences arise in experimental situations, as for example the interference between electric quadrupole and magnetic dipole reaction channels in photon absorption. The
state after the transition, if pure, is

\[ \Psi = \sum_{n,l} a_{nl} \psi_{nl} \]

with

\[ \psi_{nl} = \chi_n \phi_l(\theta, \varphi) \]

where \( \phi_l \) describes the emitted particle with \( \theta, \varphi \) the direction of emission, and with \( \chi_n \) describing the in general unobserved recoiling particle.

Assume that the incoming photon is polarized with polarization \( p \), and denote the photon density matrix by \( \sigma_{p'p''} \). Further assume that the final state of the recoil reached in the transition (transition operator denoted \( T \)) depend on the polarization. Thus the state index \( n \) splits into two classes, say \( u \) for the “up” polarization (which may stand for “cat alive”) and \( d \) for the “down” polarization (“cat dead”). Let us further denote the components of the initial state of the recoil by \( \Phi_k \), and its density matrix by \( \beta_{k'k''} \).

The (not normalized) final-state density matrix, \( \rho_{n'n',n''n''} \), is given by the transition matrix elements together with the initial-state density matrix:

\[ \rho_{n'n',n''n''} = \sum_{p'k'p''k''} \langle n'l' | T | p'k' \rangle \sigma_{p'p''} \beta_{k'k''} \langle p''k'' | T | n''l'' \rangle . \]  

This equation shows which interferences are possible. Thus the \( u - d \) interference is possible only if the density matrix \( \sigma \) has non-vanishing off-diagonal elements. Similarly, the possibility of any other interference is determined by the non-vanishing of the corresponding off-diagonal elements of the relevant \textit{initial state} density matrix.

Further limitations arise from the measurement operator. The general form of this operator interacting with the emitted particle is

\[ M = \kappa_{u',u'';d',d''} \tau_{p'p''} \delta(\theta - \theta') \delta(\varphi - \varphi') \delta(\theta - \theta'') \delta(\varphi - \varphi'') \]

The fact that the measuring operator does not interact with the recoiling particle leads to overlap of the wave functions \( \chi_n \) in the evaluation of the measurement matrix element, which enforces \( n' = n'' \). Evidently any possibility of \( u - d \) interference hinges on the form of the operator \( \tau \): it must connect the \( u \) and the \( d \) states. If instead it is of the form \( \tau_{p'p''} = \delta_{p',p''} \) the
result of the measurement is

\[ P(\theta, \varphi) = \sum_{n', n''} \langle \psi_{n'l'} | M | \psi_{n''l''} \rangle \rho_{n''l'', n'l'} \delta_{n', n''} \]

\[ = \sum_{n, l', l''} \phi_{l'}(\theta, \varphi)^* \phi_{l''}(\theta, \varphi) \rho_{nl, nl} \delta_{n', n''}. \]  

This way, not performing a measurement results in the density matrix losing the corresponding off-diagonal elements; here \( \rho_{n'l', n''l''} \) becomes \( \rho_{nl, nl} \), i.e., diagonal in \( n \). Further, interferences between the different states, specified by the quantum numbers \( l', l'' \), arise only if the corresponding off-diagonal elements of the final state density matrix, Eq. (65), are non-zero, which again is possible only if \( \sigma \) has non-vanishing off-diagonal elements. The latter condition is fulfilled in electromagnetic transitions since in the incoming photon plane wave all multipoles are coherent.

If the initial state of the system is a mixture of non-interfering states, i.e., if the density matrix \( \beta_{k', k''} \) is diagonal, the final state would consist of non-interfering blocks of states; however, the states within each of these blocks of a given \( l' \) could exhibit full interference.

Returning to our example of the initial interaction in the measurement chain, it is obvious that no macroscopic classical apparatus, no matter how small, will have the simplicity of the wave function \( \psi_{nl} \), Eq. (64). The split into the classes \( u \) and \( d \) therefore will take place early in the chain; probably at its first link. As discussed in the previous Section, decoherence is a very rapid process. Even if the orthogonality of the states of the reaction products, the equivalent to \( \chi \), Eq. (64), is incomplete, or generally, even if these states could exhibit interference, full decoherence will arrive after a very few links of the chain of events leading to the pointer position. Hence the density matrix of the input to the last link before the pointer, the analogue to the product \( \sigma \beta \) of Eq. (65), will be strictly diagonal, which is the form a classical object should have. Consequently no interference as in Eq. (62) is possible; and the pointer behaves like a classical system. Thus, Schrödinger’s cat will never exhibit “the suspended animation”; it will be either dead or alive, independently of our state of mind.

9 Summary

A full explanation of the evolution, as well as of a measurement, of a quantum system requires both the particle and the wave aspects: the first to provide the “yes-no” decision, the second
to provide the “how much” of the measurement. More precisely, if the particle aspect gives
the “yes” decision, the wave aspect provides the probability of the particular outcome, of the
particular reaction branch. Measurement or no measurement, a fully deterministic description of
the evolution is not possible. The attention by an experimentalist plays no role, except that he
can influence the further evolution of the system, for example by switching off the apparatus.

The question of the interference in the results of a quantum measurement, i.e. the question
of the meaning of the superposition of states describing different pointer positions, is answered
by investigating the meaning of a classical object in the quantum formulation. It results in the
statement that classical objects do not exhibit interference effects. They can not be described
by wave functions; their description requires the use of density matrices; more particularly, of
diagonal or block-diagonal density matrices. The reasons for the conversion of the outcome of a
measurement from the quantum to the classical realm are the inevitable dissipation associated
with the quantum measurement process, and the strength of the process of decoherence.

Since the particle aspects are not available within quantum mechanics they must be supplied
“by hand” when attempting a description within that framework. This artifact is called “the col-
lapse of the wave function”. Quantum theory does not separate the particle and the wave aspects.
The quantum physics field can, however, be factorized into these aspects, cf. Eq. (6). The wave
parts of this factorization are the wave functions of quantum mechanics; hence the correctness of
the predictions of quantum mechanics. Still, in agreement with Einstein’s observation, quantum
mechanics itself is incomplete.

Appendix A: Density Matrices

We begin with quantum mechanics. A system which can be described by a wave function is
said to be in a “pure” state. The pure state can be also defined as exhibiting full interference.
The probability of finding the particle at the point \( x \) is

\[
P(x, t) = \int d^3 x' \psi^*(x', t) \delta^3(x - x') \psi(x', t)
\]

The most general pure state is given by a wave function which is the superposition of the eigen-
functions of the Hamiltonian:

\[
\psi(x, t) = \sum_j C_j \psi_j(x) e^{-iE_j t}
\]
with \( \psi_j \) corresponding to energy level \( E_j \). Inserting (A.2) in (A.1) one sees that at every fixed position \( x \) the probability fluctuates with a superposition of energy differences \( \cos(E_j - E_{j'})t \), which is not the behavior of a classical system, as observed in our World: a system localized in space and time, e.g., a book on a table, is not an eigenstate of energy or momentum, but nonetheless it does not exhibit fluctuations. These fluctuations do not arise for a fully impure system, for which the state is described by a diagonal density matrix. The general matrix element of the density-operator is

\[
D_{kj}(x,t) = \int d^3x' \psi_j(x') e^{-iE_j t} \delta^3(x - x') \psi_k^*(x') e^{iE_k t} ; \quad (A.3)
\]

for a pure state the state density matrix is

\[
\varrho_{jk} = C_j C_k^* , \quad (A.4)
\]

while for a fully impure (also called “mixed”) state the state density matrix is

\[
\varrho_{jk} = \delta_{j,k} |C_j|^2 . \quad (A.5)
\]

The probability is computed by tracing the product of the density operator and the state density matrix. Thus Eq. (A.1) then reads

\[
P(x,t) = \sum_{j,k} \varrho_{jk} D_{kj}(x,t) \quad (A.6)
\]

which for a fully impure state indeed shows no interference between the components making up the state.

This way, a classical system cannot be represented by a wave function. It must be represented by a density matrix, which, in particular, must be diagonal. Classical systems are of necessity mixed states.

We now define the QP state vectors appropriate to the density matrix formalism. We shall call them “density state vectors”. To that end we must augment the QM wave functions with appropriate Fock operators. Thus the field operator is

\[
\Psi(x,t) = \sum_j b_j \psi_j(x) e^{-iE_j t} , \quad (A.7)
\]

and the density state vector

\[
\Omega_{jk} = b_j^\dagger |V\rangle \varrho_{jk} \langle V| b_k , \quad (A.8)
\]
which is bi-linear in the Fock operators. The expectation value of an operator, $O$, then is

$$\langle O \rangle = \sum_{j,k} \langle V|b_j \Psi(x,t) O(x,y) \Psi(y,t) b_k^\dagger|V\rangle \omega_{jk}$$

$$= \text{Tr} \; \Psi(x,t) O(x,y) \Psi(y,t) \Omega_{jk} \quad . \quad (A.9)$$

Here the $\text{Tr}$ includes tracing over the indices $j, k$ and re-ordering the Fock operators cyclically, without introducing commutator phases.

**Appendix B: More on Decoherence in Quantum Physics**

In this appendix we will illustrate the decoherence of a measured system after interacting with a macroscopic measuring device. Both of the measured and the measuring are treated as quantum systems, but classically mixed states will be realized after their interaction. Our decoherence result requires nothing more than the framework of quantum physics advocated in this paper; i.e. no extra projection or collapse of the wave function postulate. This is possible because quantum physics provides us the mechanism of the chain amplifying the first interaction between the measured and the macroscopic device through the many microscopic constituents of the device.

Explicitly, we will make use of two ingredients: (i) quantum physics detector functions, Eqs. (21,24,34,35,53), and (ii) the many degrees of freedom of the measuring device that are unobserved or unobservable.

To begin with we will derive a result for later use. In the spirit of (6) we expand the field operator either in terms of the set of c-number functions $\{\psi_\alpha(x,t)\}$ or another set $\{\phi_\beta(x,t)\}$, which satisfy the same boundary condition but arbitrary otherwise (i.e. corresponding to a different set of quantum numbers $\beta$).

$$\Psi(x,t) = \sum_\alpha a_\alpha \psi_\alpha(x,t) \quad , \quad (B.1)$$

$$= \sum_\beta b_\beta \phi_\beta(x,t) \quad . \quad (B.2)$$

We can take these two sets to be orthonormal without any loss of generality. Thus,

$$a_\alpha = \sum_\beta C_{\alpha\beta} b_\beta \quad , \quad (B.3)$$

$$\left[ a_\alpha, b_\beta^\dagger \right]_+ = C_{\alpha\beta} \quad , \quad (B.4)$$
where

\[ C_{\alpha\beta} \equiv \int \psi_\alpha^* \phi_\beta. \] (B.5)

We now choose an arbitrary state \(|\alpha\rangle = a_\alpha^\dagger |V\rangle\), say, and perform the interrogation

\[
\langle V|\Psi|\alpha\rangle = \psi_\alpha
= \sum_\beta C_{\alpha\beta}^* \phi_\beta, \tag{B.6}
\]

if the expansions (B.1) and (B.2) respectively are employed for \(\Psi\). Then from the orthonormality of \(\psi\)'s and \(\phi\)'s it follows that \(\sum_\beta |C_{\alpha\beta}|^2 = 1\), which implies

\[ |C_{\alpha\beta}| \leq 1. \] (B.7)

(The last result can also be obtained simply by considering \([a_\alpha, a_\alpha^\dagger] + = 1 = [b_\beta, b_\beta^\dagger] +\) with the substitution of \(\text{(B.3)}\).)

Coming back to the decoherence problem, let the measured system be in the pure state before the measurement, thus its density operator is given by (for simplicity, we restrict the number of states to 2)

\[
\rho = \sum_{i,j=1,2} \rho_{ij} b_i^\dagger |V\rangle \langle V| b_j. \tag{B.8}
\]

The density operator of the whole system is the direct product of those of the measured and the device \(g\).

We now go directly to the end of the measurement process, leapfrogging the measurement chain. The detector function, Eq. (21), is then replaced by the effective detector function

\[
D = \eta_1 B_1^\dagger b_1 + \eta_2 B_2^\dagger b_2, \tag{B.9}
\]

here \(B_j^\dagger\) creates the final observable states of the apparatus, i.e. the “pointer position”, and

\[
\eta_{1,2} = \prod_{m=1}^{N} d_m^{(1,2)\dagger}, \tag{B.10}
\]

where \(d_m^\dagger\) are the creation operators for the (unobserved/unobservable) degrees of freedom in the chain, and \(m = 1, \ldots, N\). Thus the density operator after the measurement becomes

\[
\rho \otimes g \rightarrow D (\rho \otimes g) D^\dagger. \tag{B.11}
\]
The reduced density operator is then obtained by tracing over the unobserved (or unobservable) dynamical degrees of freedom of the apparatus

\[
\rho_{\text{(after)}} = \text{Tr}' \left( D (\rho \otimes \varrho) D^\dagger \right) = E_1 P_1 B_1^\dagger |V\rangle \langle V| B_1 + E_2 P_2 B_2^\dagger |V\rangle \langle V| B_2 + \text{interference term.} \quad (B.12)
\]

In this expression, \( P_i = \rho_{ii}, \ i = 1, 2, \) is the probability that the system is found to be in state \( i \); and \( E_i = \text{Tr}' \left( \eta_i \varrho \eta_i^\dagger \right) \) is a measure of the efficiency of the detector in detecting the state \( i \). The interference term of (B.12) contains quantities that are explicitly proportional to the trace \( \text{Tr}' \left( \eta_i \varrho \eta_j^\dagger \right) \), where \( i \neq j \). And with the representation (B.10) for \( \eta_i \), a typical term of this trace has the form

\[
\text{Tr}' \left( \eta_i \varrho \eta_j^\dagger \right) \sim \prod_{m=1}^{N} \prod_{n=1}^{N} \langle |d_j^{(m)} a_i^{(n)}| \rangle + \text{similar terms} , \quad (B.13)
\]

where \( | = d^\dagger \cdots d^\dagger |V\rangle \) represent the initial states of the many constituents of the detector.

(Anti-)commuting the \( d \)-annihilation operators in (B.13) to the right and using the results in (B.4.B.7), which now translate to a strict inequality because \( i \neq j \),

\[
\left[ d_j^{(m)}, a_i^{(n)} \right] < 1 , \quad (B.14)
\]

we finally obtain the result that the off-diagonal elements of the reduced density operator vanish

\[
\text{Tr}' \left( \eta_i \varrho \eta_j^\dagger \right) \xrightarrow{N \to \infty} 0 , \ i \neq j , \quad (B.15)
\]

as the terms on the right hand side of (B.13) are products of complex numbers of magnitudes less than one. That is, with a macroscopic measuring device, we have complete decoherence

\[
\rho_{\text{(after)}} \xrightarrow{N \to \infty} E_1 P_1 B_1^\dagger |V\rangle \langle V| B_1 + E_2 P_2 B_2^\dagger |V\rangle \langle V| B_2 . \quad (B.16)
\]

The estimation for the time required for complete decoherence is more complicated but may be calculated with the full use of quantum field theory or its non-relativistic equivalent.

**Appendix C: Preparation of State**

A conceptual and semantic confusion exists between the similar but distinct processes of “measurement” and “preparation of a system in a specific state”. Namely, “the preparation of
the initial state for an experiment” consists in allowing the desired state to remain and rejecting all other states. This supposedly is achieved by an appropriate measurement. The selection of the desired state then in quantum mechanics is described as the “collapse of the wave function”: the “measurement puts the state into an eigenstate of the measuring apparatus”. The prototype of such a setup is a Stern-Gerlach apparatus. Indeed, the atoms emerge from the source in a beam in which they are a mixture of all possible states, of which, say, there are N. At the output of the Stern-Gerlach apparatus then emerge N separated beams, one each for the different states. Now one simply must supply a collimator such that only the beam containing the desired state is transmitted; all other beams are absorbed. No interaction with the transmitted particles has taken place; this process is non-dissipative. The statement that “the measurement puts the system into an eigenstate of the apparatus” in fact is not accurate since actually no measurement has been performed on the particle. We only know that if a particle emerges in that output beam, then it has that particular polarization – except for background effects arising from collimator scattering etc.

Another possibility for preparing the system in a specified state is the technique called “tagging”, which is of the kind of the EPR setup \[14\]: measurement of the characteristics of one partner in a correlated pair of particles provides information on the state of the other partner. A well-known example is that of the tagged bremsstrahlung photons: if the detector, which is an electron detector, registers, i.e., measures the characteristics of, a recoiling electron then one “knows” that the other particle, here the bremsstrahlung photon, is in a well-defined state having definite energy, direction of propagation, and polarization. Again, no measurement has been performed on the tagged photon, which is the particle of interest.

Overall, since in the preparation of the state the system of interest has actually been left alone, this process should not be denoted as “a measurement.” A more precise term would be that the preparation is “a filtering action.” In quantum physics the filtering is accomplished either directly by absorbing the particles being in the undesired states or by performing a measurement on another part of the system, in an EPR-type arrangement. Any measurement directly on the system of interest would be of the kind described above in the context of the two-slit experiment when checking for the passage of the particle through the slit, and involves an interaction of the system with an apparatus. The re-emitted system then inescapably is in a new state, and has
new not necessarily known properties.

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