Abstract

This proposes a new theory of Quantum measurement: a state reduction theory in which reduction is to the elements of the number operator basis of a system, triggered by the occurrence of annihilation or creation (or lowering or raising) operators in the time evolution of a system. It is from these operator types that the acronym ‘LARC’ is derived. Reduction does not occur immediately after the trigger event; it occurs at some later time with probability $P_t$ per unit time, where $P_t$ is very small. Localisation of macroscopic objects occurs in the natural way: photons from an illumination field are reflected off a body and later absorbed by another body. Each possible absorption of a photon by a molecule in the second body generates annihilation and raising operators, which in turn trigger a probability per unit time $P_t$ of a state reduction into the number operator basis for the photon field and the number operator basis of the electron orbitals of the molecule. Since all photons in the illumination field have come from the location of the first body, wherever that is, a single reduction leads to a reduction of the position state of the first body relative to the second, with a total probability of $m \tau_L$, where $m$ is the number of photon absorption events. Unusually for a reduction theory, the larc theory is naturally relativistic.

1 The Measurement Problem

Quantum Mechanical time evolution is Hamiltonian time evolution. But quantum measurement, on the face of things, appears to need a second law, unique to it. There are two reasons for this apparent need. The first is that without a second law, measurements do not produce definite results. The second is that Hamiltonian evolution is deterministic, but measurement outcomes are probabilistic; the probabilistic element of quantum measurement is provided by the second law. The most primitive form of second law is the ‘quantum jump’: a sudden, probabilistic, change of state that occurs during a measurement. Unfortunately, no-one has been able to convincingly specify the physical property of measurement that invokes this special process. The measurement problem consists in the apparent inability of pure Hamiltonian time evolution to describe
measurements correctly, combined with our inability to give any special process
a sound physical basis.

When using quantum mechanics for everyday calculation, physicists don’t
notice this problem, because they have learned to recognise physical arrange-
ments that constitute measurement instruments, and at what point measure-
ment can be said to have occurred, by applying the laws of quantum theory
with what Bell referred to as ‘good taste’[1].

A solution to the measurement problem needs to either: 1) explain mea-
surement without recourse to special processes, or 2) specify the special processes
as physical laws, most importantly providing a law to explain when the pro-
cess occurs, since ‘measurement’ is not physically well-defined. This is a basis
for a broad classification of solutions to the measurement problem: type 1 solu-
tions explain measurement without invoking any fundamental physical processes
outside Hamiltonian time evolution; type 2 solutions specify new fundamental
physical laws that allow measurement to be explained.

Everett’s relative state interpretation is an example of a type 1 solution [4].
Type 2 solutions can themselves be said to fall into two broad classes: those
that retain the bulk of the quantum theoretical formalism, only adding the
minimum needed to address the measurement problem itself; and those that
derpart more or less widely from the quantum formalism, producing effectively
a new theory. The Ghirardi-Rimini-Weber [2] theory falls into the former class,
while hidden-variable theories such as the Bohm [3] theory fall into the latter.

This present will propose a solution to the measurement problem of type 2,
and of the former class of adding only enough to quantum theory to (with luck)
explain the behaviour of measurement processes. Maudlin[5] has given a very
succinct summary of the system of arguments that leads some to believe that
the solution to the measurement problem must be of type 2.

We are going to need at least one new law which must do two things. It must
specify when the state reduction process occurs, including what states of affairs
trigger the process, and it must specify how the Hilbert space basis in which the
reduction occurs is chosen. Somewhere along the way, the theory must explain
why this new process appears to occur only during measurements and not at
other times. Since there is no physical property possessed by measurement that
distinguishes it from other processes, this is inherently tricky.

2 Outline of the Theory

Up to now, proposed type 2 solutions to the measurement problem have usually
kept within nonrelativistic quantum mechanics. They have often, even when
adequate at the nonrelativistic level, met with difficulty in developing into a
relativistic form [2] [3]. The event that will trigger state reduction in this the-
ory, while it appears in nonrelativistic quantum mechanics, is more naturally
associated with relativistic quantum mechanics and quantum field theories.

The ability of the number of particles, or of field quanta, to vary in quantum
field theory is one of its fundamental properties. Changes in the number of
field quanta have operators associated with them, known as the annihilation ($\hat{a}$), and creation ($\hat{a}^\dagger$) operators. Processes in which a particle of a given type is annihilated and replaced by the creation of particles of other types are routine in quantum field theory. Conservation laws demand that particle annihilation must be followed by particle creation. (‘Followed’ here means in the temporal sense. Formally, the creation operators fall to the left of the annihilation operators.)

The concepts behind annihilation and creation operators also apply to the less esoteric case of a bound system in a certain energy state. The classic case is the harmonic oscillator, for which there exist lowering ($\hat{a}$) and raising ($\hat{a}^\dagger$) operators that move the oscillator to the next lower or next higher energy state. The harmonic oscillator is the classic example because it is one of the very few cases for which the mathematical form of the operators can be solved exactly. However, in principle, corresponding operators exist for the electronic states of hydrogen atoms, or more complex molecules. Lowering and raising operators mix with annihilation and creation operators, following the rule that a lowering or annihilation operator (or group of them) must be followed temporally by a raising or creation operator (or group of them) so that conservation laws are obeyed. I’ll refer to a well-formed cluster of lowering/annihilation - raising/creation operators as a larc.

One example of a larc would be pair creation: a photon annihilation operator followed by an electron creation operator and a positron creation operator. Another example would be absorption of a photon by a molecule, described by a photon annihilation operator, followed by a raising operator for the electronic states of the molecule.

The larc theory postulates that the triggering event for a state reduction is the occurrence of a larc in the description of a process. This does not mean that a state reduction happens at every larc event: a larc triggers a probability of a state reduction, specifically a probability per unit time. If this probability per unit time is small enough, there will be virtually zero chance of a state reduction occurring during a process involving small numbers of larc events, which will be the case, for example, in high-energy particle experiments.

A state reduction consists of a lopping off of branches. In most cases a larc event occurs as a branching: the larc event for absorption of a photon by a molecule will be a branch superposed with the alternative event of the photon passing through the molecule without being absorbed. Each of these branches has an amplitude. The presence of a larc in this process creates a probability per unit time that one of the branches will be lopped off, the probability for each branch to be the survivor is the square modulus of its amplitude.

In the larc theory, the basis in which the reduction occurs is the number basis for the particles or field quanta defined by the lowering/annihilation and raising/creation operators involved in each larc event.

In summary, the elements of the larc theory of quantum measurement are: 1. State reduction occurs following a larc. After a larc occurs there is a probability per unit time of a state reduction, which consists of a lopping off of the branches not chosen. ‘Lopping off’ means a zeroing of the amplitudes of those branches. 2. The basis in which the reduction occurs is the number basis defined by the
operators in the larc.

3 Localisation of Macroscopic Bodies

So far, the larc theory doesn’t appear to include anything that will solve the classic problem of quantum measurement: the localisation of macroscopic bodies. Macroscopic body localisation is important for two reasons: first, it explains why the moon, or a cricket ball, is in a definite location (and is still there even when nobody looks). Second, it ensures that the pointers of measuring devices must show definite outcomes. Since pretty much any measuring device can be designed to show its output as the position of a pointer, or some equivalent, such as the position of macroscopic patches of colour on a display device, localisation of macroscopic objects is effectively a general solution to the measurement problem, at least to a first approximation.

That macroscopic objects are going to be localised by the larc theory may not be immediately obvious. I’ve postulated a process that has nothing to do with position localisation, and occurs extremely rarely to boot. In fact, the fundamental machinery required is already in place. The completion of the argument to localisation requires a further property of quantum mechanics I will refer to as amplification by correlation (abc). As far as I know, this mechanism is due to Ghirardi, Rimini, and Weber[2].

In outline, abc acts whenever a composite state contains a large number of quanta that are in states correlated with one another. The correlation means that if any one of the quanta gets reduced to a single definite state, all the other quanta in the composite state will be reduced to their corresponding definite states. Since every quantum’s state will be reduced if any quantum’s state becomes reduced, the probability of reduction of the whole composite state becomes roughly the sum of the probabilities of each individual quantum being reduced. In this way, the probability of reduction is amplified, potentially by many orders of magnitude, by the correlated state.

Body B begins in a superposition of positions $x_1$ and $x_2$. Larc-abc is capable of localising a body whose position is unknown in three dimensions, but it is sufficient to show that it can localise B. The space where B exists can be illuminated. Initially, the light is off. At $t = t_i$ the light will turn on.

The final element of the thought experiment is a pinhole camera, C. This is positioned in the plane, facing towards the line we know B is confined to. C consists of a box with a small hole in the side facing B (not small compared to the wavelength of the illumination, of course). At the back of the camera is an old-fashioned photographic plate.
Localisation of body B by pinhole camera C

Let’s start with the simplest possible case; B in a superposition of two possible locations, \( x_1 \) and \( x_2 \).

\[
\psi_B = a_1 \delta(x - x_1) + a_2 \delta(x - x_2) \quad (1)
\]

With \( |a_1|^2 + |a_2|^2 = 1 \). At time \( t_i \), the light turns on

\[
a_1 \delta(x - x_1) I(x, t_i) + a_2 \delta(x - x_2) I(x, t_i) \quad (2)
\]

Where \( I(x, t_i) \) represents the electromagnetic field radiation expanding from point \( x \), at time \( t_i \).

The photographic plate is composed of individual molecules capable of absorbing visible photons and, as a consequence, undergoing a change which will later be visible to the unaided human eye, provided enough molecules absorb photons. At time \( t_f \) the plate has been illuminated by \( n \) photons, where \( n \) is presumed to be large enough to ensure that the result will be visible.

Again to simplify the mathematics, and without loss of generality, we assume that the camera is positioned equidistant from \( x_1 \) and \( x_2 \), so that the number of photons illuminating the plate is the same for either possible position of B. If the body B is at \( x_1 \), then a patch of the photographic plate will be illuminated, call this patch \( \xi_1 \). If the body is at \( x_2 \), a different patch will be illuminated, call it \( \xi_2 \). The illumination landing on \( \xi_i \) at time \( t_f \) will be represented by \( I(\xi_i, t_f, n) \): \( n \) is the number of photons that will fall on the patch \( \xi_i \).

As \( I \) passes through each molecule it generates a factor representing possible absorption of a photon by a molecule at the photographic plate. A very simple representation of this factor is

\[
a_1 b_1 \delta(x - x_1) \hat{a}_\xi I(\xi_1, t_f, n) \hat{a}_\xi \psi(\xi_1, i) \psi(\xi_2, i) \\
+ a_1 b_2 \delta(x - x_1) I(\xi_1, t_f, n) \hat{a}_\xi \psi(\xi_1, i) \psi(\xi_2, i) \\
+ a_2 b_1 \delta(x - x_2) \hat{a}_\xi I(\xi_2, t_f, n) \hat{a}_\xi \psi(\xi_1, i) \psi(\xi_2, i) \\
+ a_2 b_2 \delta(x - x_2) I(\xi_2, t_f, n) \hat{a}_\xi \psi(\xi_1, i) \psi(\xi_2, i) \quad (3)
\]
Where $|b_1|^2 + |b_2|^2 = 1$, $b_1$ being the amplitude for a photon to be absorbed by a light-sensitive molecule, while $b_2$ is the amplitude for the photon to pass through without being absorbed (in a given time). $\psi(\xi_1, i)$ is the initial electron orbital state of a light-sensitive molecule in area $\xi_1$, the area on the photographic plate that will be illuminated if B is at $x_1$. $\hat{a}_\gamma$ is a photon annihilation operator such that $\hat{a}_\gamma I(\xi_1, t_f, n) = n^{1/2} I(\xi_1, t_f, n - 1)$. $\hat{a}_e^\dagger$ is a raising operator acting on the electron orbital of the molecule, raising it to the energy state that leads to a visible change, so that $\hat{a}_e^\dagger \psi(\xi_1, i) = \psi(\xi_1, f)$.

When states of the form (3) exist the proposed state reduction mechanism becomes active. The mechanism has three elements.

1. **Trigger.** Once the state contains a larc there is a probability per unit time of a reduction. Call the time constant for this $P_t$. In this reduction branches other than the one selected will be lopped off, their amplitudes set to zero.

2. **Basis choice.** The basis in which this reduction occurs is the number basis consistent with the annihilation and raising operators that appear in the larc used to describe the absorption of the photon by the molecule. In this case the relevant number operators are $\hat{n}_\gamma = \hat{a}_\gamma^\dagger \hat{a}_\gamma$, corresponding to the number of photons in the incoming electromagnetic field, $\hat{n}_{ei}$ the number of electrons in the initial electron orbital state, and $\hat{n}_{ef}$, the number of electrons in the final electron orbital state.

I will express these combinations of number operator eigenvalues in the form of a 3-tuple $(n, i, f)$, where $n$ is the number of photons, $i$ the number of electrons in the initial state, and $f$ the number of electrons in the final state. $i$ and $f$ can each only be zero or one, because the electron is a fermion, and $i + f = 1$, because the number of electrons is conserved. The initial state is $(n, 1, 0)_1$, where the subscript indicates which of the two regions, $\xi_1$ or $\xi_2$, is in question. There are three possible final combinations: if the photon is absorbed the final state is $(n - 1, 0, 1)_1$, which corresponds to a zeroing of amplitude $b_2$; if the photon is not absorbed the state is $(n, 1, 0)_1$, which corresponds to a zeroing of amplitude $b_1$. In each of these cases it is inherent that $a_2$ is also zeroed.

There remains a third possible final combination: $(n - 1, 1, 0)_1$. This corresponds not to the photon being absorbed by the molecule, but to a removal of one photon from the electromagnetic field $I(\xi_1, t_f, n)$ without its being absorbed by the molecule at $\xi_1$. On the face of things, this outcome violates energy conservation, since the photon disappears but its energy does not get transferred to the electron orbital of the molecule. However, recall that (3) is a superposition of two different positions for B, and consequently, two different illumination states. If a photon disappears from one of the two superposed illumination states, then it still exists in the other. This outcome corresponds with a zeroing of $a_1$.

This third way of reducing the state is necessary to the success of the whole enterprise. Imagine an alternative version of the pinhole camera, in which there is film in only the half of the camera containing $\xi_1$, the half containing $\xi_2$ being open, so that any photons that would have been absorbed at $\xi_2$ now pass through empty space without being absorbed at all. If in this situation only the first two combinations were allowed, then with probability one, B would be at $x_1$. It
would be possible to determine where B is simply by choosing whether to detect the illumination beam at $\xi_1$ or $\xi_2$. Therefore, combination 3 is necessary if the larc-abc mechanism is to produce the correct Born rule probabilities.

(3) State Reduction. The final result of the state reduction process is one of three possible states. The first in which the photon is definitely absorbed, is that corresponding to $(n-1,0,1)$

$$\delta(x - x_1)\hat{a}_\gamma I(\xi_1, t_f, n)\hat{a}_\alpha^\dagger \psi(\xi_1, i)\psi(\xi_2, i)$$

which occurs with probability $|a_1b_1|^2$. Of course, this state is equal to

$$\delta(x - x_1)n^{1/2}I(\xi_1, t_f, n - 1)\psi(\xi_1, f)\psi(\xi_2, i)$$

The second, in which the photon is definitely not absorbed, corresponding to $(n,1,0)$

$$\delta(x - x_1)I(\xi_1, t_f, n)\psi(\xi_1, i)\psi(\xi_2, i)$$

which occurs with probability $|a_1b_2|^2$.

The third case is a little harder to describe neatly. One photon is removed from the illumination field in the $\xi_1$ region, without being absorbed. This situation corresponds to $(n-1,1,0)$

$$b_1\delta(x - x_2)\hat{a}_\gamma I(\xi_2, t_f, n)\psi(\xi_1, i)\hat{a}_\alpha^\dagger \psi(\xi_2, i) + b_2\delta(x - x_2)I(\xi_2, t_f, n)\psi(\xi_1, i)\psi(\xi_2, i)$$

which occurs with probability $|a_2|^2$.

4 Amplification By Correlation

As exposure goes on, we get large numbers of factors like $\hat{a}_\gamma$ representing possible absorption of photons by the molecules in regions $\xi_1$ and $\xi_2$. Assuming that the photographic plate’s coating is opaque, all the photons will eventually be absorbed. After the entire illumination front has passed the camera, the state at the photographic plate will look something like

$$a_1\delta(x - x_1)\prod_{j=1}^{m_1} [b_2\psi_j(\xi_1, i) + b_1\hat{a}_\gamma\psi_j(\xi_1, i)\hat{a}_\alpha] I(\xi_1, t_f, n) \times \prod_{j=1}^{m_1}\psi_j(\xi_1, i) + a_2\delta(x - x_2)\prod_{j=1}^{m_2} [b_2\psi_j(\xi_2, i) + b_1\hat{a}_\gamma\psi_j(\xi_2, i)\hat{a}_\alpha] I(\xi_2, t_f, n) \times \prod_{j=1}^{m_2}\psi_j(\xi_2, i)$$

Where $m_1$ and $m_2$ represent the number of absorbing molecules in regions 1 and 2 respectively.

As before there are three elements to the state reduction process, and three possible end states.
(1) Trigger. The state described by equation (8) contains a large number of larc events. Each one creates a probability per unit time of a state reduction. The probability per unit time is \( P_t \) for each larc event. The total number of larc events in (8) is \( m_1 + m_2 \), which means that, roughly, the probability per unit time for a reduction is \( P_t(m_1 + m_2) \).

(2) Basis Choice. The basis choice for any individual larc event is the same as that for equation (3). Let us say, without loss of generality, that the larc event associated with molecule \( q \) in patch \( \xi_1 \) is the first to trigger a state reduction. The basis will then be that associated with the photon number operator, the number operator for the initial state of molecule 1; \( q \); and the number operator for the final state of molecule 1; \( q \). The possible outcomes, characterised as above, are \( (n−1,0,1)_{1,q}, (n,1,0)_{1,q}, \) and \( (n−1,1,0)_{1,q} \).

(3) State Reduction. Again, one of three possible states results from the process. The first, \( (n−1,0,1)_{1,q} \), corresponds to the photon being definitely absorbed at 1; \( q \).

\[
\delta(x-x_1)\hat{a}_q\psi_q(\xi_1,i)\hat{a}_q \prod_{\substack{j=1 \\text{j.q}}\, m_1} [b_2\psi_j(\xi_1,i) + b_1\hat{a}_q\psi_j(\xi_1,i)\hat{a}_\gamma] I(\xi_1,t_f,n) \\
\times \prod_{j=1}^{m_2} \psi_j(\xi_2,i) \tag{9}
\]

with probability \( |a_1b_1|^2 \).

The core of the larc-abc mechanism is now clear. The crucial property of the state described by (8) is this: the photons are all correlated. If any one photon is absorbed in region \( \xi_1 \), then every photon must be absorbed in region \( \xi_1 \) and none in region \( \xi_2 \). This is clear in (9), which is the result of a single photon definitely absorbed in region \( \xi_1 \). All terms in which a photon could be absorbed in region \( \xi_2 \) have vanished. Since the region in which a photon is absorbed is correlated with the position of B, any photon absorbed in \( \xi_1 \) means the position state of B is \( \delta(x-x_1) \), which corresponds to a zeroing of \( a_2 \). Again, this is clear in (9): all the terms with \( \delta(x-x_2) \) are gone.

The second possible end state after a state reduction is \( (n,1,0)_{1,q} \), corresponding to a photon being definitely not absorbed at 1; \( q \).

\[
\delta(x-x_1)\psi_q(\xi_1,i) \prod_{\substack{j=1 \\text{j.q}}\, m_1} [b_2\psi_j(\xi_1,i) + b_1\hat{a}_q\psi_j(\xi_1,i)\hat{a}_\gamma] I(\xi_1,t_f,n) \\
\times \prod_{j=1}^{m_2} \psi_j(\xi_2,i) \tag{10}
\]

with probability \( |a_1b_2|^2 \).
The third end state is \((n - 1, 1, 0)_{1:q}\), corresponding to the photon being removed from the illumination field at \(\xi_1\) without being absorbed at \(1:q\).

\[
\delta(x - x_2) \prod_{j=1}^{m_2} \left[ b_2 \psi_j(\xi_2, i) + b_1 \hat{a}_1^\dagger \psi_j(\xi_2, i) \hat{a}_\gamma \right] I(\xi_2, t_f, n) \\
\times \prod_{j=1}^{m_1} \psi_j(\xi_1, i)
\]

(11)

with probability \(|a_2|^2\).

Effectively, B has been localised to \(\delta(x - x_2)\) before any photon has been definitely absorbed, or definitely not absorbed, at \(\xi_2\). The form of equation (11) is determined by the correlation of photons in the illumination field \(a_1 I(\xi_1, t_f, n) + a_2 I(\xi_2, t_f, n)\). The photons are correlated such that they are either all at \(\xi_1\), or all at \(\xi_2\).

So, equations (9), (10), and (11), are the final end results of the measurement process. In each of them, body B has become definitely localised. As I have said, if even a single photon undergoes a state reduction, it carries all the photons with it, as well as the body B with whose position they are all in turn correlated. This means that even if the probability per unit time of a reduction is very small for each larv event, the enormous number of larv events involved implies that the probability of B becoming localised may be virtually one. This is the effect I have dubbed amplification by correlation (abc): a very small probability of a state reduction \((P_t)\) for each photon is amplified by correlation of large enough numbers of photons \((m_1 + m_2)\), into a very large probability of reduction of the position of body B \((P_t(m_1 + m_2))\). Hence (8) implies that B becomes localised with probability virtually one.

It is worth being clear about what is and is not correlated in (8). If any photon is definitely absorbed at \(\xi_1\), then no photon can be absorbed at \(\xi_2\), all must ultimately be absorbed at \(\xi_1\). However, while the dropping of one molecule into a definite state forces the position of B into a definite state, it does not force any of the other photons to decide whether they have been definitely absorbed. As a result, it is possible that when a person looks at the plate, there will not be any definite fact of the matter of exactly which molecules have been changed and which have not. Some molecules may still be in a superposition of having absorbed or not absorbed a photon. But this is not as serious as it might sound. The differing states that are superposed are not distinguishable to the human observer because they all lie within one region. Furthermore, it is likely that photon absorption and amplification processes that occur within the visual system and brain will raise the probability of these superposed states becoming definite.

It is clear that the above argument can be generalised to a body whose position is unknown in one, two or three dimensions, using two or three pinhole cameras.
5 Human-independent localisation

The argument so far has relied on the pinhole camera, which is clearly a human artifact. Can we expect that the larc-abc mechanism described above will lead to localisation by other measuring devices, or even without any human involvement? The answer is yes, for more than one reason.

First, the involvement of humans, as observers, is irrelevant to the localisation process. The only involvement of humans as observers (as opposed to artificers) in the above argument is the specification that the state reached by a molecule when it absorbs a photon, $\hat{a}_e^\dagger \psi(\xi_1, i) = \psi(\xi_1, f)$, is visibly different, to the human eye, when compared with the initial state, $\psi(\xi_1, i)$. But this assumption plays no role in the argument leading to localisation. Localisation would occur in exactly the same way if the two molecular states were not visibly different to the human eye, and hence does not depend on the camera’s use as a measuring instrument, but only on its basic physical characteristics.

Second, localisation will occur even if the ‘pinhole camera’ is just a natural cavity in the surface of a second body. After all, real macroscopic bodies are not smooth surfaced; real bodies have surfaces which have many ridges and valleys, hills and cavities. At the bottoms of these valleys and cavities will be places where illumination from another body can reach only if it is in the right relative position, so the above analysis based on the ‘pinhole camera’ will still apply.

Third, and more generally, when a photon is absorbed by a molecule or crystal lattice it transfers information about the direction of its source in the form of a momentum kick. Imagine a single molecule, M, part of a macroscopic body positioned somewhere off the line $x$, which is struck by the illumination field coming from body B. The illumination is a superposition of illumination fields coming from two different directions. Direction one corresponds to a photon coming from position $x_1$, which will give the molecule a kick $p_1$. Direction two corresponds to a photon coming from position $x_2$, which will give the molecule a kick $p_2$. Once the illumination field has passed through the molecule the state will be

$$a_1 b_1 \delta(x - x_1) \hat{a}_e I(p_1, t_f, n) \hat{a}_e^\dagger \psi(i) \hat{a}_p \phi(p_i) + a_1 b_2 \delta(x - x_1) I(p_1, t_f, n) \psi(i) \phi(p_i) + a_2 b_1 \delta(x - x_2) \hat{a}_e I(p_2, t_f, n) \hat{a}_e^\dagger \psi(i) \hat{a}_p \phi(p_i) + a_2 b_2 \delta(x - x_2) I(p_2, t_f, n) \psi(i) \phi(p_i) \tag{12}$$

where $p_1$ ($p_2$) represents the momentum of a photon travelling from $x_1$ ($x_2$) towards M. It is no longer the position of the molecule M that is correlated with the position of B; the position of B is now correlated with the direction of the momentum striking a single molecule. $\phi(p_i)$ is the centre of mass momentum state for the molecule M, with an initial momentum of $p_i$. $\hat{a}_p$ is a raising operator affecting the centre of mass momentum state of the molecule and adding momentum $p_1$ to it; hence $\hat{a}_p \phi(p_i) = \phi(p_i + p_1)$. 


Localisation of body B by momentum transferred to molecule M

Decomposing the momentum kick into components $p_1 = p_{1x} + p_{1y}$, we have

$$
\begin{align*}
& a_1 b_1 \delta(x - x_1) \hat{a}_\gamma I(p_{1x}, p_{1y}, t_f, n) \hat{a}_\delta^\dagger \psi(i) \hat{a}_{p_{1x}}^\dagger \hat{a}_{p_{1y}}^\dagger \phi(p_{ix} + p_{iy}) \\
& + a_1 b_2 \delta(x - x_1) I(p_{1x}, p_{1y}, t_f, n) \psi(i) \phi(p_i) \\
& + a_2 b_1 \delta(x - x_2) \hat{a}_\gamma I(p_{2x}, p_{2y}, t_f, n) \hat{a}_\delta^\dagger \psi(i) \hat{a}_{p_{2x}}^\dagger \hat{a}_{p_{2y}}^\dagger \phi(p_{ix} + p_{iy}) \\
& + a_2 b_2 \delta(x - x_2) I(p_{2x}, p_{2y}, t_f, n) \psi(i) \phi(p_i)
\end{align*}
$$

(13)

The larc in equation (13) create a probability per unit time $P_t$ of a state reduction which will leave only one term of equation (13) behind. If we consider all the molecules exposed to the illumination field $I(n)$ we will end up with a state like (8), but one in which the direction of the momentum transferred to each molecule is correlated, rather than the position in which the transfer takes place, which means that the abc mechanism will apply to the momenta transferred by many photons to a macro body: the directions of these momenta will be correlated, and just one reduction to a definite direction will carry all the others with it through this correlation. Localisation of body B occurs without any human involvement, and with no assumptions about the structure of the other body(s).

6 Some properties of the theory

A significant property of the theory is that while it localises macroscopic objects, it can only produce localisation relative to other bodies. The larc theory is in principle incapable of producing absolute position localisation. I regard this as a strength of the theory. Most physical theories that are not explicitly relativistic are agnostic about the relativity of position: position variables can be interpreted as either relative or absolute without any effect on observable physics. The larc-abc theory is a rare example of a theory that permits only of relative positions.
Type 2 theories of quantum measurement are usually formulated initially in nonrelativistic terms, and they often have problems extending to the relativistic case. Both the Bohm and GRW theories have difficulties of this kind. The larc theory, by contrast, analyses the situation the same way it would be done in quantum electrodynamics. Larc clusters are already present in quantum field theory, so they do not disrupt the relativistic invariance of the analysis. The branches in the larc number operator basis for bound states effectively define the energy basis, which again is consistent with a relativistic theory. If each branch is relativistic, then lopping off any one branch, or all but one, will leave behind a system that remains relativistic.

It is a vital property of the larc-abc process that it leads to position localisation in appropriate circumstances. But this localisation can only occur relative to some other body. Therefore, the theory cannot produce any problem of there being a preferred reference frame. Likewise, the temporal constant $P_t$ for the probability of a state reduction also belongs to a well-defined system: that to which the relevant larc cluster belongs, such as the molecule where the photon is absorbed, so again there is no need for a preferred reference frame. Taken altogether, larc theory should be as relativistic as quantum field theory itself.

Similar arguments show that the theory conserves energy. The emission of photons from B conserves energy, the absorption of photons, either in the pinhole camera, or in molecule M, also conserves energy. Each branch in the superposition conserves energy, so if the amplitudes of all but one branch are zeroed, whichever branch remains conserves energy.

Larc theory has a property which could be considered a further advantage over the GRW theory. GRW theory has two free parameters: the probability per unit time for a spontaneous localisation, and the parameter defining the width of the localised state. In fact, the aspect of the theory that is free to be adjusted is much larger than two parameters, because the position distribution after spontaneous localisation is free; a wide range of arbitrary localised distributions can be chosen.

Larc theory does not have this freedom: the distribution of the localised state is determined by the detail of the mechanism that establishes the correlation between relative position and the bound electron states within molecules. In fact, the distribution of the localised state is determined in exactly the same way it would have been in standard 'good taste’ quantum theory.

The time distribution for the spontaneous localisation in GRW is also free, but in GRW a constant probability per unit time is natural given that every particle may localise at any time. The only event that could form an anchor for a distribution not constant in time would be each particle's own localisations, or possibly those of particles spatially correlated with the first. In this exposition I have assumed that the Larc theory also produces a constant probability per unit time. Although the justification for this is less natural, because Larc theory does have a trigger event relative to which the probability distribution could be defined. Nevertheless, constant probability per unit time remains a natural expectation in the Larc theory.

So Larc theory contains only one free parameter, where GRW has two, plus
significant freedom in the position distribution.

I have not made any attempt to resolve one point. Larc clusters occur also in virtual processes. There is therefore the question of whether virtual larc clusters should be considered to trigger state reductions just as real ones do. If they do, it is also not clear whether the temporal constant should be the same in both cases. I have no basis on which to offer an opinion on whether virtual larc clusters should be considered as triggering state reductions. The only comment I have to make is that, if they do, this will certainly have an effect on the calculation of virtual processes, but with results that may be significant only in those of high complexity.

7 Empirical tests

Since the core purpose of a measurement theory is to match the well-known behaviour of measurements, without changing the known behaviour of other types of quantum systems, empirical verification of a measurement theory is inherently difficult. Nevertheless, as in any other case where one has changed the fundamental laws underlying a theory, one expects that there will be at least one situation where the predictions of the new theory diverge from those of the old. This is what we expect for all type 2 solutions to the measurement problem. Measurement is not a physically well-defined concept, so it is next to impossible that any physical law could exactly match its behaviour. At some point, there will be a situation where the new law operates detectably when there is no measurement occurring.

The problem of verification is made worse by the phenomenon usually called environment-induced decoherence. This phenomenon mimics an aspect of the behaviour of measurements by causing the branches corresponding to the possible outcomes of measurements to lose their coherence. A measurement necessarily requires the measured system to be put into correlation with the measuring apparatus. If one considers the measuring apparatus by itself, tracing out the measured system, its possible outcome states become decohered. As the total state of the system and apparatus remains a superposition this does not solve the measurement problem, but it does mean that interference effects that would be destroyed by a successful measurement theory are destroyed in any case by decoherence, making straightforward tests impossible.

Nevertheless, the possibility of processes that generate large enough numbers of larc to produce detectable loss of interference, without its being masked by environment-induced decoherence, remains.

If larc-triggered decoherence occurs during virtual processes, this may also produce detectable deviation from behaviours predicted by standard quantum field theory.
8 Conclusion

In 1990, Shimony[7] provided a heuristic justification for a theory in which state reductions result from sudden jumps to energy eigenstates. Technically, larc theory is not such a theory, since in it jumps are to eigenstates of number operators defined by larc operators, not energy operators. Nevertheless, since the number operator states for electron orbitals correspond to bound state energy eigenstates, larc theory is essentially a theory of the type Shimony called for. Shimony’s argument therefore may constitute support for the larc-abc theory.

In my case, at least, the core intuition behind the desire for a measurement theory involving jumps to energy eigenstates is that if bound electrons jumped to energy eigenstates after absorbing incoming particles, they would carry anything correlated with them, such as the centres of mass of other bodies, into suitably classical states. For bound states, the energy eigenstates seem to form a natural solution to the basis problem, where they do not for unbound states. A severe barrier to achieving a ‘jumps to energy eigenstates’ theory has been the expectation that it would be impossible to treat the bound states separately from the other bodies correlated with them, and therefore one would be forced to adopt a theory where the entire correlated state would need to jump to an energy eigenstate, even though this would include the centre of mass states of free bodies.

From this point of view, the crucial element of larc theory is that the creation and annihilation operators in larc events provide a barrier that separates the bound energy eigenstates from other elements of the total system, so that it is possible for bound electrons to jump to energy eigenstates, and carry other systems that are correlated with them into states that need not themselves be energy eigenstates. Nor does the total state of the whole system need to be an energy eigenstate.

The measurement problem is one of the oldest open questions in quantum theory. The list of attempted solutions to this problem is long. While many have their good points, most have failed, or at least failed to be convincing. Some remain in play, but none that can claim to be a clear solution to the problem.

The theory described here has a number of advantages that I believe have not been seen before in a single theory. It solves the problem as a matter of straightforward physics, without requiring conceptual contortions in attempting to generate the Born rule probabilities, as the Everett theory does. It produces localisation of macroscopic bodies in the most physically natural way, as a direct result of illumination reflected from one body being absorbed by other bodies. And it is that rarity, a reduction theory which is naturally relativistic.

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