# Models of Wave-function Collapse, Underlying Theories, and Experimental Tests 

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#### Abstract

Quantum mechanics is an extremely successful theory that agrees with every experiment. However, the principle of linear superposition, a central tenet of the theory, apparently contradicts a commonplace observation: macroscopic objects are never found in a linear superposition of position states. Moreover, the theory does not really explain as to why during a quantum measurement, deterministic evolution is replaced by probabilistic evolution, whose random outcomes


[^0]we review an experimentally falsifiable phenomenological proposal, known as Continuous Spontaneous Collapse: a stochastic nonlinear modification of the Schrödinger equation, which resolves these problems, while giving the same experimental results as quantum theory in the microscopic regime. Two underlying theories for this phenomenology are reviewed: Trace Dynamics, and gravity induced collapse. As one approaches the macroscopic scale, the predictions of this proposal begin to differ appreciably from those of quantum theory, and are being confronted by ongoing laboratory experiments that include molecular interferometry and optomechanics. These experiments, which essentially test the validity of linear superposition for large systems, are reviewed here, and their techni-
cal challenges, current results, and future prospects summarized. We conclude that it is likely that over the next two decades or so, these experiments can verify or rule out the proposed stochastic modification of quantum theory.

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## I. INTRODUCTION

Quantum theory has been extremely successful in explaining results of experiments, ranging from spectrum of black-body radiation, atomic spectra, molecular chemistry, atomic interferometry, quantum electrodynamics and nuclear physics, to properties of lasers, superconductivity, semi-conductor physics, Bose-Einstein condensation, Josephson junctions, nano-technology, applications in cosmology, and much more. The theory is not contradicted by any experiment. Yet, there is one apparently innocuous observed phenomenon the theory seems unable to explain, and in fact seems to contradict with. This is the observed absence of superposition of different position states in a macroscopic system. Quantum theory, by virtue of the principle of linear superposition, predicts that a microscopic object such as the electron can be in more than one position at the same time, and this is of course observed, for example in the famous double-slit interference experiment. Moreover, the theory in principle makes no distinction between microscopic objects and macroscopic ones, and predicts that large objects can also be in more than one place at the same time. But this is not what we observe. A table for example, unlike the electron, is never observed to be 'here' and 'there' simultaneously.

Why should this be so? The present review article is devoted to discussing one possible proposed resolution, known as Continuous Spontaneous Collapse, which is experimentally falsifiable. Namely that, although quantum theory is extremely successful in the microscopic domain, it is an approximation to a more general theory. This general theory is capable of explaining the observed absence of macroscopic superpositions. It goes over to quantum mechanics in the microscopic limit, and to classical mechanics in the macroscopic limit, but differs from both quantum and classical mechanics in the intermediate [mesoscopic] regime which marks the transition from the micro- to the macroworld. A large number of experiments worldwide are operating or are being planned, to test the validity of linear superposition in the mesoscopic domain, and in this article we will review the proposed modification to quantum mechanics, and the laboratory experiments which can falsify this proposal.
A. The relation
retween
relan-
classical mechanics

The classical dynamics of a system of particles having a Hamiltonian $H$ is described in phase space $\left(q_{i}, p_{i}\right)$ by Hamilton's equations
of motion

$$
\begin{equation*}
\dot{q}_{i}=\frac{\partial H}{\partial p_{i}}, \quad \dot{p}_{i}=-\frac{\partial H}{\partial q_{i}} \tag{1}
\end{equation*}
$$

or via Poisson brackets

$$
\begin{equation*}
\dot{q}_{i}=\left\{q_{i}, H\right\}, \quad \dot{p}_{i}=\left\{p_{i}, H\right\} . \tag{2}
\end{equation*}
$$

The state of the system at an time initial $t_{0}$ is a point in the phase space, and the equations of motion determine the location of the system point at a later time. An equivalent description of the dynamics is through the Hamilton-Jacobi equation

$$
\begin{equation*}
-\frac{\partial S}{\partial t}=H\left(q_{i}, \frac{\partial S}{\partial q_{i}}\right) \tag{3}
\end{equation*}
$$

where $S$ is the action of the system (Landau and Lifshitz, 1976).

In contrast, the quantum dynamics of this system is described by first converting the $q_{i}$ and $p_{i}$ to operators $\mathbf{q}_{\mathbf{i}}, \mathbf{p}_{\mathbf{i}}$ satisfying the commutation relations $\left[\mathbf{q}_{\mathbf{i}}, \mathbf{p}_{\mathbf{i}}\right]=i \hbar$ and then proposing that the operators evolve via the Heisenberg equations of motion

$$
\begin{equation*}
\dot{\mathbf{q}}_{\mathbf{i}}=\left[\mathbf{q}_{\mathbf{i}}, \mathbf{H}\right], \quad \dot{\mathbf{p}}_{\mathbf{i}}=\left[\mathbf{p}_{\mathbf{i}}, \mathbf{H}\right] . \tag{4}
\end{equation*}
$$

Quantum dynamics is equivalently described by the time evolution of the system's wavefunction $\psi$, which is a normalized element of a Hilbert space and obeys the normpreserving Schrödinger equation

$$
\begin{equation*}
i \hbar \frac{\partial \psi}{\partial t}=H \psi, \quad \int d q \psi^{*} \psi=1 \tag{5}
\end{equation*}
$$

In the Heisenberg picture the relation between quantum and classical mechanics is expressed by setting the position-momentum commutator to zero, replacing operators by ordinary numbers, and the commutators in the equations of motion by Poisson brackets. A more insightful comparison is obtained in the Schrödinger picture, and for the purpose of illustration it is adequate to consider the case of a single particle of mass $m$ moving in one dimension, for which the Schrödinger equation can be written in the position representation, after defining $\psi \equiv e^{i S / \hbar}$, as

$$
\begin{equation*}
-\frac{\partial S}{\partial t}=\frac{1}{2 m}\left(\frac{\partial S}{\partial q}\right)^{2}+V(q)-\frac{i \hbar}{2 m} \frac{\partial^{2} S}{\partial q^{2}} \tag{6}
\end{equation*}
$$

In the approximation in which the last term in (6) can be neglected, this equation reduces to the classical Hamilton-Jacobi equation (1)

$$
\begin{equation*}
-\frac{\partial S}{\partial t}=\frac{1}{2 m}\left(\frac{\partial S}{\partial q}\right)^{2}+V(q) \tag{7}
\end{equation*}
$$

provided the quantity $S$ is assumed to be real and identified with the action of the system. This essentially corresponds to the limit $S \gg$ $\hbar$. [We will not consider the more precise treatment where $S$ is separated into its real and imaginary parts, as it is not crucial to the present discussion].

There is thus a well-defined sense in which the Schrödinger equation goes over to the Hamilton-Jacobi equation in the limit, and a description of the dynamics in Hilbert space
gets replaced by a description in terms of evolution of position and momentum coordinates in phase space. Yet, there is a profound aspect which gets lost in the limiting process. The Schrödinger equation is linear : if $\psi_{1}$ and $\psi_{2}$ are two solutions of (5) then the linear superposition $c_{1} \psi_{1}+c_{2} \psi_{2}$ is also a solution, where $c_{1}$ and $c_{2}$ are complex coefficients. On the other hand the Hamilton-Jacobi equation (1) is non-linear : if $S_{1}$ is a solution corresponding to one space-time trajectory, and $S_{2}$ is a solution corresponding to another spacetime trajectory, then clearly $a_{1} S_{1}+a_{2} S_{2}$ is not a solution of this equation.

In particular, if $\psi_{1}$ is a wave-packet which is peaked around one classical solution and $\psi_{2}$ is a wave-packet peaked around another classical solution, quantum mechanics predicts that the sum of these two wave-packets is also a solution, and in principle such solutions should be observed in nature. However, according to classical mechanics, such a superposition is not a solution of the equations of motion, nor is it observed in the macroscopic world around us. Naively, we believe that classical mechanics, which applies to macroscopic systems, is a limiting case of quantum mechanics, and hence quantum mechanics should apply to large systems as well. Why then we do not observe macroscopic superpositions [such as a table being 'here' and 'there' at the same time]?

One might argue that even though the Hamilton-Jacobi equation is non-linear, its non-linearity cannot be used to deduce the observed absence of macroscopic superpositions, because the classical theory is after all an approximation. The last term in Eqn. (6), howsoever small, is always non-zero and present, and can be used to transform back to the linear Schrödinger equation. At a fundamental level, the description of the dynamics, even for a macroscopic classical object, is in terms of the wave-function of the quantum state, and not in terms of the action which appears in the Hamilton-Jacobi equation. Hence superpositions must be observed. Nonetheless, one is left wih the discomforting feeling that the prediction of the HamiltonJacobi equation regarding position superpositions seems to be at variance with quantum theory, and in accord with what is actually observed. Thus one needs to explain the following : why is it that macroscopic objects which obey the rules of classical mechanics are not found in superposition of different position states, in spite of quantum theory suggesting otherwise? There is no unique universally accepted answer to this question. In this sense this is an unsolved problem.

The absence of macroscopic superpositions is of course at the heart of the so-called quantum measurement problem. Suppose a quantum system which is in a superposition
of two eigenstates $\psi_{1}$ and $\psi_{2}$ of a physical observable $\mathbf{O}$ interacts with a classical measuring apparatus $A$. Let us say that the state $\psi_{1}$ of the quantum system corresponds to a pointer position state $A_{1}$ of the apparatus [meaning that if the system had been in the state $\psi_{1}$ and interacted with the apparatus, the pointer would result in the position $A_{1}$ and we would interpret that the observable had the value $\left.O_{1}\right]$. Similarly, the pointer position $A_{2}$ corresponds to the system state $\psi_{2}$ and a value $O_{2}$ for the observable $\mathbf{O}$. Immediately after interaction, the combined state of the system and apparatus is

$$
\begin{equation*}
\psi=c_{1} \psi_{1} A_{1}+c_{2} \psi_{2} A_{2} \tag{8}
\end{equation*}
$$

where $c_{1}$ and $c_{2}$ are complex coefficients proportional to the relative amplitudes for the system to be in the two states $\psi_{1}$ and $\psi_{2}$.

According to quantum mechanics, this state $\psi$ of Eqn. (8) should evolve linearly by way of Schrödinger evolution, and the linear superposition of the two parts should be preserved. But that is not what is observed in a quantum measurement. The outcome of the measurement is either pointer at position $A_{1}$ (and hence system is driven to state $\psi_{1}$ ) or pointer at position $A_{2}$ (system is driven to state $\psi_{2}$ ). Repeated measurements on the same initial quantum state yield outcome $\psi_{1}$ or $\psi_{2}$ with relative probability $\left|c_{1}\right|^{2}:\left|c_{2}\right|^{2}$. This is the Born probability rule. The pro-
cess of measurement destroys linear superposition of the initial states $\psi_{1}$ and $\psi_{2}$. This indeed has to do with the fact that the apparatus [which is a macroscopic object] is never simultaneously observed in a linear superposition of pointer position states $A_{1}$ and $A_{2}$. To the extent that we do not undersand why macroscopic objects are not found in superposed states, we do not understand why the measurement process breaks superposition.

Perhaps even more remarkable is the emergence of probabilities. The Schrödinger evolution is deterministic, and so is the classical evolution according to the HamiltonJacobi equation. In our discussion above, on the transition from the Schrödinger equation to the Hamilton-Jacobi equation, nowhere did we encounter probabilities. And for good reason. Because the initial state is always exactly specified [including at the start of the measurement process, as in Eqn. (8)], unlike in classical probability theory, where probabilities arise because of uncertainty in our knowledge of the initial state of the system. Thus the status of probabilities in quantum theory is absolutely unique, and besides explaining the absence of macroscopic superpositions one must also explain why during a measurement probabilities arise, in violation of deterministic linear superposition, and the quantum system is driven to one or the other outcome in accordance with the Born rule.

Another important and related unsolved problem is the following : when do we call a physical system a quantum system and when we do call it a classical measuring apparatus? In other words, where is the quantumclassical divide? How much mass, or degrees of freedom [say number of nucleons] should an object have, before it qualifies to be an apparatus? Of course, in order for it to be called an apparatus, different pointer positions should never be simulaneously realized, but one does not know at what mass scale this transition from micro to macro [and the concurrent breakdown of superposition] takes place. Interferometry experiments have shown that quantum theory, and hence linear superposition, holds for molecules at least as large as those having about a thousand atoms [hence a molecular mass of $10^{-21}$ grams]. Efforts are afoot to push this test limit up to objects of about a million atoms [ $10^{-18}$ grams]. On the other end, classical behavior [absence of position superpositions] is known to hold out down to about a microgram [ $10^{18}$ atoms]. There is thus an enormous desert of some fifteen orders of magnitude, where linear quantum superposition yet remains to be tested experimentally. Does quantum mechanics hold at all scales, including macroscopic scales, and is there a way to understand the absence of macroscopic superpositions while staying within the frame-
work of quantum theory? Or is it that somewhere in the grand desert modifications to quantum theory start becoming significant, so that linear superposition becomes more and more of an approximate principle as the size of a system is increased, until for large objects position superposition is no longer a valid principle? What exactly is the nature of the quantum-to-classical transition? A large number of ongoing and planned experiments worldwide are gearing up to address this question.

The paradoxical issue of deterministic evolution followed by a peculiar probabilistic evolution during a measurement was of course well-appreciated by the founding fathers of quantum mechanics. Over the last eightyfive years or so since the discovery of the Schrödinger equation, extraordinary theoretical effort has been invested in trying to find an answer to what is generally known as the measurement problem in quantum mechanics Albert, 1992; Bell, 1987; Ghirardi, 2005; Leggett, 2002, 2005; Maudlin, 2011; Wheeler and Zurek, 1983). In the next subsection we give a brief overview of a few of the major categories of the explanations, keeping in mind that the modern outlook is to discuss this problem not in isolation, but in conjunction with the question of lack of macroscopic superpositions, and as a part of the much broader investigation of the exact nature of
the quantum-to-classical transition.
Our review of the measurement problem will be almost exclusively confined to the context of non-relativistic quantum mechanics, as the relativistic version seems not within reach at the moment. Whether this is symptomatic of a deep incompatibility of modified quantum mechanics and relativity is perhaps hard to say at present. Thus we will not discuss issues raised by the instantaneous nature of wave-function collapse, such as the EPR paradox, whether this 'violates the spirit of relativity' or whether there is a need for a radical change in our ideas about space-time structure.

## B. Proposed resolutions for the quan-

 tum measurement problem and for the observed absence of macroscopic superpositions
## 1. The Copenhagen Interpretation

The Copenhagen interpretation $\overline{\mathrm{Bohr}}$, 1928) [reprinted in Wheeler and Zurek, 1983)] postulates an artificial divide between the micro-world and the macro-world, without quantitatively specifying at what mass scale the divide should be. Microscopic objects obey the rules of quantum theory [superposition holds] and macroscopic objects obey the rules of classical mechanics [super-
position does not hold]. During a measurement, when a micro-system interacts with a macro-system, the wave-function of the micro-system 'collapses' from being in a superposition of the eigenstates of the measured observable, to being in just one of the eigenstates. This collapse is postulated to happen in accordance with the Born probability rule, and no dynamical mechanism is specified to explain how the collapse takes place, in apparent contradiction wth the linearity of the Schrödinger equation.
von Neumann gave a more precise form to this interpretation by explicitly stating that evolution in quantum theory takes place in two ways : (i) deterministic evolution according to the Schrödinger equation before a measurement, as well as after a measurement, and (ii) non-deterministic, probabilistic evolution [the projection postulate] during a measurement (von Neumann, 1955).

At a pragmatic level, this can be taken to be a perfectly valid set of rules, in so far as the goal is to apply quantum theory to sub-atomic, atomic and molecular systems, and to compare with experiments the predictions based on theoretical calculations. However, the interpretation bypasses the questions raised in the previous section, by simply raising unresolved issues to the level of postulates. The interpretation creates an illdefined micro-macro separation, which is be-
ing challenged by modern experiments which are verifying superposition for ever larger systems. There is no precise definition as to which systems classify to serve as a 'classical measuring apparatus'. Even though there is a sense in which the Hamilton-Jacobi equation is a limit of the Schrödinger equation, no attempt is made to explain the apparently different predictions of the two theories with regard to absence of macroscpic position superpositions. At a fundamental level one should prescribe a physical mechanism which causes the so-called collapse of the wave-function.

The Copenhagen interpretation does not solve the quantum measurement problem, nor does it explain the absence of macroscopic superpositions.

## 2. Decoherence

The phenomenon of decoherence, which is observed in laboratory experiments, highlights the role played by the environment when a quantum system interacts with a measuring apparatus, during the process of measurement. By environment is meant the system of particles which surround the apparatus. More precisely one could define the environment as the collection of particles which is present within a radius $c T$ of the apparatus, where $T$ is the duration of the measure-
ment : these are hence particles which can causally interact with and influence the apparatus during a measurement.

To illustrate the effect of decoherence, we assume that the system on which measurement is to be made is a two state system initially in the state

$$
\begin{equation*}
\psi(t=0)=c_{1} \psi_{1}+c_{2} \psi_{2} \tag{9}
\end{equation*}
$$

Denoting the initial state of the apparatus by $\phi_{0 A}$ and the initial state of the environment by $\phi_{0 E}$, we can write the net initial state as the direct product

$$
\begin{equation*}
\Phi_{0}=\psi(t=0) \phi_{0 A} \phi_{0 E} \tag{10}
\end{equation*}
$$

Over time, as a result of interaction, this state evolves into the state

$$
\begin{equation*}
\Phi(t)=c_{1} \psi_{1} \phi(t)_{E A 1}+c_{2} \psi_{2} \phi(t)_{E A 2} \tag{11}
\end{equation*}
$$

Here, $\phi(t)_{E A 1}$ and $\phi(t)_{E A 2}$ denote macroscopically distinguishable entangled states of the apparatus and the environment.

As demonstrated below, during measurement, the process of decoherence operates in such a way that very quickly, the inner product

$$
\begin{equation*}
\left\langle\phi(t)_{E A 1} \mid \phi(t)_{E A 2}\right\rangle \rightarrow 0 \tag{12}
\end{equation*}
$$

starting from the value unity at $t=0$. The final state is reduced to a statistical mixture of states with relative weights $\left|c_{1}\right|^{2}:\left|c_{2}\right|^{2}$.

This by itself does not explain why during product a measurement

$$
\begin{equation*}
\Phi(t) \rightarrow \psi_{1} \phi(t)_{E A 1} \quad \text { or } \quad \Phi(t) \rightarrow \psi_{2} \phi(t)_{E A 2} . \tag{13}
\end{equation*}
$$

Decoherence destroys interference amongst alternatives, which is what Eqn. (12) signifies, but because it operates within the framework of linear quantum mechanics, it cannot destroy superposition. Since loss of superposition is what is seen during a measurement, decoherence does not explain the measurement process. What (12) implies is that decoherence forces quantum probability distributions to appear like classical probabilities (weighted sums of alternatives); however this is neither necessary nor sufficient to explain the outcome of an individual measurement.

The loss of interference can be understood as a consequence of the interaction of the very large number of particles of the environment with the apparatus. Assuming that the measurement starts at $t=0$, the product $\left\langle\phi(t)_{E A 1} \mid \phi(t)_{E A 2}\right\rangle$, which is one at $t=0$, rapidly goes to zero. To see this, one notes that in general a particle of the environment, say the $i$ th particle, will be scattered by the state $A_{1}$ of the apparatus to a final state different from the one to which it will be scattered from the apparatus state $A_{2}$. Thus, the
$\left\langle E_{1} \mid E_{2}\right\rangle(t)=\Pi_{i}{ }_{i}\langle E(t=0)| S_{A 1} S_{A 2}|E(t=0)\rangle_{i}$
is made up of an ever increasing number of quantities, each of which is smaller than one, $S_{A 1}$ and $S_{A 2}$ being scattering matrices describing the action of the apparatus on the environment. Hence this product can be written as $\exp (-\Lambda t)$ and goes to zero for large $t, \Lambda$ being the decoherence rate. Because the environment has a very large number of particles, this cross-product between the two environment states is very rapidly suppressed, and is responsible for the emergence of the property described by Eqn. (12). The decoherence time-scale $\Lambda^{-1}$ is much smaller than the duration $T$ of the measurement.

The above discussion is partly based on the article by Adler (Adler, 2003) where a more detailed description of decoherence in the context of measurement can be found. There is a vast literature on decoherence, including the experiments and models by (Brune et al., 1996; Gerlich et al., 2007; Harris and Stodolsky, 1981), books by (Breuer and Petruccione, 2000; Joos et al., 2003; Schlosshauer, 2007) and the seminal papers (Joos and Zeh, 1985; Zeh, 1970) and reviews (Schlosshauer, 2005; Vacchini and Hornberger, 2009, Zurek, 1991, 2003), and (Bacciagaluppi, 2007).

## 3. Many-Worlds interpretation

The Many-worlds interpretation was invented by Everett (Everett III, 1957) to counter the Copenhagen interpretation. According to Everett, evolution during a measurement is also Schrödinger evolution, and there is no such thing as a non-deterministic probabilistic evolution during the measurement. Thus in this interpretation the state (8) evolves during a measurement according to the Schrödinger equation. Why then does it appear as if only one of the two outcomes has been realized? The answer is that the state continues to be of the form

$$
\begin{equation*}
\Psi=c_{1} \psi_{1} A_{1} O_{1}+c_{2} \psi_{2} A_{2} O_{2} \tag{15}
\end{equation*}
$$

where $O_{1}\left(O_{2}\right)$ is the state of the observer where the observer detects the system and apparatus in state one (two). The two parts of this state exist in two different branches of the Universe.

Despite appearances, there is no logical inconsistency in the above interpretation, as has been argued by Everett : it is merely the assertion that Schrödinger evolution is universally valid at all scales, and the breakdown of superposition during a measurement is only apparent, not real. The hard part is to explain the origin of probabilities and the Born probability rule. If the evolution is deterministic through and through, why
should there be a definite probability associated with an outcome? In our opinion, despite extensive investigation, this problem remains unsolved in the many-worlds interpretation (Deutsch, 1998; DeWitt and Graham, 1973; Hsu, 2011; Kent, 1990; Tegmark, 2007, Vaidman, 2002; Wallace, 2003).

## 4. Decoherence + Many-Worlds

Decoherence by itself does not solve the measurement problem, because it does not destroy superposition. However, one could propose that the decohered alternatives both continue to co-exist in different branches of the Universe, in the sense of the many-worlds interpretation, and these branches do not interfere with each other because decoherence is operating. While this merger helps both the decoherence and the many-worlds picture of a measurement, the origin of the Born probability rule continues to be not understood, and as of now is essentially added as a postulate.

This is perhaps today the 'establishment view', wherein one believes that one does not need to modify quantum theory in order to explain measurement. Its major weakness though is that it is not experimentally falsifiable. What experiment can one perform in order to find out whether the other branches of the many-worlds exist or not? In the ab-
sence of such an experiment, we have at hand another interpretation of the same quantum theory, an interpretation which cannot be experimentally distinguished from the Copenhagen interpretation.

For discussions on decoherence in the context of many-worlds see Bacciagaluppi, 2001).

## 5. Bohmian Mechanics

It is proposed that there are additional degrees of freedom, labelled let us say by the set $Q_{k}$, which carry more information about the quantum system than is provided by the wave-function $\psi$ and the Schrödinger equation which describes the evolution of this wave-function. This additional information is such that the outcome of a measurement is known beforehand, and can be predicted deterministically, without having to assign an ad hoc probability to the outcome. The additional degrees of freedom $Q_{k}$ are distributed over a parameter space in such a way that the different outcomes are realized in accordance with the Born rule.

The degrees of freedom $Q_{k}$ are assumed to describe the coordinates of hidden "particles" which move in configuration space. The wave-function $\psi$, besides depending on the usual configuration space variables $\left(q_{1}, q_{2}, q_{3}, \ldots, q_{N}\right)$ also depends on the coordi-
nates $Q_{k}$, whose velocity $v_{k}$ obeys the equation

$$
\begin{align*}
v_{k} & =\frac{d Q_{k}}{d t} \\
& =\frac{\hbar}{m_{k}} \operatorname{Im} \nabla_{Q_{k}} \log \psi\left(Q_{1}, Q_{2}, Q_{3}, \ldots, Q_{N}, t\right) . \tag{16}
\end{align*}
$$

Bohmian mechanics is another interpretation of quantum mechanics which while being logically consistent, does not seem to be experimentally falsifiable, since it makes the same predictions as the standard theory.

For literature on Bohmian mechanics see (Bohm, 1952a b; Bohm and Bub, 1966; Bub, 1997; Dürr et al., 1992)
6. Quantum theory is an approximation to a more general theory

It is proposed here that the measurement problem and the apparent inability of quantum theory to explain the absence of macroscopic superpositions are a consequence of trying to apply the theory in a domain where it is not valid. It is proposed that there is a universal dynamics, to which quantum theory and classical mechanics both are approximations. In the domain of a quantum measurement, the universal dynamics differs from quantum dynamics, and operates in such a way that interaction of the quantum system with the apparatus causes a collapse of the wave-function from a superposition to one of
the eigenstates. The collapse is a physical, dynamical process, and hence the universal dynamics provides a physical explanation for the ad hoc collapse postulated by the Copenhagen interpretation. Furthermore, the collapse is shown to obey the Born probability rule. The universal dynamics is stochastic : the outcome of a measurement is random and unpredictable, but the mathematical structure of the dynamics is such that repeated measurements on an ensemble of identically prepared quantum systems are shown to yield different outcomes, in relative frequencies which obey the Born rule.

The universal dynamics must be nonlinear, in order to allow for the breakdown of superposition during a measurement. Yet, the non-linearity must be extremely negligible in the microscopic domain, so that the experimentally observed linear superposition in microscopic quantum systems is reproduced. The new dynamics must be stochastic; but once again, stochasticity must be negligible for microscopic systems, so that the deterministic Schrödinger evolution prevails. Thirdly, as one progresses from microscopic to macroscopic systems, the universal dynamics must allow for non-unitary [but norm-preserving] evolution : this is essential so that stochastic evolution can cause all but one outcomes to decay exponentially, something which would not be permitted during
unitary evolution. Again, non-unitarity must be utterly negligible for microscopic systems. Thus, the universal dynamics possesses a set of parameters, whose effective values are determined for the system under study in such a way that for microscopic systems these parameters take values so that the dynamics is experimentally indistinguishable from quantum dynamics. Similarly, for macroscopic systems, the parameters take effective values such that the dynamics coincides with classical dynamics. For systems that are mesoscopic [neither micro nor macro] the dynamics differs from both classical and quantum, and is hence experimentally distinguishable from quantum theory. The properties of nonlinearity, stochasticity and non-unitarity also ensure position localization for macroscopic objects and hence dynamically explain the observed absence of macroscopic superpositions. It is clear that the universal dynamics is not tied to or invented for explaining just the measurement process or absence of macroscopic superpositions - these two phenomena just happen to be special cases where the new dynamics plays a vital role in understanding the phenomenon. We say that the universal dynamics, which describes the behaviour of micro, meso and macro objects, is intrinsically non-linear, stochastic, and nonunitary.

Over the last two decades or so there has
been a significant progress in developing phenomenological models of such a universal dynamics. At the same time, one would like to know if there are underlying theoretical reasons [new symmtery principles for instance] which compel us to consider a generalization of quantum theory of the kind mentioned above, thereby lending an inevitability to the phenomenological models that have been proposed. There has been important progress on this front too. Thirdly, there have been important technological advances which are now permitting a host of experiments to be carried out to test these phenomenological models, and verify their predictions against those of quantum theory. Needless to add, all these three facets are best described as 'work currently in progress'. The purpose of the present review is to present a state-of-the-art description of (i) the phenomenological models for the universal dynamics, (ii) the underlying theories, and (iii) ongoing experiments which aim to test these models and theories. It is our hope that a review of this nature will further stimulate the cross-talk between phenomenologists, theorists and experimentalists in this field, thereby helping the community to sharply focus on those aspects of phenomenology and experimentation which might be most directly accessible and feasible in the near future.

Phenomenological models of modified

## quantum mechanics

From early times, an aspect which has received considerable attention, is possible nonlinear modifications of quantum theory, and this is not necessarily because of the measurement problem. Most fundamental differential equations which describe physical phenomena are non-linear, with linearity being a convenient approximation in some appropriate limiting cases. Why then should an equation as fundamental as the Schrödinger equation be a singular exception to this rule? [It is of course known that there are very strong bounds on non-linearity in the atomic domain, see for instance the experiment described in (Bollinger et al., 1989)]. Nonlinear quantum theories may be classified as deterministic non-linear, and stochastic nonlinear. For discussions on detrministic nonlinear quantum mechanics the reader is referred to the works by Weinberg Weinberg, 1989a b), by Goldin (Doebner and Goldin, 1992; Goldin, 2000), and Bialynicki-Birula and Mycielski, 1976). It has often been suggested, and demonstrated, though perhaps not universally so, that deterministic nonlinear modifications result in superluminal propagation (Gisin, 1990; Polchinski, 1991). This, coupled with the fact that stochasticity appears to be an essential ingredient for explaining the origin of probabilities, has meant that investigations of a universal dynam-
ics have tended to focus on stochastic nonlinearities; see for instance (Diósi, 1988a b; Gisin, 1981, 1984, 1989, Gisin and Rigo, 1995; Weinberg, 2011).

With regard to the application of stochastic non-linearity to explain measurement, the pioneering paper is due to Pearle (Pearle, 1976) - the paper is aptly titled "Reduction of the state vector by a non-linear Schrödinger equation". In hindsight, it is perhaps surprising that an idea such as this one should appear on the scene so late in the day. On the other hand, a delay of this kind is testimony to the adherence to the Copenhagen interpretation. Pearle proposed to replace the Schrödinger equation by a non-linear one, during measurement, and that certain variables which take random values just after the quantum system interacts with the apparatus, drive the system to one or the other outcomes, thus breaking superposition. For the choice of these random variables he suggested the phases of the state vectors immediately after the measurement. An appropriate assignment of the probability distribution of these phases over the allowed parameter space leads to the Born rule. It is noteworthy that this assignment of the probability distribution is something which has to be put in by hand, keeping in mind what probability rule one wants to emerge. This is one aspect where phenomenology and under-
lying theories need to do better even today: there should be a fundamental reason for the probability distribution over the stochastic variables, which inevitably implies the Born rule. Further investigations by Pearle were reported in (Pearle, 1979, 1982, 1984, 1989) and reviewed in (Pearle, 1999a).

The next major advance came from Ghirardi, Rimini and Weber Ghirardi et al., 1986) in a seminal paper titled "Unified dynamics for microscopic and macroscopic systems" and the model has come to be known as the GRW model. There were two guiding principles for this dynamical reduction model (also known as QMSL: Quantum Mechanics with Spontaneous Localization):

1. The preferred basis - the basis on which reductions take place - must be chosen in such a way as to guarantee a definite position in space to macroscopic objects.
2. The modified dynamics must have little impact on microscopic objects, but at the same time must reduce the superposition of different macroscopic states of macrosystems. There must then be an amplification mechanism when moving from the micro to the macro level.

The reduction is achieved by making the following set of assumptions:

1. Each particle of a system of $n$ distinguishable particles experiences, with a mean rate $\lambda_{i}$, a sudden spontaneous localization
process.
2. In the time interval between two successive spontaneous processes the system evolves according to the usual Schrödinger equation.

In their model, GRW introduced two new fundamental constants of nature, assumed to have definite numerical values, so as to reproduce observed features of the microscopic and macroscopic world. The first constant, $\lambda^{-1} \sim 10^{16}$ seconds, alluded to above, determines the rate of spontaneous localization (collapse) for a single particle. For a composite object of $n$ particles, the collapse rate is $(\lambda n)^{-1}$ seconds. The second fundamental constant is a length scale $r_{C} \sim 10^{-5} \mathrm{~cm}$ which is related to the concept that a widely spaced wave-function collapses to a length scale of about $r_{C}$ during the localization.

A gravity based implementation of the GRW model was studied by Diósi Diósi, 1989) and generalized by (Ghirardi et al., 1990a).

The GRW model has the limitation that it does not preserve symmetry of the wavefunction under particle exchange, and has been improved into what is known as the CSL (Continuous Spontaneous Localization) model by Ghirardi, Pearle and Rimini (Ghirardi et al., 1990c). In CSL a randomly fluctuating classical field couples with the particle number density operator of a quantum
system to produce collapse towards its spatially localized eigenstates. The narrowing of the wavefunction amounts to an increase in the energy of the particle, and actually amounts to a tiny violation of energy conservation.

An outstanding open question with regard to the dynamical reduction models is the origin of the random noise, or the randomly fluctuating classical scalar field, which induces collapse.

The current status of the Spontaneous Collapse models is discussed in detail in Section II.

A modern approach to stochastic reduction is to describe it using a stochastic nonlinear Schrödinger equation, an elegant simplified example of which is the following oneparticle case [known as QMUPL : Quantum Mechanics with Universal Position Localization (Diósi, 1989)] [See Section II for details]:

$$
\begin{align*}
d \psi(t)= & {\left[-\frac{i}{\hbar} H d t+\sqrt{\lambda}\left(q-\langle q\rangle_{t}\right) d W_{t}-\right.} \\
& \left.\frac{\lambda}{2}\left(q-\langle q\rangle_{t}\right)^{2} d t\right] d \psi(t) \tag{17}
\end{align*}
$$

$q$ is the position operator, $\langle q\rangle_{t}$ is its expectation value, and $\lambda$ is the rate constant alluded to above. $W_{t}$ is a Wiener process which describes the impact of stochasticity on the dynamics. This equation can be used to explain the collapse of wave-function during a measurement, the emergence of the Born rule, the absence of macroscopic superpositions, and
the excellent matching of the linear theory with experiments for microscopic systems.

Various studies and arguments suggest that the structure of this equation is very rigid and tightly controlled, once one assumes [as is true here] that the evoution is norm-preserving, and secondly, superluminal propagation is not possible Adler, 2004; Gisin, 1989). There is then a unique relation between the coefficient $\sqrt{\lambda}$ of the diffusion [stochastic] term and the coefficient $-\lambda / 2$ of the drift term : Drift Coefficient $=-2$ (Diffusion Coefficient) ${ }^{2}$. This is the well-known martingale structure for a stochastic differential equation.

In QMUPL, stochastic fluctuations take place only in the time direction and hence there is only one free parameter, i.e. $\lambda$. In contrast, in the CSL model the stochastic fluctuations exist over space too, and hence there is a second free parameter $r_{C}$ (analogous to GRW) which defines the scale of spatial localization. Of course, in the QMUPL and CSL models the stochastic process acts continously, unlike in GRW, wherein the stochastic jumps are discontinuous and discrete. In fact the QMUPL model can be understood as a scaling limit of the GRW process (Dürr et al., 2011) [the collapse frequency goes to infinity and the spread $r_{C}$ goes to zero in such a way that their product remains a constant].

Part of the experimental effort on testing quantum mechanics, discussed in detail in Section IV, is devoted to testing the validity of equations such as (17) above, and measuring / setting bounds on the rate constant $\lambda$ and the length scale $r_{C}$.

## Underlying Theories

Phenomenological models of dynamical wave-function collapse propose an ad hoc modification of quantum mechanics, albeit retaining certain features such as normpreservation and no superluminal propagation. In principle, there should be strong underlying theoretical reasons which make a compelling case for a modified quantum theory, rendering the phenomenological models inevitable. Here we mention three different theoretical developments in this connection, two of which arise from attempts to remove one or the other fundamental incompleteness in the formulation of quantum theory, and the third investigates how gravity might play an effective role in wave-vector reduction.

## Trace Dynamics

Classical mechanics is supposed to be a limiting case of quantum theory. And yet, in its canonical formulation, quantum theory assumes a prior knowledge of classical dynamcs! In order to 'quantize' a system, one should know the classical configuration variables and their conjugate momenta, and one should first know the Hamiltonian or
the action [for a path-integral formulation] of the classical system. This is unsatisfactory. In the canonical formulation, one then proposes canonical commutation relations such as $[\mathbf{q}, \mathbf{p}]=i \hbar$ in an ad hoc manner. Why should these be the relations, unless one already knows that they lead to results which match with experiments? It would be desirable to derive quantum theory from a starting point which is not classical mechanics, and then obtain classical mechanics as an approximation [and explain quantum measurement in the process]. The theory of Trace Dynamics developed by Adler and collaborators does well in progressing towards this goal Adler, 1994, 2004, Adler and Millard, 1996).

Trace Dynamics [TD] assumes that the underlying theory is a classical dynamics of Grassmann matrices, living on a given spacetime. However, this classicality does not mean that TD is a 'hidden variables' theory for the eventual description is at an averaged level, where no reference is made to the matrices which have been coarse-grained over. The matrices satisfy the standard Lagrangian and Hamiltonian Dynamics, but as a consequence of global unitary invariance, the theory possesses a remarkable additional conserved charge, not present in point-particle mechanics. This is the Adler-Millard charge
(Adler and Millard, 1996)

$$
\begin{equation*}
\tilde{C}=\sum_{i}\left[q_{i}, p_{i}\right]-\sum_{j}\left\{q_{j}, p_{j}\right\} \tag{18}
\end{equation*}
$$

where the first sum is over commutators of bosonic matrices, and the second sum is over anti-commutators of fermionic matrices. [See Section III for details]. This conserved charge, which has the dimensions of action, plays a central role in the emergence of quantum theory at a coarse-grained level.

Assuming that these matrix degrees of freedom are at a level sufficiently 'microscopic' [e.g. at the Planck scale] that we do not observe them in our routine laboratory experiments, a statistical thermodynamics of this matrix dynamics is constructed. An equipartition theorem for the thermodynamically averaged quantities is derived, which results in the Adler-Millard charge being uniformly distributed across the averaged commutators, each of which is assumed to equal Planck's constant. This is the origin of the quantum commutation relations. As a consequence of the assumed invariance of thermodynamic averages under constant shifts in phase space, a Ward identity is derived, which under suitable assumptions shows that the thermally averaged $q$ 's and $p$ 's satisfy Heisenberg equations of motion. A relativistic quantum field theory is arrived at, and a non-relativistic Schrödinger equation holds in the finite particle limit. Thus quantum the-
ory is shown to emerge as the thermodynamic approximation to an underlying classical dynamics of Grassmann matrices possessing a global unitary invariance.

Perhaps the greatest asset of TD is to be able to go beyond this stage and address the quantum measurement problem in a natural manner. Quantum theory emerges in the thermodynamic approximation of the statistical mechanics of the underlying matrix mechanics. Next, it is pertinent to consider the impact of Brownian motion fluctuations - remarkably these modify the Schrödinger equation and provide the necessary stochastic element for the collapse process to operate, and for the origin of probabilities. Subject to certain crucial assumptions for which one would eventually like to find a theoretical basis, the modified Schrödinger equation is a nonlinear and non-unitary [but norm-preserving] stochastic equation of the type used in the CSL model. In this way, Trace Dynamics, through its thermodynamic limit and the associated statistical fluctuations provides a theoretical underpinning for the phenomenological collapse models.

TD is perhaps the most well-developed underlying theory one has at present for collapse phenomenology. Hence in Section III we give a detailed presentation of the physics and mathematics of TD, leading to wavevector reduction, and we also point out the
open problems of TD which remain to be addressed.

Quantum theory without classical spacetime
Quantum theory requires an external classical time for describing evolution. This is of course so obvious and essential that it is almost never stated explicitly! However this dependence on an external classical time is perhaps the greatest incompleteness of quantum theory. Such a time is part of a classical spacetime geometry which is produced by classical matter fields according to the laws of general relativity. But classical matter fields are a limiting case of quantum fields. If there were no classical fields in the Universe, but only fields subject to quantum fluctuations, there will be no definite metric available to describe the spacetime geometry. An argument due to Einstein, known as the Einstein hole argument (Christian, 1998) then implies that if the metric is subject to quantum fluctuations, there is no longer available an underying classical spacetime manifold. It is then not possible to describe quantum evolution.

We see once again that via its dependence on external time, quantum theory depends on its classical limit [the required presence of a Universe dominated by classical matter]. This is unsatisfactory from a fundamental point of view, and hence there must exist an equivalent reformulation of quantum
theory which does not refer to classical time. Such a reformulation can be shown to be the limiting case of a non-linear theory, with the non-linearity becoming important at the Planck mass scale. The non-linearity is possibly stochastic, and could have implications for resolution of the quantum measurement problem. Tentative heuristic discussions towards this investigation have been given in (Singh, 2006, 2009). A possible intriguing implication is that the Planck constant $\hbar$ is no longer strictly a constant, but runs with the mass / number of degrees of freedom in the system under study. $\hbar$ is the zero-mass limit, and as the mass of the object being studied goes to infnity, the effective Planck's constant $\hbar_{\text {eff }}$ goes to zero, one possible form of the mass-dependence being

$$
\begin{equation*}
\hbar_{e f f}=\frac{\hbar}{1+m / m_{P l}} \tag{19}
\end{equation*}
$$

where $m_{P l}$ is the Planck mass. It is interesting that a laboratory experiment has been proposed (Pikovski et al., 2012) which can test quantum commutation relations directly and hence in effect the possible departure of Planck's consant from its bare value $\hbar$.

A detailed systematic program to develop a formulation of quantum theory without classical time and to study its impact on quantum measurement has recently been begun, and is qualitatively described in (Singh, 2011). The key symmetry principle here is
that basic laws should be invariant under coordinate transformations of non-commuting coordinates. The motivation being that if quantum fluctuations destroy a classical spacetime manifold, a possible replacement for ordinary spacetime could be a noncommutative spacetime. This approach proposes to generalize Trace Dynamics by raising time, and space, to the level of matrices [operators]. This has been done in Lochan and Singh, 2011) and it has been shown that by defining a non-commutative space-time metric as a Trace over the space-time operators, a Poincaré invariant dynamics can be constructed. We call this a generalized Trace Dynamics. Evolution is described with respect to the scalar constructed by taking Trace over the non-commutative metric - this is the ana$\log$ of the ordinary proper time.

The next step is to contstruct, a la TD, a statistical mechanics for this generalized matrix dynamics and obtain the equilibrium thermodynamic approximation - this yields a generalized quantum theory which has an energy-time commutation relation and a generalized Schrödinger equation with an operator time as one of the configuration variables. This is the sought for reformulation of quantum theory which does not refer to an external classical time (Lochan et al., 2012). If the Universe is dominated by macroscopic objects, the consideration of Brownian motion
fluctuations should yield position localization and the concurrent emergence of a classical space-time. This is the classical Universe, dominated by classical macroscopic objects and in possession of a classical spacetime. This Universe has a 'sprinkling' of quantum fields and non-relativistic quantum systems. On the backdrop of this classical Universe one can postulate standard quantum theory [now that an external time is given] and then proceed to implement the program of Trace Dynamics to derive quantum dynamics from matrix mechanics, for this 'sprinkling' of quantum matter fields on the classical spacetime backgound, and to resolve the attendant measurement problem.

The program described here aims to address a limitation of Trace Dynamics - a matrix treatment for matter fields while leaving spacetime as having point structure, thus leaving spacetime untouched. We regard such a limitation as one which should be addressed - in the process we see that removing time from quantum theory drives us to a starting point [generalized Trace Dynamics] whose eventual outcome is a possible resolution for the measurement problem. We thus wish to assert that there is a deep connection between the problem of time in quantum theory and the measurement problem in quantum theory. Addressing the former will possibly compel us to consider a modification
of quantum theory and that modification will have a bearing on the measurement problem.

Since this work is at present in an early stage of development, we will not discuss it any further in the remainder of the review.

## Gravity induced wave-function collapse

The fact that the fundamental mass scale, Planck mass $M_{P l}=(\hbar c / G)^{1 / 2} \sim 10^{-5}$ grams is not far from the scale where the micro to macro transition takes place has often intrigued some physicists. Mass seems to have something to do with deciding which objects are quantum, and which are classical, and mass also produces gravity. Could gravity thus play some role in causing wave-function collapse, and in localization of macro- objects? For instance, Feynman (Feynman et al., 1995) wrote
" I would like to suggest that it is possible that quantum mechanics fails for large distances and large objects. Now, mind you, I do not say that quantum mechanics does fail at large distances, I only say that it is not inconsistent with what we do know. If this failure of quantum mechanics is connected with gravity, we might speculatively expect this to happen for masses such that $G M^{2} / \hbar c=1$, or $M$ near $10^{-5}$ grams, which corresponds to some $10^{18}$ particles."

The idea that gravity might somehow be responsible for wave-function collapse has been seriously pursued by Karolyhazy and
collaborators (Karolyazy, 1966; Karolyhazy et al., 1986), by (Diósi, 1987), and by Penrose (Penrose, 1996). Penrose's proposal is also the subject of an important ongoing experiment aimed at testing it (Marshall et al., 2003). These issues will be discussed in Section III.

## Experimental Tests

The Copenhagen interpretation was a need of the times when it was proposed: pioneering experiments were being carried out for atomic systems. The measuring apparatus was a classical object, and the Born probability rule had to be invoked to explain the random outcomes of measurements. For some, this dual aspect of quantum theory - unitary evolution followed by wavepacket reduction - was the 'truth' in quantum theory; this is how nature is. For others, this was completely unacceptable, and reinterpretations and new mathematical formulations such as many-worlds, Bohmian mechanics, and decoherent histories, were developed. However, the idea that quantum theory may be an approximation to a holistic theory which better explains both the unitary and reductionist aspects as limits of a unified mathematical description has taken shape only over the last three deacdes or so. And yet, none other than Einstein himself (Schilpp, 1949) saw it this way early on, and
had this to say about quantum theory:
'... it would, within the framework of future physics, take an approximately analogous position to that of statistical mechanics within the framework of classical mechanics'.

In the light of the theory of Trace Dynamics and models of spontaneous collapse, these words are prophetic. These modern ideas suggest the emergence of probabilities as a consequence of thermodynamic averaging in a deterministic theory, and the related significance of stochastic fluctuations. Above all, their predictions for results of experiments differ from the predictions of quantum theory. The difference will be far too small to be detectable for an atomic system, but starts becoming significant as the size of the system is increased. The best example of an experiment which could detect such a difference is double slit interference. If an object of mass $m$ is directed at a suitably prepared double slit, with appropriate slit width and separation between the slits, quantum theory predicts that an interference pattern will be seen on the screen no matter what the value of $m$. Not so, say collapse models. According to these models, the superposition state created after the object has passed the slits lasts only for a finite time $\tau$, where $\tau$ decreases with increasing $m$, and its value can be calculated precisely from a given theoretical model. Thus, according to these mod-
els, if the time of travel from the slits to the screen is greater than $\tau$, suerposition will break down before the screen is reached, and no interference pattern will be seen. This is perhaps the cleanest confrontation that spontaneous collapse and gravity collapse models make with experiment. A successful diffraction experiment in the right mass domain will irrefutably confirm or rule out these models.

One should of course stay cautioned against assuming that quantum theory will be successful through and through, and that interference will be seen for all values of $m$. The fact that a theory is extremely successful in one part of the parameter space should not be taken as guarantee that it will continue to be successful in a different part of the parameter space - in the present instance the absence of macroscopic superpositions already provides reason for caution. And there are historical examples of long-standing successful theories eventually turning into approximations to more general theories when their extrapolation into a new part of the parameter space failed to be confirmed by experiment: (i) classical mechanics became an approximation to special relativity at speeds close to the speed of light, (ii) quantum dynamics took over from classical dynamics in the atomic domain, and (iii) Newton's inverse square law of gravitation was replaced by the laws of general relativity for strong gravita-
tional fields.
Interference experiments with matter have a fascinating history, with a quest developing over decades to test superposition using larger and larger objects. Over eighty years have passed since the classic experiment by Davisson and Germer in 1927 where interference was demonstrated with electrons. Landmarks on the way included cofirmation of interference for Helium [1930] and neutrons [1988]. A great modern breakthrough came in 1999 when interference was demonstrated for $C_{60}$ in the famous fullerene experiment (Arndt et al., 1999). [There seems to be prevalent a popular belief in some quarters that the discovery of quantum superposition in a molecule as 'large' as fullerene means the end of all theories which predict breakdown of superposition for large systems. This of course is not true - breakdown of superposition and the quantum-to-classical transition is expected around $10^{6} \mathrm{amu}$ to $10^{9} \mathrm{amu}$.] This opened the door for larger molecules, and today, a decade later, interference has been demonstrated for molecules with 7,000 nucleons (Gerlich et al., 2011). Proposed future interferometry experiments plan to push the limit to macromolecules with a million nucleons and beyond, going up to molecules with a 100 million nucleons. Doing so involves overcoming great technological challenges (Hornberger et al., 2011), and there are many or-
ders of magnitudes in the mass scale yet to be covered. But we certainly live in exciting times where predictions of collapse models and gravity based models are being testsed by these experiments, and constraints are being put on model parameters. See Section IV for details.

Also, this is perhaps a good place to clear another misconception regarding the domain over which quantum mechanics has been tested. Various remarkable macroscopic internal states have been achieved experimentally, in which an enormous collection of internal degrees of freedom behave as a collective one-particle coherent state. We have in mind of course systems such as superconductors, superfluids and Bose-Einstein condensates. The existence of such states however does not explain why macroscopic objects are not found in superposition of position states. Quantum mechanics may yet have to be modified so that the modified theory can explain the absence of position superposition, but the modified theory will certainly continue to successfully explain a collective phenomenon such as superconductivity. In other words, the discovery of superconductivity does not solve / trivialize the Schrödinger cat paradox!

Apart from direct laboratory experiments, collapse model parameters are also constrained by their effect on known measurements. Section IV discusses the various
experimental tests of the phenomenological models.

## C. Plan and outline of the article

Sections II, III and IV are the main parts of the review. Sec. II reviews the phenomenological models of Spontaneous Collapse which explain wave-function collapse and absence of macroscopic superpositions, via a stochastic non-linear modification of the Schrödinger equation. Sec. III gives a review of Trace Dynamics and gravity-inducedcollapse as possible underlying theories for the phenomenology discussed in Sec. II. Sec. IV is a review of the techniques and results of ongoing and planned experiments which are testing the proposed phenomenological models. Sec. V provides a critique of the current understanding on the theoretical and experimental front, and lists open problems. The Appendix in Sec. VI provides a pedagogical introduction to stochastic processes and stochastic differential equations.

Sec. II begins by introducing spontaneous collapse, and recalls the various collapse models that have been proposed. The original GRW model is then introduced. This is then followed by a detailed review of the QMUPL model, which is applied to show how stochasticity induces collapse, and how the Born probability rule is derived. The simplest col-
lapse models have been generalized to include dissipation and colored noise, and two subsections are devoted to describing these generalizations. The possible fundamental origin of the noise field is briefly discussed. The next sub-section discusses the most important collapse model, i.e. the CSL model, and its generalizations. Lastly, the current understanding of the numerical values of the two parameters of the collapse model is reviewed.

Sec. III reviews Adler's Trace Dynamics as a candidate fundamental theory for Spontaneous Localization. The fundamental matrix degrees of freedom of the theory are introduced, and their dynamics described. The conserved charges of the theory, including the all-important Adler-Millard charge, are derived. This is followed by the construction of the statistical mechanics and the canonical ensemble for thermodynamic equilibrium for the theory. Following this, an important Ward identity, which is an analog of the equipartition theorem, is proved. It is shown how the commutation relations for quantum theory, and the Schrödinger equation, emerge at this coarse-grained level, from the microscopic theory. Finally, consideration of fluctuations described by Brownian motion leads to generalization from the Schrödinger equation to the stochastic non-linear Schrödinger equation, which makes contact with the CSL model. Subsequent sections describe the
gravity based models for collapse, based on the work of Karolyhazy et al., Diósi, and the work of Penrose.

Sec. IV on Experimental Tests starts by discussing the basics of the collapse theory necessary for performing and interpreting the diffraction experiments with macromolecules. Matter-wave interferometry and optomechanics experiments with mechanical cantilevers are reviewed in detail. Cavity optomechanics with micro-spheres and nanoparticles is discussed, followed by a review of new developments which combine optical tweezing techniques with near-field matter-wave interferometry. The challenges proposed to these experiments by various kinds of decoherence are considered. The current bounds on collapse model parameters coming from the diffraction experiments and from other measurement processes are summarized.

## II. SPONTANEOUS COLLAPSE MODELS

## A. Introducing spontaneous collapses

Quantum Mechanics, in its standard textbook formulation, refers only to the outcomes of measurements, but it has nothing to say about the world as it is, independently of any measurement or act of observation. This
is a source of serious difficulties, which have been clearly elucidated e.g. by J. Bell (Bell, 1990): It would seem that the theory is exclusively concerned about 'results of measurements', and has nothing to say about anything else. What exactly qualifies some physical systems to play the role of 'measurer'? Was the wavefunction of the world waiting to jump for thousands of millions of years until a single-celled living creature appeared? Or did it have to wait a little bit longer, for some better qualified system ... with a Ph.D.?

Measuring devices, like photographic plates and bubble chambers, are very sophisticated and highly structured physical systems, which anyhow are made of atoms; we then expect them to be ultimately described in quantum mechanical terms by means of the Schrödinger equation. What else should we expect, taking into account that physicists are trying to describe even the entire universe quantum mechanically? But if we describe measurements in this way, then the theory does not predict any definite outcome, at the end of the process. The Schrödinger equation is linear, the superposition principle holds, and it does so in such a way that all possible outcomes are there simultaneously in the wave function, but none of them is selected as the one which actually occurs. Yet, if we perform a measurement, we always get a definite outcome. So we have a problem
with Quantum Mechanics.
Continuing quoting Bell: If the theory is to apply to anything but highly idealized laboratory operations, are we not obliged to admit that more or less 'measurement-like' processes are going on more or less all the time, more or less everywhere? Do we not have jumping then all the time?

The basic idea behind the dynamical reduction program is precisely this: spontaneous and random collapses of the wave function occur all the time, for all particles, whether isolated or interacting, whether they form just an atom or a complex measuring device. Of course, such collapses must be rare and mild for microscopic systems, in order not to alter their quantum behavior as predicted by the Schrödinger equation. At the same time, their effect must add up in such a way that, when thousands of millions of particles are glued together to form a macroscopic system, a single collapse occurring to one of the particles affects the global system. We then have thousands of millions of such collapses acting very frequently on the macro-system, which together force its wave function to be very rapidly well-localized in space.

On the mathematical level, the program is accomplished by modifying the Schrödinger evolution, introducing new terms having the following properties:

- They must be non-linear: The new dynamics must break the superposition principle at the macroscopic level and guarantee the localization of the wave function of macro-objects.
- They must be stochastic because, when describing measurement-like situations, the dynamics must explain why the outcomes occur randomly; more than this, it must explain why they are distributed according to the Born probability rule.
- There must be an amplification mechanism according to which the new terms have negligible effects on the dynamics of microscopic systems but, at the same time, their effect becomes very strong for large many-particle systems such as macroscopic objects, in order to recover their classical-like behavior.
- They must not allow for superluminal signalling, as one would like to preserve the causal structure of spacetime.

Looking carefully at these requirements, one soon realizes that they are very demanding: there is no reason beforehand, that they can be consistently fulfilled. One of the greatest merits of collapse models is to have shown that this program can be implemented in a consistent and satisfactory way.

## B. The zoo of collapse models

In the literature, different collapse models have been proposed. A first characterization depends on the choice of the collapse operators, i.e. on the basis on which the wave function is localized. Some models induce the collapse in the energy basis (Adler, 2002, 2004; Adler et al., 2001; Adler and Brun, 2001; Adler and Horwitz, 2000), others in the momentum basis (Benatti et al., 1988), or the spin basis (Bassi and Ippoliti, 2004). However, only models which collapse in the position basis make sure that different macroscopic superpositions rapidly collapse towards localized states. To understand this, one can think of a superposition of two spatially separated states of a macroscopic object, which have the same (or very similar) energy. In this case, an energy-based collapse model would not be able to collapse the superposition fast enough, because the superposition in energy is null or negligible. Such a model would not be able to guarantee that macro-objects always occupy a definite position in space. Only space collapse models make sure that macroscopic objects always behave classically, and therefore we will consider only them in the following.

Space collapse models can be conveniently grouped depending on the properties of the noise, which is responsible for the collapse.

A first distinction is between white and nonwhite models. In white-noise models, the collapse-noise is assumed to be a Wiener process in time, and the resulting evolution is Markovian. All frequencies of the noise contribute to the collapse with the same weight. Examples of collapse models of this type are the GRW model (Ghirardi et al., 1986), the CSL model (Ghirardi et al., 1990c), the QMUPL model (Bassi, 2005; Diósi, 1989). In non-white noise models, the collapse-noise is taken to be a generic Gaussian noise, with mean equal to zero, and a generic correlation function. The corresponding dynamics turns out to be non-Markovian, and they are more difficult to analyze. Models of this kind are the non-Markovian QMUPL model (Bassi and Ferialdi, 2009a b), and the non-Markovian CSL model (Adler and Bassi, 2007, 2008). General non-Markovian collapse models have been discussed in (Bassi and Ghirardi, 2002; Diósi et al., 1998; Pearle, 1993, 1996).

A second distinction is between infinite temperature and finite temperature models. In the first type of models, the collapsenoise acts like a reservoir at infinite temperature. The wave function collapses, but at the same time the energy of the quantum system increases steadily; no dissipative effects are taken into account. This is a well-known feature of collapse models. Mathematically,
these models are characterized by the fact that the wave function and the collapse-noise are coupled through the position operator only. The GRW model, the (Markovian and non-Markovian) CSL model, the (Markovian and non-Markovian) QMUPL model all belong to this group. In the finite temperature models instead, the collapse-noise behave like a reservoir at finite temperature. The wave function still collapses, but now dissipative terms are included (through a position and momentum coupling between the wave function and the noise), which thermalize any quantum systems to the temperature of the noise. The only such model so far available is the non-dissipative QMUPL model (Bassi et al., 2005b), though also the other models can be generalized in this sense. Recently, the QMUPL model has been generalized in order to include both non-Markovian and dissipative effects (Ferialdi and Bassi, 2011).

A final distinction is between first quantized models and second quantized models. Models of the first type consider only system of distinguishable particles; the GRW model, the QMUPL model and their non-Markovian and/or dissipative generalization belong to this group. Models of the second type are formulated in the language of quantum field theory and include systems of identical particles. The Tumulka-GRW model (Tumulka, 2006a) and the CSL model belong to this group.

Some comments are in order. The first one is that all space-collapse model are qualitatively equivalent: they all induce the collapse of the wave function in space, and the collapse is faster, the larger the system. Of course, they can differ also in a significant way in the technical details, as we will see. The second comment refers to the nature of the stochastic character of the collapse process. One way to look at itwhich corresponds to the original attitude towards these models-is that nature is intrinsically stochastic, therefore stochastic differential equations are the natural type of equations for describing the dynamics of microscopic systems. A new way to look at it is to assume that there is a random field, filling space, which couples to quantum matter in a non-standard way, and is responsible for the collapse of the wave function. The new terms in the modified Schrödinger equation are meant to describe such a coupling. Since this noise fills the whole space, most likely it has a cosmological origin. According to this scenario, the physically most reasonable collapse model is a quantum field theoretical model, where the collapsing field is "cosmologically reasonable", e.g. it has a typical cosmological correlation function and a typical cosmological temperature. This could be the case for the colored-noise and dissipative CSL model, which however has not been formu-
lated yet. What one can do, is to extrapolate predictions from the other models already available. Therefore, in the following we will focus our attention on two of the above mentioned models: the CSL model, the one that more closely resembles the physically most reasonable model; and the QMUPL model, which is less physical, but has already been generalized in order to include dissipations, besides colored noises, and is relatively easy to analyze mathematically.

A third comment is about the origin of the noise field. The important thing to bear in mind, is that this field cannot be a standard quantum field, otherwise we would fall back in the realm of standard quantum mechanics, with the superposition principle and the measurement problem. This field couples to quantum matter though an anti-hermitian and non-linear coupling. The most intriguing guess is that this noise has a gravitational origin. In fact, a gravitational background is part of the standard cosmological scenario; gravity is non-linear; gravity has not been successfully quantized yet, and we do not know today what shape a quantum theory of gravity will eventually take. Gravity-induced collapse models have been formulated in the literature, and we will discuss them in Sec. III.B.

A fourth comment is about the relativistic extension of collapse models. All models pre-
viously listed are non-relativistic. Their generalization to relativistic quantum field theories has not been successful. The reason is very simple to understand: the collapse of the wave function is an instantaneous process, or at least faster than light. This is a necessary requirement, in order to reproduce non-local quantum correlations encoded in Bell inequalities ( $\overline{\mathrm{Bell}}$, 1987), which have been verified experimentally. An instantaneous collapse process is not welcome in a relativistic framework, hence the difficulty in formulating relativistic collapse models. We will discuss this issue in Sec. II.I

## C. The GRW model

In order to appreciate how collapse models work, and what they are able to achieve, we briefly review the GRW model, the first consistent model proposed. Though it is not expressed in terms of a compact stochastic differential equation, it has the advantage of being physically very intuitive.

Let us consider a system of $N$ particles which, only for the sake simplicity, we take to be scalar; the GRW model is defined by the following postulates:

States. The state of the system is represented by a wave function $\psi\left(\mathbf{x}_{1}, \mathbf{x}_{2}, \ldots \mathbf{x}_{N}\right)$ belonging to the Hilbert space $\mathcal{L}^{2}\left(\mathbf{R}^{3 N}\right)$.

Dynamics. At random times, the wave function experiences a sudden jump of the form:

$$
\begin{align*}
& \psi_{t}\left(\mathbf{x}_{1}, \mathbf{x}_{2}, \ldots \mathbf{x}_{N}\right) \longrightarrow \\
&  \tag{20}\\
& \frac{L_{n}(\mathbf{x}) \psi_{t}\left(\mathbf{x}_{1}, \mathbf{x}_{2}, \ldots \mathbf{x}_{N}\right)}{\left\|L_{n}(\mathbf{x}) \psi_{t}\left(\mathbf{x}_{1}, \mathbf{x}_{2}, \ldots \mathbf{x}_{N}\right)\right\|},
\end{align*}
$$

where $\psi_{t}\left(\mathbf{x}_{1}, \mathbf{x}_{2}, \ldots \mathbf{x}_{N}\right)$ is the state vector of the whole system at time $t$, immediately prior to the jump process. $L_{n}(\mathbf{x})$ is a linear operator which is conventionally chosen equal to:

$$
\begin{equation*}
L_{n}(\mathbf{x})=\frac{1}{\left(\pi r_{C}^{2}\right)^{3 / 4}} e^{-\left(\mathbf{q}_{n}-\mathbf{x}\right)^{2} / 2 r_{C}^{2}} \tag{21}
\end{equation*}
$$

where $r_{C}$ is a new parameter of the model which sets the width of the localization process, and $\mathbf{q}_{n}$ is the position operator associated to the $n$-th particle of the system; the random variable $\mathbf{x}$ corresponds to the place where the jump occurs. Between two consecutive jumps, the state vector evolves according to the standard Schrödinger equation.

The probability density for a jump taking place at the position $\mathbf{x}$ for the $n$-th particle is given by:

$$
\begin{equation*}
p_{n}(\mathbf{x}) \equiv\left\|L_{n}(\mathbf{x}) \psi_{t}\left(\mathbf{x}_{1}, \mathbf{x}_{2}, \ldots \mathbf{x}_{N}\right)\right\|^{2} \tag{22}
\end{equation*}
$$

and the probability densities for the different particles are independent.

Finally, it is assumed that the jumps are distributed in time like a Poissonian process with frequency $\lambda$; this is the second new parameter of the model.

The standard numerical values for $r_{C}$ and $\lambda$ are:

$$
\begin{equation*}
\lambda \simeq 10^{-16} \mathrm{sec}^{-1}, \quad r_{C} \simeq 10^{-5} \mathrm{~cm} . \tag{23}
\end{equation*}
$$

We will come back on the issue of numerical value of these parameters in Sec. II.J.

Ontology. In order to connect the mathematical formalism with the physical world, one needs to provide an ontology, which is rather straightforward for collapse models. Let $m_{n}$ be the mass associated to the $n$-th "particle" of the system (one should say: to what is called "a particle", according to the standard terminology); then the function:

$$
\begin{aligned}
\rho_{t}^{(n)}\left(\mathbf{x}_{n}\right) \equiv & m_{n} \int d^{3} x_{1} \ldots d^{3} x_{n-1} d^{3} x_{n+1} \\
& \ldots d^{3} x_{N}\left|\psi_{t}\left(\mathbf{x}_{1}, \mathbf{x}_{2}, \ldots \mathbf{x}_{N}\right)\right|^{2}(24)
\end{aligned}
$$

represents the density of mass Ghirardi et al., 1995) of that "particle" in space, at time $t$.

These are the axioms of the GRW model: as we see, words such as 'measurement', 'observation', 'macroscopic', 'environment' do not appear. There is only a universal dynamics governing all physical processes, and an ontology which tells how the physical world is, according to the model, independently of any act of observation.

The GRW model, as well as the other dynamical reduction models which have appeared in the literature, has been extensively studied (see (Bassi and Ghirardi, 2003)
and (Pearle, 1999a) for a review on this topic); in particular-with the numerical choice for $\lambda$ and $r_{C}$ given in (23) - the following three important properties have been proved, which we will state in more quantitative terms in the following section:

- At the microscopic level, quantum systems behave almost exactly as predicted by standard Quantum Mechanics, the differences being so tiny that they can hardly be detected with present-day technology.
- At the macroscopic level, wave functions of macro-objects are almost always very well localized in space, so well localized that their centers of mass behave, for all practical purposes, like point-particles moving according to Newton's laws.
- In a measurement-like situation, e.g. of the von Neumann type, the GRW model reproduces-as a consequence of the modified dynamics-both the Born probability rule and the standard postulate of wave-packet reduction.

In this way, models of spontaneous wave function collapse provide a unified description of all physical phenomena, at least at the nonrelativistic level, and a consistent solution to the measurement problem of Quantum Mechanics.

It may be helpful to stress some points about the world-view provided by the GRW model, and collapse models in general. According to the ontology given by the third axiom, there are no particles at all in the theory! There are only distributions of masses which, at the microscopic level, are in general quite spread out in space. An electron, for example, is not a point following a trajectory-as it would be in Bohmian Mechanics-but a wavy object diffusing in space. When, in a double-slit experiment, it is sent through the two apertures, it literally goes through both of them, as a classical wave would do. The peculiarity of the electron, which qualifies it as a quantum system, is that when we try to locate it in space, by making it interact with a measuring device, e.g. a photographic film, then, according to the collapse dynamics, its wave function very rapidly shrinks in space till it gets localized to a spot, the spot where the film is impressed and which represents the outcome of the position measurement. Such a behavior, which is added $a d$ hoc in the standard formulation of Quantum Mechanics, is a direct consequence of the universal dynamics of the GRW model.

Also macroscopic objects are waves; their centers of mass are not mathematical points, rather they are represented by some function defined throughout space. But macro-objects have a nice property: according to the GRW
dynamics, each of them is always almost perfectly located in space, which means that the wave functions associated to their centers of mass are appreciably different from zero only within a very tiny region of space (whose linear extension is of order $10^{-14} \mathrm{~m}$ or smaller, as we shall see), so tiny that they can be considered point-like for all practical purposes. This is the reason why Newton's mechanics of point particles is such a satisfactory theory for macroscopic classical systems.

Even though the GRW model contains no particles at all, we will keep referring to micro-systems as 'particles', just as a matter of convenience.

Though the collapse dynamics is expressed entirely in terms of the wave function, not of the density matrix, in order to eliminate any possible ambiguity about the nature of the collapse, it is often convenient to look at the collapse dynamics for the density matrix, to analyze specific features of the model. The 1particle master equation of the GRW model takes the form (Ghirardi et al., 1986):

$$
\begin{equation*}
\frac{d}{d t} \rho(t)=-\frac{i}{\hbar}[H, \rho(t)]-T[\rho(t)] \tag{25}
\end{equation*}
$$

where $H$ is the standard quantum Hamiltonian of the particle, and $T[\cdot]$ represents the effect of the spontaneous collapses on the particle's wave function. In the position repre-
sentation, this operator becomes:
$\langle\mathbf{x}| T[\rho(t)]|\mathbf{y}\rangle=\lambda\left[1-e^{-(\mathbf{x}-\mathbf{y})^{2} / 4 r_{C}^{2}}\right]\langle\mathbf{x}| \rho(t)|\mathbf{y}\rangle$.

As expected, the effect of the spontaneous collapse is to suppress the off-diagonal elements of the density matrix, with a rate proportional to $\lambda$, depending also on the distance between the off-diagonal elements: larger superpositions are suppressed faster than closer ones.

The many particle master equation is the generalization of Eqn. (25), where an operator $T_{i}[\cdot], i=1,2, \ldots N$ appears for each particle. For ordinary matter-and with a good approximation-one can separate the center-of-mass motion from the internal motion (Ghirardi et al., 1986). The reduced density matrix for the internal motion obeys the standard Schrödinger equation, while that for the center of mass is equivalent to Eqn. (25), where now the collapse rate entering the definition of the operator $T[\cdot]$ is $N \lambda$, with $N$ the total number of particles making up the object. This is a manifestation of the amplification mechanics, perhaps the most important feature of collapse models: the wave function of an object collapses with a rate which is proportional to the size of the system. This is the mathematical reason why collapse models can accommodate both the quantum dynamics of microscopic systems (negligible collapse
rate) and the classical dynamics of macroscopic systems (fast collapse) within one unified dynamical principle.

## D. The QMUPL model

We now focus our attention on the QMUPL (Quantum Mechanics with Universal Position Localizations) model. As previously anticipated, the reason is that this model has the virtue of being both physically realistic, though very simplified compared to the more realistic CSL model, and mathematically simple enough to be analyzed in great detail. The axioms defining this model are the same as those of the GRW model, with the only difference that the dynamics is described by a stochastic differential equation. The one-particle equation takes the form:

$$
\begin{align*}
d \psi_{t}= & {\left[-\frac{i}{\hbar} H d t+\sqrt{\lambda}\left(q-\langle q\rangle_{t}\right) d W_{t}\right.} \\
& \left.-\frac{\lambda}{2}\left(q-\langle q\rangle_{t}\right)^{2} d t\right] \psi_{t} \tag{27}
\end{align*}
$$

where $q$ is the position operator of the particle, $\langle q\rangle_{t} \equiv\left\langle\psi_{t}\right| q\left|\psi_{t}\right\rangle$ is the quantum expectation, and $W_{t}$ is a standard Wiener process. For simplicity, we work in only one spatial dimension, the generalization to three dimensions being straightforward. The collapse constant $\lambda$ sets the strength of the collapse mechanics, and it is chosen proportional to the mass $m$ of the particle according to the
formula:

$$
\begin{equation*}
\lambda=\frac{m}{m_{0}} \lambda_{0} \tag{28}
\end{equation*}
$$

where $m_{0}$ is the nucleon's mass and $\lambda_{0} \simeq$ $10^{-2} \mathrm{~m}^{-2} \mathrm{~s}^{-1}$ is the reference value for the collapse strength (Bassi, 2005). We will come back on the numerical values of the collapse parameter in Sec. II.J. The generalization for a many-particle system can be easily obtained by considering the position operator $q_{i}$ of every particle, each coupled to a different Wiener process $W_{t}^{(i)}$. The structure remains the same, with a sum to include the contribution to the collapse coming from each particle.

As we expect, Eqn. (27) contains both non-linear and stochastic terms, which are necessary to induce the collapse of the wave function. In order to see this, let us consider a free particle $\left(H=p^{2} / 2 m\right)$, and a Gaussian state:

$$
\begin{equation*}
\psi_{t}(x)=\exp \left[-a_{t}\left(x-\bar{x}_{t}\right)^{2}+i \bar{k}_{t} x+\gamma_{t}\right] \tag{29}
\end{equation*}
$$

It is not too difficult to show that $\psi_{t}(x)$ is solution of Eqn. (27), provided that the parameters solve appropriate stochastic differential equations (Bassi, 2005). In particular, the equations for $a_{t}$ which control the spread both in position and momentum, for the mean position $\bar{x}_{t}$ and the mean momen-
tum $\bar{k}_{t}$ ar ${ }^{1}$

$$
\begin{align*}
d a_{t} & =\left[\lambda-\frac{2 i \hbar}{m}\left(a_{t}\right)^{2}\right] d t,  \tag{30}\\
d \bar{x}_{t} & =\frac{\hbar}{m} \bar{k}_{t} d t+\frac{\sqrt{\lambda}}{2 a_{t}^{\mathrm{R}}} d W_{t},  \tag{31}\\
d \bar{k}_{t} & =-\sqrt{\lambda} \frac{a_{t}^{\mathrm{I}}}{a_{t}^{\mathrm{R}}} d W_{t} . \tag{32}
\end{align*}
$$

Eqn. (30) is deterministic and easy to solve. The spread in position and momentum:

$$
\begin{align*}
\sigma_{q}(t) & =\frac{1}{2} \sqrt{\frac{1}{a_{t}^{\mathrm{R}}}} \\
\sigma_{p}(t) & =\hbar \sqrt{\frac{\left(a_{t}^{\mathrm{R}}\right)^{2}+\left(a_{t}^{\mathrm{T}}\right)^{2}}{a_{t}^{\mathrm{R}}}} \tag{33}
\end{align*}
$$

are given by the following analytical expressions:

$$
\sigma_{q}=\sqrt{\frac{\hbar}{m \omega} \frac{\cosh \left(\omega t+\varphi_{1}\right)+\cos \left(\omega t+\varphi_{2}\right)}{\sinh \left(\omega t+\varphi_{1}\right)+\sin \left(\omega t+\varphi_{2}\right)}}
$$

$$
\begin{equation*}
\sigma_{p}=\sqrt{\frac{\hbar m \omega}{2} \frac{\cosh \left(\omega t+\varphi_{1}\right)-\cos \left(\omega t+\varphi_{2}\right)}{\sinh \left(\omega t+\varphi_{1}\right)+\sin \left(\omega t+\varphi_{2}\right)}} \tag{34}
\end{equation*}
$$

with:

$$
\begin{equation*}
\omega=2 \sqrt{\frac{\hbar \lambda_{0}}{m_{0}}} \simeq 10^{-5} \mathrm{sec}^{-1} \tag{36}
\end{equation*}
$$

The two parameters $\varphi_{1}$ and $\varphi_{2}$ are functions of the initial condition. Also, note that setting $\lambda_{0}=0$ will give the same results as one will get by using the Schrödinger equation instead of Eqn. (27).

[^2]Eqns. (34) and (35) tell that the spread in position and momentum do not increase in time, but reach an asymptotic final value give by:

$$
\begin{equation*}
\sigma_{q}(\infty)=\sqrt{\frac{\hbar}{m \omega}} \simeq\left(10^{-15} \sqrt{\frac{\mathrm{Kg}}{m}}\right) \mathrm{m} \tag{37}
\end{equation*}
$$

and:

$$
\begin{equation*}
\sigma_{p}(\infty)=\sqrt{\frac{\hbar m \omega}{2}} \simeq\left(10^{-19} \sqrt{\frac{m}{\mathrm{Kg}}}\right) \frac{\mathrm{Kg} \mathrm{~m}}{\mathrm{sec}} \tag{38}
\end{equation*}
$$

such that:

$$
\begin{equation*}
\sigma_{q}(\infty) \sigma_{p}(\infty)=\frac{\hbar}{\sqrt{2}} \tag{39}
\end{equation*}
$$

which corresponds to almost the minimum allowed by Heisenberg's uncertainty relations. As we see, the spread in position does not increase indefinitely, but stabilizes to a finite value. For microscopic systems, this value is still large, such as to guarantee that in all standard experiments - in particular, diffraction experiments - one observes interference effects. For macroscopic objects instead, the spread is very small, so small that for all practical purposes the wave function behaves like a point-like system. Once again, this is how collapse models are able to accommodate both the "wavy" nature of quantum systems and the "particle" nature of classical objects, within one single dynamical framework. One should also note that, as a byproduct of the collapse in position, one has an almost perfect collapse in momentum, compatibly with Heisenberg's uncertainty relations.

Eqn. (32) says that the mean momentum undergoes a diffusion process. For microscopic systems, such a diffusion is appreciably large: the wave function is kicked back and forth by the collapse noise. For larger objects instead, the diffusion becomes weaker and weaker, to the point that at the macroscopic level it is almost entirely negligible. The same is true for the mean in position, according to Eqn. (31). In this way, collapse models can explain both the stochastic nature of quantum phenomena and the (apparently) deterministic nature of classical ones. Moreover, the average momentum $\mathbb{E}\left[\langle p\rangle_{t}\right]$ is constant $\left(\langle p\rangle_{t}=\hbar k_{t}\right)$, while the average position is given by $\mathbb{E}\left[\langle q\rangle_{t}\right]=\mathbb{E}\left[\langle p\rangle_{t}\right] / m$ : the particle, on the average, moves along a straight line, depending on its initial momentum.

Two comments are in order. The first one is that the above results refer only to the special case of Gaussian wave functions, like that of Eqn. (29). However, in Bassi and Duerr, 2008; Bassi et al., 2010b) a remarkable result has been proven: with probability one, any initial state converges asymptotically to a Gaussian wave function, having a fixed spread both in position and in momentum, given by Eqn. (37) and (38) respectively. The collapse process not only localizes wave functions, but also smoothens all their bumps and eventually shapes them as Gaussian functions. The second comment is that
the above results refer only to a free particle. Also, the harmonic oscillator can be treated in a fully analytical way, but more general potentials require perturbative approaches, which have not been explored so far.

To conclude this section, let us consider the many-particle equation:

$$
\begin{align*}
d \psi_{t}= & {\left[-\frac{i}{\hbar} H d t+\sum_{i=1}^{N} \sqrt{\lambda_{i}}\left(q_{i}-\left\langle q_{i}\right\rangle_{t}\right) d W_{t}^{(i)}\right.} \\
& \left.-\frac{1}{2} \sum_{i=1}^{N} \lambda_{i}\left(q_{i}-\left\langle q_{i}\right\rangle_{t}\right)^{2} d t\right] \psi_{t} \tag{40}
\end{align*}
$$

where $H$ is the quantum Hamiltonian of the composite system, the operators $q_{i}(i=$ $1, \ldots N)$ are the position operators of the particles of the system, and $W_{t}^{(i)}(i=1, \ldots N)$ are $N$ independent standard Wiener processes.

As often in these cases, it is convenient to switch from the particles' coordinates $\left(x_{1}, x_{2}, \ldots x_{N}\right)$ to the center-of-mass $(R)$ and relative $\left(\tilde{x}_{1}, \tilde{x}_{2}, \ldots \tilde{x}_{N}\right)$ coordinates:

$$
\left\{\begin{array}{l}
R=\frac{1}{M} \sum_{i=1}^{N} m_{i} x_{i} \quad M=\sum_{i=1}^{N} m_{i},  \tag{41}\\
x_{i}=R+\tilde{x}_{i} ;
\end{array}\right.
$$

let $Q$ be the position operator for the center of mass and $\tilde{q}_{i}(i=1 \ldots N)$ the position operators associated to the relative coordinates. It is not difficult to show that-under the assumption $H=H_{\text {См }}+H_{\text {rel }}$-the dynamics for the center of mass and that for
the relative motion decouple; in other words, $\psi_{t}(\{x\})=\psi_{t}^{\mathrm{CM}}(R) \otimes \psi_{t}^{\mathrm{rel}}(\{\tilde{x}\})$ solves Eqn. (40) whenever $\psi_{t}^{\mathrm{CM}}(R)$ and $\psi_{t}^{\text {rel }}(\{\tilde{x}\})$ satisfy the following equations:

$$
\begin{align*}
d \psi_{t}^{\mathrm{rel}}= & {\left[-\frac{i}{\hbar} H_{\mathrm{rel}} d t+\sum_{i=1}^{N} \sqrt{\lambda_{i}}\left(\tilde{q}_{i}-\left\langle\tilde{q}_{i}\right\rangle_{t}\right) d W_{t}^{(i)}\right.} \\
& \left.-\frac{1}{2} \sum_{i=1}^{N} \lambda_{i}\left(\tilde{q}_{i}-\left\langle\tilde{q}_{i}\right\rangle_{t}\right)^{2} d t\right] \psi_{t}^{\mathrm{rel}}, \tag{42}
\end{align*}
$$

and:

$$
\begin{align*}
d \psi_{t}^{\mathrm{CM}}= & {\left[-\frac{i}{\hbar} H_{\mathrm{CM}} d t+\sqrt{\lambda_{\mathrm{CM}}}\left(Q-\langle Q\rangle_{t}\right) d W_{t}\right.} \\
& \left.-\frac{\lambda_{\mathrm{CM}}}{2}\left(Q-\langle Q\rangle_{t}\right)^{2} d t\right] \psi_{t}^{\mathrm{CM}} \tag{43}
\end{align*}
$$

with:

$$
\begin{equation*}
\lambda_{\mathrm{CM}}=\sum_{n=1}^{N} \lambda_{n}=\frac{M}{m_{0}} \lambda_{0} \tag{44}
\end{equation*}
$$

The first of the above equations describes the internal motion of the composite system: it basically says that the internal structure of the system behaves quantum mechanically, modulo tiny modifications given by the collapse process. The second equation describes the center-of-mass evolution, and here we can see once again the most important feature of collapse models: the amplification mechan$i c s$. The collapse strength of the center of mass is proportional to the size (i.e. the number of constituents) of the system. For microscopic systems, $\lambda_{\mathrm{CM}}$ is similar to $\lambda_{0}$, i.e. very weak; in these cases, the collapse is almost negligible. For macroscopic objects, $\lambda_{\text {CM }}\left(\sim N \lambda_{0}\right.$, with $\left.N \sim 10^{24}\right)$ can be very
strong, implying a rapid and efficient collapse of the wave function. It is precisely because of the amplification mechanics that, with a single choice of $\lambda_{0}$, one can describe both quantum and classical phenomena at the same time.

## 1. The quantum formalism derived

Collapse models contain a unique and universal dynamics, which applies to all physical situations. Measurements play no special role in collapse models. It becomes then interesting and important to show how the entire phenomenology about quantum measurements emerges from the universal dynamics of collapse models. To do this, we use the QMUPL model, because of its relatively simple mathematical structure. We will now show that measurements always have a definite outcome, are randomly distributed according to the Born rule, and that at the end of the measurement process, the wave function of the micro-system collapses according to the von Neumann projection postulate. All these features are included in Eqn. (27), without any need for extra axioms.

The measurement setup we consider consists of a microscopic system $\mathcal{S}$ interacting with a macroscopic system $\mathcal{A}$, which acts like a measuring apparatus; both systems are described in quantum mechanical terms. We as-
sume that the measurement includes a finite set of outcomes. Accordingly, we assume that the microscopic system $\mathcal{S}$ can be described by a finite-dimensional complex Hilbert space. For the sake of simplicity, and without loss of generality, we can consider the simplest case: $\mathcal{H}_{\mathcal{S}}=\mathbb{C}^{2}$, because the generalization of what follows to $\mathbb{C}^{n}$ is quite straightforward. Since the most general self-adjoint operator $O$ acting on $\mathbb{C}^{2}$ can be written as

$$
\begin{equation*}
O=o_{+}|+\rangle\langle+|+o_{-}|-\rangle\langle-|, \tag{45}
\end{equation*}
$$

where $|+\rangle$ and $|-\rangle$ are the eigenstates of $O$, while $o_{+}$and $o_{-}$are its two real eigenvalues, for definiteness and with no loss of generality, in what follows we will take $o_{ \pm}= \pm \hbar / 2$ and $O$ to be the $z$-component of the spin, $S_{z}$, of a spin $1 / 2$ particle.

We take the following model for the measuring apparatus $\mathcal{A}$, which is general enough to describe all interesting physical situations: we assume that the apparatus consists of a fixed part plus a pointer moving along a graduate scale, in such a way that different positions of the pointer along the scale correspond to different possible outcomes of the measurement. To simplify the analysis, we study the evolution of the center of mass of the pointer only, and disregard all other macroscopic and microscopic degrees of freedom; accordingly, the pointer will be treated like a macroscopic quantum particle
of mass $m$ moving in one dimension only, whose state space is described by the Hilbert space $\mathcal{H}_{\mathcal{A}}=\mathrm{L}^{2}(\mathbb{R})$.

We assume that the pointer of $\mathcal{A}$ undergoes a spontaneous collapse mechanism according to Eqn. (27), while the microscopic system $\mathcal{S}$ evolves according to the standard Schrödinger equation, since, as typical of dynamical reduction models, the stochastic collapse terms have negligible effects on microscopic quantum systems. For definiteness, let us consider a pointer of mass $m=1 \mathrm{~g}$ (i.e., a pointer made of an Avogadro number of nucleons).

We take the total Hamiltonian $H$ to be of the form: $H=H_{\mathcal{S}}+H_{\mathcal{A}}+H_{\mathrm{INT}}$. The first term is the quantum Hamiltonian for the microscopic system: we assume that the time scale of the free evolution of the microscopic system is much larger than the characteristic time scale of the experiment ("instantaneous measurement" assumption); accordingly we take $H_{\mathcal{S}}$ to be the null operator. The second term is the quantum Hamiltonian of the pointer, which we take equal to that of a nonrelativistic free quantum particle of mass $m$ : $H_{\mathcal{A}}=p^{2} /(2 m)$, where $p$ is the momentum operator. Finally, we assume the interaction term $H_{\text {INT }}$ between the two systems to be of the von Neumann type, and devised in such
a way as to measure the $\operatorname{spin} S_{z}$ :

$$
\begin{equation*}
H_{\mathrm{INT}}(t)=\kappa \Delta_{t}^{T} S_{z} \otimes p, \tag{46}
\end{equation*}
$$

where $\kappa$ is a coupling constant and $\Delta^{T}: t \mapsto$ $\Delta_{t}^{T}$ is a $T$-normalized ${ }^{2}$ non-negative, real valued, function of time, identically equal to zero outside a given interval of the form $\left(t_{0}, t_{0}+T\right)$, i.e., outside the time interval of length $T$, say $T=1 \mathrm{sec}$, during which the experiment takes place; we choose the time origin in such a way that the experiment begins at $t_{0}=0$. As is well known in standard Quantum Mechanics, $H_{\text {INT }}$ generates the following type of evolution, depending on the initial state of the micro-system $\mathcal{S}$ :

$$
\begin{align*}
& {\left[c_{+}|+\rangle+c_{-}|-\rangle\right] \otimes \phi_{0}} \\
& \quad \mapsto c_{+}|+\rangle \otimes \phi_{+}+c_{-}|-\rangle \otimes \phi_{-} \tag{47}
\end{align*}
$$

where $\phi_{ \pm}$are final pointer states spatially translated with respect to the initial state $\phi_{0}$ by the quantity $\pm(\hbar / 2) \kappa T$. We will see how collapse models modify this linear evolution.

The strength of the coupling constant $\kappa$ has to be chosen in such a way that the distance $\hbar \kappa T$ between the initial state $\phi_{0}$ of the

$$
\begin{aligned}
& { }^{2} \text { By a } T \text {-normalized function, we simply mean } \\
& \qquad \int_{-\infty}^{+\infty} \Delta_{t}^{T} d t=\int_{t_{0}}^{t_{0}+T} \Delta_{t}^{T} d t=T
\end{aligned}
$$

Note that $\Delta_{t}^{T}$ depends also on the initial time $t_{0}$; we will avoid to indicate it explicitly, when no confusion arises.
pointer and any of the two final states $\phi_{ \pm}$is macroscopic; for definiteness, let us choose $\hbar \kappa=1 \mathrm{~cm} \mathrm{~s}^{-1}$, so that $\hbar \kappa T=1 \mathrm{~cm}$.

We take the initial states of the microscopic system $\mathcal{S}$ and of the macroscopic apparatus $\mathcal{A}$ to be completely uncorrelated, as it is customary and appropriate for the description of a measurement process. Accordingly, we assume the initial state of the total system $\mathcal{S}+\mathcal{A}$ to be:

$$
\begin{equation*}
\left[c_{+}|+\rangle+c_{-}|-\rangle\right] \otimes \phi_{0}, \tag{48}
\end{equation*}
$$

where $\phi_{0}$ describes the "ready" state of the macroscopic apparatus $\mathcal{A}$.

Regarding the initial state $\phi_{0}$ of the pointer, some considerations have to be done. In the previous section we showed that, according to Eqn. (27), the wave function for the centre of mass of an isolated quantum system reaches asymptotically (and very rapidly, for a macro-object) a Gaussian state of the form

$$
\begin{equation*}
\phi_{t}^{\mathrm{G}}(x)=\sqrt[4]{\frac{1}{2 \pi \sigma_{q}^{2}}} \exp \left[-\frac{1-i}{4 \sigma_{q}^{2}}\left(x-\bar{x}_{t}\right)^{2}+i \bar{k}_{t} x\right] \tag{49}
\end{equation*}
$$

(modulo a time-dependent global phase factor) with $\sigma_{q}$ defined as in Eq. (37), taking the value: $\sigma_{q} \simeq 4.6 \times 10^{-14} \mathrm{~m}$ for $m=1 \mathrm{~g}$. The dispersion of the Gaussian function of Eqn. (49) in momentum space is $\sigma_{p} \simeq 1.6 \times$ $10^{-21} \mathrm{~kg} \mathrm{~m} \mathrm{~s}^{-1}$, as described in Eqn. (38).

In our measurement model, we assume that the pointer is isolated for the time prior
to the experiment; during this time its wave function converges rapidly towards a state close to (49), which we therefore assume to be the initial state of the pointer. To summarize, we take as the initial state of the composite system $\mathcal{S}+\mathcal{A}$ the ket:

$$
\begin{equation*}
\Psi_{0}=\left[c_{+}|+\rangle+c_{-}|+\rangle\right] \otimes \phi^{\mathrm{G}} . \tag{50}
\end{equation*}
$$

We choose the natural reference frame where the pointer is initially at rest, so that $\bar{k}_{0}=0$ $\mathrm{m}^{-1}$, with the origin set up in such a way that $\bar{x}_{0}=0 \mathrm{~m}$.

We are now ready to solve the collapse equation. It is not difficult to show that, for the given initial condition, the solution takes the form

$$
\begin{equation*}
\psi_{t}=|+\rangle \otimes \phi_{t}^{+}+|-\rangle \otimes \phi_{t}^{-}, \tag{51}
\end{equation*}
$$

where, after defining the $c_{ \pm}$in terms of $\gamma^{ \pm}$, the wave functions $\phi_{t}^{+}$and $\phi_{t}^{-}$have the form:
$\phi_{t}^{ \pm}(x)=\exp \left[-\alpha_{t}\left(x-\bar{x}_{t}^{ \pm}\right)^{2}+i \bar{k}_{t}^{ \pm} x+\gamma_{t}^{ \pm}+i \theta_{t}^{ \pm}\right]$
whose parameters $\alpha_{t} \in \mathbb{C}$, and $\bar{x}_{t}^{ \pm}, \bar{k}_{t}^{ \pm}, \gamma_{t}^{ \pm}$, $\theta_{t}^{ \pm} \in \mathbb{R}$ satisfy a complicated set of nonlinear stochastic differential equations (Bassi) and Salvetti, 2007).

In order to extract the relevant physical information from Eqns. (51) and (52), let us consider the differences $X_{t}:=\bar{x}_{t}^{+}-\bar{x}_{t}^{-}$ and $K_{t}:=\bar{k}_{t}^{+}-\bar{k}_{t}^{-}$, which represent the distance in position and (modulo $\hbar$ ) momentum
space between the centres of the two Gaussian functions $\phi_{t}^{+}$and $\phi_{t}^{-}$. One can easily prove that $X_{t}$ and $K_{t}$ satisfy a set of linear and deterministic equations Bassi and Salvetti, 2007):

$$
\frac{d}{d t}\left[\begin{array}{l}
X_{t}  \tag{53}\\
K_{t}
\end{array}\right]=\left[\begin{array}{cc}
-\omega & \hbar / m \\
-2 \lambda & 0
\end{array}\right]\left[\begin{array}{l}
X_{t} \\
K_{t}
\end{array}\right]+\left[\begin{array}{c}
\hbar \kappa \Delta_{t}^{T} \\
0
\end{array}\right]
$$

where both the non-linear and the stochastic terms cancel out. The solution of the above system depends of course on the specific choice for the function $\Delta_{t}^{T}$; a simple reasonable choice is the following:

$$
\Delta_{t}^{T}= \begin{cases}1 & t \in[0, T]  \tag{54}\\ 0 & \text { else }\end{cases}
$$

which-in a standard quantum scenariomeans that during the measurement each term of the superposition moves with a constant speed towards the left and towards the right, respectively. According to this choice, $X_{t}$, given the initial condition $X_{0}=0 \mathrm{~m}$, evolves in time as follows:

$$
X_{t}=\left\{\begin{array}{r}
\frac{2 \hbar \kappa}{\omega} e^{-\omega t / 2} \sin \frac{\omega}{2} t \quad \text { for } 0 \leq t \leq T  \tag{55}\\
\frac{2 \hbar \kappa}{\omega} e^{-\omega t / 2}\left[\sin \frac{\omega}{2} t-e^{\omega T / 2} \sin \frac{\omega}{2}(t-T)\right] \\
\text { for } t \geq T
\end{array}\right.
$$

Since $\omega^{-1} \simeq 2.0 \times 10^{4} \mathrm{sec}$ is a very long time compared to the measurement-time, we can meaningfully expand Eqn. (55) to first order
in $\omega t$ :
$X_{t} \simeq \begin{cases}\hbar \kappa t & \text { for } 0 \leq t \leq T=(\hbar \kappa)^{-1}=1 \mathrm{~s}, \\ 1 \mathrm{~cm} & \text { for } T \leq t \ll \omega^{-1} \simeq 2.0 \times 10^{4} \mathrm{~s}\end{cases}$

As we see, the distance between the two peaks increases almost linearly in time, reaching its maximum ( 1 cm ) at the end of the measurement process, as predicted by the standard Schrödinger equation; after this time, their distance remains practically unaltered for extremely long times, and only for $t \simeq 2.0 \times 10^{4} \mathrm{~s}$ it starts slowly to decrease, eventually going to 0 . Note that such a behavior, being determined by $\omega$, does not depend on the mass of the pointer, thus a larger pointer will not change the situation. The moral is that $X_{t}$ behaves as if the reduction mechanism were not present (as if $\lambda_{0}=0$ ) so we have to look for the collapse somewhere else.

As we shall see now, the collapse occurs because, in a very short time, the weight of one of the two Gaussian wave functions ( $\phi_{t}^{+}$or $\phi_{t}^{-}$) becomes much smaller than the weight of the other component; this implies that one of the two components practically disappears, and only the other one survives, the one which determines the outcome of the experiment. Of course, this process is random and, as we shall prove, it occurs with a probability almost equivalent to the Born probability rule.

The relative damping between the two Gaussian components of Eqn. (51) is measured by the stochastic process $\Gamma_{t}=\gamma_{t}^{+}-\gamma_{t}^{-}$: if, at the end of the measurement process, it occurs that $\Gamma_{t} \gg 1$, it means that $\phi_{t}^{-}$is suppressed with respect to $\phi_{t}^{+}$, so that the initial state (50) practically evolves to $|+\rangle \otimes \phi_{t}^{+}$; the opposite happens if $\Gamma_{t} \ll-1$.

In (Bassi and Salvetti, 2007) it is shown that $\Gamma_{t}$ satisfies the nonlinear stochastic differential equation:

$$
\begin{equation*}
d \Gamma_{t}=\lambda X_{t}^{2} \tanh \Gamma_{t} d t+\sqrt{\lambda} X_{t} d W_{t} \tag{57}
\end{equation*}
$$

to be solved with initial condition $\Gamma_{0}=$ $\ln \left|c_{+} / c_{-}\right|$. To proceed further with the analysis, it is convenient to perform the following time change:

$$
\begin{equation*}
t \quad \longrightarrow \quad s_{t}:=\lambda \int_{0}^{t} X_{t}^{2} d t^{\prime} \tag{58}
\end{equation*}
$$

which allows us to describe the collapse process in terms of the dimensionless quantity $s$ that measures its effectiveness. Using Eqn. (55), one can solve exactly the above integral and compute $s$ as a function of $t$. Such a function however cannot be inverted in order to get $t$ from $s$. Therefore, we use the simplified expression (56) in place of the exact formula (55) to compute the integral, an expression which, as we have seen, represents a very good approximation to the time evolution of $X_{t}$ throughout the whole time during which the experiment takes place. Accord-
ingly, we have:

$$
\begin{align*}
& s \equiv s_{t} \simeq \frac{\lambda \hbar^{2} \kappa^{2}}{3} t^{3} \simeq 2.0 \times 10^{17}(t / \mathrm{s})^{3} \\
& \quad \text { for } 0 \leq t \leq T=1 \mathrm{~s}  \tag{59}\\
& t \equiv t_{s} \simeq \sqrt[3]{\frac{3}{\lambda \hbar^{2} \kappa^{2}} s} \simeq\left(1.7 \times 10^{-6} \sqrt[3]{s}\right) \mathrm{s} \\
& \quad \text { for } 0 \leq s \leq \lambda \hbar^{2} \kappa^{2} / 3=2.0 \times 10^{17} . \tag{60}
\end{align*}
$$

Note that, according to the above equations, the physical time $t$ depends on $s$ through the inverse cubic-root of $\lambda$, i.e. on the inverse cubic-root of the mass of the pointer; this time dependence of $t$ on $\lambda$ is important since, as we shall see, it will affect the collapse time. We do not study the functional dependence between $s$ and $t$ for $t \geq T$ since, as we shall soon see and as we expect it to be, the collapse occurs at times much smaller than $T$.

Written in terms of the new variable $s$, Eqn. (57) reduces to:

$$
\begin{equation*}
d \Gamma_{s}=\tanh \Gamma_{s} d s+d W_{s} \tag{61}
\end{equation*}
$$

this equation belongs to a general class of stochastic differential equations whose properties are known in detail Gikhman and Skorokhod, 1972). Here we report the main results.

Collapse time. The collapse time is the time when $\left|\Gamma_{s}\right| \geq b$, with $b \gg 1$, i.e. the time when one of the two terms of the superposition becomes dominant with respect to the other:

$$
\begin{equation*}
S_{\mathrm{COL}} \equiv \inf \left\{s:\left|\Gamma_{s}\right| \geq b\right\} \tag{62}
\end{equation*}
$$

This is a random variable (for each run of the experiment, the collapse time slightly changes), whose mean and variance can be exactly computed (Gikhman and Skorokhod, 1972). In particular, if we start with an equal-weight superposition $\left(\Gamma_{0}=0\right)$, then: $\mathbb{E}_{\mathbb{P}}\left[S_{\mathrm{COL}}\right] \simeq b$ and $\mathbb{V}_{\mathbb{P}}\left[S_{\mathrm{COL}}\right] \simeq b$, where $\mathbb{E}_{\mathbb{P}}[\cdot]$ and $\mathbb{V}_{\mathbb{P}}[\cdot]$ denote the mean and variance, respectively. If we move back from $s$ to the physical time $t$, we have the following estimate for the collapse time (Bassi and Salvetti, 2007):

$$
\begin{equation*}
T_{\mathrm{COL}} \simeq 1.5 \times 10^{-4} \mathrm{~s} \tag{63}
\end{equation*}
$$

The collapse occurs within a time interval smaller than the perception time of a human observer. Moreover, as previously argued, $T_{\mathrm{COL}}$ is proportional to the inverse cubic-root of the mass of the pointer: therefore, the bigger the pointer, the shorter the collapse time. With our choice for $\lambda_{0}$, even for a $1-\mathrm{g}$ pointer the reduction occurs practically instantaneously.

It is important to note that, at time $T_{\mathrm{COL}} \simeq 1.5 \times 10^{-4} \mathrm{~s}$, the distance between the two Gaussian components is approximately $X_{T_{\mathrm{COL}}} \simeq 1.5 \times 10^{-4} \mathrm{~cm}$ : this means that, with very high probability, the collapse occurs before the two components have enough time to spread out in space to form a macroscopic superposition. This means that, from the physical point of view, there is no col-
lapse of the wave function at all, since it always remains perfectly localized in space at any stage of the experiment.

Collapse probability. Let us call $P_{+}$the probability that $\Gamma_{s}$ hits the point $+b$ before the point $-b$, i.e. the probability that $\phi_{s}^{+}$ survives during the collapse process so that the outcome of the measurement is " $+\hbar / 2$ ". Such a probability turns out to be equal to (Bassi and Salvetti, 2007):

$$
\begin{equation*}
P_{+}=\frac{1}{2} \frac{\tanh b+\tanh \Gamma_{0}}{\tanh b} ; \tag{64}
\end{equation*}
$$

while the probability $P_{-}$that $\Gamma_{s}$ hits the point $-b$ before the point $+b$, i.e. that the outcome of the experiment is " $-\hbar / 2$ ", is:

$$
\begin{equation*}
P_{-}=\frac{1}{2} \frac{\tanh b-\tanh \Gamma_{0}}{\tanh b} . \tag{65}
\end{equation*}
$$

By taking into account that $\tanh b \simeq 1$, since we have assumed that $b \gg 1$, we can write, with extremely good approximation:

$$
\begin{align*}
P_{+} & \simeq \frac{1}{2}\left[1+\tanh \Gamma_{0}\right]=\frac{e^{\Gamma_{0}}}{e^{\Gamma_{0}}+e^{-\Gamma_{0}}} \\
& =\frac{e^{2 \gamma_{0}^{+}}}{e^{2 \gamma_{0}^{+}}+e^{2 \gamma_{0}^{-}}}=\left|c_{+}\right|^{2}  \tag{66}\\
P_{-} & \simeq \frac{1}{2}\left[1-\tanh \Gamma_{0}\right]=\frac{e^{-\Gamma_{0}}}{e^{\Gamma_{0}}+e^{-\Gamma_{0}}} \\
& =\frac{e^{2 \gamma_{0}^{-}}}{e^{2 \gamma_{0}^{+}}+e^{2 \gamma_{0}^{-}}}=\left|c_{-}\right|^{2} . \tag{67}
\end{align*}
$$

We see that the probability of getting one of the two possible outcomes is practically equivalent to the Born probability rule. On the one hand, this is an entirely expected result, since collapse models have been designed precisely in order to solve the measurement problem and in particular to reproduce
quantum probabilities; on the other hand, it is striking that a very general equation like Eqn. (27), which is meant to describe both quantum systems as well as macroscopic classical objects (i.e. all physical situations, at the non-relativistic level), when applied to a measurement situation, provides not only a consistent description of the measurement process, but also reproduces quantum probabilities with such a good precision.

State vector after the collapse. At time $t \geq T_{\mathrm{COL}}$ the state of the composite system is:

$$
\begin{equation*}
\Psi_{t}=\frac{|+\rangle \otimes \tilde{\phi}_{t}^{+}+\epsilon_{t}|-\rangle \otimes \tilde{\phi}_{t}^{-}}{\sqrt{1+\epsilon_{t}^{2}}} \tag{68}
\end{equation*}
$$

where $\epsilon_{t} \equiv e^{-\left(\gamma_{t}^{+}-\gamma_{t}^{-}\right)}$and the normalized Gaussian states $\tilde{\phi}_{t}^{ \pm}$are defined as follows:

$$
\begin{align*}
& \tilde{\phi}_{t}^{ \pm}=\sqrt[4]{\frac{1}{2 \pi \sigma_{q}^{2}}} \exp \left[-\frac{1-i}{4 \sigma_{q}^{2}}\left(x-\bar{x}_{t}^{ \pm}\right)^{2}\right. \\
&\left.+i \bar{k}_{t}^{ \pm} x+i \theta_{t}^{ \pm}\right] \tag{69}
\end{align*}
$$

Let us assume that the collapse occurred in favor of the positive eigenvalue, i.e. in such a way that $\Gamma_{t} \geq b$ for $t \geq T_{\mathrm{COL}}$; it follows that:

$$
\begin{equation*}
\epsilon_{t} \leq e^{-b} \simeq 0 \quad \text { if } b \gg 1 \tag{70}
\end{equation*}
$$

and we can write, with excellent accuracy:

$$
\begin{equation*}
\Psi_{t} \simeq|+\rangle \otimes \tilde{\phi}_{t}^{+} \tag{71}
\end{equation*}
$$

We recover in this way the postulate of wave packet reduction of standard quantum mechanics: at the end of the measurement process, the state of the micro-system reduces
to the eigenstate corresponding to the eigenvalue which has been obtained as the outcome of the measurement, the outcome being defined by the surviving Gaussian component ( $\tilde{\phi}_{t}^{+}$in this case). Note the important fact that, according to our model, the collapse acts directly only on the pointer of the measuring apparatus, not on the microsystem; however, the combined effect of the collapse plus the von Neumann type of interaction is that the microscopic superposition of the spin states of the micro-system is rapidly reduced right after the measurement.

Note finally that, after the collapse, the states of the micro-system and of the pointer are de facto factorized: as such, after the measurement process one can, for all practical purposes, disregard the pointer and focus only on the micro-system for future experiments or interactions with other systems, as is the custom in laboratory experiments.

To conclude, we have seen how collapse models can describe quantum measurements in a precise way, without ambiguities and paradoxes. We have seen that the standard recipe for quantum measurements (definite outcomes, the Born rule, the postulate of wave function collapse) derives from the dynamical equation (27) and need not be postulated in an ad hoc way. But there is something more: it can be shown (Bassi et al. 2007) that also the Hilbert space op-
erator formalism - according to which observable quantities are represented by selfadjoint operators - can also be derived from Eqn. (27). In other words, in collapse models there is only the wave function and a collapse equation like Eqn. (27): everything else can be derived from it.

## E. The dissipative QMUPL model

One of the characteristic features of collapse models is the violation of energy conservation for isolated systems. In fact, a consequence of Eqn. (27) is that the mean energy $E_{t} \equiv \mathbb{E}\left[\langle H\rangle_{t}\right]$ of a free particle $\left(H=p^{2} / 2 m\right)$ changes in time as follows:

$$
\begin{equation*}
E_{t}=E_{0}+\frac{\lambda \hbar^{2}}{2 m} t \tag{72}
\end{equation*}
$$

This linear increase of the mean kinetic energy is common not only to the QMUPL model, but also to the GRW model (Ghirardi et al. 1986) and the CSL model Ghirardi et al., 1990c). The reason is quite easy to understand. If one goes back to the analysis of Sec. II.E, one can see that a collapse in position induces indirectly also a collapse in momentum: the wave function is almost perfectly localized both in position and momentum, compatibly with Heisenberg's uncertainty relations. However, the wave function diffuses both in position and momentum, and the fluctuations become larger and
larger. The diffusion in momentum forces the system to pick up higher and higher components, hence the energy increase. For typical values of the parameters, such an increase is very small and undetectable with present day technology Adler, 2007, Adler and Bassi, 2009); still, one would wish to restore the principle of energy conservation within collapse models.

The way to solve this problem is to include dissipative effects into the system-noise interaction. This is achieved by the dissipative QMUPL model (Bassi et al., 2005b). The modified Schrödinger equation is:

$$
\begin{align*}
d \psi_{t}= & {\left[-\frac{i}{\hbar} H d t+\sqrt{\lambda}\left(A-r_{t}\right) d W_{t}\right.} \\
& \left.-\frac{\lambda}{2}\left(A^{\dagger} A-2 r_{t} A+r_{t}^{2}\right) d t\right] \psi_{t} \tag{73}
\end{align*}
$$

with:

$$
\begin{equation*}
r_{t}=\frac{1}{2}\left\langle\psi_{t}\right|\left(A+A^{\dagger}\right)\left|\psi_{t}\right\rangle . \tag{74}
\end{equation*}
$$

The collapse operator $A$ is taken equal to:

$$
\begin{equation*}
A=q+i \frac{\alpha}{\hbar} p \tag{75}
\end{equation*}
$$

where $p$ is the momentum operator. Moreover, we define the operator $H$ as follows:

$$
\begin{equation*}
H=H_{0}+\frac{\lambda \alpha}{2}\{q, p\} \tag{76}
\end{equation*}
$$

where $H_{0}$ is the standard quantum Hamiltonian. As we see, the model is defined in terms of the two constants $\lambda$ and $\alpha$; the first constant has been first introduced in Eqn. (28); the second one is defined as follows:

$$
\begin{equation*}
\alpha=\frac{m_{0}}{m} \alpha_{0} \tag{77}
\end{equation*}
$$

where $\alpha_{0}$ is a fixed constant, which has been takes equal to: $\alpha_{0} \simeq 10^{-18} \mathrm{~m}^{2}$. As shown in (Bassi et al., 2010a, 2005b), with this numerical choice for $\alpha_{0}$ one can show that all the collapse properties remain unaltered: the wave function collapses basically in the same way as in the original QMUPL model. The only important difference is that now the system's diffusion contains friction terms which constrain the fluctuations. As a consequence, the mean kinetic energy of a free particle evolves in time as follows:

$$
\begin{equation*}
E_{t}=\left(E_{0}-\frac{\hbar^{2}}{8 m \alpha}\right) e^{-4 \lambda \alpha t}+\frac{\hbar^{2}}{8 m \alpha} \tag{78}
\end{equation*}
$$

The energy still changes in time and thus is not conserved; however it does not diverge for $t \rightarrow+\infty$, but reaches the asymptotic finite value:

$$
\begin{equation*}
E_{\infty}=\frac{\hbar^{2}}{8 m \alpha}=\frac{\hbar^{2}}{8 m_{0} \alpha_{0}} \tag{79}
\end{equation*}
$$

corresponding to a temperature

$$
\begin{equation*}
T=\frac{\hbar^{2}}{4 m_{0} \alpha_{0} k_{B}} \simeq 10^{-1} \mathrm{~K} \tag{80}
\end{equation*}
$$

which is independent of the mass of the particle. Note that the time evolution of $E_{t}$ is very slow, the rate of change being equal to $4 \lambda \alpha=4 \lambda_{0} \alpha_{0} \simeq 10^{-20} \sec ^{-1}$ which means that, with very high accuracy, the mean energy remains constant for very long times.

The above equations imply that the stochastic process acts like a dissipative medium which, due to friction, slowly thermalizes all systems to the temperature $T$ by
absorbing or transferring energy to them depending on their initial state. Note that according to Eq. (80) a very "cold" medium is enough to guarantee the localization in space of the wave functions of macroscopic objects. Note also that one recovers the QMUPL model by setting $\alpha \rightarrow 0$ which corresponds to the high temperature limit $T \rightarrow+\infty$. This calculation shows that the reason why the energy increases in the original collapse models and eventually diverges is that the noise acts like an infinite-temperature stochastic medium with no friction.

The above discussion suggests that the model can be further developed by promoting $W_{t}$ to a real physical medium with its own equations of motion, having a stochastic behavior to which a temperature $T$ can be associated and such that, with good accuracy, can be treated like a Wiener process. This would imply not only that the medium acts on the wave function by changing its state according to Eqn. (73), but also that the wave function acts back on the medium according to equations which still have to be studied. The above suggestion opens the way for the possibility that by taking into account the energy of both the quantum system and the stochastic medium, one can restore perfect energy conservation not only on the average but also for single realizations of the stochastic process. A similar proposal has been con-
sidered by P. Pearle (Pearle, 2000) and by S . Adler (Adler, 2004).

## F. The colored-noise QMUPL model

The models so far considered involve a Wiener process $W_{t}$ which, though being mathematically very convenient since the corresponding stochastic differential equations are relatively easy to handle, are not physically realistic. The reason is the following. Using a mathematically less precise, but physically more intuitive, notation, Eqn. (27) can be written as follows:

$$
\begin{align*}
\frac{d}{d t} \psi_{t}= & {\left[-\frac{i}{\hbar} H+\sqrt{\lambda}\left(q-\langle q\rangle_{t}\right) w_{t}\right.} \\
& \left.-\frac{\lambda}{2}\left(q-\langle q\rangle_{t}\right)^{2}\right] \psi_{t} \tag{81}
\end{align*}
$$

where $w_{t} \equiv d W_{t} / d t$ is a white noise, i.e. a stochastic process with zero mean and correlation function $\mathbb{E}\left[w_{t} w_{s}\right]=\delta(t-s)$. The white noise then has a flat spectrum, where all frequencies appear with the same weight. While this is alright when describing rapidly fluctuating fields, in general it is a very idealized situation. If one thinks of the collapse noise as a real random field, possibly having a cosmological origin, then very likely its spectrum has a more complicated structure, with a cut-off at high frequencies. It becomes then important to analyze models where the noise has a generic correlation function $D(t, s)$, and
see how the collapse depends on its properties.

This has been done in (Bassi and Ferialdi, 2009a b) for the QMUPL model, in (Ferialdi and Bassi, 2011) for the dissipative QMUPL model and in Adler and Bassi, 2007, 2008) for the CSL model. The calculations are much more difficult, since Eq. (81) changes into a non-Markovian integro-differential equation. The integral covers the past history and is weighted by the time correlator $D(t, s)$; in the white noise case $D(t, s)=\delta(t-s)$ the integral vanishes and one recovers the original equations. Here we do not enter into the details of the calculations, which are not easy to report; it is sufficient to say that all the main features of the collapse mechanism are preserved. On the other hand, some of the predictions may radically change, more specifically all those which depend on the behavior of the noise at high frequencies Adler and Ramazanoğlu, 2007).

## G. On the origin of the noise field

In the dissipative and colored-noise QMUPL models, the noise field acquires a more physical character: it can be assigned a finite temperature, and its spectrum is arbitrary; moreover, this noise is assumed to fill space. It becomes then natural to con-
sider whether it can have a cosmological origin. At present it is too early to answer such a question, though some work has already been done (Adler and Bassi, 2008) and some people have already suggested that it could have a gravitational (Diósi, 1989; Feynman et al. 1995; Karolyhazy et al., 1986; Penrose, 1996) or pre-quantum (Adler, 2004) nature. Moreover, it is still not clear why it has an antiHermitian coupling to matter, which is necessary to ensure the collapse ${ }^{3}$. However, one can meaningfully ask whether a noise with 'typical' cosmological properties (in terms of temperature and correlation function) can induce an efficient collapse of the wave function, where by 'efficient' we mean that the collapse is fast enough to avoid the occurrence of macroscopic superpositions.

Let us consider a Gaussian state as in Eqn. (29), whose time evolution can be analytically unfolded in all models so far considered. The quantity we are interested in, is its spread in space $\sigma_{t}=\left(2 \sqrt{\alpha_{t}^{\mathrm{R}}}\right)^{-1 / 2}$. This is plotted in Fig. 1 and Fig. 2. The first figure refers to the QMUPL model. It shows how the spread of a wave function with initial spread $\sigma=5 \times 10^{-7} \mathrm{~m}$ changes af-

[^3]

FIG. 1 Wave function spread $\sigma$ after a time $t=10^{-2} \mathrm{~s}$, as a function of particle number $n$ and collapse parameter $\lambda$, according to the QMUPL model. The initial value $\sigma_{0}=5 \times 10^{-7} \mathrm{~m}$ reproduces the geometry of the macromolecule diffraction experiments of Arndt et al. 1999 Nairz et al. 2001. For few particles, and for small values of $\lambda$, the spread increases just like in the standard quantum case. Increasing $\lambda$, the collapse becomes stronger. The black line marks where $\sigma$ has reduced to half its initial value, after the considered time. This can be taken as the threshold from quantum (no collapse) to classical (collapse) regime. In the case of the macromolecule diffraction experiments marked by the white strip, values of $\lambda \lesssim 10^{-5} \mathrm{~s}^{-1}$ imply that collapse models' predictions agree with standard quantum predictions. This is consistent with the analysis of Adler 2007, for the CSL model. The proposed theoretical bound, represented by the white region, shows that with an increase of the particle number of $\sim 2-3$ orders of magnitude, these experiments would represent a crucial test for macroscopic quantum superpositions.
ter $t=10^{-2} \mathrm{~s}$, as a function of the collapse strength $\lambda$ and of the number of particles. The numerical values for $\sigma$ and $t$ mimic the setup of the macromolecule diffraction experiments (Arndt et al., 1999; Nairz et al., 2001), which can be considered the most straightforward tests of quantum linearity, and therefore of collapse models. As the picture shows, in order for these experiments to become crucial tests of quantum linearity, the mass of the molecules has to be increased by $\sim 2-3$ orders of magnitude. Recent analysis (Gerlich et al. 2011) shows that such an improvement
is already within reach.
Fig. 2. (top row) shows the difference in the evolution of the spread as given by the dissipative QMUPL model and by the original QMUPL model. Two temperatures have


FIG. 2 Difference between the spread $\sigma$ predicted by the QMUPL model with that given by the dissipative QMUPL model (QMUPLd), for $\mathrm{T}=2.73 \mathrm{~K}$ (top row left) and $\mathrm{T}=2.73 \times 10^{-3} \mathrm{~K}$ (top row right); and by the colored-noise model (QMUPLen), with cutoff at $10^{10} \mathrm{~Hz}$ (bottom row left) and $10^{2} \mathrm{~Hz}$ (bottom row right). As the color bars on the right show, the whiter the region, the greater the difference in the spreads. The initial spread $\sigma_{0}$ and the elapsed time $t$ are the same as in Fig. [...]. At lower temperatures or lower cutoffs the wave function tends to collapse more slowly, which results in a bigger difference with respect to the QMUPL model. Regarding the plots in the top row, the discrepancy manifests in the lower left corner of the plot for $T=2.73 \times 10^{-3} \mathrm{~K}$ which disappears for $T=2.73 \mathrm{~K}$. Regarding the plots in the bottom row, it is manifest in the diagonal strip for $\gamma=10^{2} \mathrm{~Hz}$ which decreases for $\gamma=10^{10} \mathrm{~Hz}$. This diagonal feature exactly lies on the ridge between the quantum and the classical regime as displayed in Fig. [...]. The large discrepancy there is due to the missing high frequencies in the noise spectrum of the colored-noise model. Without these high frequencies the colored-noise model is not able to reproduce the sharpness of the ridge predicted by the QMUPL model.
been considered: $T=2.73 \mathrm{~K}$ (top row right), as for the CMBR and $T=2.73 \times 10^{-3} \mathrm{~K}$ (top row left). In both cases the difference is negligible. This means that even a rather cold thermal field, according to cosmological standards, can induce an efficient collapse of the wave function, as efficient as with the standard QMUPL model.

Fig. 2 (bottom row) shows the difference in the evolution of the spread as given by the QMUPL model, and by the colored-noise model, with a noise having a frequency cutoff. No high-frequency cutoff affects the collapsing properties of the model in an appreciable way. These results can be compared with the behavior of typical cosmological fields such as the CMBR, the relic neutrino background and the relic gravitational background. The spectrum of the first two have a cutoff (measured or expected) at $\sim 10^{11} \mathrm{~Hz}$, while the spectrum of the third one probably lies at $\sim 10^{10} \mathrm{~Hz}$ Grishchuk, 2010). All these cutoffs as well as that of $\sim 10^{15} \mathrm{~Hz}$ proposed in (Bassi and Ghirardi, 2003) for $w(t, \mathbf{x})$ ensure a rapid collapse of the wave function. While the collapse is robust over a large range of cutoffs, other effects, such as the emission of radiation from charged particles, highly depend on the spectrum of the noise correlator (Adler and Ramazanoğlu, 2007).

The message which can be drawn is that a cosmological field with 'typical' properties can induce an efficient collapse. A great challenge is to test the existence of such a field.

## H. The CSL model

The QMUPL model has the advantage of allowing for quite a rigorous mathematical analysis of the main features of collapse mod-
els, as was shown in the previous sections. However it does not seem physically realistic, for two main reasons. The first one is that it is built for systems of distinguishable particles, and its generalization to identical particles does not seem straightforward. The second reason is that the noise field depends only on time, not on space, thus it cannot be immediately identified with a random field of Nature. The CSL model (Ghirardi et al., 1990c) overcomes the above difficulties, and so far remains the most advanced collapse model. In its mass proportional version (Pearle and Squires, 1994), it is defined by the following stochastic differential equation in the Fock space:

$$
\begin{align*}
d \psi_{t} & =\left[-\frac{i}{\hbar} H d t\right.  \tag{82}\\
& +\frac{\sqrt{\gamma}}{m_{0}} \int d \mathbf{x}\left(M(\mathbf{x})-\langle M(\mathbf{x})\rangle_{t}\right) d W_{t}(\mathbf{x}) \\
& \left.-\frac{\gamma}{2 m_{0}^{2}} \int d \mathbf{x}\left(M(\mathbf{x})-\langle M(\mathbf{x})\rangle_{t}\right)^{2} d t\right] \psi_{t}
\end{align*}
$$

as usual, $H$ is the standard quantum Hamiltonian of the system and the other two terms induce the collapse of the wave function in space. The mass $m_{0}$ is a reference mass, which as usual is taken equal to that of a nucleon. The parameter $\gamma$ is a positive coupling constant which sets the strength of the collapse process, while $M(\mathbf{x})$ is a smeared mass
density operator:

$$
\begin{align*}
& M(\mathbf{x})=\sum_{j} m_{j} N_{j}(\mathbf{x})  \tag{83}\\
& N_{j}(\mathbf{x})=\int d \mathbf{y} g(\mathbf{y}-\mathbf{x}) \psi_{j}^{\dagger}(\mathbf{y}) \psi_{j}(\mathbf{y}) \tag{84}
\end{align*}
$$

$\psi_{j}^{\dagger}(\mathbf{y}, s), \psi_{j}(\mathbf{y}, s)$ being, respectively, the creation and annihilation operators of a particle of type $j$ in the space point $\mathbf{y}$. The smearing function $g(\mathbf{x})$ is taken equal to

$$
\begin{equation*}
g(\mathbf{x})=\frac{1}{\left(\sqrt{2 \pi} r_{c}\right)^{3}} e^{-\mathbf{x}^{2} / 2 r_{c}^{2}} \tag{85}
\end{equation*}
$$

where $r_{c}$ is the second new phenomenological constant of the model. $W_{t}(\mathbf{x})$ is an ensemble of independent Wiener processes, one for each point in space. [In the original, i.e. 'not mass proportional', CSL model, the integrals are proportional to the number density operator, instead of the mass density operator].

As one can see from Eqn. 82), in the CSL model the collapse operators are the density number operator $\psi_{j}^{\dagger}(\mathbf{y}) \psi_{j}(\mathbf{y})$, which means that superpositions containing different numbers of particles in different points of space are suppressed. This is equivalent to collapsing the wave function in space, in a secondquantized language.

The collapse occurs more or less like in the QUMPL model, though it is more difficult to unfold: an easier and more handy way to look at the collapse is through the density matrix, in particular how its off-diagonal elements decay in time. Since ordinary matter is made
just of electrons and nucleons and, according to Eqns. (82) and (83), the collapse effect on electrons is negligible with respect to the effect on nucleons, we focus our attention only on nucleons.

According to Eqn. (82), the decay of the off-diagonal elements of the density matrix $\rho_{t} \equiv \mathbb{E}\left[\left|\psi_{t}\right\rangle\left\langle\psi_{t}\right|\right]$ (where $\mathbb{E}[\cdot]$ denotes the stochastic average) of a many-nucleons system, in the position basis, is Ghirardi et al., 1990c):

$$
\begin{equation*}
\frac{\partial}{\partial t}\left\langle\overline{\mathbf{x}}^{\prime}\right| \rho_{t}\left|\overline{\mathbf{x}}^{\prime \prime}\right\rangle=-\Gamma\left(\overline{\mathbf{x}}^{\prime}, \overline{\mathbf{x}}^{\prime \prime}\right)\left\langle\overline{\mathbf{x}}^{\prime}\right| \rho_{t}\left|\overline{\mathbf{x}}^{\prime \prime}\right\rangle, \tag{86}
\end{equation*}
$$

where $\overline{\mathbf{x}}^{\prime} \equiv \mathrm{x}_{1}^{\prime}, \mathrm{x}_{2}^{\prime}, \ldots \mathrm{x}_{N}^{\prime}$ (and similarly for $\left.\overline{\mathbf{x}}^{\prime \prime}\right)$. In the above equation, we have neglected the standard quantum evolution. The decay function $\Gamma$ is:

$$
\begin{align*}
\Gamma & =\frac{\gamma}{2} \sum_{i, j}\left[G\left(\mathbf{x}_{i}^{\prime}-\mathbf{x}_{j}^{\prime}\right)+G\left(\mathbf{x}_{i}^{\prime \prime}-\mathbf{x}_{j}^{\prime \prime}\right)\right. \\
& \left.-2 G\left(\mathbf{x}_{i}^{\prime}-\mathbf{x}_{j}^{\prime \prime}\right)\right] \tag{87}
\end{align*}
$$

where the indices $i, j$ run over the $N$ nucleons of the system, and:

$$
\begin{equation*}
G(\mathbf{x})=\frac{1}{\left(4 \pi r_{C}^{2}\right)^{3 / 2}} e^{-\mathbf{x}^{2} / 4 r_{C}^{2}} \tag{88}
\end{equation*}
$$

As a first observation, in the case of a single nucleon $\Gamma$ reduces to:
$\Gamma\left(\mathbf{x}^{\prime}, \mathbf{x}^{\prime \prime}\right)=\frac{\gamma}{\left(4 \pi r_{C}^{2}\right)^{3 / 2}}\left[1-\exp -\frac{\left|\mathbf{x}^{\prime}-\mathbf{x}^{\prime \prime}\right|}{4 r_{c}^{2}}\right]$,
which is precisely the GRW 1-particle collapse term (see Eqn. (26)). This allows us to make the following identification between
the collapse rate $\lambda$ of the GRW model and the collapse strength $\gamma$ of the CSL model:

$$
\begin{equation*}
\lambda=\frac{\gamma}{\left(4 \pi r_{C}^{2}\right)^{3 / 2}} \tag{90}
\end{equation*}
$$

In this way, the two models give similar predictions regarding the collapse effects on systems containing just a few particles, while for many-particle systems important differences emerge, as we will see soon. In (Ghirardi et al., 1990c) the following choice for $\gamma$ was made:

$$
\begin{equation*}
\gamma \sim 10^{-30} \mathrm{~cm}^{3} \mathrm{~s}^{-1} \tag{91}
\end{equation*}
$$

corresponding to $\lambda \sim 2.2 \times 10^{-17} \mathrm{~s}^{-1}$.
Several useful approximate formulas can be obtained from Eqn. 87). The first one is for large distances. When the particles in a superposition are displaced by a distance $\ell=\left|\mathbf{x}^{\prime}-\mathbf{x}^{\prime \prime}\right| \ll r_{C}$, then according to Eqn. (88) their contribution to $\Gamma$ is negligibly small. Thus, only superpositions with $\ell \geq r_{C}$ contribute to $\Gamma$ and trigger the collapse of the wave function. In such a case, the formula tells that for groups of particles separated (in each term of the superposition) by less than $r_{C}$, the rate $\Gamma$ increases quadratically with the number of particles while for groups of particles separated by more than $r_{C}$ it increases linearly. Thus we have the following simplified formula for the collapse rate Adler, 2007):

$$
\begin{equation*}
\Gamma=\lambda n^{2} N \tag{92}
\end{equation*}
$$

where $n$ is the number of particles within a distance $r_{C}$, and $N$ is the number of such clusters. We note that the quadratic dependence of $\Gamma$ on the number of particles - which is absent in the original GRW model-is a direct effect of the identity of particles. This means that the identity of particles works in favor of the collapse.

An estimate for small distances can be obtained by Taylor expanding $G(\mathbf{x})$ as follows:

$$
\begin{equation*}
G(\mathbf{x}) \simeq \frac{1}{\left(4 \pi r_{C}^{2}\right)^{3 / 2}}\left[1-\frac{\mathbf{x}^{2}}{4 r_{C}^{2}}\right] \tag{93}
\end{equation*}
$$

which leads to:

$$
\begin{equation*}
\Gamma\left(\mathbf{x}^{\prime}, \mathbf{x}^{\prime \prime}\right) \simeq \frac{\lambda}{2 r_{C}^{2}}\left(\sum_{i}\left(\mathbf{x}_{i}^{\prime}-\mathbf{x}_{i}^{\prime \prime}\right)\right)^{2} \tag{94}
\end{equation*}
$$

As we can see - and as expected-the collapse strength grows quadratically with the superposition distance for small distances, like in the GRW and QMUPL models, the important difference here being that that there is a quadratic dependence also in the number of particles. In both cases (large- and smalldistance approximation), we see the amplification mechanism: the collapse rate increases with the size of the system.

Another useful formula can be obtained for macroscopic rigid systems, for which the mass distribution can be expressed by a density function $\mathrm{D}(\mathrm{x})$, averaging the contributions of the single nucleons. In such a case the decay function $\Gamma$ takes the simpler ex-
pression (Ghirardi et al., 1990c):

$$
\begin{align*}
\Gamma\left(\mathbf{X}^{\prime}, \mathbf{X}^{\prime \prime}\right) & =\gamma \int d \mathbf{x}\left[D^{2}(\mathbf{x})\right. \\
& \left.-D(\mathbf{x}) D\left(\mathbf{x}+\mathbf{X}^{\prime}-\mathbf{X}^{\prime \prime}\right)\right] \tag{95}
\end{align*}
$$

where $\mathbf{X}^{\prime}$ and $\mathbf{X}^{\prime \prime}$ are the positions of the center of mass of the object, in the two terms of the superposition. The physical meaning of Eqn. (95) can be understood by making reference to a homogeneous macroscopic body of constant density $D$. Then the decay rate becomes:

$$
\begin{equation*}
\Gamma=\gamma D n_{\mathrm{OUT}} \tag{96}
\end{equation*}
$$

where $n_{\text {out }}$ is the number of particles of the body when the center-of-mass position is $\mathbf{X}^{\prime}$, which do not lie in the volume occupied by the body when the center of mass position is $\mathrm{X}^{\prime \prime}$.

Further properties of the CSL model are discussed in (Ghirardi et al., 1990c) and (Bassi and Ghirardi, 2003).

1. On the nature of the noise field of the CSL model

As anticipated, contrary to the QMUPL model, in the CSL model the noise field can be given a straightforward physical interpretation. In order to see this, it is convenient to rewrite the CSL dynamical equation (82)
in the following equivalent form:

$$
\begin{align*}
d \psi_{t}= & {\left[-\frac{i}{\hbar} H d t\right.}  \tag{97}\\
+ & \frac{\sqrt{\gamma}}{m_{0}} \int d \mathbf{x}\left(M(\mathbf{x})-\langle M(\mathbf{x})\rangle_{t}\right) d \bar{W}_{t}(\mathbf{x}) \\
- & \frac{\gamma}{2 m_{0}^{2}} \int d \mathbf{x} d \mathbf{y}\left(M(\mathbf{x})-\langle M(\mathbf{x})\rangle_{t}\right) \\
& \left.\quad G(\mathbf{x}-\mathbf{y})\left(M(\mathbf{y})-\langle M(\mathbf{y})\rangle_{t}\right) d t\right] \psi_{t}
\end{align*}
$$ speculation. Moreover, one has to justify the non-Hermitian coupling and the non-linear character of the collapse equations, which are necessary in order to obtain the effective collapse dynamics.

2. Generalizations of the CSL model: dissipative, non-Markovian and relativistic models

Like the QMUPL model, the CSL model can also be generalized in several directions. The first generalization one would make, is to include dissipative terms, in order to cure the problem of energy non-conservation. In fact, in the CSL model also the energy increases at a steady rate, though such an increase is negligible for all practical purposes. This can be easily seen by noting that the 1-particle master equation of the CSL model coincides with the 1-particle equation of the GRW model. This type of generalization should not present particular problems, however it has not been worked out so far.

The second generalization consists in replacing the white noise field with a more general Gaussian noise. As for the QMUPL model, the equations become non-Markovian, therefore difficult to analyze mathematically. The analysis has been carried out to the leading perturbative order-with respect to the
collapse parameters $\gamma$-in Adler and Bassi, 2007, 2008). The result of the analysis is the expected one: the collapse qualitatively occurs with the same modalities as in the white noise case, the rate depending on the correlation function of the noise. In particular, the rate is robust against changes of the correlation functions, while other predictions are very sensitive to the form of the time correlator (Adler and Ramazanoğlu, 2007).

The great challenge of the dynamical reduction program is to formulate a consistent model of spontaneous wave function collapse for relativistic quantum field theories; many attempts have been proposed so far, none of which is as satisfactory as the non-relativistic CSL model.

The first attempt (Ghirardi et al., 1990b) aimed at making the CSL model relativistically invariant by replacing Eqn. (82) with a Tomonaga-Schwinger equation of the type:

$$
\begin{align*}
\frac{\delta \psi(\sigma)}{\delta \sigma(x)} & =\left[-\frac{i}{\hbar} \mathcal{H}(x)\right.  \tag{99}\\
& +\sqrt{\gamma}(\mathcal{L}(x)-\langle\mathcal{L}(x)\rangle) w(x) \\
& \left.-\frac{\gamma}{2}(\mathcal{L}(x)-\langle\mathcal{L}(x)\rangle)^{2}\right] \psi(\sigma)
\end{align*}
$$

where now the wave function is defined on an arbitrary space-like hypersurface $\sigma$ of spacetime. The operator $\mathcal{H}(x)$ is the Hamiltonian density of the system ( $x$ now denotes a point in space-time), and $\mathcal{L}(x)$ is a local density of the fields, on whose eigenmanifolds one decides to localize the wave function. The $c$ -
number function $w(x)$ is a random field on space-time with mean equal to zero, while the correlation function-in order for the theory to be Lorentz invariant in the appropriate stochastic sense (Ghirardi et al., 1990b)— must be a Lorentz scalar. And here the problems arise.

The simplest Lorentz invariant choice for the correlation function is:

$$
\begin{equation*}
\mathbb{E}[w(x) w(y)]=\delta^{(4)}(x-y), \tag{100}
\end{equation*}
$$

which however is not physically acceptable as it causes an infinite production of energy per unit time and unit volume. The reason is that in Eqn. (99) the fields are locally coupled to the noise which, when it is assumed to be white, is too violent, so to speak, and causes too many particles to come out of the vacuum. To better understand the situation, let us go back to the non-relativistic Eqn. (82): also there we basically have a white-noise process, which however is not coupled locally to the quantum field $a^{\dagger}(s, \mathbf{y}) a(s, \mathbf{y})$, the coupling being mediated by the smearing Gaussian function appearing in the definition of $N(\mathbf{x})$. One can compute the energy increase due to the collapse mechanism, which turns out to be proportional to $\alpha$. Now, if we want to have a local coupling between the quantum field and the noise, we must set $\alpha \rightarrow+\infty$, in which case the energy automatically diverges also for finite times.

The simplest way out one would think of, in order to cure this problem of Eqn. (99), is to replace the local coupling between the noise and the quantum field by a non-local one, as in the CSL equation (82); this procedure would essentially amount to replacing the white noise field with a non-white one. In both cases we need to find a Lorentz invariant function which either smears out the coupling or replaces the Dirac- $\delta$ in the definition of the correlation function (100). This however is not a straightforward task, for the following reason.

One of the reasons why the third term $(\gamma / 2)(\mathcal{L}(x)-\langle\mathcal{L}(x)\rangle)^{2}$ appears in Eqn. (99) is to guarantee that the collapse mechanism occurs with the correct quantum probabilities (for those who are experts in stochastic processes, the third term is such that the equation embodies an appropriate martingale structure); if we change the noise, we then have to change also the third term, and it turns out that we have to replace it with a non-local function of the fields (Myrvold et al., 2009; Nicrosini and Rimini, 2003). But, having a non-local function of the fields jeopardizes the entire (somehow formal) construction of the theory based on the Tomanaga-Schwinger formalism, as the integrability conditions are not automatically satisfied. More analysis is required, however it is very likely that the model will turn out

What we have briefly described is the major obstacle to finding a relativistic dynamical reduction model. We want to briefly mention three research programs which try to overcome such an impasse.
P. Pearle has spent many years in trying to avoid the infinite energy increase of relativistic spontaneous collapse models, e.g. by considering a tachyonic noise in place of a white noise as the agent of the collapse process (Pearle, 1999b), obtaining suggestive results. Unfortunately, as he has recently admitted (Myrvold et al., 2009), the program so far did not succeed.

Dowker and Henson have proposed a spontaneous collapse model for a quantum field theory defined on a $1+1$ null lattice (Dowker and Henson, 2004, Dowker and Herbauts, 2004), studying issues like the non-locality of the model and the no-faster-than-light constraint. More work needs to be done in trying to apply it to more realistic field theories; in particular, it would be important to understand if, in the continuum limit, one can get rid of the divergences which plague the relativistic CSL model.

In a recent paper (Tumulka, 2006b), generalizing a previous idea of Bell (Bell, 1987), Tumulka has proposed a discrete, GRW-like, relativistic model, for a system of $N$ noninteracting particles, based on the multi-time
formalism with $N$ Dirac equations, one per particle; the model fulfills all the necessary requirements, thus it represents a promising step forward in the search for a relativistic theory of dynamical reduction. Now it is important to understand whether it can be generalized in order to include also interactions.

It is rather discomforting that, after so many years and so many efforts, no satisfactory model of spontaneous wave function collapse for relativistic quantum field theories exists. And some have started to wonder whether there is some fundamental incompatibility between the dynamical reduction program and relativity. In this regard, we mention the analysis of (Ghirardi, 2000), where a toy model of spontaneous wave function collapse is analyzed: the collapse mechanism is supposed to occur instantaneously along all spacelike hypersurfaces crossing the center of the jump process; in spite of this superluminal effect, the whole picture is perfectly Lorentz invariant, it agrees with quantum mechanical predictions, it does not lead to any contradiction, e.g. it does not allow faster-than-light signaling and, moreover, different inertial observers always agree on the outcomes of experiments. Unfortunately, the missing piece (which would make the toy model a real physical model) is the dynamics for the reduction mechanism; in any case, this model suggests that there is no reason of
principle forbidding the relativistic reduction program.

Recently, a completely different perspective towards relativistic collapse models emerged Adler, 2004). If one assumes that the random field causing the collapse of the wave function is a physical field filling space and possibly having a cosmological origin, then there is no need to make the equations relativistically invariant. The noise would select a privileged reference framein pretty much the same way in which the CMBR identifies a preferred frame. Then the collapse equations, being phenomenological equations, need not be relativistic invariant, being dependent on the noise. The underlying theory, out of which these equations would emerge at an appropriate coarsegrained level, should respect the appropriate symmetries (Lorentz invariance or a possible generalization of it). The theory would explain the origin of the noise field which, because of the initial conditions, would break the relevant symmetry. This is only speculation at the moment. However it is a reasonable program, though difficult to carry out.

## I. Choice of the parameters

The choice (23) for $\lambda$ and $r_{C}$ makes sure that collapse models agree with all observational evidence about the quantum nature of
microscopic systems, while macro-objects are always localized in space and behave according to Newton's laws (within experimentally testable limits). It sets a quantum-classical threshold at $\sim 10^{13}$ nucleons (Bassi and Ghirardi, 2003; Ghirardi et al., 1990c).

Recently, a much stronger bound on $\lambda$ has been proposed (Adler, 2007), namely:

$$
\begin{equation*}
\lambda \simeq 2.2 \times 10^{-8 \pm 2} \mathrm{~s}^{-1}, \tag{101}
\end{equation*}
$$

corresponding to a threshold of $\sim 10^{5}$ nucleons. The underlying motivation is that the collapse should be effective in all measurement processes, also those involving only a small number of particles as it happens in the process of latent image formation in photography, where only $\sim 10^{5}$ particles are displaced more than $r_{C}$. In order for the collapse to be already effective at this scale, one has to increase the conventional CSL value for $\lambda$ by $\sim 10^{9 \pm 2}$ orders of magnitude. Eqn. (101) is also the value one has to take in order to make sure that a superposition of 6 photons $\mathbb{4}^{4}$ reaching the human eye, collapses within the eye itself (Bassi et al., 2010a).

Both values $\lambda \simeq 10^{-16} \mathrm{~S}^{-1}$ and $\lambda \simeq 2.2 \times$ $10^{-8 \pm 2} \mathrm{~s}^{-1}$ are compatible with known experimental data (Adler and Bassi, 2009). However, such a large discrepancy of $\sim 9$ orders of magnitude shows that there is no general consensus on the strength of the collapse process

[^4]and consequently on the scale at which violations of quantum linearity can be expected to manifest. Experiments will decide which value - if any - is the correct one.

## III. UNDERLYING THEORIES

It is beyond doubt that if Continuous Spontaneous Collapse models are the right way to resolve the problems of quantum theory, these models must derive from an underlying physical theory based on new symmetry principles. Here, we review two such possibilities. One, known as Trace Dynamics, is a theory due to Stephen Adler and collaborators, which bears the same relation to quantum theory that statistical mechanics bears to thermodynamics and the kinetic theory of gases. The other, which is not quite a theory yet, is the idea that gravity has a role to play in bringing about collapse of the wavefunction.

## A. Trace Dynamics, Quantum Theory,

 and Spontaneous CollapseIn the process of canonical quantization the operator algebra or the commutation rule is arrived at starting from the Poisson bracket in the Classical theory. This may be somewhat unsatisfactory, since the Classical theory is supposed to be a limiting case of the
fundamental quantum theory. In order to know the fundamental theory we need not know its limit. The quantum commutations should be achieved in a more fundamental manner and this is what Adler's scheme of Trace Dynamics sets out to do. However, eventually the theory goes beyond deriving quantum theory from a more fundamental framework, and provides a possible resolution of the quantum measurement problem which is experimentally testable.

The physics of Trace Dynamics might appear formidable, but is in essence quite straightforward, and can be described in the following three well-laid out steps :
(i) The classical theory, which is the Newtonian dynamics of Grassmann-valued noncommuting matrices; and which as a consequence of global unitary invariance possesses a unique non-trivial conserved charge of great significance [Sections 1.1.1-1.1.4 below].
(ii) The statistical thermodynamics of this theory, the construction of the canonical ensemble and the derivation of equilibrium. The derivation of an all important Ward identity, as a cosequence of assumed invariance under constant shifts in phase space, from which there emerge, as thermodynamic averages, the canonical commutation relations of quantum theory, the Heisenberg equations of motion and the equivalent Schrödinger equation of quantum theory
[Sections 1.1.5-1.1.7].
(iii) The consideration of Brownian motion fluctuations around the above thermodynamic approximation, the consequent non-linear stochastic modification of the Schrödinger equation, the resolution of the quantum measurement problem and derivation of the Born probability rule [Section 1.1.8].

The review below is based on the book by Adler (Adler, 2004). The interested reader should consult Adler's book and the references therein for further details.

## 1. The fundamental degrees of freedom

In this scheme the fundamental degrees of freedom are matrices living on a background spacetime with complex Grassmann numbers (or more precisely elements of a graded algebra $\mathcal{G}_{\mathcal{C}}$ of complex Grassmann numbers) as elements. Grassmann numbers have the following properties:

$$
\begin{array}{r}
\theta_{i} \theta_{j}+\theta_{j} \theta_{i}=0 \\
\theta_{i}^{2}=0 \\
\chi=\theta_{R}+i \theta_{I m} \\
\left\{\chi_{r}, \chi_{s}\right\}=0
\end{array}
$$

Thus we have a matrix field with the help of which to each spacetime point we can asso-
ciate a matrix

$$
M(x)=\left(\begin{array}{llll}
\chi_{11}(x) & \chi_{12}(x) & . . & . . \\
\chi_{21}(x) & \chi_{22}(x) & . . & . . \\
. . & . . & . . & . . \\
. . & . . & . . & . .
\end{array}\right)
$$

where $\chi_{11}(x), .$. etc. are the matrix elements at space-time point $x$. Some further important properties of these Grassmann (matrix-) elements are the following:
(i) Product of even number of Grassmann elements commutes with all the elements of Grassmann algebra.
(ii) Product of odd number of Grassmann elements anti-commutes with any other odd number product.

Therefore we have two disjoint sectors :
Bosonic Sector- B: Consists of the identity and the Even Grade elements of the algebra : $B \equiv\left\{I, \chi_{a} \chi_{b}, \chi_{a} \chi_{b} \chi_{c} \chi_{d}, \ldots.\right\}$
Fermionic Sector-F: Consists of the Odd Grade elements from the algebra : $F \equiv$ $\left\{\chi_{a}, \chi_{a} \chi_{b} \chi_{c}, \ldots.\right\}$.
Therefore, the fundamental d.o.f. of the trace dynamics theory are the matrices made out of elements from these sectors

$$
\begin{gathered}
B_{I} \in\left\{M ; M_{i j} \in B\right\} \\
\chi_{I} \in\left\{M ; M_{i j} \in F\right\} \\
B_{I}, \chi_{I} \in \mathcal{G}_{M} .
\end{gathered}
$$

A general matrix can be decomposed as

$$
M=\left(\begin{array}{llll}
\chi_{11}^{B}(x) & \chi_{12}^{B}(x) & . . & \ldots \\
\chi_{21}^{B}(x) & \chi_{22}^{B}(x) & . . & \ldots \\
. . & . . & . . & . . \\
. . & . . & . . & . .
\end{array}\right)
$$

## $+$

$$
\left(\begin{array}{llll}
\chi_{11}^{F}(x) & \chi_{12}^{F}(x) & . . & . . \\
\chi_{21}^{F}(x) & \chi_{22}^{F}(x) & . . & \ldots \\
. . & . . & . . & . . \\
. . & . . & . . & . .
\end{array}\right)
$$

into Bosonic and Fermionic sectors.
Here we note that being matrices the d.o.f. of this scheme are non-commutative in nature. Dimensionality of the matrices can be arbitrary, however we work with finite dimensional matrices with the assumption that everything done subsequently can be extended to infinite dimensional matrices as well.

Next, we define an operation Trace on this matrix field as follows

$$
\operatorname{Tr}: \mathcal{G}_{M} \rightarrow \mathcal{G}_{C},
$$

which is a map from the space of matrices [ $\left.\mathcal{G}_{M}\right]$ to the field of complex Grassmann numbers $\left[\mathcal{G}_{C}\right]$ and is given by the sum of the diagonal elements of a given matrix (for the time being we specialize to finite dimensional matrices).
There are some nice Trace properties satisfied by the d.o.f. of the theory

$$
\begin{gather*}
\operatorname{Tr} B_{1} B_{2}=\Sigma_{m, n}\left(B_{1}\right)_{m n}\left(B_{2}\right)_{n m}= \\
\Sigma_{m, n}\left(B_{2}\right)_{n m}\left(B_{1}\right)_{m n}=\operatorname{Tr} B_{2} B_{1}  \tag{102}\\
\operatorname{Tr} \chi_{1} \chi_{2}=\Sigma_{m, n}\left(\chi_{1}\right)_{m n}\left(\chi_{2}\right)_{n m}= \\
-\Sigma_{m, n}\left(\chi_{2}\right)_{n m}\left(\chi_{1}\right)_{m n}=-\operatorname{Tr} \chi_{2} \chi_{1}  \tag{103}\\
\operatorname{Tr} B \chi=\operatorname{Tr} \chi B \tag{104}
\end{gather*}
$$

One also has some interesting Trace trilinear cyclic identities

$$
\begin{array}{r}
\operatorname{Tr} B_{1}\left[B_{2}, B_{3}\right]=\operatorname{Tr} B_{2}\left[B_{3}, B_{1}\right]= \\
\operatorname{Tr} B_{3}\left[B_{1}, B_{2}\right] \\
\operatorname{Tr} B_{1}\left\{B_{2}, B_{3}\right\}=\operatorname{Tr} B_{2}\left\{B_{3}, B_{1}\right\}= \\
\operatorname{Tr} B_{3}\left\{B_{1}, B_{2}\right\} \\
\operatorname{Tr} B\left[\chi_{1}, \chi_{2}\right]=\operatorname{Tr} \chi_{1}\left[\chi_{2}, B\right]= \\
\operatorname{Tr} \chi_{2}\left[\chi_{1}, B\right] \\
\operatorname{Tr} \chi_{1}\left\{B, \chi_{2}\right\}=\operatorname{Tr}\left\{\chi_{1}, B\right\} \chi_{2}= \\
\operatorname{Tr}\left[\chi_{1}, \chi_{2}\right] B \\
\operatorname{Tr} \chi\left\{B_{1}, B_{2}\right\}=\operatorname{Tr} B_{2}\left\{\chi, B_{1}\right\}= \\
\operatorname{Tr} B_{1}\left\{B_{2}, \chi\right\} \\
\operatorname{Tr} \chi\left[B_{1}, B_{2}\right]=\operatorname{Tr} B_{2}\left[\chi, B_{1}\right]= \\
\operatorname{Tr} B_{1}\left[B_{2}, \chi\right] \\
\operatorname{Tr} \chi_{1}\left[\chi_{2}, \chi_{3}\right]=\operatorname{Tr} \chi_{2}\left[\chi_{3}, \chi_{1}\right]= \\
\operatorname{Tr} \chi_{3}\left[\chi_{1}, \chi_{2}\right] \\
\operatorname{Tr} \chi_{1}\left\{\chi_{2}, \chi_{3}\right\}=\operatorname{Tr} \chi_{2}\left\{\chi_{3}, \chi_{1}\right\}= \\
\operatorname{Tr} \chi_{3}\left\{\chi_{1}, \chi_{2}\right\} \cdot( \tag{105}
\end{array}
$$

where $g_{i}$ is the grade (odd/even) of the matrix $O$. The anti-commutative feature of matrix elements induces non-triviality in the adjointness properties as seen above.

The matrices with the above properties play the role of 'particles' of the theory, which obey standard classical dynamics; a dynamics which we now examine. A little later, we will construct the statistical mechanics of a gas of such 'particles', and we will find that the equations of quantum theory are nothing but thermodynamic identities, valid in the thermodynamic limit of this underlying theory.

## 2. Classical dynamics of these degrees of free-

 domWe can construct a polynomial $P$ from these non-commuting matrices (say $O$ ) and obtain the trace of the polynomial (from now on bold letters will signify a trace)

$$
\mathbf{P}=\operatorname{Tr} P
$$

We define trace derivative of $\mathbf{P}$ w.r.t. variable $O$ (any general $O$ can be expanded in terms of $B_{I}, \chi_{I}$ as shown above) as

$$
\delta \mathbf{P}=\operatorname{Tr} \frac{\delta \mathbf{P}}{\delta O} \delta O
$$

Next, we can verify that the matrices have the following adjoint rule

$$
\left(O_{1}^{g_{1}} \ldots O_{n}^{g_{n}}\right)^{\dagger}=(-1)^{\sum_{i<j} g_{i} g_{j} g_{j}} O_{n}^{g_{n} \dagger} \ldots . . O_{1}^{g_{1} \dagger}
$$

i.e. $\delta$ - variation in $\mathbf{P}$ should be written in such a way that resulting $\delta$ - variation of $O$ in each monomial sits on the right, and the term
coming on the left of $\delta O$ is defined as trace derivative. It should be mentioned that one always constructs the polynomial $\mathbf{P}$ to be an even graded element of Grassmann algebra. Moreover, $\delta O$ and $\delta \mathbf{P}$ are also taken to be of same type (Bosonic/Fermionic) as $O$ and $\mathbf{P}$ respectively. Thus $\frac{\delta \mathbf{P}}{\delta O}$ will be of the same type as $O$.
EXAMPLE
For example, let us start with a polynomial

$$
P=A O B O C
$$

where $A, B, C$ are operators which in general do not commute with each other or with the variable $O$. Operator here stands for matrix. Then,

$$
\delta P=A(\delta O) B O C+A O B(\delta O) C
$$

Therefore,

$$
\begin{array}{r}
\delta \operatorname{Tr} P=\delta \mathbf{P}=\operatorname{Tr}\left(\epsilon_{A O} B O C A(\delta O)+\right. \\
\left.\epsilon_{C} C A O B(\delta O)\right) \tag{106}
\end{array}
$$

using trace properties (102), (103), (104) and (105) given above. Hence, the trace derivative will be

$$
\frac{\delta \mathbf{P}}{\delta O}=\epsilon_{A O} B O C A+\epsilon_{C} C A O B
$$

Above, $\epsilon_{M}= \pm 1$ depending on whether the matrix $M$ is Bosonic or Fermionic.
:: Lagrangian and Hamiltonian Dynamics::

Armed with these tools we write the $L a$ grangian of a theory as a Grassmann even polynomial function of bosonic/fermionic operators $\left\{q_{r}\right\}$ and their time derivatives $\left\{\dot{q}_{r}\right\}$ with $\left\{q_{r}, \dot{q}_{r} \in \mathcal{G}_{M}\right\}$.
We go on to define the Trace Lagrangian

$$
\mathbf{L}\left[\left\{q_{r}\right\},\left\{\dot{q}_{r}\right\}\right]=\operatorname{Tr} L\left[\left\{q_{r}\right\},\left\{\dot{q}_{r}\right\}\right],
$$

and subsequently the Trace Action

$$
\mathbf{S}=\int d t \mathbf{L}
$$

Using the trace derivative we obtain the equation of motion by extremizing the action w.r.t. variation in $q_{r}$ using the differentiation technique just described above

$$
\begin{equation*}
\frac{\delta \mathbf{L}}{\delta q_{r}}=\frac{d}{d t} \frac{\delta \mathbf{L}}{\delta \dot{q}_{r}} \tag{107}
\end{equation*}
$$

Above is a matrix equation, in component form we can write down the equation of motion as

$$
\left(\frac{\delta \mathbf{L}}{\delta q_{r}}\right)_{i j}=\frac{\partial \mathbf{L}}{\partial\left(q_{r}\right)_{j i}}
$$

thus having $N^{2}$ Euler-Lagrange equations of motion if $N$ is the dimensionality of matrices $q_{r}, \dot{q}_{r}$. Further we define conjugate momenta as

$$
p_{r} \equiv \frac{\delta \mathbf{L}}{\delta \dot{q}_{r}}
$$

Since the Lagrangian is Grassmann even, momentum will be of same type (Bosonic/Fermionic) as $q_{r}$. The Trace Hamiltonian is obtained in the usual way

$$
\mathbf{H}=\sum_{r} p_{r} \dot{q}_{r}-\mathbf{L}
$$

Using the trace derivative and the definition of momentum, we get

$$
\begin{equation*}
\delta \mathbf{H}=\operatorname{Tr} \sum_{r}\left(\epsilon_{r} \dot{q_{r}} \delta p_{r}-\dot{p_{r}} \delta q_{r}\right) . \tag{108}
\end{equation*}
$$

Therefore,

$$
\begin{align*}
& \frac{\delta \mathbf{H}}{\delta q_{r}}=-\dot{p_{r}}, \\
& \frac{\delta \mathbf{H}}{\delta p_{r}}=\epsilon_{r} \dot{q}_{r}, \tag{109}
\end{align*}
$$

where $\epsilon_{r}= \pm 1$, depending upon whether $r$ is a bosonic or fermionic degree of freedom.

Note that in general the coordinates do not commute with each other, the momenta do not commute with each other, and the coordinates and momenta do not commute either, for these are all arbitrary matrices.

## 3. Hamiltonian dynamics

We define a generalized Poisson Bracket over this phase space

$$
\{\mathbf{A}, \mathbf{B}\}=\operatorname{Tr} \sum_{r} \epsilon_{r}\left(\frac{\delta \mathbf{A}}{\delta q_{r}} \frac{\delta \mathbf{B}}{\delta p_{r}}-\frac{\delta \mathbf{B}}{\delta q_{r}} \frac{\delta \mathbf{A}}{\delta p_{r}}\right) .
$$

Writing the phase space variables as $x$-s, i.e. $x_{1}=q_{1}, x_{2}=p_{1}, \ldots, x_{2 D-1}=q_{D}, x_{2 D}=p_{D}$, the generalized Poisson Bracket looks like

$$
\{\mathbf{A}, \mathbf{B}\}=\operatorname{Tr} \sum_{r, s=1}^{2 D}\left(\frac{\delta \mathbf{A}}{\delta x_{r}} \omega_{r s} \frac{\delta \mathbf{B}}{\delta x_{s}}\right)
$$

This generalized Poisson Bracket satisfies the Jacobi identity
$\{\mathbf{A},\{\mathbf{B}, \mathbf{C}\}\}+\{\mathbf{C},\{\mathbf{A}, \mathbf{B}\}\}+\{\mathbf{B},\{\mathbf{C}, \mathbf{A}\}\}=0$.

$$
\mathbf{H}=\operatorname{Tr} \sum_{r, s \in F}\left(p_{r} q_{s} B_{1 r s}+p_{r} B_{2 r s} q_{s}\right)
$$

For $\mathbf{A}\left[\left\{q_{r}\right\},\left\{p_{r}\right\}, t\right]$ one can easily verify that,

$$
\begin{equation*}
\dot{\mathbf{A}}=\frac{\partial \mathbf{A}}{\partial t}+\{\mathbf{A}, \mathbf{H}\} \tag{110}
\end{equation*}
$$

We observe that the matrix dynamics obtained above is non-unitary in general. For operators that do not have explicit time dependence, (110) does not show a unitary evolution of type

$$
\dot{x}_{r}(t)=i[G, x(t)] .
$$

However, for Weyl-ordered Hamiltonians, trace-dynamics evolution and Heisenberg unitary time evolution can be shown to be equivalent.

## 4. Conserved parameters

As an obvious result of (110), the trace Hamiltonian itself is conserved,

$$
\begin{equation*}
\dot{\mathbf{H}}=\{\mathbf{H}, \mathbf{H}\}=0 . \tag{111}
\end{equation*}
$$

Moreover, for Trace Hamiltonian restricted to bilinear form in Fermionic sector with self-adjoint kinetic part
+bosonic, (112)
the quantity Trace Fermion number

$$
\mathbf{N}=\frac{1}{2} i \operatorname{Tr} \sum_{r \in F}\left[q_{r}, p_{r}\right]
$$

is conserved, i.e. $\dot{\mathbf{N}}=0$.
This conserved charge corresponds to $U(1)$ gauge transformations of Fermionic d.o.f. :

$$
q_{r} \rightarrow \exp \{i \alpha\} q_{r}: p_{r} \rightarrow \exp \{i \alpha\} p_{r}
$$

for real and constant $\alpha$ and $r \in F$.
For such a Hamiltonian (112), the trace Lagrangian is

$$
\begin{align*}
\mathbf{L}=\operatorname{Tr} L & =\operatorname{Tr} \sum_{r \in F} p_{r} \dot{q}_{r}-\operatorname{Tr} \sum_{r, s \in F}\left(p_{r} q_{s} B_{1 r s}\right. \\
& \left.+p_{r} B_{2 r s} q_{s}\right)+ \text { bosonic. } \tag{113}
\end{align*}
$$

If we take

$$
\begin{equation*}
p_{r}=q_{r}^{\dagger} \tag{114}
\end{equation*}
$$

then,

$$
\left(p_{r} \dot{q}_{r}\right)^{\dagger}=p_{r} \dot{q}_{r}-\frac{d}{d t}\left(q_{r}^{\dagger} q_{r}\right)
$$

making the kinetic term in the trace Lagrangian (113) self-adjoint up to a total derivative,

$$
\begin{align*}
\mathbf{L}=\operatorname{Tr} L & =\operatorname{Tr} \sum_{r \in F} q_{r}^{\dagger} \dot{q}_{r}-\operatorname{Tr} \sum_{r, s \in F}\left(p_{r} q_{s} B_{1 r s}\right. \\
& \left.+p_{r} B_{2 r s} q_{s}\right)+ \text { bosonic } \tag{115}
\end{align*}
$$

and hence the conserved charge corresponding to $U(1)$ gauge transformations of Fermionic d.o.f. is

$$
\begin{align*}
\mathbf{N} & =\frac{1}{2} i \operatorname{Tr} \sum_{r \in F}\left[q_{r}, p_{r}\right]=-i \operatorname{Tr} \sum_{r \in F} p_{r} q_{r} \\
& =-i \operatorname{Tr} \sum_{r \in F} q_{r}^{\dagger} q_{r} \tag{116}
\end{align*}
$$

which resembles the number operator for Fermionic degrees of freedom.

If we demand self-adjointness of $L$ and $H$ as well, then we have to live with

$$
B_{1 r s}=-B_{1 r s}^{\dagger} ; B_{2 r s}=-B_{2 r s}^{\dagger}
$$

There is a Nöether charge corresponding to the global unitary invariance of the Trace Lagrangian (and Hamiltonian) i.e.

$$
\begin{aligned}
\mathbf{L}\left[\left\{U^{\dagger} q_{r} U\right\},\left\{U^{\dagger} \dot{q}_{r} U\right\}\right] & =\mathbf{L}\left[q_{r}, \dot{q}_{r}\right], \\
\mathbf{H}\left[\left\{U^{\dagger} q_{r} U\right\},\left\{U^{\dagger} \dot{q}_{r} U\right\}\right] & =\mathbf{H}\left[q_{r}, \dot{q}_{r}\right],(118)
\end{aligned}
$$

for some constant unitary matrix $U$. From (118) and (109) we get

$$
\frac{d}{d t} \tilde{C}=0
$$

for the important quantity

$$
\begin{equation*}
\tilde{C}=\sum_{r \in B}\left[q_{r}, p_{r}\right]-\sum_{r \in F}\left\{q_{r}, p_{r}\right\} \tag{119}
\end{equation*}
$$

known as the Adler-Millard charge. This charge will play a fundamental role in the emergence of quantum theory. In fact it is this charge, having the dimensions of action, and which is trivially zero in point particle mechanics, which makes all the difference between Trace Dynamics, and ordinary classical mechanics of point particles where all position and momenta commute with each other. Note that in (119) the individual commutators/anti-commutators take arbitrary values in time; yet the particular combination shown in this equation remains conserved.

Now, if the Fermionic d.o.f. have the adjointness property of (114), and the Bosonic d.o.f. are self-adjoint (or anti-self adjoint) then the conserved charge is anti-self adjoint

$$
\begin{equation*}
\tilde{C}=-\tilde{C}^{\dagger} \tag{120}
\end{equation*}
$$

and traceless

$$
\begin{equation*}
\operatorname{Tr} \tilde{C}=0 \tag{121}
\end{equation*}
$$

Since $\tilde{C}$ is Nöether charge corresponding to global unitary invariance of the matrix model, it can be used to construct generator of global unitary transformation

$$
\mathbf{G}_{\boldsymbol{\Lambda}}=\operatorname{Tr} \Lambda \tilde{C}
$$

The algebra of the generator is

$$
\begin{equation*}
\left\{\mathbf{G}_{\boldsymbol{\Lambda}}, \mathbf{G}_{\boldsymbol{\Sigma}}\right\}=\mathbf{G}_{[\boldsymbol{\Lambda}, \boldsymbol{\Sigma}]} . \tag{122}
\end{equation*}
$$

Now we consider those canonical transformations which have global unitary invariant generator G. For those canonical transformations clearly,

$$
\begin{equation*}
\left\{\mathbf{G}, \mathbf{G}_{\boldsymbol{\Lambda}}\right\}=0 \tag{123}
\end{equation*}
$$

Alternatively we can interpret 123 by saying that $\mathbf{G}_{\boldsymbol{\Lambda}}$ is invariant under the action of G. Then, along these lines it can be shown that $\tilde{C}$ is Poincaré invariant, where Poincaré transformations are generated by trace functional Poincaré generators. These generators are global unitary invariant. Hence, we can make use of this charge $\tilde{C}$ in Poincaré invariant theories.

If we consider a Lagrangian which has fermionic Kinetic part as

$$
\begin{equation*}
\mathbf{L}_{k i n}=\operatorname{Tr} \sum_{r, s \in F} q_{r}^{\dagger} A_{r s} \dot{q}_{s} \tag{124}
\end{equation*}
$$

where $A_{r s}$ is a constant matrix having the property (for real trace Lagrangian)

$$
A_{r s}=A_{s r}^{\dagger}
$$

then $\tilde{C}$ is still conserved but now it can have a self-adjoint part as well,

$$
\begin{equation*}
\tilde{C}+\tilde{C}^{\dagger}=-\sum_{r, s \in F}\left[q_{s} q_{r}^{\dagger}, A_{r s}\right] \tag{125}
\end{equation*}
$$

Generically, for a continuous spacetime based trace Lagrangian written in terms of trace Lagrangian density $\mathcal{L}\left(\left\{q_{l}(x)\right\},\left\{\partial_{\mu} q_{l}(x)\right\}\right)$ which is invariant under the following symmetry transformations

$$
\begin{array}{r}
q_{l}(x) \rightarrow q_{l}(x)+\alpha(x) \Delta_{l}(x) \\
\partial_{\mu} q_{l}(x) \rightarrow \partial_{\mu} q_{l}(x)+\alpha(x) \partial_{\mu} \Delta_{l}(x) \\
+\partial_{\mu} \alpha(x) \Delta_{l}(x), \tag{126}
\end{array}
$$

there is a local trace current

$$
\mathbf{J}^{\mu}=\operatorname{Tr} \sum_{l} \frac{\delta \mathcal{L}}{\delta \partial_{\mu} q_{l}(x)} \Delta_{l}(x)
$$

for which

$$
\partial_{\mu} \mathbf{J}^{\mu}=0
$$

suggesting that there is a conserved charge

$$
Q=\int d^{3} x \mathbf{J}^{0}(x)
$$

For a global unitary and Poincaré invariant theory the conserved charges are the following

$$
\begin{array}{r}
\tilde{C}=\int d^{3} x \sum_{l}\left(\epsilon_{l} q_{l} p_{l}-p_{l} q_{l}\right) \\
\mathbf{P}^{\mu}=\int d^{3} x \mathcal{T}^{0 \mu} \\
\mathbf{M}^{\mu \nu}=\int d^{3} x \mathcal{M}^{0 \mu \nu} \tag{129}
\end{array}
$$

where the momentum conjugate to $q_{l}(x)$ is

$$
\begin{equation*}
p_{l}(x)=\frac{\delta \mathcal{L}}{\delta \partial_{0} q_{l}(x)} ; \tag{130}
\end{equation*}
$$

trace energy momentum density is

$$
\begin{equation*}
\mathcal{T}^{\mu \nu}=\eta^{\mu \nu} \mathcal{L}-\operatorname{Tr} \sum_{l} \frac{\delta \mathcal{L}}{\delta \partial_{\mu} q_{l}} \partial^{\nu} q_{l} \tag{131}
\end{equation*}
$$

$\eta^{\mu \nu}=\operatorname{diag}(1,1,1,-1)$ is Minkowski metric and
$\mathcal{M}^{\lambda \nu \mu}=x^{\mu} \mathcal{T}^{\lambda \nu}-x^{\nu} \mathcal{T}^{\lambda \mu}+\operatorname{Tr} \sum_{l m} \frac{\delta \mathcal{L}}{\delta \partial_{\lambda} q_{l}} \chi_{l m}^{\nu \mu} q_{m}$
is the trace angular momentum density, with $\chi_{l m}$ being the matrix characterizing the intrinsic spin structure of field $q_{l}$ such that under four space rotation

$$
\begin{array}{r}
x_{\mu} \rightarrow x_{\mu}^{\prime}=x_{\mu}+\omega_{\mu \nu} x^{\nu} \\
q_{l}\left(x^{\prime}\right)=q_{l}(x)+(1 / 2) \omega_{\mu \nu} \sum_{m} \chi_{l m}^{\mu \nu} q_{m}(x), \tag{133}
\end{array}
$$

for antisymmetric infinitesimal rotation parameter $\omega_{\mu \nu} . \quad \mathbf{P}^{0}$ is the conserved trace Hamiltonian. $\mathbf{P}^{\mu}$ and $\mathbf{M}^{\mu \nu}$ together form complete set of Poincaré generators.

## 5. Phase space

Matrix operator phase space is well behaved. We can define on this phase space a measure

$$
\begin{align*}
d \mu & =\prod d \mu^{A} \\
d \mu^{A} & =\prod_{r, m, n} d\left(x_{r}\right)_{m n} \tag{134}
\end{align*}
$$

where $A=0,1$ for

$$
\left(x_{r}\right)_{m n}=\left(x_{r}\right)_{m n}^{0}+i\left(x_{r}\right)_{m n}^{1} .
$$

This measure is invariant under canonical transformation (Louiville's theorem)

$$
d \mu\left[\left\{x_{r}+\delta x_{r}\right\}\right]=d \mu\left[\left\{x_{r}\right\}\right] .
$$

Further, bosonic and fermionic measures can be separated

$$
d \mu=d \mu_{B} d \mu_{F}
$$

which are separately invariant under above assumed adjointness properties. Clearly this measure is invariant under dynamic evolution of the system as time evolution is a canonical transformation generated by $d t \mathbf{H}$.

At this stage, we have in hand a description of the classical dynamics [called Trace Dynamics because of the introduction of the trace derivative] of matrices. In essentially all respects but one, it is analogous to ordinary point-particle dynamics. The exception is the presence of the Adler-Millard charge (119) in Trace Dynamics, which makes all the
difference in the following discussion, which demonstrates the emergence of quantum theory. We will construct a statistical mechanics, assuming that we are interested, not in the microscopic description of Trace Dynamics, but only in a coarse-grained average.

## 6. Canonical Ensemble

With the matrix equations of motion 109 for time evolution in trace dynamics theory we study the evolution of phase space distribution. We assume that a large enough system rapidly forgets its initial distribution and the time averages of physical quantities are equal to the statistical averages over an equilibrium ensemble which is determined by maximizing combinatoric probability subject to conservation laws. If

$$
d P=d \mu\left[\left\{x_{r}\right\}\right] \rho\left[\left\{x_{r}\right\}\right]
$$

is the probability of finding the system in operator phase space volume element $d \mu\left[\left\{x_{r}\right\}\right]$, then,

$$
\int d P=1
$$

For a system in statistical equilibrium, phase space density distribution is constant

$$
\dot{\rho}\left[\left\{x_{r}\right\}\right]=0 .
$$

Hence, $\rho$ depends only upon conserved operators, conserved trace functionals and constant parameters. By going to a frame where
the system is not translating, accelerating or rotating the charges associated with Poincaré symmetry can be put to zero. In that case

$$
\rho=\rho(\tilde{C}, \mathbf{H}, \mathbf{N})
$$

In addition the distribution function of dynamical variables can depend on constant parameters.

$$
\rho=\rho(\operatorname{Tr} \tilde{\lambda} \tilde{C} ; \mathbf{H}, \tau ; \mathbf{N}, \eta)
$$

where $\tilde{\lambda}, \tau$ and $\eta$ are the Lagrange multipliers conjugate to $\tilde{C}, \mathbf{H}$ and $\mathbf{N}$ respectively. One important aspect to note is that while $\mathbf{H}$ and $\mathbf{N}$ belong to $\mathcal{G}_{C}, \tilde{C} \in \mathcal{G}_{M}$. Hence, $\tau, \eta \in \mathcal{G}_{C}$ while $\tilde{\lambda} \in \mathcal{G}_{M}$. The dependence of $\rho$ on $\operatorname{Tr} \tilde{\lambda} \tilde{C}$ is motivated from global unitary invariance. If $\tilde{C}$ has a self-adjoint part as well, one can break it into its self-adjoint ( $s a$ ) and anti-self adjoint (asa) parts,

$$
\rho=\rho\left(\operatorname{Tr} \tilde{\lambda}^{s a} \tilde{C}^{s a}, \operatorname{Tr} \tilde{\lambda}^{a s a} \tilde{C}^{a s a} ; \mathbf{H}, \tau ; \mathbf{N}, \eta\right)
$$

Next, we define ensemble average of an operator $O$ as

$$
\langle O\rangle=\int d \mu \rho O
$$

which is a map from $\mathcal{G}_{M}$ to $\mathcal{G}_{M}$.
This ensemble average has nice properties
(i) When $O$ is constructed only from phase space variables $\left\{x_{r}\right\}$ then this ensemble average depends only on the constant parameters $\bar{\lambda}=\{\tilde{\lambda}, \tau, \eta\}$

$$
\langle O\rangle=F_{O}(\bar{\lambda})
$$

(ii) Since the integration measure is unitary invariant and $O$ is made up of only $\left\{x_{r}\right\}$ using $c$ - number coefficients, under global unitary transformation

$$
F_{O}(\bar{\lambda})=U F_{O}\left(U^{-1} \bar{\lambda} U\right) U^{-1}
$$

(iii) As a consequence $F_{O}(\bar{\lambda})$ depends explicitly on $\bar{\lambda}$ only and hence commutes with $\tilde{\lambda}$

$$
[\bar{\lambda},\langle O\rangle]=0
$$

Taking a specific case when $O=\tilde{C}$

$$
[\bar{\lambda},\langle\tilde{C}\rangle]=0
$$

$\bar{\lambda}$ being \{anti- $\}$ self-adjoint can be diagonalized by unitary transformation, hence so will be $\langle\tilde{C}\rangle$. Again specializing to anti-self-adjoint $\bar{\lambda}$ and $\langle\tilde{C}\rangle$ we get
(i) for a real, non-negative, diagonal magnitude operator $D_{\text {eff }}$ and unitary diagonal phase-operator $i_{e f f}$,

$$
\langle\tilde{C}\rangle=i_{e f f} D_{e f f}
$$

(ii) Since, $\tilde{C}$ is traceless,

$$
\operatorname{Tr}\left(i_{e f f} D_{e f f}\right)=0
$$

(iii) anti-self adjointness of $\tilde{C}$ is ensured with

$$
i_{e f f}=-i_{e f f}^{\dagger}
$$

and

$$
\left[i_{e f f}, D_{e f f}\right]=0
$$

(iv) As a consequence of this decomposition

$$
i_{e f f}^{2}=-I
$$

For an ensemble symmetric in Hilbert space basis, i.e. the ensemble does not prefer any state, the averaged operator should have identical entries as eigenvalues. Therefore,

$$
D_{e f f}=k I
$$

Clearly, $D_{\text {eff }}$ is determined by a single real number with dimension of action

$$
\begin{equation*}
\langle\tilde{C}\rangle=i_{e f f} \hbar \tag{135}
\end{equation*}
$$

The constant $\hbar$ will eventually be identified with Planck's constant.

The traceless $\langle\tilde{C}\rangle$ implies

$$
\operatorname{Tr}_{i}{ }_{e f f}=0
$$

The above mentioned properties of $i_{e f f}$ alongwith property (iv) above forces the dimension of Hilbert space to be even and uniquely fixes $i_{e f f}$ to

$$
i_{e f f}=i[\operatorname{diag}(1,-1,1,-1, \ldots, 1,-1)]
$$

Next, we obtain the functional form of $\rho$ through maximizing entropy defined as

$$
\begin{equation*}
S=\int d \mu \rho \log \rho \tag{136}
\end{equation*}
$$

subject to the following constraints

$$
\begin{aligned}
\int d \mu \rho & =1 \\
\int d \mu \rho \tilde{C} & =\langle\tilde{C}\rangle
\end{aligned}
$$

$$
\int d \mu \rho \mathbf{H}=\langle\mathbf{H}\rangle
$$

$$
\int d \mu \rho \mathbf{N}=\langle\mathbf{N}\rangle
$$

which on some calculations gives

$$
\begin{aligned}
\rho & =Z^{-1} \exp (-\operatorname{Tr} \tilde{\lambda} \tilde{C}-\tau \mathbf{H}-\eta \mathbf{N}) \\
Z & =\int d \mu \exp (-\operatorname{Tr} \tilde{\lambda} \tilde{C}-\tau \mathbf{H}-\eta \mathbf{N})
\end{aligned}
$$

Using this distribution and partition function we can evaluate ensemble averages

$$
\begin{align*}
\langle\tilde{\lambda}\rangle & =-\frac{\delta \log Z}{\delta \tilde{\lambda}}  \tag{137}\\
\langle\mathbf{H}\rangle & =-\frac{\partial \log Z}{\partial \tau}  \tag{138}\\
\langle\mathbf{N}\rangle & =-\frac{\partial \log Z}{\partial \eta} \tag{139}
\end{align*}
$$

and the mean square fluctuations

$$
\begin{gather*}
\Delta_{\tilde{P} \tilde{C}}^{2} \equiv\left(\operatorname{Tr} \tilde{P} \frac{\delta}{\delta \tilde{\lambda}}\right)^{2} \log Z  \tag{140}\\
\Delta_{\mathbf{H}}^{2} \equiv \frac{\partial^{2} \log Z}{(\partial \tau)^{2}}  \tag{141}\\
\Delta_{\mathbf{N}}^{2} \equiv \frac{\partial^{2} \log Z}{(\partial \eta)^{2}} \tag{142}
\end{gather*}
$$

as analogous to statistical mechanics, with $\tilde{P}$ being any arbitrary fixed self-adjoint operator.

We can further introduce a book-keeping matrix source term $j_{r}$ for each matrix variable $x_{r}$ of same type $(B / F)$ and adjointness that can be varied and set to zero, s.t.

$$
\begin{gather*}
\rho_{j}=Z^{-1} \exp (-\operatorname{Tr} \tilde{\lambda} \tilde{C}- \\
\left.\tau \mathbf{H}-\eta \mathbf{N}-\sum_{r} \operatorname{Tr} j_{r} x_{r}\right)  \tag{143}\\
Z_{j}=\int d \mu \exp (-\operatorname{Tr} \tilde{\lambda} \tilde{C}- \\
\left.\tau \mathbf{H}-\eta \mathbf{N}-\sum_{r} \operatorname{Tr} j_{r} x_{r}\right) \tag{144}
\end{gather*}
$$ coefficients. Since $D_{\text {eff }}$ is constant times identity,

$$
\begin{equation*}
\tilde{\lambda}=\lambda i_{e f f} \tag{146}
\end{equation*}
$$

for a real c-number $\lambda$. Now for a unitary matrix $U_{\text {eff }}$ that commutes with $i_{e f f}$

$$
\begin{equation*}
\left[U_{e f f}, \tilde{\lambda}\right]=0 \Rightarrow U_{e f f} \tilde{\lambda} U_{e f f}^{\dagger}=\tilde{\lambda} \tag{147}
\end{equation*}
$$

Clearly, presence of $\operatorname{Tr} \tilde{\lambda} \tilde{C}$ term in the partition function breaks the global unitary invariance since under,

$$
q_{r} \rightarrow U^{\dagger} q_{r} U, p_{r} \rightarrow U^{\dagger} p_{r} U, \tilde{C} \rightarrow U^{\dagger} \tilde{C} U
$$

$$
\begin{equation*}
\operatorname{Tr} \tilde{\lambda} \tilde{C}=\lambda \operatorname{Tr} i_{e f f} \tilde{C} \rightarrow \lambda \operatorname{Tr} U i_{e f f} U^{\dagger} \tilde{C} \tag{148}
\end{equation*}
$$

Thus, presence of this term breaks the global unitary invariance to $\left\{U_{e f f}\right\}$. With this residual invariance in canonical ensemble we define an integration measure as

$$
d \mu=d\left[U_{e f f}\right] d \hat{\mu},
$$

with $d\left[U_{e f f}\right]$ as the Haar measure on the residual symmetry group and $d \hat{\mu}$ is the measure over operator phase space when overall global unitary transformation $U_{\text {eff }}$ is kept fixed.

In this case, for a polynomial $R_{e f f}$ which is a function of $i_{e f f}$ and dynamic variables $\left\{x_{r}\right\}$,

$$
\begin{equation*}
R_{e f f A V} \equiv \frac{\int d\left[U_{e f f}\right] d \hat{\mu} \rho R_{e f f}}{\int d\left[U_{e f f}\right] d \hat{\mu} \rho} \tag{149}
\end{equation*}
$$

Now if we fix the $U_{\text {eff }}$ rotation

$$
q_{r}=U_{e f f}^{\dagger} \hat{q}_{r} U_{e f f}, \quad p_{r}=U_{e f f}^{\dagger} \hat{p}_{r} U_{e f f}
$$

resulting in,

$$
R_{e f f}=U_{e f f}^{\dagger} \hat{R}_{e f f} U_{e f f}
$$

and hence,

$$
R_{e f f A V} \equiv \frac{\int d\left[U_{e f f}\right] U^{\dagger} R_{e f f A V} U_{e f f}}{\int d\left[U_{e f f}\right]}
$$

In above equation,

$$
R_{e f f A \hat{}} \equiv \frac{\int d \hat{\mu} \hat{\rho} \hat{R}_{e f f}}{\int d \hat{\mu} \hat{\rho}}
$$

where $\hat{R}_{e f f}$ and $\hat{\rho}$ are obtained from $R_{e f f}$ and $\rho$ by replacing $q, p$ therein by $\hat{q}, \hat{p}$ as defined above.

Now, a general matrix $M$ can be decomposed in the form

$$
\begin{align*}
M= & \frac{1}{2}\left(\sigma_{0}+\sigma_{3}\right) M_{+}+\frac{1}{2}\left(\sigma_{0}-\sigma_{3}\right) M_{-}+ \\
& \sigma_{1} M_{1}+\sigma_{2} M_{2} \tag{150}
\end{align*}
$$

where $\sigma_{i}, i=1,2,3$ are $2 \times 2$ Pauli matrices and $\sigma_{0}=1_{2}$.
This can further be re-written in terms of

$$
M_{e f f}=\frac{1}{2}\left(\sigma_{0}+\sigma_{3}\right) M_{+}+\frac{1}{2}\left(\sigma_{0}-\sigma_{3}\right) M_{-}
$$

and

$$
M_{12}=\sigma_{1} M_{1}+\sigma_{2} M_{2}
$$

These new matrices satisfy

$$
\begin{array}{r}
{\left[i_{e f f}, M_{e f f}\right]=0} \\
\left\{i_{e f f}, M_{12}\right\}=0 \\
2 i_{e f f} M_{e f f}=\left\{i_{e f f}, M_{e f f}\right\} . \tag{153}
\end{array}
$$

In irreducible systems unitary fixing can be done by fixing the global unitary rotation of one canonical pair of dynamic variables. With the restricted measure we need to know the restricted canonical average $\tilde{C}_{\hat{A V}}$. Since unitary fixing does not disturb $i_{e f f}$, for any operator $O$ made up of $\left\{x_{r}\right\}$ using $c$-number coefficients,

$$
\left[\tilde{\lambda}, O_{\hat{A V}}\right]=0
$$

When $O=\tilde{C}$,

$$
\left[\tilde{\lambda}, \tilde{C}_{\hat{A V}}\right]=0
$$

Using properties 153, most general $\tilde{C}_{A V}$ commuting with $\tilde{\lambda}$ is
$\tilde{C}_{\hat{A V}}=i_{e f f} \hbar+\frac{1}{2}\left(\sigma_{0}+\sigma_{3}\right) \Delta_{+}+\frac{1}{2}\left(\sigma_{0}-\sigma_{3}\right) \Delta_{-}$.

Thus, the ensemble average in unitary fixed system is different from the canonical ensemble average (135). However, since the unitary fixing has been done by restricting only one canonical pair, in systems involving large number of canonical pairs, the restricted average should be close to unrestricted average. Therefore, $\Delta_{ \pm}$should be small.

The canonical ensemble that has been constructed here broadly resembles the construction that would follow from the statistical thermodynamics of an ordinary classical system, though once again with the exception of the Adler-Millard charge $\tilde{C}$, and the emergence of its canonical average 135 .
7. General Ward Identity and Emergence of Quantum Theory

Thus far we have progressed from the classical theory of Trace Dynamics, to developing a statistical thermodynamics of this theory. We now have in hand the tools necessary to describe the emergence of quantum theory in this thermodynamic approximation. The first step is the derivation of a crucial general Ward identity. This identity should be thought of as an analog of the
equipartition theorem in statistical mechanics, and its implications in the present context are deeply connected with the existence of the Adler-Millard charge and its canonical average (135). With the restricted measure, average is defined as

$$
\begin{equation*}
\langle O\rangle_{A V}=\int d \hat{\mu} \rho_{j} O \tag{155}
\end{equation*}
$$

with
$\rho_{j}=Z^{-1} \exp \left(-\operatorname{Tr} \tilde{\lambda} \tilde{C}-\tau \mathbf{H}-\eta \mathbf{N}-\sum_{r} \operatorname{Tr} j_{r} x_{r}\right)$
$Z_{j}=$
$\int d \hat{\mu} \exp \left(-\operatorname{Tr} \tilde{\lambda} \tilde{C}-\tau \mathbf{H}-\eta \mathbf{N}-\sum_{r} \operatorname{Tr} j_{r} x_{r}\right)$.
Therefore,

$$
\langle O\rangle_{\hat{A V}}=\langle O\rangle_{\hat{A V}, j=0} .
$$

Under constant shift of any matrix variable apart from the restricted pair

$$
x_{r} \rightarrow x_{r}+\delta x_{r}, r \neq R, R+1
$$

$$
\begin{equation*}
\int d \hat{\mu} \delta_{x_{r}}\left(\rho_{j} O\right)=0 \tag{156}
\end{equation*}
$$

Using this and the definition of average given above for an $O=\left\{\tilde{C}, i_{e f f}\right\} W$

$$
\begin{array}{r}
Z_{j}\left\langle\operatorname{Tr}\left\{\tilde{C}, i_{e f f}\right\} W\right\rangle_{j}=\int d \hat{\mu} \exp (-\operatorname{Tr} \tilde{\lambda} \tilde{C}-\tau \mathbf{H} \\
\left.-\eta \mathbf{N}-\sum_{r} \operatorname{Tr} j_{r} x_{r}\right) \operatorname{Tr}\left\{\tilde{C}, i_{e f f}\right\} W
\end{array}
$$

after taking trace. Using (156) we get

$$
\begin{align*}
& \int d \mu \delta_{x_{s}}[\exp (-\operatorname{Tr} \tilde{\lambda} \tilde{C}-\tau \mathbf{H}-\eta \mathbf{N} \\
& \left.\left.-\sum_{r} \operatorname{Tr} j_{r} x_{r}\right) \operatorname{Tr}\left\{\tilde{C}, i_{e f f}\right\} W\right]=0
\end{align*}
$$

Using chain rule we have

$$
\begin{array}{r}
\int d \mu \exp (-\operatorname{Tr} \tilde{\lambda} \tilde{C}-\tau \mathbf{H}-\eta \mathbf{N} \\
\left.-\sum_{r} \operatorname{Tr} j_{r} x_{r}\right)\left[\left(-\operatorname{Tr} \tilde{\lambda} \delta_{x_{s}} \tilde{C}-\tau \delta_{x_{s}} \mathbf{H}\right.\right. \\
\left.-\eta \delta_{x_{s}} \mathbf{N}-\operatorname{Tr} j_{s} \delta x_{s}\right)\left\{\tilde{C}, i_{e f f}\right\} W \\
\left.+\delta_{x_{s}}\left\{\tilde{C}, i_{e f f}\right\} W\right]=0 .(1 \tag{158}
\end{array}
$$

Evaluating term by term in (158) we have,

$$
\begin{array}{r}
\operatorname{Tr} \tilde{\lambda} \delta_{x_{s}} \tilde{C}=\operatorname{Tr}\left[\tilde{\lambda}, \sum_{r} \omega_{r s} x_{r}\right] \delta x_{s} \\
\delta_{x_{s}} \mathbf{H}=\sum_{r} \omega_{r s} \operatorname{Tr} \dot{x_{r}} \delta x_{s} \\
\delta_{x_{s}} \mathbf{N}=\sum_{r} \tilde{\omega}_{r s} \operatorname{Tr} x_{r} \delta x_{s} \\
\delta_{x_{s}}\left\{\tilde{C}, i_{e f f}\right\} W=\operatorname{Tr}\left(\left\{i_{e f f}, W\right\} \delta_{x_{s}} \tilde{C}+\right. \\
\left.\left\{\tilde{C}, i_{e f f}\right\} \delta_{x_{s}} W\right), \tag{159}
\end{array}
$$

with, further, for a polynomial $W$

$$
\delta_{x_{s}} W=\sum_{l} W_{s}^{L l} \delta x_{s} \sum_{l} W_{s}^{R l}
$$

where $l$ labels each monomial in the polynomial. $W_{s}^{L l}$ and $W_{s}^{R l}$ are the left and right fractions of a monomial.
For example, for

$$
W=A O B O C O
$$

for constant matrices $A, B, C$,

$$
\begin{align*}
\delta_{O} W=A \delta O B O C O & +A O B \delta O C O \\
& +A O B O C \delta O \tag{160}
\end{align*}
$$

Therefore,

$$
W^{L 1}=A, W^{R 1}=B O C O
$$

$$
W^{L 2}=A O B, W^{R 2}=C O
$$

$$
W^{L 3}=A O B O C, W^{R 3}=1
$$

Collecting above terms and plugging back in (158) with some manipulations leads to the generalized WARD IDENTITY

$$
\left\langle\Lambda_{u_{e f f}}\right\rangle_{j}=0
$$

where,

$$
\begin{aligned}
& \Lambda_{u_{e f f}}=\left(-\tau \dot{x}_{u e f f}+i \eta \xi_{u} x_{u e f f}-\Sigma_{s} \omega_{u s} j_{\text {seff }}\right) \\
& \\
& \quad \operatorname{Tr} \tilde{C} i_{e f f} W_{e f f}+\left[i_{e f f} W_{e f f}, x_{u e f f}\right]+ \\
& \quad \sum_{s, l} \omega_{u s} \epsilon_{l}\left(W_{s}^{R l} \frac{1}{2}\left\{\tilde{C}, i_{e f f}\right\} W_{s}^{L l}\right) \\
& \text { with } \\
& \qquad \\
& \qquad=\operatorname{diag}\left(\Omega_{B}, \ldots, \Omega_{B}, \Omega_{F}, \ldots, \Omega_{F}\right)
\end{aligned}
$$

and,

$$
\tilde{\omega}=\operatorname{diag}\left(0, \ldots, 0, \Omega_{B}, \ldots, \Omega_{B}\right),
$$

where

$$
\Omega_{B}=\left(\begin{array}{ll}
0 & 1 \\
-1 & 0
\end{array}\right), \Omega_{F}=-\left(\begin{array}{ll}
0 & 1 \\
1 & 0
\end{array}\right)
$$

with,

$$
2 i_{e f f} M_{e f f}=\left\{i_{e f f}, M\right\}
$$

$\xi_{u}=1(-1)$ for fermionic $q(p)$ and zero for bosonic $x_{u}$. As a result

$$
\begin{gather*}
\sum_{s} \omega_{u s} \omega_{r s}=\delta_{u r}  \tag{161}\\
\sum_{s} \omega_{u s} \tilde{\omega}_{r s}=-\xi_{u} \delta_{u r} \tag{162}
\end{gather*}
$$

In more compact form the Ward Identity can be written as

$$
\begin{equation*}
\left\langle\mathcal{D} x_{u e f f}\right\rangle_{j}-\sum_{s} \omega_{u s} j_{\text {seff }}\left\langle\operatorname{Tr} \tilde{C} i_{e f f} W_{e f f}\right\rangle_{j}=0 \tag{163}
\end{equation*}
$$

where,

$$
\begin{gathered}
\mathcal{D} x_{u e f f}=\left(-\tau x_{u e f f}+i \eta \xi_{u} x_{u e f f}\right) \\
\operatorname{Tr} \tilde{C} i_{e f f} W_{e f f}+\left[i_{e f f} W_{e f f}, x_{u e f f}\right]+ \\
\sum_{s, l} \omega_{u s} \epsilon_{l} W_{s}^{R l} \frac{1}{2}\left\{\tilde{C}, i_{e f f}\right\} W_{s}^{L l} \cdot(1
\end{gathered}
$$

It can be shown from (163) that for a polynomial $S$ made up of $x_{r e f f}$ with $c$-number coefficients,

$$
\begin{equation*}
\left\langle S_{L}\left(x_{t e f f}\right)\left(\mathcal{D} S\left(x_{r e f f}\right)\right) S_{R}\left(x_{t e f f}\right)\right\rangle_{A V, 0}=0 \tag{165}
\end{equation*}
$$

for left and right decompositions $S_{L}\left(x_{t e f f}\right), S_{R}\left(x_{t e f f}\right) \quad$ of the polynomial $S$.
We now make the following realistic and important assumptions regarding (163), which lead to a drastic simplification
(i) Support properties of $\dot{x}_{u e f f}$ and $\tilde{C}_{e f f}$ are such that

$$
-\tau \dot{x}_{u e f f} \operatorname{Tr} \tilde{C} i_{e f f} W_{e f f}
$$

in (164) can be neglected.
(ii) Chemical potential $\eta$ is very small, such that the term

$$
i \eta \xi_{u} x_{u e f f} \operatorname{Tr} \tilde{C} i_{e f f} W_{e f f}
$$

in (164) can be neglected. In fact, for bosonic d.o.f. this term vanishes and is taken to be
small for fermionic d.o.f.
(iii) When number of degrees of freedom is large, $\tilde{C}$ can be replaced by its zero-source ensemble average

$$
\left\langle\tilde{C}_{e f f}\right\rangle_{\hat{A V}}=i_{e f f} \hbar
$$

With these assumptions the identity (163) simplifies to
$\mathcal{D} x_{u e f f}=i_{e f f}\left[W_{e f f}, x_{u e f f}\right]-\hbar \sum_{s} \omega_{u s}\left(\frac{\delta \mathbf{W}}{\delta x_{s}}\right)_{e f f}$
Important physical consequences emerge for different choices of $W$. If we consider $W=H$ in the Ward identity, we obtain

$$
\mathcal{D} x_{u e f f}=i_{e f f}\left[H_{e f f}, x_{u e f f}\right]-\hbar \dot{x}_{e f f}
$$

which gives the effective Heisenberg equation of motion for the dynamics when sandwiched between $S_{L}\left(x_{t e f f}\right)$ and $S_{R}\left(x_{t e f f}\right)$ and averaged over zero-source ensemble.
For an arbitrary polynomial function $P_{\text {eff }}$ made up of $x_{r e f f}$

$$
\begin{array}{r}
\left\langle S_{L}\left(x_{t e f f}\right) \dot{P}_{e f f} S_{R}\left(x_{t e f f}\right)\right\rangle_{\hat{A V, 0}} \\
=\left\langle S_{L}\left(x_{t e f f}\right) i_{e f f} \hbar^{-1}\left[H_{e f f}, P_{e f f}\right] S_{R}\left(x_{t e f f}\right)\right\rangle_{\hat{A V}, 0}, \tag{166}
\end{array}
$$

suggesting that within our assumptions $H_{e f f}$ is a constant of motion. Next, for $W=\sigma_{v} x_{v}$ for some $c$-number parameter $\tilde{\sigma_{v}}$

$$
\begin{equation*}
i_{e f f} \mathcal{D} x_{u e f f}=\left[x_{u e f f}, \tilde{\sigma_{v}} x_{v e f f}\right]-i_{e f f} \hbar \omega_{u v} \tilde{\sigma}_{v} \tag{167}
\end{equation*}
$$

Thus, when multiplied with $S_{L}\left(x_{t e f f}\right)$ and $S_{R}\left(x_{t e f f}\right)$ and averaged over zero source ensemble EFFECTIVE CANONICAL COMMUTATORS EMERGE

$$
\begin{align*}
\left\{q_{u e f f}, q_{v e f f}\right\} & =\left\{p_{u e f f}, p_{v e f f}\right\}=0 \\
\left\{q_{u e f f}, p_{v e f f}\right\} & =i_{e f f} \hbar \delta_{u v} \tag{168}
\end{align*}
$$

for $u, v$ fermionic, and

$$
\begin{gather*}
{\left[q_{u e f f}, q_{v e f f}\right]=\left[p_{u e f f}, p_{v e f f}\right]=0,} \\
{\left[q_{u e f f}, p_{v e f f}\right]=i_{e f f} \hbar \delta_{u v}} \tag{169}
\end{gather*}
$$

for $u, v$ bosonic.

It is important to emphasize that these commutation relations emerge only upon statistical averaging, as a consequence of there being the conserved Adler-Millard charge. At the level of the underlying theory of Trace Dynamics, the commutators/anticommutators amongst the above operators are arbitrary.

Next, let $W=G$ be a self-adjoint operator such that $\mathbf{G}$ generates canonical transformation

$$
\begin{equation*}
\hbar^{-1} \mathcal{D} x_{u e f f}=i \hbar^{-1}\left[G_{e f f}, x_{e f f}\right]-\delta x_{e f f} \tag{170}
\end{equation*}
$$

Thus, on sandwiching between $S_{L}\left(x_{t e f f}\right)$ and $S_{R}\left(x_{t e f f}\right)$ that do not contain $x_{u}$, and averaging we see that infinitesimal canonical transformations at the ensemble level and within the above-mentioned assumptions are generated by unitary transformations

$$
U_{\text {can eff }}=\exp \left(i_{e f f} \hbar^{-1} G_{e f f}\right)
$$

Therefore, we have at hand the essential features of quantum field theory. The commutator/ anti-commutator structure, time evolution in Heisenberg picture and unitary generation of canonical transformations emerge when we do the statistical thermodynamics of the matrix variables.

Now we make following correspondences between operator polynomials in trace dynamics and operator polynomials in quantum field theory

$$
S\left(\left\{x_{r e f f}\right\}\right) \Leftrightarrow S\left(\left\{X_{r e f f}\right\}\right)
$$

with $X_{\text {reff }}$ being quantized operators in quantum field theory. Here, $i_{e f f}$ acts as two blocks of $i$ and $-i$. With the assumptions
(a) in the continuum limit the trace Lagrangian is Poincaré invariant,
(b) the Hamiltonian $H_{e f f}$ is bounded from below by magnitude of the corresponding effective three-momentum operator $\vec{P}_{e f f}$, and there is a unique eigenvector $\psi_{0}$ with lowest eigenvector of $H_{\text {eff }}$ and zero eigenvalue of $\vec{P}_{e f f}$,
we also have a correspondence between trace dynamics canonical averages and Wightman functions in emergent quantum field theory, $\psi_{0}^{\dagger}\left\langle S\left(\left\{x_{r e f f}\right\}\right)\right\rangle_{\hat{A V}} \psi_{0}=\langle v a c| S\left(\left\{X_{r e f f}\right\}\right)|v a c\rangle$.

Using (167), 168) and (169) and the correspondence proposed we have at the quantum level

$$
\begin{equation*}
\left[X_{u e f f}, \tilde{\sigma}_{v} X_{v e f f}\right]=i_{e f f} \hbar \omega_{u v} \tilde{\sigma}_{v} \tag{171}
\end{equation*}
$$

which gives the commutators at the quantum and, level

$$
\begin{align*}
\left\{Q_{u e f f}, Q_{v e f f}\right\} & =\left\{P_{u e f f}, P_{v e f f}\right\}=0 \\
\left\{Q_{u e f f}, P_{v e f f}\right\} & =i_{e f f} \hbar \delta_{u v} \tag{172}
\end{align*}
$$

for $u, v$ fermionic,

$$
\begin{align*}
{\left[Q_{u e f f}, Q_{v e f f}\right] } & =\left[P_{u e f f}, P_{v e f f}\right]=0, \\
{\left[Q_{u e f f}, P_{v e f f}\right] } & =i_{e f f} \hbar \delta_{u v} \tag{173}
\end{align*}
$$

for $u, v$ bosonic.
Time evolution is given by

$$
\begin{equation*}
\dot{X}_{u e f f}=i_{e f f} \hbar^{-1}\left[H_{e f f}, X_{u e f f}\right] \tag{174}
\end{equation*}
$$

or equivalently for a polynomial $S_{\text {eff }}$ of $\left\{X_{e f f}\right\}$

$$
\begin{equation*}
\dot{S}_{u e f f}=i_{e f f} \hbar^{-1}\left[H_{e f f}, S_{u e f f}\right] \tag{175}
\end{equation*}
$$

For fermionic anti-commutator to be operator equation, following adjointness assignment is required,

$$
\begin{gather*}
\psi_{r e f f}=q_{r e f f} \leftrightarrow Q_{r e f f}=\Psi_{r e f f}, \\
\psi_{r e f f}^{\dagger}=p_{r e f f} \leftrightarrow P_{r e f f}=i_{e f f} \Psi_{r e f f}^{\dagger}, \tag{176}
\end{gather*}
$$

such that

$$
\begin{equation*}
\left\{\Psi_{u e f f}, \Psi_{v e f f}^{\dagger}\right\}=\hbar \delta_{u v} \tag{177}
\end{equation*}
$$

For the bosonic sector we obtain creation and annihilation operators $A_{\text {reff }}, A_{\text {reff }}^{\dagger}$ such that,

$$
\begin{array}{r}
Q_{u e f f}=\frac{1}{\sqrt{2}}\left(A_{r e f f}+A_{r e f f}^{\dagger}\right) \\
P_{u e f f}=\frac{1}{\sqrt{2} i_{e f f}}\left(A_{r e f f}-A_{r e f f}^{\dagger}\right) \tag{178}
\end{array}
$$

$$
\begin{gather*}
{\left[A_{u e f f}, A_{v e f f}\right]=\left[A_{u e f f}^{\dagger}, A_{v e f f}^{\dagger}\right]=0} \\
{\left[A_{u e f f}, A_{v e f f}^{\dagger}\right]=\hbar \delta_{u v}} \tag{179}
\end{gather*}
$$

Thus, we have correct commutation/anticommutation rules for bosonic/fermionic d.o.f. in both the $i_{e f f}= \pm i$ sectors. Once we have Heisenberg equation of motion (174) and (175), we make transition to Schrödinger picture as usual without making any reference to the background trace dynamics theory.
When the effective Hamiltonian has no time dependence, we define

$$
\begin{equation*}
U_{e f f}(t)=\exp \left(-i_{e f f} \hbar^{-1} t H_{e f f}\right) \tag{180}
\end{equation*}
$$

so that,

$$
\begin{align*}
\frac{d}{d t} U_{e f f}(t) & =-i_{e f f} \hbar^{-1} H_{e f f} U_{e f f}(t) \\
\frac{d}{d t} U_{e f f}^{\dagger}(t) & =i_{e f f} \hbar^{-1} U_{e f f}^{\dagger}(t) H_{e f f} \tag{181}
\end{align*}
$$

From Heisenberg picture time-independent state vector $\psi$ and time dependent operator $S_{\text {eff }}(t)$, in Schrödinger picture we do the construction

$$
\begin{aligned}
\psi_{S c h r}(t) & =U_{e f f}(t) \psi \\
S_{e f f ~ S c h r} & =U_{e f f}(t) S_{e f f}(t) U_{e f f}^{\dagger}(t),
\end{aligned}
$$

which gives,

$$
\begin{align*}
i_{e f f} \hbar \frac{d}{d t} \psi_{S c h r}(t) & =H_{e f f} \psi_{S c h r}(t) \\
\frac{d}{d t} S_{e f f ~ S c h r} & =0 \tag{183}
\end{align*}
$$

For obtaining the Schrödinger equation, we make contact with space-time by taking the label $r$ as $\vec{x}$. In that case, fermionic anticommutator becomes

$$
\begin{equation*}
\left\{\Psi_{e f f}(\vec{x}), \Psi_{e f f}^{\dagger}(\vec{y})\right\}=\hbar \delta^{3}(\vec{x}-\vec{y}) . \tag{184}
\end{equation*}
$$

We have assumed that $H_{\text {eff }}$ is bounded from below having vacuum state as the lowest eigenvalue state and $\Psi_{e f f}$ should annihilate it,

$$
\Psi_{e f f}|v a c\rangle=0
$$

Therefore,

$$
\begin{align*}
\langle v a c|\left\{\Psi_{e f f}(\vec{x}), \Psi_{e f f}^{\dagger}(\vec{y})\right\}|v a c\rangle & =\hbar \delta^{3}(\vec{x}-\vec{y}) \\
\langle v a c| \Psi_{e f f}(\vec{x}) \Psi_{e f f}^{\dagger}(\vec{y})|v a c\rangle & =\hbar \delta^{3}(\vec{x}-\vec{y}) \tag{185}
\end{align*}
$$

Similarly, bosonic operator $A_{\text {eff }}$ should annihilate $|v a c\rangle$ resulting in

$$
\begin{equation*}
\langle v a c| A_{e f f}(\vec{x}) A_{e f f}^{\dagger}(\vec{y})|v a c\rangle=\hbar \delta^{3}(\vec{x}-\vec{y}) . \tag{186}
\end{equation*}
$$

Thus any description of will apply on (186) as well. Therefore, we have

$$
\begin{array}{r}
\sum_{n}\langle v a c| \Psi_{e f f}(\vec{x})|n\rangle\langle n| \Psi_{e f f}^{\dagger}(\vec{y})|v a c\rangle= \\
\hbar \delta^{3}(\vec{x}-\vec{y}) \tag{187}
\end{array}
$$

Next, defining,

$$
\hbar^{\frac{1}{2}} \Psi_{n}(\vec{x})=\langle v a c| \Psi_{e f f}(\vec{x})|n\rangle
$$

results in

$$
\begin{equation*}
\sum_{n} \Psi_{n}(\vec{x}) \Psi_{n}^{*}(\vec{y})=\delta^{3}(\vec{x}-\vec{y}) \tag{188}
\end{equation*}
$$

thus modifying (191) into a Schrödinger equation

$$
\begin{equation*}
i \hbar \frac{d}{d t} \Psi_{n}(\vec{x})=\mathcal{H}_{e f f}(\vec{x}) \Psi_{n}(\vec{x}) \tag{192}
\end{equation*}
$$

Thus we have a Schrödinger evolution at the effective level. Starting from a fundamental trace dynamics theory we have obtained quantum mechanics from the thermodynamics of the fundamental degrees of freedom.

However, in deriving (192) we have made certain approximations valid at equilibrium. More explicitly, we replaced $\tilde{C}$ by its canonical average. If we also consider the fluctuations about the average quantities we have possibilities of obtaining a stochastic equation of evolution by adding stochastic nonlinear terms to the Schrödinger equation.

Herein perhaps lies the greatest virtue of Trace Dynamics. By treating quantum theory as a thermodynamic approximation to a statistical mechanics, the theory opens the door for the ever-present statistical fluctuations to play the desired role of the non-linear stochasticity which impacts on the measurement problem.
If we consider fluctuation in $\tilde{C}$ as

$$
\begin{array}{r}
\Delta \tilde{C} \simeq \tilde{C}-i_{e f f} \hbar \\
\frac{1}{2}\left\{\tilde{C}, i_{e f f}\right\}=-\hbar+\frac{1}{2}\left\{\Delta \tilde{C}, i_{e f f}\right\} \\
\frac{1}{2}\left\{\Delta \tilde{C}, i_{e f f}\right\}=-\hbar(\mathcal{K}+\mathcal{N}) \tag{195}
\end{array}
$$

with fluctuating $c$-number $\mathcal{K}$ and fluctuating matrix $\mathcal{N}$, we obtain the modified linear Schrödinger equation

$$
\begin{gather*}
|\dot{\Phi}\rangle=\left[-i \hbar^{-1} H_{e f f}+\mathcal{K}_{0}(t) i \hbar^{-1} H_{e f f}-\right. \\
\left.\mathcal{K}_{1}(t) \hbar^{-1} H_{e f f}+\frac{1}{2} i \mathcal{M}_{0}(t)-\frac{1}{2} \mathcal{M}_{1}(t)\right]|\Phi\rangle, \tag{196}
\end{gather*}
$$

where 0,1 label the real and imaginary part of $\mathcal{K}$ and

$$
\mathcal{M}(t)=\sum_{r, l} m_{r} \mathcal{N}(t)_{r, l} .
$$

In above equations $m_{r}$ is the rest mass of $r$-th species such that the Hamiltonian is

$$
\begin{equation*}
H=\sum_{r} \sum_{l} \frac{1}{2} i m_{r}\left[\psi_{r l}^{\dagger}, \psi_{r l}\right]+\text { const } \tag{197}
\end{equation*}
$$

with $l$ labeling a general complete basis set and after using the correspondence proposed
between trace-dynamics and quantum mechanics,

$$
|\Phi\rangle=\prod_{r, l} \Psi_{r l e f f}^{\dagger}|v a c\rangle
$$

As we will see next, with the added assumption of norm conservation the above equation (196) is modified to a CSL type nonlinear stochastic equation. This is how Trace Dynamics connects to models of continuous spontaneous localization. The weak link in the chain, at the present stage of our understanding, is the assumption that norm is conserved. Rather than being an assumption, this should follow from the underlying theory, and hopefully with improved understanding this will become possible in the future. On the other hand, the presence of antiHermitean modifications in the Schrödinger equation is inevitable from the Trace Dynamics viewpoint, since it is possible for $\tilde{C}$ to have a self-adjoint part as well.

## 8. Spontaneous collapse from stochastic mod-

ification of Schrödinger equation

We further motivate that the fluctuations can be described by linear superposition of white noise terms owing to the hierarchy between the length scale associated with the $\tilde{C}$ fluctuation and the length scale characterizing the emergent quantum degrees of freedom, much like in the case of Brownian mo-
tion fluctuations. A Brownian motion is described by a stochastic process $d W_{t}^{n}$ satisfying the following Itô table [see Appendix]

$$
\begin{array}{r}
\left(d W_{t}^{n}\right)^{2}=\gamma_{n} d t \\
d W_{t}^{n} d W_{t}^{m}=0, m \neq n \\
d W_{t}^{n} d t=d t^{2}=0 \tag{200}
\end{array}
$$

For our case we make following identifications
(i) For $c$ - number fluctuations $\mathcal{K}_{0,1}$

$$
\begin{align*}
i \hbar^{-1} \mathcal{K}_{0} d t & =i \beta_{I} d W_{t}^{I}  \tag{201}\\
-\hbar^{-1} \mathcal{K}_{1} & =\beta_{R} d W_{t}^{R} \tag{202}
\end{align*}
$$

with following Itô table

$$
\begin{equation*}
\left(d W_{t}^{R}\right)^{2}=\left(d W_{t}^{I}\right)^{2}=d t, d W_{t}^{R} d W_{t}^{I}=0 \tag{203}
\end{equation*}
$$

(ii) For the fluctuating matrix $\mathcal{M}_{0,1}$ having spatially correlated noise structure

$$
\begin{align*}
\frac{1}{2} i \mathcal{M}_{0} d t & =i \int d^{3} x d W_{t}^{I}(\vec{x}) \mathcal{M}_{t}^{I}(\vec{x})  \tag{204}\\
-\frac{1}{2} \mathcal{M}_{1} & =\int d^{3} x d W_{t}^{R}(\vec{x}) \mathcal{M}_{t}^{R}(\vec{x}),(2
\end{align*}
$$

> of superluminal signaling, after some calcu- lation at length, gives

$$
\begin{align*}
& d \hat{\rho}=i \hbar^{-1}\left[\hat{\rho}, H_{e f f}\right] d t-\frac{1}{2}\left(\beta_{R}^{2}+\beta_{I}^{2}\right)\left[H_{e f f},\left[H_{e f f}, \hat{\rho}\right]\right] d t- \\
& \begin{array}{r}
\frac{\gamma}{2} \int d^{3} x\left(\left[\mathcal{M}^{R}(\vec{x}),\left[\mathcal{M}^{R}(\vec{x}), \hat{\rho}\right]\right]+\left[\mathcal{M}^{I}(\vec{x}),\left[\mathcal{M}^{I}(\vec{x}), \hat{\rho}\right]\right]\right) d t \\
+\beta_{R}\left[\hat{\rho},\left[\hat{\rho}, H_{e f f}\right]\right] d W_{t}^{R}+i \beta_{I}\left[H_{e f f}, \hat{\rho}\right] d W_{t}^{I}
\end{array} \\
& +\int d^{3} x\left(\left[\hat{\rho},\left[\hat{\rho}, \mathcal{M}^{R}(\vec{x})\right]\right] d W_{t}^{R}(\vec{x})+i\left[\mathcal{M}^{I}(\vec{x}), \hat{\rho}\right] d W_{t}^{I}(\vec{x})\right)
\end{align*}
$$

corresponding to

$$
\begin{array}{r}
d|\Psi\rangle=\left[-i \hbar^{-1} H_{e f f} d t-\frac{1}{2} \beta_{I}^{2} H_{e f f}^{2} d t+\right. \\
\beta_{R}^{2}\left(H_{e f f}-\left\langle H_{e f f}\right\rangle\right)^{2} d t-\frac{\gamma}{2} d t \int d^{3} x \mathcal{M}^{I}(\vec{x})^{2}- \\
\frac{\gamma}{2} d t \int d^{3} x\left(\mathcal{M}^{R}(\vec{x})-\left\langle\mathcal{M}^{R}(\vec{x})\right\rangle\right)^{2}+i \beta_{I} H_{e f f} d W_{t}^{I}+ \\
\left.\left.\beta_{R}\left(H_{e f f}-\left\langle H_{e f f}\right\rangle\right)^{2} d W_{t}^{R}+i \int d^{3} x \mathcal{M}^{I}(\vec{x})\right]\right] d W_{t}^{I}(\vec{x})+ \\
\left.\int d^{3} x\left(\mathcal{M}^{R}(\vec{x})-\left\langle\mathcal{M}^{R}(\vec{x})\right\rangle\right)^{2} d W_{t}^{R}(\vec{x})\right]|\Psi\rangle . \tag{214}
\end{array}
$$

This equation (214) is a stochastic non-linear Schrödinger equation which has the martingale structure of spontaneous collapse models, and is capable of state vector reduction. In this sense, Trace Dynamics is an underlying theory for spontaneous collapse models. be larger for larger systems remains to be Of course, at the present stage of understanding, it cannot pick out one collapse model out of the many discussed, nor provide a theoretical origin for the values of the CSL parameters $\lambda$ and $r_{C}$. Nonethless, one cannot escape the profound and natural hypothesis that on
one hand thermodynamic equilibrium corresponds to quantum theory, and on the other hand fluctuations around equilibrium correspond to stochastic modifications of quantum theory. Why the effect of stochasticity must understood. Nor is it understood why norm should be preserved during evolution in Trace Dynamics : one should not have to put this in as an assumption into the theory, but rather have it come out of the underlying theory as a consequence.

For explicit demonstration of the collapse of the wave-function induced by stochasticity, we study a simplified version of (214)

$$
\begin{align*}
d|\Psi\rangle & =\left(-i \hbar^{-1} H_{\text {eff }}-\frac{1}{2}\left[\beta_{R}^{2}(A-\langle A\rangle)^{2}\right.\right. \\
& \left.\left.+\beta_{I}^{2} A^{2}\right]\right)|\Psi\rangle d t+\beta_{R}(A-\langle A\rangle)|\Psi\rangle d W_{t}^{R} \\
& +i \beta_{I} A|\Psi\rangle d W_{t}^{I}, \tag{215}
\end{align*}
$$

with

$$
\begin{gather*}
d \hat{\rho}=i \hbar^{-1}\left[\hat{\rho}, H_{e f f}\right] d t-\frac{1}{2}|\beta|^{2}[A,[A, \hat{\rho}] d t+ \\
\beta_{R}[\hat{\rho},[\hat{\rho}, A]] d W_{t}^{R}+i \beta_{I}[A, \hat{\rho}] d W_{t}^{I} \tag{216}
\end{gather*}
$$

Defining $E[]$ as expectation with respect to the stochastic process, $\left(E\left[d W_{t}^{R}\right]=0=\right.$ $\left.E\left[d W_{t}^{I}\right]\right)$ and variance of $A$,

$$
V=\left\langle(A-\langle A\rangle)^{2}\right\rangle=\operatorname{Tr} \hat{\rho} A^{2}-(\operatorname{Tr} \hat{\rho} A)^{2}
$$

and using the Itô product rules gives

$$
\begin{equation*}
d E[V]=E[d V]=-4 \beta_{R}^{2} E\left[V^{2}\right] d t \tag{217}
\end{equation*}
$$

Therefore,

$$
\begin{equation*}
E[V(t)]=E[V(0)]-4 \beta_{R}^{2} \int_{0}^{t} d s E\left[V(s)^{2}\right] \tag{218}
\end{equation*}
$$

Using the inequality

$$
0 \leq E\left[(V-E[V])^{2}\right]=E\left[V^{2}\right]-E[V]^{2}
$$

Non-negativity of variance suggests that $E[V(\infty)]=0$ and again as $V(t)$ is not supposed to be negative anywhere this will enforce

$$
V[\infty] \rightarrow 0
$$

As the variance in expectation of $A$ goes to zero asymptotically, the system in this way
results in one of the eigenstates of $A$. The demonstration of collapse using a systemapparatus interaction in the QMUPL model in Sec. II is a specific explicit application of this general analysis.

Also, we obtain from (218)

$$
E[V(t)] \leq \frac{V[0]}{1+4 \beta_{R}^{2} V[0] t}
$$

and hence a time scale of reduction as $\Gamma=$ $4 \beta_{R}^{2} V[0]$.

We can also see that in such a reduction scheme the Born probability rule follows for the outcomes. To see that, let us take $\Pi_{a}$ as the projector into $a$-th eigenstate of operator A,

$$
\Pi_{a}=|a\rangle\langle a| .
$$

Now for any operator $G$ commuting with $H_{\text {eff }}$ and $A$

$$
\begin{align*}
E[d\langle G\rangle] & =\operatorname{Tr}\left(-i \hbar^{-1}\left[G, H_{e f f}\right] E[\hat{\rho}]\right. \\
& \left.-\frac{1}{2}|\beta|^{2}[G, A][A, \hat{\rho}]\right) d t=0 . \tag{219}
\end{align*}
$$

If the initial state of the system is

$$
\left|\Psi_{i}\right\rangle=\sum_{a} p_{i a}|a\rangle
$$

at $t=0$ when the stochastic evolution has not started,

$$
\begin{equation*}
E\left[\left\langle\Pi_{a}\right\rangle\right]_{t=0}=\left\langle\Pi_{a}\right\rangle_{t=0}=\left|p_{i a}\right|^{2} \tag{220}
\end{equation*}
$$

As we have argued that when evolution is driven by $A$, the system lands in a particular eigenstate $|f\rangle$ with some probability $P_{f}$.

Then, for $a$-th eigenstate,

$$
\begin{equation*}
E\left[\left\langle\Pi_{a}\right\rangle\right]_{t=\infty}=\sum_{f}\langle f| \Pi_{a}|f\rangle P_{f}=P_{a} \tag{221}
\end{equation*}
$$

Now, since $A$ was taken to be commuting with $H_{e f f}$, we can have their simultaneous eigenstates, which we can call $|a\rangle$. Therefore, operators $\Pi_{a}$ constructed from these eigenstates will commute with $H_{e f f}$ and $A$, resulting in time-independence of $E\left[\left\langle\Pi_{a}\right\rangle\right]$ as evident from (219). Therefore,

$$
\begin{equation*}
E\left[\left\langle\Pi_{a}\right\rangle\right]_{t=0}=\left\langle\Pi_{a}\right\rangle_{t=0}=E\left[\left\langle\Pi_{a}\right\rangle\right]_{t=\infty} \tag{222}
\end{equation*}
$$

giving $P_{a}=\left|p_{i a}\right|^{2}$. Thus, we have obtained the Born probability rule.

We have seen that when treated as a fluctuation around the thermodynamic limit of trace dynamics theory, the emergent nonlinear equation captures the essential features of CSL, and in a sense, can possibly be a theoretical motivation for the phenomenological CSL equation of evolution. Of course at the present stage of understanding, trace dynamics makes no definite prediction for the actual numerical values of the CSL parameters, and this remains a challenge for the theory.

## B. Gravity induced wave-function collapse

The general theory of relativity dictates that gravity is the curvature of space-time. This curvature is produced by classical material bodies. However, even the motion of
classical bodies possesses intrinsic quantum fluctuations, and these fluctuations imprint a small uncertainty on spacetime structure. When one considers the motion of quantum mechanical objects on such a fluctuating spacetime, the coherence of the quantum state can be lost, providing a possible mechanism for wave-function collapse in the macroscopic domain, while leaving microphysics untouched by gravity. Counterintuitive though it may seem, gravity possibly plays a profound role in bringing about wavevector reduction, as the studies described below indicate.

## 1. The model of Karolyhazy [K-model]

The proposal of Karolyhazy (Karolyazy, 1966; Karolyhazy et al., 1986) deals with the smearing of space-time which results from the fundamental uncertainty in quantum theory being forced upon space-time structure. It starts with a viewpoint that nature 'somehow' tries to reconcile classical general relativity with quantum mechanics as much as possible. Space-time has in general, a fairly definite metric structure mainly determined by classical massive objects with fairly definite positions. However, the metric should not be completely sharp, and must have an in-built haziness to avoid contradiction with the fundamental quantum aspect of massive
objects (spread in position and momenta). Even a macroscopic massive body will have to satisfy

$$
\delta x \delta v \geq \frac{\hbar}{2 m}
$$

where $m$ is the mass of the body. The resulting haziness in the metric produced by the body leads to a stochastic correction in the evolution of state-vectors in quantum theory.

The basic idea of the approach is that when a wave packet of the center of mass of a body, sufficiently narrow in the beginning, spreads out in the Schrödinger evolution, into a space domain larger than a critical value (characteristic to the system), the coherence between distinct parts of the wave function gets destroyed, owing to space-time haziness. This is interpreted as a signal for stochastic reduction of the extended wave function to one of its smaller, coherent parts.

Quantum imprecision introduced in spacetime by quantum theory:

Let us consider a world-line segment $s=$ $c T$, in a flat space-time. We wish to estimate the precision with which we can realize this segment. Thus, the segment of $t$-axis is to be realized by the narrowest possible tube formed by a standing wave packet. Let, at the start (i.e. at the bottom of the worldline segment) the width of the wave packet be $\triangle x_{0}$. For mass $M$ of the wave packet the
velocity spread is

$$
\triangle V=\frac{\hbar}{2 M \triangle x_{0}}
$$

The corresponding spread at the end (i.e. at the top of line segment) will be

$$
\begin{equation*}
\triangle x=\triangle V T=\frac{\hbar}{2 M \triangle x_{0} c} c T \tag{223}
\end{equation*}
$$

The uncertainties $\triangle x$ and $\triangle x_{0}$ are the uncertainties in the top and bottom of the segments as well as in the length of the segments. Minimum amount of uncertainty in the length of segment will be introduced if we choose

$$
\begin{equation*}
\triangle x=\triangle x_{0} \tag{224}
\end{equation*}
$$

Clearly the uncertainty in the length of segment decreases with increasing $M$ and pointlike description becomes progressively more valid. Now, the gravitational radius of the mass $M$ is bounded by the fact that it should not be greater than the spread $\triangle x$,

$$
\begin{equation*}
\Delta x \approx G M / c^{2} \tag{225}
\end{equation*}
$$

From Eqns. (223), (224) and (225), the uncertainty in the length of the segment is given by

$$
\begin{array}{r}
(\triangle s)^{2}=(\triangle x)^{2}=\frac{\hbar}{2 M c} c T= \\
\frac{\hbar}{2 M c} s=\frac{G \hbar}{2 \triangle s c^{3}} s . \\
(\triangle s)^{2}=\left(\frac{G \hbar}{2 c^{3}}\right)^{2 / 3} s^{2 / 3} . \tag{226}
\end{array}
$$

This relation is often known as the Karolyhazy uncertainty relation. It gives the minimum amount of uncertainty in space-time
structure, which can be argued just from the demand of compatibility of quantum theory and metric structure of space-time. Therefore, we should be careful in using classical space-time considerations once the length of the segment starts approaching its uncertainty value; this providing a critical length scale for the system.

Next we consider a physical space-time domain of nearly Minkowski metrics with a corresponding smear structure as argued in (Karolyazy, 1966; Karolyhazy et al., 1986). We introduce a family $\left\{g_{\mu \nu}^{\beta}\right\}$ of matter-free metrics very close to the Minkowski metric

$$
\begin{gathered}
g_{00}=-1 ; g_{\mu \nu}=0 \Leftrightarrow \mu \neq \nu, \\
g_{11}=g_{22}=g_{33}=1,
\end{gathered}
$$

where different $\beta$ mark different members (hence different metrics) of the family. The proper length $s=c T$ between two world points $x_{1}$ and $x_{2}$ will be defined as the mean value of the lengths $s_{\beta}$ corresponding to different $g_{\mu \nu}^{\beta}$,

$$
s=\bar{s}_{\beta},
$$

with the bar describing average over $\beta$. The uncertainty in the line segment is defined as

$$
\Delta s=\left[\overline{\left(s-s_{\beta}\right)^{2}}\right]^{1 / 2}
$$

In the family of metrics $\beta=0$ gives the Minkowski metric. In the present analysis attention will be confined to the case in which
we do not have macroscopic bodies moving relatively to each other with a velocity near that of light. The co-ordinate system will therefore be assumed to be one relative to which all macroscopic bodies move slowly. This will enable us to confine our use of the set $\left\{s_{\beta}\right\}$ to non-relativistic many-particle wave equations in spite of the fact that by invoking curved manifolds we are employing the language of general relativity.

Since we are considering only slowly moving particles, $v \ll c$, only $\left(g_{00}\right)_{\beta}$ part will be required for the analysis. The general form of the metric in the family is of the form

$$
\left(g_{00}\right)_{\beta}(x)=-1+\gamma_{\beta}(x) ; \quad(\beta \neq 0)
$$

Since the space-time is matter-free apart from the test particle, we have

$$
\square \gamma_{\beta}=0
$$

Now, the idea is to fix the set $\gamma_{\beta}$ in such a way that the length of the world-line

$$
\begin{equation*}
s_{\beta}=\int d t\left[g_{\mu \nu}^{\beta} \frac{d x^{\mu}}{d t} \frac{d x^{\nu}}{d t}\right]^{1 / 2} \tag{227}
\end{equation*}
$$

is averaged to

$$
s=\bar{s}_{\beta}
$$

and the uncertainty obtained from

$$
\triangle s=\left[\overline{\left(s-s_{\beta}\right)^{2}}\right]^{1 / 2},
$$

is the same as obtained in 226). We thus do not regard the functions $\gamma_{\beta}$ as dynamical
variables, rather we represent physical spacetime by the whole set $\left\{g_{\mu \nu}^{\beta}\right\}$ at once.

In this spirit we construct $\gamma \mathrm{s}$ through their Fourier series,

$$
\begin{array}{r}
\gamma_{\beta}(x)=\frac{1}{L^{3 / 2}} \sum_{\vec{k}} \\
\left(c_{\beta}(\vec{k}) \exp [i(\vec{k} \cdot \vec{x}-\omega t)]+c . c\right), \tag{228}
\end{array}
$$

where $L$ is the length of an arbitrarily chosen large box (for normalization),

$$
\vec{k}=\frac{2 \pi}{L} \vec{n}
$$

and,

$$
\omega=c|\vec{k}| .
$$

We now choose an integer

$$
N_{\vec{k}}>2
$$

for each $\vec{k}$ and introduce a random variable $\alpha(\vec{k})$, such that

$$
\alpha(\vec{k}) \in \frac{2 \pi}{N_{\vec{k}}}\left[0,1,2, \ldots, N_{\vec{k}}-1\right] .
$$

For a particular $\alpha(\vec{k})$ a particular Fourier coefficient $c_{\beta}(\vec{k})$ is given as

$$
\begin{equation*}
c_{\beta}(\vec{k})=f(k) \exp [i \alpha(\vec{k})] . \tag{229}
\end{equation*}
$$

The unknown function $f(k)$ is obtained from the scheme proposed above, and is found out to be

$$
\begin{equation*}
f(k)=\left(\frac{G \hbar}{2 c^{3}}\right)^{1 / 3} k^{-5 / 6} \tag{230}
\end{equation*}
$$

using (226). The contribution to $f_{k}$ for large values of $k$ comes from the requirement that

Eqn. (226) should be valid even if $s$ is very small. Clearly, 226 ) is not meaningful in the limit $s \rightarrow 0$ and a cut-off is assumed: $f(k)=$ 0 for $k>10^{13} \mathrm{~cm}^{-1}, s<10^{-13} \mathrm{~cm}$. It is asserted that details of the cut-off are not important, and only long-wave components are relevant. This has been contested in Diósi and Lukacs, 1993) where it has been claimed that this cut-off is at a very high physically unacceptable value of $k$ and leads to absurd situations such as neutron star scale densities all over space. However, it seems that this objection can possibly be avoided by working entirely in real space, without going to Fourier space (Frenkel, 2002). The analysis of Karolyhazy has been repeated by Frenkel, accoring to whom some of the Fourier sums diverge in some intermediate expressions, but "in the formulae for physical quantities these sums are convegent". In this work, the impact of the Karolyhazy uncertainty relation is realized, not by introducing a family of metrics, but by introducing a local time operator, and a corresponding phase operator in the wave-function describing the quantum state. The final results on wave-vector reduction are the same as those described below.

Wave Propagation in a smeared space-time:
For considering wave propagation (Schrödinger type evolution) we generalize the quantum description of evolution in this 'hazy' space-time. We introduce a
family $\left\{\psi_{\beta}\right\}$ of wave-functions corresponding to the metric family $\left\{g_{\mu \nu}^{\beta}\right\}$. For a single scalar elementary particle, via the relativistic Klein-Gordon equation (alternatively, the Dirac equation for spinors)

$$
\begin{equation*}
\frac{1}{\sqrt{-g_{\beta}}} \frac{\partial}{\partial x^{\mu}}\left(\sqrt{-g_{\beta}} g_{\beta}^{\mu \nu} \frac{\partial \phi}{\partial x^{\nu}}\right)-\left(\frac{m c}{\hbar}\right)^{2} \phi=0 \tag{231}
\end{equation*}
$$

we obtain the non-relativistic generalization

$$
\begin{equation*}
i \hbar \frac{\partial}{\partial t} \psi_{\beta}=\left(-\frac{\hbar^{2}}{2 m} \nabla^{2}+V_{\beta}\right) \psi_{\beta} \tag{232}
\end{equation*}
$$

The small perturbation $V_{\beta}$ is given as

$$
V_{\beta}(\vec{x}, t)=\frac{m c^{2} \gamma_{\beta}(\vec{x}, t)}{2}
$$

More interesting is the case for many particles: the multi-particle equation, where $V_{\beta}$ is replaced by

$$
\begin{array}{r}
U_{\beta}(\{\vec{X}\}, t)=\sum_{i} \frac{m_{i} c^{2} \gamma_{\beta}\left(\vec{x}_{i}, t\right)}{2} \\
\{\vec{X}\}=\left\{\vec{x}_{i}\right\} \tag{233}
\end{array}
$$

To see the qualitative effect of such a smearing we start with an initial 'composite wave-function' $\Psi_{0}(\{\vec{X}\}, 0)$ for all the metrics $\left\{g_{\beta}^{\mu \nu}\right\}$. After the evolution different $\Psi_{\beta}(\{\vec{X}\}, t)$ will become different. We can write to a good approximation,

$$
\begin{equation*}
\Psi_{\beta}(\{\vec{X}\}, t) \approx \Psi_{0}(\{\vec{X}\}, t) e^{i \phi_{\beta}(\{\vec{X},\} t)}, \tag{234}
\end{equation*}
$$

with

$$
\begin{equation*}
\phi_{\beta}(\{\vec{X}\}, t)=-\frac{1}{\hbar} \int_{0}^{t} d t^{\prime} U_{\beta}(\{\vec{X}\}, t) . \tag{235}
\end{equation*}
$$

Let us choose and fix an $\vec{X}_{1}$ and an $\vec{X}_{2}$, and calculate the difference in phase between
these two points in configuration space for different $\beta$. The answer will depend on $\beta$ and on time. The root mean square spread in the phase (average is over $\beta$ )

$$
\left\{\overline{\left\{\phi_{\beta}\left(\left\{\vec{X}_{1}\right\}, t\right)-\phi_{\beta}\left(\left\{\vec{X}_{2}\right\}, t\right)\right]^{2}}\right\}^{1 / 2}
$$

can be estimated as a function of $\left\{\vec{X}_{1}, \vec{X}_{2}\right\}$ and time $t$. The uncertainty in the relative phase depends only on the separation between the two points in configuration space, and for a sufficiently large separation can reach the value $\pi$. Therefore, the set $\Psi_{\beta}$ can be of particular importance once we study microscopic and macroscopic systems.

## Microscopic and Macroscopic behavior:

For a single quantum particle of mass $M$ for small values of $a \equiv\left|x_{1}-x_{2}\right|$ the spread in the phase

$$
\triangle(a) \ll \pi
$$

and only for a large critical value $a_{c}$

$$
\triangle(a) \approx \pi
$$

will be achieved. The spread in the phase and the separation for which the critical value is reached can be calculated from the formalism described above. We next discretize the space in cells of dimension $a_{c}$, called 'coherence cells'. If initially the particle is confined to a single cell, then Schrödinger evolution will try to spread the wave packet, resulting in the wave function extending over to different
cells and the set $\left\{\psi_{\beta}\right\}$ will no longer behave as a single coherent wave function. When the original coherent set develops incoherent parts of comparable weights, it is taken as signal for stochastic reduction of $\left\{\psi_{\beta}\right\}$ to a single cell. Therefore, this stochastic reduction scheme is governed by Schrödinger evolution and stochastic part coming through smearing of space-time metric. This process provides us with a description of a physical phenomenon taking place regardless of the presence of any observer. Still this formalism indicates towards but does not provide any formal embedding of idea of stochastic jumps into evolution in a consistent mathematical framework. It can be heuristically argued that microscopc quantum particles will take astronomically large time before their wavefunctions "spill over" a single coherence shell, making the possibility of stochastic reduction very remote.

For an elementary particle of mass $m$, it can be shown that (Frenkel, 2002)

$$
\begin{equation*}
a_{c} \approx \frac{\hbar^{2}}{G} \frac{1}{m^{3}} \approx\left(\frac{L}{L_{p}}\right)^{2} L \quad ; L \approx \frac{\hbar}{m c}, \tag{236}
\end{equation*}
$$

and the critical time of reduction can be shown to be

$$
\begin{equation*}
\tau_{c} \approx \frac{m a_{c}^{2}}{\hbar} \tag{237}
\end{equation*}
$$

For a proton one finds

$$
a_{c} \approx 10^{25} \mathrm{~cm}, \quad \tau_{c} \approx 10^{53} \mathrm{~s}
$$

thus showing that one can never observe wave-packet reduction for a proton. The origin of the expression for the reduction time lies in the fact that according to the Schrödinger equation, a wave-packet initially spread over $a_{c}$ will spread to a size $2 a_{c}$ over time $\tau_{c}$. When this happens, we could take that as an indicator of loss of coherence, and hence stochastic reduction. The dynamics thus consists of cycles of deterministic Schrödinger evolution followed by stochastic jumps - something fully reminiscent of the GRW model; a comparison to which we will return shortly. In fact, $\tau_{c}$ is analogous to $\lambda^{-1}$ in GRW, and $a_{c}$ is analogous to $r_{c}$.

For more complex systems, such as a macroscopic body, one works with the center of mass co-ordinate. However, care is needed because the gravitational perturbation described by the multi-particle potential (233) depends on extended region of space. Still it can be shown that only the phase of the wave-function of the center of mass is affected, and the already introduced concepts of coherence cells and coherence length $a_{c}$ can be applied to the center of mass coordinate. In such cases not only mass $M$ of the system but the size $R$ also enters in the expression for $a_{c}$ (as arbitrariness in metric will be experienced throughout the size). It can be shown
that
$a_{c} \approx\left(\frac{\hbar^{2}}{G}\right)^{1 / 3} \frac{R^{2 / 3}}{M}=\left(\frac{R}{L_{p}}\right)^{2 / 3} L ; \quad L=\frac{\hbar}{m c}$

The reduction time is again given by (237). For a ball of $R=1 \mathrm{~cm}$ and for terrestrial densities this gives $a_{c} \approx 10^{-16} \mathrm{~cm}$, and $\tau_{c} \approx 10^{-4} \mathrm{~cm}$. The wave-function undergoes $10^{4}$ expansion-reduction cycles per second, and at the end of each cycle the momentum performs a jump $\Delta p_{c}$ of the order $\hbar / a_{c}$ which corresponds to a velocity shift of the order $a_{c} / \tau \sim 10^{-12} \mathrm{~cm} / \mathrm{sec}$. These repeated kicks amount to an anomalous Brownian motion and a tiny associated energy nonconservation of the order $\hbar^{2} / M a_{c}^{2}$, another feature in common with spontaneous collapse models.

One can try to understand the transition region from micro- to macro- behavior. We have seen that for $R \approx 1 \mathrm{~cm}$ we have $a_{c} \ll R$. Furthermore, the expressions (238) and 236 become the same when $a_{c}=R$. So one can now classify
(i) $a_{c} \gg R$ (i.e. $\left.\hbar^{2} / G \gg M^{3} R\right)$ microbehavior regime
(ii) $a_{c} \approx R \quad$ (i.e. $\left.\hbar^{2} / G \approx M^{3} R\right) \quad$ transition region
(iii) $a_{c} \ll R \quad$ (i.e. $\left.\hbar^{2} / G \gg M^{3} R\right)$ macrobehavior regime.

If $a_{c} \gg R$ it can be shown that Frenkel, 2002) the expression (236) continues to hold,
for a micro-object having an extended linear size $R$.

Setting $a_{c}=R$ in (238) and assuming density to be about 1 gram/cc for terrestrial bodies gives for the transition region
$a^{t r} \approx 10^{-5} \mathrm{~cm}, \tau^{t r} \approx 10^{3} \mathrm{sec}, M^{t r} \approx 10^{-14} \mathrm{gram}$

It is significant that $a^{t r}$ coincides with the favored value for $r_{C}$ in the GRW and CSL model. The transition mass corresponds to about $10^{10} \mathrm{amu}$. Note that because Planck length and the size of the body also enter the picture, the transition occurs at a mass much lower than the simplistic but much higher Planck mass ( $10^{-5}$ grams).

Interestingly, the jump velocity $\Delta v_{c}=$ $a_{c} / \tau_{c}$ in a reduction cycle takes its maximal value in the transition region $a_{c} \approx R$ and decreases on either side away from this transition region (Karolyhazy et al., 1986).

Thus one can, in principle, elegantly argue for classical behavior of massive and large objects and provide a transition point which can be tested upon. The measurement process can be argued as interactions resulting in significant change in mass distribution of whole set up (system+surrounding) making the definite and different outcome states incoherent thereby reducing the state of set-up to a particular outcome. Still the formal mathematical framework of this idea is missing making
it a challenge to precisely calculate the characteristics of quantum state reduction.

The K-model has also been discussed in (Frenkel, 1977, 1990, 1995, 1997, 2002 Karolyhazy, 1974, 1990, 1995, Karolyhazy et al., 1982).

Comparison with GRW model
There is a fascinating similarity between the K-model and GRW model, even though there are significant differences in detail. The overall picture of Schrödinger evolution interrupted by stochastic reduction is the same. In the K-model, the origin of stochasticity lies in the intrinsic uncertainty of space-time structure, whereas in GRW the origin is left unspecified. Both models have a length scale ( $a_{c}$ in K-model, $r_{C}$ in GRW) and a timescale ( $\tau_{C}$ in K-model, and $\lambda^{-1}$ in GRW). There are no free parameters in the K-model, whereas GRW introuduce new parameters $\lambda$ and $r_{C}$. Thus it is entirely possible that gravity might provide the fundamental underpinning for models of spontaneous collapse. Of course a mathematically rigorous treatment of gravity in the K-model remains to be developed, but the physical principles and semirigorous results already obtained are highly suggestive by themselves.

An important early study comparing the K-model and GRW was made in Frenkel, 1990). It should be noted that while in both cases the reduction time decreases with in-
creasing mass, the quantitative dependence is different. In the K-model, for $a_{c} \geq R$ the reduction time falls as $1 / m^{5}$, and if $a_{c} \ll R$ it falls as $1 / m^{5 / 9}$ assuming a fixed density. In GRW, the reduction time simply falls as $1 / m$, whereas we have seen that in the CSL model the dependence is more complex. Similarly, $a_{c}$ falls with increasing mass. While $r_{c}$ in GRW does not depend on mass, the linear size to which the stochastic reduction confines an expanding wave-packet does depend on mass - this linear size is the analog of the coherence cell $a_{c}$ (Frenkel, 1990). In the light of modern experiments there is perhaps need for a more careful comparison between the quantitative predictions of the K-model and GRW/CSL. Also, a careful quantitative description of the quantum measurement process, showing the emergence of the Born rule, seems to not yet have been developed in the K-model. A time-evolution equation for the density operator in the K-model, analogous to the corresponding equation in GRW, has been discussed in (Frenkel, 1990). See also related discussions in Unturbe and SanchezGomez, 1992).

## 2. The model of Diósi

Diósi's approach (Diósi, 1987), while being similar to Karolyhazy's, is inspired by the famous work of Bohr and Rosenfeld
(Wheeler and Zurek, 1983) which investigated the principles of measuring the electromagnetic field by apparatuses obeying quantum mechanics. It was shown in (Diósi and Lukacs, 1987) that if a Newtonian gravitational field $\mathbf{g}=-\nabla \phi$ is measured by a quantum probe over a time $T$, then its average $\tilde{\mathbf{g}}(\mathbf{r}, t)$ over a volume $V$ exhibits an uncertainty which is universally bounded by

$$
\begin{equation*}
(\delta \tilde{g})^{2} \geq \hbar G / V T \tag{240}
\end{equation*}
$$

This is Diósi's analog of the Karolyhazy uncertainty relation, and the idea now is to see how how this intrinsic quantum imprecision in the space-time metric affects the Schrödinger evolution of a quantum state in quantum mechanics.

To this effect, Diósi introduces the concept of a universal gravitational white noise, by proposing that the gravitational field possesses universal fluctuations [in other words the potential $\phi(\mathbf{r}, t)$ is a stochastic variable] whose stochastic average equals, upto numerical factors of order unity, the intrinsic uncertainty given by 240

$$
\begin{equation*}
\left\langle[\nabla \tilde{\phi}(\mathbf{r}, t)]^{2}\right\rangle-[\langle\nabla \tilde{\phi}(\mathbf{r}, t)\rangle]^{2}=\mathrm{const} \times \hbar G / V T \tag{241}
\end{equation*}
$$

From here, it can be shown that, assuming $\langle\phi(\mathbf{r}, t)\rangle \equiv 0$, the correlation function of $\phi(\mathbf{r}, t)$ is given by

$$
\begin{equation*}
\left\langle\phi(\mathbf{r}, t) \phi\left(\mathbf{r}^{\prime}, t^{\prime}\right)\right\rangle=\hbar G\left|\mathbf{r}-\mathbf{r}^{\prime}\right|^{-1} \delta\left(t-t^{\prime}\right) \tag{242}
\end{equation*}
$$

The probability distribution of the stochastic variable $\phi(\mathbf{r}, t)$ is completely specified by this correlation function if the distribution is assumed to be gaussian [gaussian white noise].

Next, one asks for the effect of the stochastic fluctuations in $\phi$ on the propagation of the quantum state $\psi$ of a system whose evolution is assumed to be described by the Schrödinger equation

$$
\begin{equation*}
i \hbar \dot{\psi}=\left(\hat{H}_{0}+\int \phi \hat{f}(\mathbf{r}) d^{3} r\right) \psi(t) \tag{243}
\end{equation*}
$$

where $\hat{f}(\mathbf{r})$ stands for the operator of the local mass density of the system.
$\psi$ is now a stochastic variable, and the corresponding density operator $\hat{\rho}=\left\langle\psi(t) \psi^{\dagger}(t)\right\rangle$ obeys the following deterministic master equation for the assumed gaussian white noise

$$
\begin{align*}
& \dot{\hat{\rho}}=-\frac{i}{\hbar}\left[\hat{H}_{0}, \rho(t)\right] \\
&- \frac{G}{2 \hbar} \iint \frac{d^{3} r d^{3} r^{\prime}}{\left|\mathbf{r}-\mathbf{r}^{\prime}\right|}\left[\hat{f}(\mathbf{r}),\left[\hat{f}\left(\mathbf{r}^{\prime}\right), \hat{\rho}(t)\right]\right] \tag{244}
\end{align*}
$$

The second term on the right hand side is the damping term which represents the universal violation of quantum mechanics.

To compute the nature of violation, denote the configuration coordinates of a dynamical system by $X$, and denote the corresponding mass density at a point $\mathbf{r}$ by $f(\mathbf{r} \mid X)$. Given a pair of configurations, a characteris-
tic damping time $\tau_{d}\left(X, X^{\prime}\right)$ is defined by

$$
\begin{align*}
& {\left[\tau_{d}\left(X, X^{\prime}\right)\right]^{-1} } \\
= & \frac{G}{2 \hbar} \iint \frac{\left[f(\mathbf{r} \mid X)-f\left(\mathbf{r}^{\prime} \mid X^{\prime}\right)\right]^{2} d^{3} r d^{3} r^{\prime}}{\left|\mathbf{r}-\mathbf{r}^{\prime}\right|} \tag{245}
\end{align*}
$$

Introducing the coordinate eigenstates $|X\rangle$ the master equation can be written as

$$
\begin{gather*}
\langle X| \dot{\hat{\rho}}\left|X^{\prime}\right\rangle=-\frac{i}{\hbar}\langle X|\left[\hat{H}_{0}, \rho(t)\right]\left|X^{\prime}\right\rangle \\
-\left[\tau_{d}\left(X, X^{\prime}\right)\right]^{-1}\langle X| \hat{\rho}(t)\left|X^{\prime}\right\rangle \tag{246}
\end{gather*}
$$

Just like in decoherence and in models of spontaneous collapse, the second term on the right hand side destroys iterference between the states $|X\rangle$ and $\left|X^{\prime}\right\rangle$ over the characteristic time $\tau_{d}$, and this effect can become significant if the difference between the mass distributions $f(\mathbf{r} \mid X)$ and $f\left(\mathbf{r} \mid X^{\prime}\right)$ is significant.

To estimate the scale of the gravitationally induced violation Diósi considers a dynamical system consisting of a rigid spherical ball of homogeneously distributed mass $m$ and radius $R$, so that the configuration $X$ is represented by the center of mass coordinate $\mathbf{x}$. The characteristic damping time $\tau_{d}$ is shown to be

$$
\begin{equation*}
\tau_{d}\left(\mathbf{x}, \mathbf{x}^{\prime}\right)=\hbar\left[U\left(\left|\mathbf{x}-\mathbf{x}^{\prime}\right|\right)-U(0)\right]^{-1} \tag{247}
\end{equation*}
$$

where $U$ is the gravitational potential between two spheres, each of mass $m$ and radius $R$. The master equation can now be written
as

$$
\begin{gather*}
\frac{d}{d t}\langle\mathbf{x}| \rho\left|\mathbf{x}^{\prime}\right\rangle=\frac{i \hbar}{2 m}\left(\Delta-\Delta^{\prime}\right)\langle\mathbf{x}| \rho\left|\mathbf{x}^{\prime}\right\rangle \\
-\frac{1}{\hbar}\left[U\left(\left|\mathbf{x}-\mathbf{x}^{\prime}\right|\right)-U(0)\right]\langle\mathbf{x}| \rho\left|\mathbf{x}^{\prime}\right\rangle \tag{248}
\end{gather*}
$$

We define the coherent width $l$ of a given state as the characteristic distance $l=\left|\mathrm{x}-\mathrm{x}^{\prime}\right|$ above which the off-diagonal terms $\langle\mathbf{x}| \rho\left|\mathbf{x}^{\prime}\right\rangle$ become negligibly small. The time-scale $t_{k i n}$ over which kinetic changes are introduced due to ordinary quantum evolution given by the first term on the right hand side is of the order $m l^{2} / \hbar$. A crtitical length $l_{\text {crit }}$ is defined by equating $t_{k i n}\left(l_{c r i t}\right)$ and the damping time $\tau_{d}\left(l_{\text {crit }}\right)$

$$
\begin{equation*}
m l_{c r i t}^{2} / \hbar=\hbar\left[U\left(l_{c r i t}\right)-U(0)\right]^{-1} \tag{249}
\end{equation*}
$$

If the coherent width $l$ of the quantum state is much smaller than the critical value $l_{\text {crit }}$ then the standard quantum kinetics dominates and damping is not effective. On the other hand if $l \gg l_{\text {crit }}$ then the coherence of the state will be destroyed by the gravitational damping term in the master equation. $l_{\text {crit }}$ is the analog of the phase coherence length $a_{c}$ of the K-model, and the length parameter $r_{C}$ of GRW. Also, there clearly are analogs of $\tau_{d}\left(l_{\text {crit }}\right)$ in the other two models.

One can now show that in two limiting cases $l_{\text {crit }}$ takes the following form:

$$
\begin{align*}
l_{\text {crit }} & \sim\left(\hbar^{2} / G m^{3}\right)^{1 / 4} R^{3 / 4}, \quad \text { if } \quad R m^{3} \gg \hbar^{2} / G \\
& \sim\left(\hbar^{2} / G m^{3}\right)^{1 / 2} R^{1 / 2}, \quad \text { if } \quad R m^{3} \ll \hbar^{2} / G \tag{250}
\end{align*}
$$

These expressions are similar to, though not identical with, those in the K-model. The fact that they are similar but not identical suggests that the involvement of gravity in wave-vector reduction is strongly indicated, but the exact mathematical treatment remains to be found. Importantly, the transition $l_{\text {crit }}=R$ happens at the same value $l_{\text {crit }}=\hbar^{2} / G m^{3}$ in both the models. Notice though that for small masses $l_{\text {crit }}$ is not independent of $R$, unlike in the K-model. For a proton, taking $R$ to be the classical radius $10^{-13} \mathrm{~cm}$, Diósi estimates $l_{\text {crit }}$ to be $10^{6} \mathrm{~cm}$, which is curiously much smaller than the prediction $10^{25} \mathrm{~cm}$ for the K-model. Also, the reduction time is $10^{15} \mathrm{sec}$, much smaller than in the K-model. However, the models are in better agreement in the macro- region, and in Diósi's model too, the transition parameters are the same as that given by Eqn. (239).

Subsequently, Diósi took the inevitable step of casting the master equation in the equivalent language of a stochastic Schrödinger equation (Diósi, 1989). He called this model QMUDL (Quantum mechanics with universal density localization). It is similar to his QMUPL model, which we reviewed earlier in this article, except that the localization is not in the position operator $q$, but in the mass density operator $\hat{f}(\mathbf{r})$ introduced above. The universal free parameter $\lambda$ of QMUPL is now replaced by the gravita-
tional constant, so that the theory becomes parameter free.

As was discussed in Ghirardi et al., 1990a) the QMUDL model has certain limitations - it cannot deal with point particles (for which case it leads to divergent densities) and restricts itself to extended objects. The model parameters are such that it leads to an unacceptably high rate of energy increase during reduction. Furthermore, for microscopic dynamics the reduction and localization process can lead to unacceptable processes such as excitation or dissociation of nuclei. To avoid these problems. Ghirardi et al. proposed a CSL type modification of QMUDL, and the introduction of a new universal length parameter. It is suggested that it does not seem possible to have a parameter free theory for reduction, such as gravity induced collapse. An alternate way out of the difficulties of the otherwise very attractive model of Diósi has been suggested by Penrose - we will recall this proposal next, but find that here one is faced with a possibly new set of difficulties. Thus it would seem that at present CSL might be the best model at hand, even though its fundamental origin remains to be understood, and it yet may have a strong connection with gravity whose proper implementation remains to be achieved.

## 3. The model of Penrose

Penrose (Penrose, 1996, 1998) addressed the question of the stationarity of a quantum system which consists of a linear superposition $|\psi\rangle=a|\alpha\rangle+b|\beta\rangle$ of two well-defined states $|\alpha\rangle$ and $|\beta\rangle$, each of which would be stationary on its own, and each of the states is assumed to have the same energy $E$

$$
\begin{equation*}
i \hbar \frac{\partial|\alpha\rangle}{\partial t}=E|\alpha\rangle, \quad i \hbar \frac{\partial|\beta\rangle}{\partial t}=E|\beta\rangle \tag{251}
\end{equation*}
$$

If gravitation is ignored, as is done in standard quantum theory, the superposition $|\psi\rangle=a|\alpha\rangle+b|\beta\rangle$ is also stationary, with the same energy $E$

$$
\begin{equation*}
i \hbar \frac{\partial|\psi\rangle}{\partial t}=E|\psi\rangle \tag{252}
\end{equation*}
$$

However, the inclusion of gravitation raises a new question: what is the meaning of the Schrödinger time-evolution operator $\partial / \partial t$ ? There will be a nearly classical spacetime associated with the state $|\alpha\rangle$, and a Killing vector associated with it which represents the time displacement of stationarity. And there will be a different nearly classical spacetime associated with the state $|\beta\rangle$, and a different Killing vector associated with it which represents the associated time displacement of stationarity. The two Killing vectors can be identified with each other only if the two space-times can be identified with each other point by point. However, the prin-
ciple of general covariance in general relativity forbids that, since the matter distributions associated with the two states are different. On the other hand, unitary evolution in quantum theory requires and assumes the existence of a Schrödinger operator which applies to the superposition in the same way that it appies to the individual states, and its action on the superposition is the superposition of its action on individual states. There is thus a conflict between the demands of quantum theory and of general relativity.

A tentative resolution is to make an approximate pointwise identification between the two spacetimes, which in turn corresponds to a slight error in the identification of the Schrödinger operator for one spacetime with that for the other. This corresponds to a slight uncertainty in the energy of the superposition, for which it is possible to make an estimate in the case when the superposition amplitudes are nearly equal in magnitude. In the Newtonian approximation, this energy uncertainty $E_{G}$ is of the order of the gravitational self-energy of the mass distribution in the two superposed states. In accordance with the Heisenberg uncertainty principle, the superposition lifetime can be taken to be $\hbar / E_{G}$, beyond which time the superposition will decay. In concept and in detail, this is quite like the damping time $\tau_{d}$ in Diosi's model. It is not clear here though, as
to how the Born rule will be recovered dynamically.

Penrose notes the commonality with Diósi's ideas, the difficulties encountered by Diósi, and the resolution proposed by Ghirardi et al. by way of introducing a fundamental length scale. Penrose observes that essentially the same difficulty would arise in his own approach too, because if one were dealing with point particles, the gravitational self-energy difference can become infinitely high, implying instantaneous reduction, which is clearly unreasonable. While Ghirardi et al. avoid this problem by introducing a new length scale, Penrose proposes a different way out. The way out is in particular based on noting that one needs to specify which states are the basic [stable] states, to which superpositions of basic states decay.

It is proposed that the basic stationary states to which a general superposition will decay by state reduction are stationary solutions of the so-called Schrödinger-Newton equation (SN-equation). This equation is actually a pair of coupled differential equations which are set up as follows, for a quantum mechanical particle of mass $m$ moving in its own gravitational field

$$
\begin{align*}
& i \hbar \frac{\partial \psi}{\partial t}=-\frac{\hbar^{2}}{2 m} \nabla^{2} \Psi+m \Phi \Psi \\
& \nabla^{2} \Phi=4 \pi G m|\Psi|^{2} \tag{253}
\end{align*}
$$

This system of equations has been ana-
lyzed in (Bernstein et al., 1998; Giulini and Großardt, 2011; Harrison et al., 1998, 2003; Moroz and Tod, 1999; Ruffini and Bonazzola, 1969). These equations are closely related to the Schrödinger-Poisson equations which have been studied for much longer (Lange et al., 1995).

At this stage an important difference with the models of Karolyhazy and Diósi seems to be that that unlike in the latter two models, where an intrinsic uncertainty in spacetime structure is assumed, here the impact on the evolution of the quantum state is due to the particle's own gravitational field. Also, the system seems to be set up deterministically and the presence of a stochastic element is not evident, at least a priori. Thus one could ask as to the origin of the stochastic feature which actually drives the system to one of the stationary states, and the accompanying Born rule. Also, if the evolution is deterministic and non-linear, there appears to be present the possibility of superluminal propagation.

These issues apart, the SN system of equations yields some very intersting results. Spherically symmetric stationary solutions have been found and their stability has been investigated. A comprehensive recent analysis is given in Giulini and Großardt, 2011). Their study was motivated in response to Carlip, 2008; Salzman and Car-
lip, 2006) - the SN equation induces a gravitational suppression of expanding Gaussian wave-packets, and it was suggested by Carlip and Salzman that the suppression [and hence wave-vector reduction] becomes significant already at $m \sim 1600 \mathrm{amu}$. This surprisingly low value is at variance with the much higher estimates coming from simple analytical estimates [and also from the work of Karolyhazy and Diósi] and prompted (Giulini and Großardt, 2011) to look at the problem closely.

Various numerical studies, as well as heuristic estimates, show that the ground state energy is of the order

$$
\begin{equation*}
E \sim-\frac{1}{8} \frac{G^{2} m^{5}}{\hbar^{2}} \tag{254}
\end{equation*}
$$

The width $a$ of the mass distribution in the ground state is

$$
\begin{equation*}
a_{0} \approx \frac{2 \hbar^{2}}{G m^{3}} \tag{255}
\end{equation*}
$$

which we immediately notice coincides with the phase coherence cell length in microscopic limit of the K-model.

By introducing a length scale $l$ the SN equation can be written in terms of a dimensionless coupling constant

$$
\begin{equation*}
K=2 \frac{G m^{3} l}{\hbar^{2}}=2\left(\frac{l}{L_{p}}\right)\left(\frac{m}{m_{P}}\right)^{3} \tag{256}
\end{equation*}
$$

One considers the time-dependent SN equation for initial values given by a spherically symmetric gaussian wave-packet of width $a$

$$
\begin{equation*}
\psi(r, t=0)=\left(\pi a^{2}\right)^{-3 / 4} \exp \left(-\frac{r^{2}}{2 a^{2}}\right) \tag{257}
\end{equation*}
$$

There are thus two free parameters $a$ and $m$, and one asks for the regions in this parameter space where significant inhibitions of the usual free quantum dispersion occur. In an important analysis Giulini and Großardt, 2011) give four different analytical arguments to show that inhibition of the dispersion becomes significant when the dimensionless coupling constant $K$ of (256) becomes order unity. This conclusion coincides with that of Karolyhazy and Diosi, and we believe it leads us to an important inference:

The models of Karolyhazy, Diósi and Penrose all agree that if the width of the quantum state associated with an object of mass $m$ becomes greater than of the order $\hbar^{2} / G m^{3}$, the quantum-to-classical transition sets in.

For the experimentally interesting $a=$ $0.5 \mu \mathrm{~m}$ this gives $m$ to be about $10^{9} \mathrm{amu}$.

These results are further supported by numerical investigations of the SN equation in (Giulini and Großardt, 2011). The authors also note that the coherence time, the time beyond which collapse takes place, can be brought down by reducing the grating period in a molecule interferometry experiment. For instance, for a mass of $10^{11} \mathrm{amu}$ and grating period of $0.5 \mu \mathrm{~m}$ they report a coherence loss time of 300 ms .

It is significant that while the Penrose approach does not directly address the emergence of the Born rule, it correctly predicts
the regime where the quantum to classical transition takes place, in agreement with the other gravity models.

This essentially completes our brief review of the three well-known models of gravityinduced collapse. Other consideations of gravity induced collapse have been made in (Ellis et al., 1984) and (Percival, 1995).

In our view, gravity induced collapse is a promising physical mechanism for a physical realization of spontaneous collapse. Furthermore, Trace Dynamics and its extension to space-time structure [treating space and time as operators] provide a plausible mathematical avenue for rigorously developing the stochastic theory of gravity induced spontaneous collapse.

## IV. EXPERIMENTAL TESTS OF THE THEORETICAL PREDICTIONS

## A. Introduction

We have considered two classes of underlying theories for dynamical collapse: Trace Dynamics, and gravity induced collapse. The phenomenology of Trace Dynamics manifests itself through models of spontaneous collapse. If spontaneous collapse or gravity induced collapse is a possible explanation for the measurement problem, then the experimental predictions of these models differ from
those of standard quantum theory. Bounds can be set on the parameters of these models by requiring that their predictions should not disagree with those observations which are well-explained by the standard quantum theory. On the other hand one can perform new experiments, such as diffraction experiments with large molecules, for which the predictions of these experiments differ appreciably from those of quantum theory. The results of such experiments could vindicate the modified quantum dynamics [and specific values of the associated parameters] or rule it out. This section reviews the bounds on model parameters which come from known physical and astrophysical processes, and from diffraction experiments that have been carried out in the laboratory or are planned for the near future. Experiments are discussed through Sections IV.B to IV.G. Bounds on spontaneous collapse models will be discussed in Sec. IV.H and IV.I and those on gravity based models in Sec. IV.J.

As we saw in Sec. II, a large variety of collapse models has been proposed: QMUPL, GRW, CSL, dissipative and nondissipative, white, colored, Markovian and non-Markovian. The overall task of constraining these models is very extensive, given their large variety, and considering that a large variety of observations [laboratory and astrophysical] as well as table-top exper-
iments, has to be considered. The subject is in a state of rapid flux, and still in a developmental stage. Here we try to do as complete a job as possible, relying on the analysis in a host of important papers that have appeared in the last few years (Adler, 2007; Adler and Bassi, 2009; Adler and Ramazanoğlu, 2007; Feldmann and Tumulka, 2012; Nimmrichter et al., 2011b; Romero-Isart, 2011). It should be noted though that the CSL model has received the maximum amount of attention, and we will focus mainly on CSL.

As we have seen, the CSL model (like the GRW model from which it originates) introduces two new parameters, the rate constant $\lambda$ and the correlation length $r_{C}$. If spontaneous collapse is a correct theory of nature, the values of these parameters must follow from some underlying fundamental principles and/or be determined by experiments. As mentioned earlier, GRW chose $\lambda \simeq 10^{-16}$ $\mathrm{sec}^{-1}$ and $r_{C} \simeq 10^{-5} \mathrm{~cm}$, in order to be consistent with observations, while Adler chose $\lambda \simeq 10^{-8 \pm 2} \mathrm{sec}^{-1}$ and $r_{C} \simeq 10^{-5} \mathrm{~cm}$. However, there is room for more general considerations and for establishing the allowed part of the parameter space in the $\lambda-r_{C}$ plane. In Sec. IV.G we will consider bounds coming from cosmology and in Sec. IV.H we will summarize other physical processes which constrain $r_{C}$ and $\lambda$.

In Sec. IV.C we will summarize experi-
ments which directly test the superposition principle of quantum mechanics by means of interferometers. As such, these experiments are also tests of collapse models and gravity models. The rationale is that if we can observe superposition at mesoscopic and possibly even macroscopic scale, the quantum dynamics does not need alteration. If instead we experimentally observe a quantum to classical transition such as the collapse of the wave function while convincingly reducing all potential sources of noise, this would strongly hint that an alteration of the fundamental equations of quantum mechanics is needed. The CSL model predicts a quantum to classical transition at only two or three orders of magnitude away from present large particle matter-wave experiments. This and new proposals for optomechanics experiments with trapped bead particles [see Sec. IV.D and IV.E] bring experimental tests of CSL (amongst other proposals) within reach. The main focus of the following sections will be on those table-top matter-wave and optomechanics experiments.

## B. Possible experimental tests of CSL

 basing on quantum superpositionMatter-wave interference experiments such as molecule interference are approaching the mass limit for the quantum to
classical transition in a 'bottom-up' fashion, starting with particles, where quantum superposition is existing and pushing the limit upwards step by step. Nano- and micro-mechanical devices cooled to the quantum-mechanical ground state within opto-mechanics approach the problem from the top, starting at very massive objects namely mechanical cantilevers of hundreds of nanometer and even some micrometer size, which are sometimes even visible with the bare eye. The range where both types of experiments are hitting towards at this very moment and where both will most probably meet to combine techniques, knowhow and ideas to overcome experimental hurdles and to switch-off known decoherence mechanisms to test limits of quantum mechanics: the 10 nm to 100 nm size range [mass $\sim 10^{6}-10^{9} \mathrm{amu}$. The experimental aim is to show quantum superposition by negativity in the Wigner function of the motional states or by proving the wave-nature of such particles by single-particle interference. Interestingly, this size range is vital to test non-standard quantum theories such as CSL and gravity induced collapse.

1. Collapse theory for diffraction experiments

In order to understand how collapse models differ from standard quantum mechanics, when applied to interferometric experiments, let us consider once again the QMUPL model of Sec. II.E, due to its simplicity. The multiparticle dynamics is given by Eqn. (40). By using Itô calculus, it is easy to show that the master equation for the statistical operator $\rho_{t}=\mathbb{E}\left[\left|\psi_{t}\right\rangle\left\langle\psi_{t}\right|\right]$ is:

$$
\begin{equation*}
\frac{d}{d t} \rho_{t}=-\frac{i}{\hbar}\left[H, \rho_{t}\right]-\frac{1}{2} \sum_{i=1}^{n} \lambda_{i}\left[q_{i},\left[q_{i}, \rho_{t}\right]\right] . \tag{258}
\end{equation*}
$$

Suppose - for simplicity sake - that all particles are identical, and that during the time of measurement, the free evolution (given by $H)$ can be neglected. Then, according to the above equation, the density matrix in the position representation evolves as follows:

$$
\begin{equation*}
\rho_{t}(x, y)=\rho_{0}(x, y) e^{-\lambda N(x-y)^{2} t / 2} . \tag{259}
\end{equation*}
$$

This equation contains everything there is to know about the effect of collapse models on interferometric experiments, at least from the conceptual point of view. Different models differ only in the technical details. The above equation tells that, in order to measure a collapse effect, corresponding to a significant damping factor, the following criteria must be met: the system should be as big as possible (large $N$ ); it should be created in a
"large" superposition (large $|x-y|$ ), which is monitored after a time as large as possible (large $t$ ). This is the goal that all interferometric experiments aim at reaching, in order to test the validity of the superposition principle, and thus also of collapse models.

We now come back to the CSL model, which we are primarily interested in. In this case, the damping behavior is less trivial than that of the QMUPL model. As we have seen in Sec. II.I, for small distances there is a quadratic dependence of the decay function on the superposition distance $|x-y|$, while for large distances such a dependence disappears. The intermediate behavior is not easy to unfold, but it can be conveniently modeled by the following ansatz. Recall first that for a single constituent the master equation
$\frac{d}{d t} \rho_{t}(x, y)=-\frac{i}{\hbar}\left[H, \rho_{t}(x, y)\right]-\Gamma_{\mathrm{CSL}}(x, y) \rho_{t}(x, y)$
implies that the decay function is

$$
\begin{equation*}
\Gamma_{\mathrm{CSL}}(x)=\lambda\left[1-e^{-x^{2} / 4 r_{c}^{2}}\right] \tag{261}
\end{equation*}
$$

for one single constituent. See Fig. 3 for a plot of $\Gamma_{\mathrm{CSL}}$ vs $x$. Here we see how the two fundamental parameters of the CSL model enter into play. For a many-particle system, one makes an ansatz and assumes that the above expression for the decay function holds, except that one has to multiply $\lambda$ by the appropriate numeric factoral, as described in Sec. II.I. The numerical factor is $n^{2} N$ where
$n$ is the number of particles [called one cluster] within a volume of linear size $r_{C}$, and $N$ is the number of clusters in the many-particle system.

In all interferometric experiments so far realized, the period of the grating is comparable to $r_{C} \sim 100 \mathrm{~nm}$. Therefore, to extract the significant order of magnitude, it is sufficient to work in the regime $x \gg r_{C}$. Taking into account Eqn. (92) (Adler, 2007) and that in the case of macromolecules $N=1$ (typical molecule size being about 1 nm ), we have:

$$
\begin{equation*}
\Gamma_{\mathrm{CSL}} \simeq \lambda n^{2} \tag{262}
\end{equation*}
$$

Since no interferometry-based experiments have so far detected any spontaneous collapse effect, this implies that the damping factor $\exp \left[-\Gamma_{\mathrm{CSL}} t\right]$ must be insignificant. We then have:

$$
\begin{equation*}
\lambda \leq 1 / n^{2} t \tag{263}
\end{equation*}
$$

where $n$ measures the number of nucleons in the system, and $t$ the duration of the experiment. This is the type of bound that interferometric-experiments place on the collapse rate $\lambda$. The experiments do not provide a bound on the second parameter of the CSL model $r_{C}$, for the reasons explained here above. More general situations could be considered, but they have not been analyzed so far. It will be desirable to carry out a careful analysis of the allowed part of the $\lambda-r_{C}$


FIG. 3 Upper Panel: The CSL decay function $\Gamma$ is shown over the spatial distance $x$ for four different masses (number of particles), using $\lambda=\lambda_{C S L}$. At the points $a, b, c, d$ where the curves intersect the line $x=r_{C}, \Gamma$ takes the values $\Gamma_{a}=.022 \mathrm{~s}^{-1}, \Gamma_{b}=\Gamma_{a} / 10^{9}, \Gamma_{c}=\Gamma_{a} / 10^{11}$ and $\Gamma_{d}=\Gamma_{a} / 10^{13}$ respectively. Lower panel: Decay function for a fixed mass of $10^{4} \mathrm{amu}$ and for different values of $\lambda$. The lower panel represents the experiment of (Gerlich et al., 2011). $\Gamma>500$ is ruled out by experiment. The values $\lambda_{G R W}, \lambda_{C S L}$ and $\lambda_{A D L E R}$ are taken from (Ghirardi et al., 1986), (Ghirardi et al., 1990c) and
plane, based on the data available from experiments, and to understand what role the grating period and size of the macromolecule will eventually play in bringing experiment and theory closer.

The latest situation on the results from diffraction experiments has been discussed in (Feldmann and Tumulka, 2012; Nimmrichter et al., 2011b; Romero-Isart, 2011). The strongest current bound on $\lambda$ seems to be from the experiment of Gerlich et al., 2011) which sets $\lambda<10^{-5} \sec ^{-1}$ for $n=$ 7,000 . Adler estimates that an experiment with $n=500,000$ will confront the enhanced CSL value proposed by him, based on reduction in latent image formation (Adler, 2007). Interestingly, in the same paper he also proposes to test whether 'latent image formation' constitutes a measurement, by using a photographic emulsion as a 'which path' detector in one arm of a quantum interferometer.

Spatial or centre of mass motion superposition is needed to be demonstrated in experiments to test the quantum to classical transition. As described previously atoms are too light that even the very large areas in today's atom interferometers do not increase the chance to test CSL (see $\Gamma$ vs $x$ plot). On the other hand the very massive cantilevers do not possess a large enough spatial separation (spatial size of superposition) to be-
come good test embodiments for the quantum to classical transition. It seems that the size range of particles of 10 nm to 100 nm , which corresponds to a mass range of $10^{6} \mathrm{amu}$ to $10^{9} \mathrm{amu}$ are ideal for such tests in matterwave interference experiments.

In this section we focus on possible experimental tests of the CSL model, while similar or quite different experiments are possible to test different collapse models.

## C. Matter-wave interferometry:

## Molecule interferometry

Experiments with matter waves exist since 1927 when Davissson and Germer diffracted a beam of electrons. It was the first proof of de Broglie's hypothesis on particle-wave duality. Since then matterwave interferometry of electrons (Hasselbach, 2010), neutrons (Rauch and Werner, 2000), atoms (Cronin et al., 2009) and molecules (Hornberger et al., 2011) has a long and successful history to investigate fundamental physics, and has been applied for metrology and sensing (Arndt et al., 2011). Interestingly, a recent interpretation of atom interferometry experiments resulted in a debate on the possible detection of gravitational red shift by such tabletop experiments Müller et al., 2010).

We are here interested in centre-of-mass-
motion interferometry (or de Broglie interference) of very massive particles as these experiments are promising to test modifications of Schrödinger dynamics such as collapse models predicting a quantum to classical transition at mesoscopic length and mass scales. The appearance of a single-particle interference pattern demonstrates wave-like behaviour of the particles and can be seen as an indication for superposition. The full beauty of this particle position superposition can be seen from reconstruction of the Wigner function of the motional quantum state by tomography (Kurtsiefer et al., 1997).

Technically, to perform de Broglie interference experiments, one has to overcome challenges of preparation of intense gas-phase beams, of preparation of spatial and temporal coherence of the matter wave, and of the efficient detection of the particles. Central to all experimental demonstrations of matter wave interference are optical elements which serve to coherently manipulate wave phases, and in particular to divide the wave fronts, thus creating different possible interference paths. While bulk and surface crystals are well-adapted to diffract electrons and neutrons with de Broglie wavelengths in the range of 1 pm to 10 pm , it is impossible to use the same structures for atoms or molecules as those would stick to the surfaces. Typically beam splitters for molecules are real-
ized by gratings. Gratings are nanofabricated highly ordered periodic structures of freestanding nanowires made from metal or semiconductor materials or realized by standing light fields using the Kapitza-Dirac effect (Kapitza and Dirac, 1933). Today the tightest bound for the quantum to classical transition comes from molecule interferometry. We shall give a brief history on molecule interferometry before we describe more details of the work horse of molecule interferometry - the Talbot-Lau interferometer.

Beams of small molecules were first scattered at surfaces in the experiments by Estermann and Stern in 1930 (Estermann and Stern, 1930) followed by interferometry experiments with di-atomic molecules in the 1990s. In 1999, Matter-wave interferometry with large neutral molecules was first demonstrated with the $C_{60}$ fullerene in Vi enna (Arndt et al., 1999). Fraunhofer farfield interference was shown by using molecular diffraction at a single nano-fabricated silicon nitride grating with a grating constant of 100 nm . The beam was collimated by a series of $5 \mu \mathrm{~m}$ slits to a beam divergence smaller than the expected beam diffraction angle of about $10 \mu \mathrm{rad}$. Only very few molecules originally in the beam reached the diffraction grating and the detector and typical count rates were of only very few molecules per second with a detector efficiency of around
$10 \%$. The resulting long integration time to resolve the interference pattern makes such experiments susceptible to noise. Prospects for large particle far-field interferometry and the related Poisson spot experiments can be found elsewhere (Juffmann et al., 2012b) as well as new developments of promising techniques for far-field experiments (Juffmann et al., 2012a).

Talbot-Lau interferometer: Later molecule interferometry experiments were done with a so called Talbot-Lau interferometer (TLI) to increase the beam intensity of the diffracted beam. Technically, a multi-grating Talbot-Lau interferometer (TLI) scheme has been implemented to realize this experiment. A TLI is operating in the near-field diffraction regime described by Fresnel integrals, where the spatial period of the diffraction grating and the interference pattern are on the same size scale. The scheme has been invented by Clauser to cope with beams of low intensity and low collimation in interferometry experiments Clauser and Reinsch, 1992). An advantage of a TLI with respect to a Fraunhofer single-grating far-field interferometer is that the scaling of the distance between the gratings (Talbot length: $L_{T}$ ) is inversely proportional to the de Broglie wavelength $\lambda_{d B}$ but quadratic with the grating period $d, L_{T}=d^{2} / \lambda_{d B}$. This helps to compensate for small de Broglie
wavelength by increasing the distance between the gratings.

In more detail, the three grating TLI operates with weakly collimated molecular beams with divergence of about 1 mrad and accepts a large number of molecules in the initial beam contributing to the final interference pattern. The first grating prepares the beam coherence, while imprinting a spatial structure on the molecular beam (see illustration of the Talbot-Lau interferometer in Fig (4) acting as an absorptive mask. The second grating - the diffraction grating - is then simultaneously illuminated by $10^{4}$ individual coherent molecular beams. The second grating generates a self-image at the Talbot distance $L_{T}=d^{2} / \lambda_{d B}$, where $d$ is the grating constant and $\lambda_{d B}$ is the de Broglie wavelength. Therefore each of the $10^{4}$ initial sources will be coherently mapped to the Talbot distance after the second grating. This Talbot effect results in a self-image of the second - the diffraction - grating at half-integer multiples of $L_{T}$. While the Lau effect makes an incoherent summation of many individual coherent beam sources located at the first grating to an integrated signal. Basically, the number of molecules contributing to the final interference pattern is multiplied by the number of illuminated slits of the first grating. Therefore, all coherent beams from the $10^{4}$ source slits will be incoherently summed to


FIG. 4 Different configurations of the TalbotLau interferometer are shown. (A) Three material grating as experimentally realized in (Brezger et al., 2002). (B) Kapitza-Dirac-Talbot-Lau interferometer realized in (Gerlich et al., 2007). (C) Optical Time-Domain Ionizing Matter Interferometer [OTIMA] as proposed in (Nimmrichter et al., 2011a)
contribute to the same interference pattern. The third grating is then placed close to this Talbot position after grating two and scans over the diffraction pattern perpendicular to the molecular beam to enable integrated signal detection. This is enabled as the period of the scan grating exactly matches the period of the Talbot self-images of the diffraction grating. Talbot, Lau and Talbot-Lau effects have been nicely illustrated by recent optical experiments (Case et al., 2009). The successful implementation of different TalbotLau interferometers for molecules has been summarized in a recent review article (Hornberger et al., 2011), where more and detailed information about techniques and requirements can be found.

A recent version of the TLI is the socalled Kapitza-Dirac-Talbot-Lau interferometer (KDTLI), which has been used to demonstrate interference of a 3 nm long diazobenzene molecule (Gerlich et al., 2007). Here the second - the diffraction grating was realized by an optical phase grating, where molecules are diffracted at periodic optical potentials due to the Kapitza-Dirac effect (Kapitza and Dirac, 1933). The use of light gratings avoids the dispersive van der Waals (vdW) or Casimir-Polder (CP) attraction between molecules and gratings Hornberger et al., 2009), which is known to phase shift the interference pattern but also to re-
duce the fringe visibility (Hackermüller et al. 2003) due to dispersive effects for molecular beams with finite velocity spread. The interaction effect scales with the particle velocity and particle polarisability as well as the dielectric properties of the grating material. The details of the interaction potentials and related vdW-CP effects and how those can be investigated by molecule interferometry experiments are still under intense investigation (Buhmann et al., 2012; CanaguierDurand et al., 2012a b). Estimations show that even with improved velocity selection schemes, where the width of the selected velocity is even below one percent of the mean velocity at full width half maximum, it is expected to disable interference with particles with masses beyond $10^{5} \mathrm{amu}$. Presently, the largest de Broglie interfered particle is an about 7000 amu massive perfluoro-alkylated $C_{60}$ molecule (Gerlich et al., 2011).

The specifications of a TLI can be estimated easily. For instance the specifications for interference of $10^{6}$ amu massive particle in a Talbot-Lau type interference with gratings of period $d=100 \mathrm{~nm}$ : at a Talbot distance of $L_{T}=2.5 \mathrm{~cm}$ a particle velocity of $\mathrm{v}=1 \mathrm{~m} / \mathrm{s}$ would be needed to be constant over this distance $L_{T}$. [Grating opening on the order of 50 nm , which is close to minimum possible size for grating dimensions which can be realized at the moment by material structures
as well as by light. Also, particle size for $10^{6} \mathrm{amu}$ is on the order of 10 nm to 30 nm and the grating opening should be larger than the particle]. For a higher mass the particle would need to be slower at the same Talbot distance or alternatively the Talbot distance would need to be extended for the same particle speed. Simple estimates show that for a particle of $10^{8}$ amu we find already $L_{T}=2.5 \mathrm{~m}$ at the same speed and grating constant. However any particle traveling over that distance, even if it starts at zero velocity, will be accelerated to higher speed than $1 \mathrm{~m} / \mathrm{s}$ (namely $7 \mathrm{~m} / \mathrm{s}$ over 2.5 m ) by Earth's gravity $g$ acting over that distance on the particle. Slowing or compensation of acceleration by additional carrier fields or in space experiments would be needed to overcome this limitation while presenting a significant experimental challenge. Therefore, TLI experiments (where the speed of the particle or equivalently it's wavelength has to have a certain value between the gratings) without compensation of Earth's gravity are limited to a particle mass of around $10^{7} \mathrm{amu}$. That limit exists for all possible orientations of the interferometer to $g$. While that is true the alternative single grating far-field interferometry is not limited by $g$. Diffraction of the matter-wave at the location of the grating, the separation of maxima and minima of the interference pattern does not depend
on the speed of the particles, but only on the distance between the grating and the particle detector.

## Technical challenges for mass scal-

 ing: The quest is for new technologies which can efficiently control and manipulate the centre of mass motion of heavy particles. The mature techniques of ion manipulation and of optical tweezing are of particular interest to scale the mass up to particles of 10 nm to 1 um (mass of $10^{6} \mathrm{amu}$ to $10^{10} \mathrm{amu}$ ) in diameter. All experiments have to be performed under ultra-high vacuum conditions to avoid decoherence by collision. We will come back to this in Sec. IV.F. In particular the challenges are:- Generation of intense particle beams: Particles need to be slow if massive to keep the de Broglie wavelength within the range for experimental possibilities (not much smaller than pm ), which means the longitudinal velocity needs to be small. The ideal particle beam has a high phase space density, which means that many (ideally all) particles should propagate at the same speed. The beam needs to be highly collimated, which means that the transverse velocity needs to be as small as possible. [Very high beam collimation ( $<10 \mu \mathrm{rad}$ ) would enable the conceptually simpler far-field single grating
interferometry.] In the wave picture this means that the transverse or spatial coherence needs to be high. All this could be achieved by cooling techniques, those which effect the centre of mass motion of particles, which yet have to be developed for complex particles. Very interesting and promising approaches have been followed in the last few years. This especially includes the collisional buffer gas cooling (Maxwell et al., 2005) as well as optical cooling techniques (Shuman et al., 2010). Both techniques have so far been demonstrated for diatomic molecules. Interestingly, a feedback cooling technique has been realized for optically trapped beads of $1 \mu \mathrm{~m}$ in diameter in the field of optomechanics $(\boxed{\mathrm{Li}}$ et al., 2011). We will come back to this in Sec. IV.D.

Furthermore particles need to be structurally stable to survive launch and detection procedures. This includes techniques to generate gas-phase particles such as by thermal or laser induced sublimation, laser desorption or ablation, but also sprays of particles from solutions, etc. and the subsequent manipulation of such particles to meet the coherence requirements of matter-wave experiments.

- Beam path separation: This is the need for coherent beam splitters and other matter-wave optical elements. While many different techniques for beam splitters are known for instance for cold atoms (Cronin et al., 2009), material and optical gratings are promising candidates to act as beam splitters for very massive particles as those already work for large molecules. The challenging part is the realization of gratings with a high enough precision in periodicity. The demand on the periodicity is very high for the TLI scheme, where the average grating pitch has to be accurate within sub-nanometer scales between all gratings. This can only be realized so far by complicated optical interference lithography techniques. For far-field gratings the demand is lower and electron-beam lithography with sophisticated alignment pattern to avoid stitching errors is possible for fabrication. The ability to form laser light gratings from retro-reflection or other superposition of laser beams depends on the intensity and frequency stability of the laser. The power of the laser needs to be sufficient to form an optical potential strong enough to act as a phase grating. This is on the order of some 1 W continuous power.

The limited availability of stable and medium power UV and XUV (wavelength $<200 \mathrm{~nm}$ ) lasers limits the fabrication of grating periods by optical lithography as well as the optical grating periodicity to about 100 nm (opening of about 50 nm ). About an order of magnitude smaller grating periods can be possibly fabricated by electronbeam lithography or direct focus ion beam (including the novel $\mathrm{He}+$ direct write) milling. Another limitation is that the grating area has to fit the size of the particle beam diameter which is on the order of 1 mm . Not all fabrication techniques are capable of manufacturing precise gratings on that size scale. However, in combination with an efficient detector this dimension can be decreased.

- Efficient detection of neutral heavy particles: Ideally, we want single particle detection resolution. For example in the most recent molecule interferometry experiments detection is realized by ion counting after electron impact ionization, which is known to have a very low ionisation efficiency $\left(10^{-4}\right)$. This has to be at least kept at the same level for particles of increased size and mass. To resolve the interference pat-
tern a spatial resolution on the order of the grating period is needed for nearfield interferometry experiments, which is elegantly realized in the case of the TLI by the third grating. Also a high spatial resolution of the detection is needed if the particle beam is not velocity selected before entering the interferometer gratings. This is important to select the temporal coherence which is given by the distribution of de Broglie wavelengths of matter waves emitted by the source.

In the following, we will discuss different alternative approaches on how to possibly implement experiments to probe the quantum superposition of particles. We will summarize different proposals for such experiments.

## 1. Neutral particles vs charged particles

Quantum superposition experiments per se need to avoid any decoherence effect which is able to read out which-way information and to localize the particle. A neutral particle is a natural choice for superposition experiments as the number of possible interactions, which would enable a readout of which-way information, is reduced in comparison to charged particles. That is especially true for superposition of slow particles. Therefore all interference experiments with
molecules have been performed with neutrals. On the other hand the centre of mass motion of charged particles can be manipulated and controlled to a higher degree by external electric and magnetic fields. This would be handy to prepare coherent particle beams. Here we will discuss the benefits and possibilities for charged particle interferometry as neutrals have been covered in the previous section. From the matter wave point of view we have to achieve the same parameter values: de Broglie wavelength, periodicity of the diffractive element, etc. to observe an interference pattern. This especially means that for a given (high) mass the speed needs to be rather low: for $\mathrm{m}=10^{6} \mathrm{amu}$ requires around $\mathrm{v}=1 \mathrm{~m} / \mathrm{s}$.

Electrically trapping charged particles such as electrons or ions in Paul and Penning traps has a long and successful history (Paul, 1990). It has been used for studies of fundamental physics such as the precise evaluation of physical constants Brown and Gabrielse, 1986), for quantum information processing with one or many ions (Duan and Monroe, 2010; Leibfried et al., 2003; Singer et al. 2010), in chemical physics to investigate the kinetics and dynamics of chemical reactions on the few molecule level under controlled conditions (Kreckel et al., 2005 , Mikosch et al., 2010; Willitsch et al., 2008) such as with buffer gas cooled polyatomic
ions in multi-pole traps (Gerlich, 1995, Gerlich and Borodi, 2009). The obvious benefit of using charged particles for matterwave experiments is the higher control over the motion of the particles. Guiding, trapping, and cooling is possible even for massive ions. For instance 200 bio-molecules of 410 amu have been co-trapped with laser cooled atomic ions $\left(\mathrm{Ba}^{+}\right)$and cooled to $150 \mathrm{mK}(\mathrm{Os}-$ tendorf et al., 2006). This sympathetic cooling via Coulomb interaction of laser-cooled atomic ions with the molecular ions has been demonstrated to be efficient, however a difficulty which remains is to realize an ion trap which is stable for both species. The mass $m$ over charge $q$ ratio $m / q$ must not be too different for both particles, which demands also a high control on the ionisation technique for atom and molecule.

Most of these techniques aim to spatially fix the ion in the trap to increase interaction times for spectroscopic and collision studies or to cool the ions, while we are interested in well controlled centre of mass motion for interference. It might be difficult to achieve a coherent centre of mass motion manipulation, but seems not impossible also with respect to exciting new guiding techniques such as the recently demonstrated microwave-manipulation Hoffrogge et al., 2011), the manipulation of ions by light (Schneider et al., 2010), or multi-pole
trap techniques (Gerlich, 1995).
To scale up the mass of ions for experiments in order to test collapse models the very mature techniques of gas-phase cluster sources are available, such as sputter Magnetron sources (Haberland, 1994) or other noble gas aggregation sources with pick up cell for larger molecules (Goyal et al., 1992; Toennies and Vilesov, 1998). Beams of such sources are intense as cooled by the supersonic expanding noble gas and the mass of a single cluster can be $10^{9} \mathrm{amu}$ and beyond (von Issendorff and Palmer, 1999). In combination with quadrupole mass filters, which work very similar to ion Paul traps, metal clusters of very narrow mass distributions can be realized $\mathrm{m} / \Delta \mathrm{m}=25$ (Pratontep et al., 2005). Additional techniques will need to be realized for deceleration of such big clusters, but as long as the particle is charged a high degree of control is guaranteed.

While this is true massive ion interference has yet to be shown to work. A recent review article on ion interferometry lists that so far only electrons and the $\mathrm{He}^{+}$ion showed quantum interference (Hasselbach, 2010). Typically electrons are diffracted at bi-prisms or solid surfaces as if applied for holography (Tonomura, 1987), but also light gratings are possible utilizing the Kapitz-Dirac effect (Batelaan, 2007). Electron interferometry has been for instance used to investi-
gate the Aharonov-Bohm effect Tonomura and Batelaan, 2009). The general understanding, which is supported by experiments on for instance image charge decoherence effects, is that ions have to be very fast to not decohere via one of the multiple interaction channels with the environment. As for any other interferometry experiments any interaction which is able to read back which-way information will localize the particle and destroy quantum superposition Sonnentag and Hasselbach, 2007). The challenge will be to avoid and shield all possible interactions of ions with surrounding matter and fields, such as for instance the coupling of the ion to its own image charge in a metal surface.

As for the neutral particles in the case of a TLI the acceleration by Earth's gravitational field has to be compensated. Guiding potentials have to be extremely flat to not influence the superposition state, to not localize the particle. External electro-magnetic fields have to be shielded by a Faraday cage of the right dimensions and materials, where recent technological progress has been made for the stabilization of magnetic field in atom experiments (Gross et al., 2010). Very stable electrical power supplies will be needed for the cold Paul trap for ion beam generation and an ion guide field. Electric stray fields from patch effects of adsorbed atoms and molecules at the shielding and elsewhere
may be avoided as well as time varying electronic inhomogeneities in the shielding material. Edge fields of the guiding electrodes and other parts inside the shielding have to be carefully considered.

However, a simple estimate shows that for instance all applied voltages would have to be stabilized to the level of below $10^{-10} \mathrm{~V}$ for the time of interference which seems to be impossible to be achieved at the moment. At present only the neutrals show success for large particle centre of mass motion interference. On the other hand interference attempts with larger particles suffer from the non-existence of guiding, slowing and cooling techniques for neutrals. Therefore a clever solution for now is to try to take the best of both the worlds: manipulation of charged particles and interfering after neutralization, which we describe in the following section.
2. The compromise - combination of techniques for charged and neutral particles: OTIMA

A novel three light grating Talbot-Lau scheme in the time domain aims towards the interference of particles of up to $10^{9}$ amu as proposed by the Vienna molecule interferometry group as described in Nimmrichter et al. (Nimmrichter et al., 2011a). This interferometer is called optical time-
domain matter-wave (OTIMA) interferometer. The charged particles will be provided by a mass filtered metal cluster aggregation source as mentioned before. A further cooling/deceleration device will reduce the velocity of the big clusters which is an existing technology for charged particles. A chopper modulated particle beam can be used for mass as well as velocity selection of the clusters in combination with a time of flight mass spectrometer (TOF-MS) detector.

The main invention is a clever neutralisation/ionization scheme implemented as the interferometer. The neutralization of the clusters to enable a coherent propagation of the superposition state is planned to be achieved by light-matter effects directly at the light gratings. The scheme makes use of a sequence of three vacuum ultra-violet (VUV, $\lambda=157 \mathrm{~nm}) \mathrm{ns}$ long light pulses to realize the interferometer gratings. The energy of a single photon of about 8 eV is sufficient to ionize or neutralize by photo-detachment metal clusters (Haberland, 1994). These processes are also applicable to large bio-molecular complexes (Marksteiner et al., 2009). The light intensity pattern realized by three retroreflected laser pulses hitting the propagating particles transversely at precisely timed locations with respect to each other realize the TLI gratings with grating period of $\lambda / 2$. The formed standing wave normal mode pattern
forms on one hand the gratings but is also a spatial resolved ionisation/neutralization device: the intensity in the antinodes is sufficient to ionize or neutralize the particles while it is not in the nodes. Therefore clusters which pass through the antinode will get ionized while others in nodes will not. This in combination with electrodes to divide the beams of neutrals and ions is the realization of an absorptive grating. The first and third gratings need to be absorptive gratings, which means they need to spatially mask out parts of the cluster beam and are realized by intensity dependent ionisation of the molecules. The second grating needs to be a phase grating and will be realized by the optical dipole force acting on the particle by making use of the Kapitza-Dirac effect (Batelaan, 2007).

The charged clusters will be neutralized by the first grating, diffracted at the second phase grating and ionized again for detection at the third grating. More details about the setup and simulations of the expected interference pattern can be found in Nimmrichter et al., 2011a) and (Reiger et al., 2006). This is a promising attempt to realize matter wave experiments with very massive particles to test collapse models. More details about the OTIMA approach to test CSL can be found in (Nimmrichter et al., 2011b). The spatial size of the superposition state is estimated by
the grating constant and is on the order of the CSL parameter, namely $r_{C}=10^{-5}$. One estimate for CSL $\lambda$ is $10^{-10} \mathrm{~Hz}$ which corresponds to a mass of about $10^{6} \mathrm{amu}$.

## D. Optomechanics: Cantilever

Here we describe an experimental approach which is alternative to matter wave interferometry. While the aim to understand the limitations of quantum mechanics is as old as quantum mechanics itself, the first proposals for a table-top experimental test by using the superposition or other non-classical states of massive mesoscopic or even macroscopic mirrors have been published in the late 1990's Bose et al., 1997, 1999, Marshall et al., 2003).

The mechanical motion - the vibration of the mirror, which was later realized by a nano-mechanical or micro-mechanical cantilever, has to be cooled to the quantum mechanical ground state which is modeled by a simple harmonic oscillator $\left(k_{B} T<\hbar \omega\right)$. The readout of the vibrating mirror is done by coupling it to a sensitive optical interferometer to compare the light phase with a stabilized cavity. The conceptual idea is to first prepare the mechanical oscillator in the vibrational or phononic ground state $|0\rangle$ by cooling and then generate a coherent superposition of or with the first excited vibra-
tional state $|1\rangle$ by single photon excitations.
A high mechanical as well as a high optical quality (Q) factor is needed to reach the regime of low dissipation to strongly couple optics to mechanics and to cool the device ultimately to the ground state. While optical control of cantilevers was under investigation for quite some time it was only in 2006 that two groups reported the successful optical cooling of mechanical cantilevers (Gigan et al., 2006; Schliesser et al., 2006). Interestingly, the cooling mechanism is very similar to the optical cooling of atoms: The optical resonance - in most cases an optical cavity resonance - is slightly detuned to the cooling sideband of the mechanical resonance, which is typically in the range of 1 MHz to 10 GHz . This opens a cooling channel for the mechanics of the cantilever though the optical leakage of the cavity. Achieved temperatures corresponded still to a high phononic occupation - to many vibrational states occupied, but it boosted the rapid development of an exciting new field of research, namely opto-mechanics. This is summarized elsewhere (Aspelmeyer et al., 2010; Kippenberg and Vahala, 2008; Marquardt and Girvin, 2009). Also the first schemes on how to generate and probe the superposition state of a cantilever appeared (Kleckner et al., 2008).

Only a few years later - and that is probably one of the most exciting developments


FIG. 5 Optomechanics. Upper Panel: Mechanical Resonator Interference in a Double Slit (MERID) as proposed in (Romero-Isart, 2011). The centre of mass motion of a single optically trapped nanoparticle is first cooled and then superimposed by an optical double potential. The interference pattern evolves in free fall after switching off the trapping field. Lower Panel: Prototype of optomechanically cooled cantilever as realized in (Gigan et al., 2006). Quantum optical detection techniques enable the sensitive read out of vibrations as they couple to light fields.
in experimental physics over the last few years - ground state cooling of micrometer sized structures by optical techniques has been achieved by Cleland's group (O'Connell et al. 2010) and by the groups of Aspelmeyer and Painter (Chan et al., 2011). Advanced nanofabrication technology enabled realization of structures with both high mechanical and high optical Q factors $\left(10^{5}\right)$ in addition to clever optical or electronic readout techniques. This opens the door to many exciting quantum information processing and sensing experiments in the near future, but let us go back to our initial question if those structures can test our beloved collapse models.

These structures are very massive, $10^{6}$ amu to $10^{15} \mathrm{amu}$ depending on their size, but the vibration amplitudes when compared to the parameter $r_{C}$ of the CSL model are very small. This limits their ability to test collapse models and the parameter range to test CSL by such systems is indicated in Fig. 3. To investigate this a little further we estimate the spatial size $x_{0}$ of this position superposition state by using the size of the zero point motion of a simple harmonic oscillator $x_{0}=\sqrt{\hbar / 2 m \omega}$, where $m$ is the mass of the cantilever and $\omega$ its frequency in a harmonic potential. This spatial size of the ground state at $25 \mu \mathrm{~K}$ is $1 \cdot 10^{-15} \mathrm{~m}$ for a typical micromechanical oscillator of a mass of 50 ng resonating at about 1 MHz (Gigan et al., 2006).

However, spatial superpositions to tests CSL have to be on the order of 10 nm or larger $\left(r_{C}=10^{-5} \mathrm{~cm}\right)$, which is roughly seven orders of magnitude away from what micromechanical oscillators can achieve at the moment. Mass or frequency or combinations of both have to be improved by that amount, which is very difficult as for most materials mass and resonance frequency are coupled and depend on the spatial dimensions of the cantilever. To see vibrational state superposition larger than the quantum mechanical ground state the optomechanical device has to be driven in an extreme regime: eight orders of magnitude in mass or frequency at optical finesse of $10^{6}$. But there are very interesting systems providing a larger zero point motion such as novel carbon materials with exceptional mechanical properties (Iijima, 1991; Novoselov et al., 2004): e.g. individual single-wall carbon nanotube oscillators generate $x_{0}=1 \mathrm{pm}$, with $\mathrm{m}=8 \times 10^{-18} \mathrm{~g}$ and at $\omega=100 \mathrm{Mhz}$ at a ground state temperature of $\mathrm{T}=2.5 \mathrm{mK}$ (Sazonova et al., 2004). Such systems have been used for mass sensing with hydrogen mass resolution Chaste et al. 2012). One big challenge for such carbon materials stays which is their very small absorption and reflection cross sections. This means it is not clear how to realize the needed high optical quality factor for optomechanics. But hope is to cool via other in-
teraction channels possibly in the electronic regime (Brown et al., 2007; Chen et al., 2011; Eichler et al., 2011).

Another difficulty for the test of collapse models by cooling mechanical cantilevers to the ground state $|0\rangle$, is that the light field has to be switched on all the time. Otherwise the substrate where the cantilever is coupled to will rapidly heat back which is probably much faster than the collapse time. There is not much time for 'free propagation' of the superposition. New ideas on pulsed optomechanics may help to prepare and reconstruct quantum state of the mechanical motion (Vanner et al. (2011) faster. So ideally we would prefer to use a massive harmonic oscillator which is realized without a link to any substrate. This is what we discuss in the next section.

Interestingly, mirror stabilization ideas are in principle linked to the much larger interferometers for the detection of gravitational waves (Braginsky et al., 2002), while in a different parameter range due to the much higher mass of the mirrors in use.

## E. Micro-spheres and nanoparticles in

 optical potentialsHere we describe a new and very promising route to test collapse models by generating spatial superposition states of the centre of mass motion of very massive nanopar-
ticles and possibly even microspheres. This is a combination of optomechanics with centre of mass motion superposition states as in matter-wave interferometry. Optically trapped particles represent an almost ideal realization of a harmonic oscillator as already mentioned by Ashkin (Ashkin, 1970) and more recently re-discovered for cavity optomechanics (Chang et al., 2010; RomeroIsart et al., 2010). In comparison to mechanical cantilevers as discussed in the previous section there is no mechanical link acting as a dissipation channel for the mechanical oscillation in such systems if implemented in a vacuum chamber to avoid collisions with background gas particles. Therefore the mechanical quality factor is very large. Such trapped particles can be seen as an optomechanical system and techniques such as for cooling the oscillation - which is now the centre of mass motion of the particle in the optical trap - need to be implemented. Both criteria to test CSL, and other collapse models, are fulfilled: a high mass of the particle and a width of the spatial separation which can be comparable to $r_{C}$.

In addition these systems enable free centre of mass motion of the initially trapped particles after switching off the trapping field and after generation of the spatial superposition. This allows for combination with matter-wave interferometric techniques and
schemes. Recently, some ideas have been put forth how to perform tests of quantum superposition with so-called beads (balls of diameter 10 nm to $10 \mu \mathrm{~m}$ made of glass or polystyrene) (Romero-Isart et al., 2011b|c). The basic sequence for such experiments is to first optically dipole trap a single particle, use optical techniques to cool the centre of mass motion of the bead in the optical trapping potential. The next step is to generate a superposition state of the particle position by a double-well optical potential by single photon addressing of the first excited vibrational state, as theoretically described within cavity quantum electrodynamics (QED). Cooling must be sufficient to increase the size of the particle wave-packet to overlap with both wells, so that there is an equal probability to find the particle left or right - the coherent superposition by a measurement of a squared position observable. After switching off the trapping potential in free fall the spatial density distribution of the particle in multiple subsequent experiments can be mapped and evaluated for a quantum signature by for instance state tomography through Wigner function reconstruction (Romero-Isart et al., 2011c) or much simpler by interference pattern detection at a fixed detector position. This is basically a clever double-slit interference experiment applied to very massive objects (polystyrene
bead of about 30 nm diameter has a mass of $10^{6} \mathrm{amu}$ ). To get a significant detection statistics the single particle experiment has to be repeated many times. The connection to test collapse models is worked out in detail in (Romero-Isart, 2011) and the experiment is called MERID which is the shortcut for mechanical resonator interference in a double slit.

The manipulation of microscopic particles as silica and polystyrene sphere but also biological cells and even living organisms as viruses by optical fields has been pioneered by Ashkin and others since the 1970s and is now very broadly applied in many fields of science Ashkin, 1970; Ashkin and Dziedzic, 1987). Techniques which are typically summarized by the term optical tweezing include the broad fields investigating optical angular momentum (Allen et al., 2003), optimizing the trapping, levitating and guiding of single dielectric particles by optical gradient and scattering forces in various geometries Ashkin, 2006; Chu, 1998) including the guiding through hollow core photonic crystal fibres (Benabid et al., 2002) and optical binding (Dholakia and Zemánek, 2010). Ashkin and co-workers demonstrated already the trapping of polystyrene and glass microspheres, of viruses and bacteria and even of complete cells in solutions and high-vacuum. They developed a vacuum loading system
and they demonstrated the stable levitation of particles at a vacuum of $10^{-6}$ mbar for half an hour by a feedback stabilization technique. A very detailed summary of this field can be found in Ashkin's book (Ashkin, 2006). The particle size is typically limited to be not smaller than $1 \mu \mathrm{~m}$ to form a stable trap, while optical near-field techniques have been very recently used to trap single nanoparticles with the help of plasmonic (Juan et al. 2009) or photonic crystal structures (Rahmani and Chaumet, 2006) in solution. An application of such advanced trapping techniques in vacuum has to engineer the challenge of particle-surface van der Waals (vdW) and Casimir-Polder (CP) interactions or in turn could be used to investigate those interactions. However it has been demonstrated very recently that even 30 nm particles can be optically trapped in tightly focused free beams under vacuum conditions when gradient forces dominate scattering forces and with parametric stabilization (Gieseler et al., 2012).

Experimental challenges: Cooling is again the key for this experiment. Here one needs to cool the centre of mass motion of a bead in an optical field ideally to the ground state: Ashkin pioneered the feedback stabilization (Ashkin, 2006). The Doppler cooling using whispering gallery modes of the particle has been proposed (Barker, 2010). Re-
cently, the cooling of the centre of mass motion of a single $1 \mu \mathrm{~m}$ glass bead to 1 mK has been achieved by a fast feedback stabilization technique (Li et al., 2011) as well as the optical parametric stabilization of a single silica nanoparticle ( 30 nm ) under vacuum conditions ( $10^{-4} \mathrm{mbar}$ ) at 400 mK (Gieseler et al., 2012). These are the first promising steps to realize the proposed experiments to test superposition of such large and heavy particles. Importantly, feedback stabilization techniques will enable to trap beads under vacuum conditions to dissipate the kinetic energy of the trapped particle. All experiments - as the competing cluster and molecule interferometry experiments - have to be performed at ultra-high vacuum (UHV) conditions (p $<10^{-10} \mathrm{mbar}$ ) to avoid collisional decoherence of the superposition state Hornberger et al., 2003). A further challenge is that the interferometer has to be stable over the duration of many single particle experiments. One idea is for experiments in space while it is not clear yet, if such an environment is needed Kaltenbaek et al., 2012). Centre of mass motion trapping would also be possible with ions (Leibfried et al., 2003). The electric trapping fields would replace the optical trap, but optical fields would still be needed for the cooling. While this is true free propagation of the charged particles as after switching off the trap in the protocols for bead superposition
experiments as explained above - would not be possible. The Coulomb interaction will certainly dominate the motion of the particle - it will not be a free motion. On the other hand a recent proposal with magnetic levitated superconducting particles claims feasibility for large superpositions Romero-Isart et al., 2011a).

To avoid the difficulty of ground state cooling, one possibility is to use the TalbotLau interferometer scheme. Here, as we know from molecule interferometry, the requirements on cooling are lowered as a quantum interference effect can be observed at low spatial coherence of the matter wave. Centre of mass motion temperatures of 1 mK (for a given TLI geometry) would be sufficient to observe interference. This will work with a single particle source, but also many particles in parallel traps would be possible which could significantly reduce the operation time of the interferometer and therefore lower the stabilization criteria on the interferometer. We are looking forward to see more exciting developments in this rapidly progressing field of research.

One significant advantage of cavity optomechanics with trapped particles is the in principle very large separation of left and right for the superposition state which can be tuned by the optical field. Furthermore the optical field can be switched off and the
particle can propagate in free space - showing the signature of superposition: an interference pattern in the spatial distribution, which is not possible in cantilever optomechanical systems: if the field is switched off the quantum mechanical system does not exist any more. The size of the beads in the proposed experiments is on the order of 10 nm to 100 nm , exactly the same size and mass range where cluster matter-wave experiments such as OTIMA are heading towards. We think that in the foreseeable future collapse models will be tested in this size range and we will see which experimental strategy is more fruitful or faster. Certainly both approaches have to overcome a number of technical challenges mainly related to coherent state preparation. In the following we will discuss decoherence effects which affect both cluster matter-wave and bead optomechanics experiments and consider how those limit the parameter range for possible experiments.

## F. Environmental Decoherence

While the aforementioned and discussed collapse models can be seen as an exotic decoherence mechanism we here discuss decoherence effects of the environment interacting with the particle in superposition. Collisions with background particles and thermal radiation of the superimposed particle itself
are counted as the major processes to localize the superimposed particle. Both docohering effects affect all the different experimental schemes to perform mesoscopic quantum superposition experiments and set limits on particle (and experimental setup) temperature as well as background pressure inside the vacuum chamber depending on the size of the particle. According to decoherence theory the superposition state is destroyed and the particle is entangled with the environment whenever any interaction of the superimposed particle with the environment has the sufficient resolution to localize - to measure the position of - the particle, which-way information is read out.

We note, that a intrinsic difficulty with the test of collapse models is that it is clear how to falsify a proposed model with respect to predicted parameters. If on the other hand no interference pattern is shown by the experiments all systematic effects related to environmental decoherence have to be excluded as reason for the quantum to classical transition. Here scaling of one of the test parameters such as mass or spatial width $x$ of the superposition as well as a detailed study of the environmental decoherence effects will help to overcome this problem.

Mathematically, decoherence is described (as for the case of collapse models) by the effect on the off diagonal elements of density
matrix of the system including the particle and the environment which are reduced by the decoherence effect as given by the master very similar to Eq. 260. The effect is evaluated by the decoherence rate function $\Gamma$ as given in Eq. 261. More details on the concept and formalism of standard decoherence theory can be found in the references given in Section I. More details on estimations of decoherence effects and associated decay rates for superposition experiments can be found in Romero-Isart (Romero-Isart, 2011) and Nimmrichter et al. Nimmrichter et al., 2011b). Both processes, collision and black body photon decoherence, have been experimentally investigated and compared to theory with fullerene interference Hackermüller et al., 2004, Hornberger et al., 2003). We will here summarize the most recent estimates from the literature to give boundaries to the experiments.

## 1. Thermal decoherence

The emission, absorption, scattering of thermal - black body - radiation by the particle in superposition can localize the particle if the wavelength of that light is comparable or smaller than the size of the superposition. The emission of thermal photons is seen as the most important effect as the internal temperature of the particle is typically higher
than the temperature of the environment. As an example for $C_{70}$ fullerenes there is still full quantum contrast for emission of thermal photons by the fullerene at about 1500 K , as experimentally observed (Hackermüller et al. 2004; Hornberger et al., 2005). The interference visibility is rapidly reduced for temperatures at above 2000 K where the wavelength of the emitted photons is comparable to the size of the superposition which was about $1 \mu \mathrm{~m}$ in this experiment. For a more detailed discussion of this long and short wavelength regimes see (Chang et al., 2010) RomeroIsart, 2011).

Romero-Isart estimates in (Romero-Isart, 2011) an emission localization time, which is inverse to the superposition decay rate, of 100 ms at a temperature of 100 K for a 50 nm particle, but claims that this time is independent of the particle size. If this claim is correct it contradicts the observation of fullerene interferometry at 1500 K . An extrapolation of this relation to mesoscopic particles ( $10^{6} \mathrm{amu}-10^{8} \mathrm{amu}$, which is 10 nm to 100 nm ) gives temperatures between 800 K and 200 K (Nimmrichter et al., 2011b). This strongly hints that we need to deepen our understanding of this decoherence process by further theoretical and experimental work. In any case the predicted temperatures will have to be reached for the particle and the environment. This may require the cooling of the internal
degrees of freedom of the particle which is an experimental challenge, but buffer gas techniques are in principle applicable to any particle and cool all degrees of freedom (Gerlich and Borodi, 2009; Maxwell et al., 2005).

## 2. Collision decoherence:

Here the collision of the superimposed particle with any other particle present will localize the particle and read out which-way information. This destroys the quantum state and entangles the particle with the environment; according to standard decoherence theory off-diagonal elements of the particleenvironment density matrix vanish. Such collision decoherence processes have been studied in depth for fullerene experiments and an elaborate theory has been developed (Hornberger, 2006; Hornberger et al., 2004, 2003). Applied to the mesoscopic range of $10^{6} \mathrm{amu}$ to $10^{8} \mathrm{amu}$ particles in OTIMA this gives minimum required pressures between $10^{-8} \mathrm{mbar}$ and $10^{-11} \mathrm{mbar}$ Nimmrichter et al., 2011b).

Romero-Isart estimates for a 100 nm sized particle and collisions with ( $\mathrm{N}_{2}$ )-molecules at $10^{-11} \mathrm{mbar}$, a decoherence time of about 100ms in MERID. A more detailed parameter set is given in (Romero-Isart, 2011) and is in agreement with the above values for OTIMA. This pressure is possible to achieve in ultra-high vacuum experiments. Most sur-
face science or ion trap experiments are performed at similar conditions. The parameter set also means that single experimental sequence from preparation of the coherence through superposition and detection has to be done in 100 ms , which seems feasible in OTIMA as well as MERID. This estimate strongly depends on the mean free path of the particle under the given pressure/vacuum conditions and therefore the size of the particle.

Certainly, more work on the theoretical side is needed to investigate those decoherence effects further. This will be an important guidance for experiments. For now it seems that collision decoherence can be controlled for mesoscopic particles while maintaining extreme UHV conditions in the experiments. On the other hand thermal radiation can become a more serious issue for larger particles of $10^{8} \mathrm{amu}$.

## G. Concluding Remarks on Laboratory

## Experiments

Interference of beads suspended in optical field and Interference of large metal clusters are both promising experimental routes to test collapse models. Clearly there is a certain possibility that other experimental routes or variations and combinations of the two main proposals: OTIMA Nimmrichter
et al., 2011a) and MERID (Romero-Isart, 2011) are sucessful in observing a mesoscopic single particle superposition state. OTIMA and MERID are the most advanced experimental attempts reported in the literature at this time.

We think an experimental test of collapse models such as the weakest CSL, with a mass bound of $10^{6} \mathrm{amu}$ is in reach within the next 5 to 10 years. This will only be possible with intense research and development of new technologies for the handling of mesoscopic 10 nm to 100 nm sized - neutral and charged particles. Conditions to control environmental decoherence seem feasible to be reached in the experiments. We hope to see a scientific competition to probe this quantum to classical transition in the coming years. It will be interesting to see if quantum mechanics again survives.

## H. Cosmological Bounds

As we saw in Sec. II.F, stochastic collapse leads to a secular increase energy in the energy of a system. For a group of particles of mass $M$ the rate of energy increase is given by (Adler, 2007)

$$
\begin{equation*}
\frac{d E}{d t}=\frac{3}{4} \frac{\hbar^{2}}{r_{C}^{2}} \frac{M}{m_{N}^{2}} \tag{264}
\end{equation*}
$$

See also (Bassi and Ghirardi, 2003, Pearle and Squires, 1994). If there is no dissipa-
tion in the stochastic collapse model, such an energy deposit will heat the system and the absence of the observed heating can be used to put upper bounds on $\lambda$.

An important case is the ionized intergalactic medium (IGM), which has a temperature of about $2 \times 10^{4}$ between the redshifts of $z=2$ and $z=4$. The IGM is kept in thermal equilibrium because the cooling due to the adiabatic expansion of the Universe and the recombination of the plasma is balanced by the energy input into the IGM that comes from astrophysical processes such as supernova explosions and quasars. An upper bound on the stochastic parameter $\lambda$ can be obtained by assuming that all the heating of the IGM is from the stochastic heating of protons and this gives that $\lambda$ should be smaller than about $10^{-8}$. More detailed discussions of cosmological and astrophysical bounds can be found in Adler, 2007, Feldmann and Tumulka, 2012).

A subject that is recently beginning to draw attention (De Unanue and Sudarsky, 2008; Perez et al., 2006; Sudarsky, 2007, 2011) is the possible role of wave-function collapse in the very early Universe. A possible mechanism for the generation of primordial density fluctuations which eventually grow to form large scale structures is provided by the hypothesized inflationary epoch in the very early history of the Universe, just after the

Big Bang. Inflation may have been driven by a scalar field and the zero point fluctuations of the quantized scalar field serve as a possible source for generating the requisite density inhomeogeneities (Lyth and Liddle, 2009). But how do these quantum fluctuations become classical, as the Universe evolves? Decoherence accompanied by the many worlds interpretation has been proposed as one possible solution (Kiefer and Polarski, 2009). Another possibility is that classicality is introduced by the models of stochastic collapse reviewed here, and it will be important and interesting to understand what sort of bounds are placed on the CSL parameters by the quantum-to-classical transition of density fluctuations in the very early Universe.

## I. Bounds from other physical pro-

 cessesThe standard GRW and CSL values for the model parameters were reviewed in Sec. II.I [including the enhanced value for $\lambda$ proposed by Adler and by Bassi et al. based on latent image formaton in a photograph, and based on image formation in the eye]. Earlier in this section we discussed bounds coming from diffraction experiments and from cosmology. A few other upper bounds have been placed too, taking into account how some other processes would be affected Adler,
2007). In so far as $r_{C}$ is concerned, tentative but plausible arguments have been given that it should be in the range $10^{-5}$ to $10^{-4}$ cm .

Amongst the processes studied thus far are: (i) decay of supercurrents induced by stochastic collapse, giving $\lambda<10^{-3} \sec ^{-1}$, (ii) excitation of bound atomic and nuclear systems [cosmic hydrogen should not decay during the life-time of the Universe : $\lambda<$ $10^{-3} \mathrm{sec}^{-1}$; proton does not decay : $\lambda<10$; rate of spontaneous 11 keV photon emission from Germanium : $\lambda<10^{-3} \sec ^{-1}$, (iii) effect on the rate of radiation from free electrons : $\lambda<10^{-5} \sec ^{-1}$.

## J. Tests of gravity induced collapse

Experiments on molecule interferometry and optomechanics are per se also test of gravity based collapse models: if a violation of quantum superposition were to be observed, the next task would of course be to analyze which of the collapse models is indicated - CSL, gravity, or perhaps something entirely different. Another test, which has received considerable attention in the literature on K-model cited above, is to look for the anomalous Brownian motion induced by the stochastic reductions. Such motion, which of course could also be induced by spontaneous collapse, seems too tiny too be detectable by
present technology, but further careful investigation into current technological limitations is perhaps called for.

The optomechanical cantilever experiment proposed in (Marshall et al., 2003) and discussed above in Sec. IV.D has received particular attention with regard to gravity induced collapse. Related discussions on this experiment can be found in Adler et al., 2005; Bassi et al., 2005a; Bernad et al., 2006).

It is not clear at this stage that there is a unique experimental signature of gravity models which will distinguish it from gravityindependent models of spontaneous collapse.

## V. SUMMARY AND OUTLOOK

In the early years following the development of quantum theory in the 1920s the Copenhagen interpretation took shape. Dynamics is described by deterministic Schrödinger evolution, followed by a probabilistic evolution when the quantum system interacts with a classical measuring apparatus and quantum superposition is broken. An artificial divide was introduced between a quantum system and a classical measuring apparatus, in order for one to be able to interpret results of experiments on atomic systems. While widely accepted, even in its early years the Copenhagen interpretation had worthy detractors including Einstein and

Schrödinger, to whom it was immediately apparent that quantum theory by itself never says that it does not apply to large macroscopic objects, and a direct consquence is paradoxes such as Schrödinger's cat.

Two broad classes of attitudes developed towards the theory. One, given the extraordinary success of the theory, was to not question it at all: since no experiment to date contradicts the theory, one should accept the Copenhagen interpretation and the associated probability interpretation as a recipe for making predictions from theory and comparing them with experiment.

The other was to take serious note of the following difficulties : (i) classical macroscopic systems are also quantum systems, and the quantum-classical divide introduced by the Copenhagen interpretation is vague; (i) the observed absence of macroscopic position superpositions is in conflict with a straightforward interpretation of the quantum superposition principle; (iii) Schrödinger evolution being deterministic, it is 'paradoxical' that probabilities should show up when one tries to describe the outcome of a measurement.

As appreciation of these difficulties grew, the Copenhagen interpretation took a backseat, and today it is perhaps fair to say that the interpretation is no longer considered viable, and should be permanently put to rest,
having well served its purpose in the early phase of quantum theory.

What has emerged on the scene instead, is three classes of explanations which address the difficulties mentioned in the previous paragraph:

## [i] Do not modify quantum theory,

 but change its interpretation: This is the many-worlds interpretation. Quantum linear superposition is never broken, despite appearances. The different outcomes of a measurement are realized in 'different' universes, which do not interfere with each other because of decoherence. It seems to us that in this interpretation it is not easy to understand the origin of probabilities and the Born probability rule.[ii] Do not modify quantum theory, but change its mathematical formulation: This is Bohmian mechanics. There are additional, unobservable, degrees of freedom whose introduction implies that outcomes of measurements can in principle be predicted beforehand, and probabilities can be avoided.
[iii] Modify quantum theory: Replace quantum theory by a different theory, which agrees with quantum theory in the microscopic limit, agrees with classical mechanics in the macroscopic limit, quantitatively and dynamically explains the absence of macroscopic superpositions and the emergence of
probabilities, and whose experimental predictions differ from those of quantum theory as one approaches the mesoscopic and macroscopic regime.

In so far as the empirical situation is concerned, all three explanations are acceptable today. The many-worlds interpretation is in fact perhaps the favoured establishment viewpoint, because it involves minimal change in standard quantum theory: everything can continue to be as such, and that which is not observed is attributed to parallel branches of the Universe which cannot be observed.

In our opinion, herein lies the difficulty with many-worlds, and perhaps also with Bohmian mechanics. They make exactly the same experimental predictions as standard quantum theory, and are empirically indistinguishable from the standard theory. We have at hand an experimentally non-falsifiabile interpretation / re-formulation. It is perhaps a fair statement that no successful theory of the physical world which rested on non-falsifiable premise(s) has in the long run survived the test of time. A prime example is the unobservable ether which was invented to protect a Newtonian, pre-relativistic interpretation of Maxwell's electrodynamics. With hindsight we all know that this discomforting ether was never meant to be, and the invariance of the speed of light [which has been
experimentally verified] turned out to be a far more viable proposition. It seems to us that history is repeating itself: a reinterpretation / reformulation of standard quantum theory is analogous to the ether; it is designed to protect the traditional theory, even though in so doing unobservable features have been introduced. Of course it should also be said that unless experiment dictates to the contrary, conservatism reigns and for some it is all but natural that the traditional theory be protected.

We have here proposed that the third avenue mentioned above be pursued: modify quantum theory. What happens during a quantum measurement is a stochastic process. Even though the initial conditions and evolution for a microscopic system, successfully described by Schrödinger evolution, are completely deterministic, the outcome of a measurement is completely random! A straightforward resolution would be to face the evidence head-on and declare that in the dynamics, deterministic Schrödinger evolution competes with stochastic evolution/reduction. For microsystems Schrödinger evolution completely dominates over stochastic reduction. For macro-systems stochastic reduction dominates Schrödinger evolution, giving evolution the effective appearance of Newtonian mechanics. Somewhere between the micro-
and the macro- the Schrödinger evolution becomes comparable in strength to stochastic reduction. In this regime, which experiments are now beginning to probe, new physical phenomena are predicted, which can be explained neither by quantum theory, nor by classical mechanics. These predictions, which are vulnerable to falsification, are also the strengths of a modified quantum theory. They are benchmarks against which the domain of validity and accuracy of the standard theory can be verified in the laboratory.

To this effect, the quantitative phenomenological models of Spontaneous Collapse, such as QMUPL, GRW and CSL, and others, have been rigorously defined within the well-defined mathematical framework of stochastic dynamics. The models successfully incorporate a Schrödinger type evolution, and a stochastic evolution - the demanding requirements of non-linearity, causality, non-unitarity and norm preservation are successfully fulfilled. Two new universal parameters are introduced. One is a strength parameter which scales with mass, and ensures that stochastic reduction is negligible for microsystems, but significant for macrosystems. The other is a localization length scale which defines the linear extent of the region to which stochastic reduction localizes an expanding wave-function. While known physical and astrophysical processes put upper and
lower bounds on these parameters, there is still a large permitted part of the parameter space and it will now be up to future laboratory experiments to confirm or rule out these parameter values.

Keeping in view the phenomenological nature of these models, which have been devised especially to resolve the quantum measurement problem, it is highly desirable to search for underlying physical principles and theories for these models. Theories which emerge for reasons of their own, and which are not designed for the explicit purpose of explaining measurement. Trace Dynamics does well in this regard: its goal is to derive quantum theory from a deeper level, instead of arriving at quantum theory by 'quantizing' its own limiting case [classical dynamics]. It is an elegant structure in which Schrödinger evolution is the equilibrium thermodyamics of a 'gas' of classical matrices, and the ever-present Brownian motion fluctuations of the gas provide the stochastic process which competes with the equilibrium Schrödinger evolution. Under appropriate circumstances, the Brownian motion becomes important enough to be noticeable, and is responsible for the breakdown of quantum superposition. There perhaps could not be a more compelling representation of 'determinism + randomness' than 'statistical equilibrium + statistical fluctuations'. What is missing still is an important
piece of the puzzle: why do the Brownian motion fluctuations become more important for larger systems?

Keeping Trace Dynamics aside for a moment, one turns to investigate if gravity could couple with quantum effects and lead to an intrinsic uncertainty in space-time structure in such a way as to enable stochastic reduction in macro-systems. At first glance, this seems not possible at all: quantum gravitational effects can only be important at the Planck scale, and Planck length is too small to be of interest in laboratory physics, whereas Planck mass is too large to play a role in the quantum-classical transition. However, as more than one analysis shows, a subtle combination of linear extent of the object [measured in Planck units] and its mass [again measured in Planck units] allows gravity to bring about stochastic reduction. Gravity predicts the quantum-classical transition very much in the domain in which it is expected on other grounds.

Gravity provides a much needed physical mechanism which could underlie spontaneous collapse models. However, a proper mathematical treatment for building a gravity based theory of reduction is not yet available. It is quite possible that a generalization of Trace Dynamics that includes gravity could unify spontaneous collapse and gravity models. Doing so could also explain why
the Brownian fluctuations in Trace Dynamics and spontaneous collapse become larger for larger systems. For, we have indeed explicitly seen in gravity models that the stochastic effect increases with mass.

The need for inclusion of gravity in Trace Dynamics also stems from reasons having to do with space-time structure. With hindsight, it is apparent that only when position localization is complete for nearly all objects in the Universe, it becomes meaningful to talk of a background classical spacetime geometry. If position localization is not achieved, and quantum coherence is significant, indeed that would prevent a meaningful definition of classical spacetime. Under such circumstances, and if one does not want to use classical physics as a starting point for quantization, one would have to include in trace dynamics, a matrix structure not only for the matter degrees of freedom, but also for space-time and gravity. Doing so holds the promise that one will be naturally led to a concrete mathematical formalism for describing gravity induced collapse. Investigation and development of these ideas is currently in progress.

A big stumbling block is the construction of relativistic models of spontaneous collapse. It is difficult to say at this stage whether this block will eventually be overcome, or is an indicator of some incompatibility between dy-
namical models of wave-function collapse and special relativity. The collapse of the wavefunction is an instantaneous process, and is said to violate the 'spirit' of relativity [like in an EPR experiment]. Radical though it may seem, we should eventually not be averse to a possible modification of special relativity to make it consistent with spontaneous collapse theories. Perhaps a generalized trace dynamics in which space and time are nonclassical might have something useful to contribute here.

The development of modified quantum theory has received great impetus from the arrival of pioneering experiments on molecule interferometry and optomechanics which can test these modifications. Prime amongst these is perhaps the 1999 discovery of interference and the verification of superposition in the fullerene diffraction experiment. This paved the way for the developments that took place in the next two decades. Interference has now been observed in molecules with $7,000 \mathrm{amu}$ and tremendous effort is afoot to push this frontier to a million amu and beyond. Great ingenuity is being invested in devising new experimental techniques and technology which help advance this frontier. These experiments undoubtedly hold a place beside experiments which ushered in quantum theory a century ago: the spectrum of black-body radiation, atomic spectra, pho-
toelectric effect, and matter interferometry with electrons. A broad class of theories predict that new physics will be seen in the range $10^{6}$ amu to $10^{9} \mathrm{amu}$. Perhaps in two decades from now, this range will have been tested. If quantum theory is found to hold good through this regime, then chances are good that linear quantum theory is universally valid on all mass scales: we must then be content with many-worlds / Bohmian mechanics, lest a more convincing interpretation of the standard theory should emerge by then. If confirmation of the predicted modifications is found, this will be nothing short of a revolution; a new theory of dynamics will have been born, to which quantum theory and classical mechanics will be approximations.

## VI. APPENDIX: STOCHASTIC PROCESSES

The best known example of a stochastic process is Brownian motion : random motion of small particles suspended in a liquid, under the influence of a viscous drag, and a fluctuating force resulting from collisions with the molecules of this liquid. The quantitative explanations of Brownian motion by Einstein and Smoluchowski were simplified by Langevin, through his use of what is now called the Langevin equation. This dynami-
cal equation for the randomly evolving position of the suspended particle is equivalent to the Fokker-Planck equation for the time evolution of the probability distribution of the random variable. The Langevin equation can be put on a firm footing as a stochastic differential equation using Itô's differential calculus for stochastic variables. The brief review below introduces stochastic processes, and should be of some assistance in understanding the previous sections of ths review. For a detailed survey the reader is referred to the books by Arnold, 1971; Gardiner, 1983; Gikhman and Skorokhod, 1972; Grimmett and Stirzaker, 2001; Risken, 1996) and the review article by (Chandrasekhar, 1943).

## A. Some Basic Concepts from Proba-

## bility Theory

Let $\Omega$ denote a sample space, i.e. a set of all possible events in an experiment. Then one can define the following structures on $\Omega$ Definition 1. A collection $\mathcal{F}$ of subsets of $\Omega$ is called a $\sigma$ - field if

1. $\phi \in \mathcal{F}$.
2. if $A_{1}, A_{2} \cdots \in \mathcal{F}$ then $\cup_{i=1}^{\infty} A_{i} \in \mathcal{F}$.
3. if $A \in \mathcal{F}$ then $A^{C} \in \mathcal{F}$.

The above properties also imply that closed countable intersections are also included in $\mathcal{F}$.

Examples:
a) $\mathcal{F}=\{\phi, \Omega\}$ is the smallest $\sigma-$ field.
b) If $A \subset \Omega$ then $\mathcal{F}=\left\{\phi, A, A^{c}, \Omega\right\}$ is a $\sigma$-field.

Definition 2. A probability measure $P$ on $(\Omega, \mathcal{F})$ is a function $P: \mathcal{F} \rightarrow[0,1]$ satisfying:

1. $P(\phi)=0, P(\Omega)=1$.
2. if $A_{1}, A_{2} \ldots$ is a collection of disjoint members of $\mathcal{F}$ in that $A_{i} \cap A_{j}=\phi$ for all points $i, j$ satisfying $i \neq j$ then

$$
P\left(\cup_{i=1}^{\infty} A_{i}\right)=\sum_{i=1}^{\infty} P\left(A_{i}\right)
$$

The triple $(\Omega, \mathcal{F}, P)$ is defined to be the probability space.

## B. Random Variables

Definition 3. A random variable is a function $X: \Omega \rightarrow \mathbf{R}$ with the property that $\{\omega \in \Omega: X(\omega) \leq x\} \in \mathcal{F}$ for each $x \in \mathbf{R}$, such functions are called $\mathcal{F}$ - measurable.

## 1. Distribution of random variables

- $X$ is called a discrete random variable if it takes values in some countable subset $\left\{x_{1}, x_{2}, \ldots,\right\}$ only of $\mathbf{R}$. Then $X$ has probability mass function $f: \mathbf{R} \rightarrow$ $[0,1]$ defined by $f(X)=P(X=x)$.
- $X$ is called a continuous random variable if its distribution function $F$ :
$\mathbf{R} \rightarrow[0,1]$ given by $F(x)=P(X \leq x)$ can be expressed as

$$
F(x)=\int_{-\infty}^{x} f(u) d u x \in \mathbf{R}
$$

for some integrable function $f: \mathbf{R} \rightarrow$ $[0, \infty)$, where $f$ is the probability density function.

- Joint distribution Function : A random variable $\mathbf{X}=\left(X_{1}, X_{2}, \ldots X_{n}\right)$ on $(\Omega, \mathcal{F}, P)$ is a function $F_{\mathbf{X}}: \mathbf{R}^{n} \rightarrow[0,1]$ given by $F_{\mathbf{X}}(\mathbf{x})=P(\mathbf{X} \leq \mathbf{x})$ for $\mathbf{x} \in$ $\mathbf{R}^{n}$.


## 2. Time Dependent Random Variables

Let $\xi \equiv \xi(t)$ be a time dependent random variable. Assume an ensemble of systems, such that each system leads to a number $\xi$ which depends on time.

The outcome for one system cannot be precisely predicted, but ensemble averages exist and can be calculated. For fixed $t=t_{1}$ we define the probability density by

$$
\begin{equation*}
W_{1}\left(x_{1}, t\right)=\left\langle\delta\left(x_{1}-\xi\left(t_{1}\right)\right)\right\rangle \tag{265}
\end{equation*}
$$

where the angular brackets denote the ensemble average. The probability to find the random variable $\xi\left(t_{1}\right)$ in the interval $x_{1} \leq$ $\xi\left(t_{1}\right) \leq x_{1}+d x_{1}$ is given by $W_{1}\left(x_{1}, t\right) d x_{1}, \ldots$, in interval $x_{n} \leq \xi\left(t_{n}\right) \leq x_{n}+d x_{n}$ is given
by $W_{n}\left(x_{n}, t_{n} ; \ldots ; x_{1}, t_{1}\right) d x_{n} d x_{(n-1)} \ldots d x_{1}$, where,
$W_{n}\left(x_{n}, t_{n} ; \ldots ; x_{1}, t_{1}\right)=\left\langle\delta\left(x_{1}-\xi\left(t_{1}\right)\right) \ldots \delta\left(x_{n}-\xi\left(t_{n}\right)\right)\right\rangle$
Given $W_{n}$ for all $n$, for every $t_{i}$ in the interval $t_{0} \leq t_{i} \leq t_{0}+T$ the time dependence of the process described by $\xi(t)$ in the interval $\left[t_{0}, t_{0}+T\right]$ can be known completely.

One can obtain probability densities with lower number of variables from those of higher number of variables by integrating as follows. For $i<r$

$$
\begin{align*}
& W_{i}\left(x_{1}, x_{2}, \ldots, x_{i}\right)= \\
& \int \ldots \int W_{r}\left(x_{1}, x_{2}, \ldots, x_{i}, x_{i+1}, \ldots, x_{r}\right) \\
& d x_{i+1} \ldots d x_{r} \tag{266}
\end{align*}
$$

Conditional Probability density is defined as:

$$
P\left(x_{1} \mid x_{2} \ldots x_{r}\right)=\frac{W_{r}\left(x_{1} \ldots x_{r}\right)}{W_{r-1}\left(x_{2}, \ldots x_{r}\right)}
$$

where $P\left(x_{1} \mid x_{2} \ldots x_{r}\right)$ denotes the probability density of $x_{1}$ given the $x_{2}, \ldots, x_{r}$.

## 3. Stationary Processes

If the probability densities do not change by replacing $t_{i}$ by $t_{i}+T$ ( $T$ arbitrary) the process is called a stationary process. In such a case $W_{1}$ does not depend on $t$ and $W_{2}$ can depend on the time difference $t_{2}-t_{1}$.

The Wiener-Khinchin Theorem : According to this theorem, the spectral density
is the Fourier Transform of the correlation function for stationary processes.

Instead of the random variable $\xi(t)$ one may consider its Fourier Transform

$$
\tilde{\xi}(\omega)=\int_{-\infty}^{\infty} \exp (-i \omega t) \xi(t) d t
$$

For a stationary process $\left\langle\xi(t) \xi^{*}\left(t^{\prime}\right)\right\rangle$ is a function only of the difference $t-t^{\prime}$ i.e.

$$
\left\langle\xi(t) \xi^{*}\left(t^{\prime}\right)\right\rangle=\left\langle\xi\left(t-t^{\prime}\right) \xi^{*}(0)\right\rangle
$$

Introducing the new variables

$$
\begin{gathered}
\tau=t-t^{\prime} \\
t_{0}=\left(t+t^{\prime}\right) / 2
\end{gathered}
$$

we have

$$
\begin{aligned}
& \left\langle\tilde{\xi}(\omega) \tilde{\xi}^{*}\left(\omega^{\prime}\right)\right\rangle=\int_{-\infty}^{\infty} \exp \left(-i\left(\omega-\omega^{\prime}\right) t_{0}\right) d t_{0} \\
& \int_{-\infty}^{\infty} \exp \left(-i\left(\omega^{\prime}+\omega\right) \tau / 2\right)\left\langle\xi(\tau) \xi^{*}(0)\right\rangle d \tau(267)
\end{aligned}
$$

which gives

$$
\left\langle\tilde{\xi}(\omega) \tilde{\xi}^{*}\left(\omega^{\prime}\right)\right\rangle=\pi \delta\left(\omega-\omega^{\prime}\right) S(\omega)
$$

where

$$
S(\omega)=2 \int_{-\infty}^{\infty} \exp (i \omega \tau)\left\langle\xi(\tau) \xi^{*}(0)\right\rangle d \tau
$$

is the spectral density.

## C. Stochastic Processes and their Classification

Stochastic processes are systems which evolve probabilistically, i.e. systems in which a time-dependent random variable exists.

The stochastic processes described by a random variable $\xi$ can be classified as follows

Purely Random Process: If $P_{n}(n \geq 2)$ does not depend on the values $x_{i}=\xi\left(t_{i}\right),(i<$ $n)$ at $t_{i}<t_{n}$, then,

$$
P\left(x_{n}, t_{n} \mid x_{n-1}, t_{n-1} ; \ldots ; x_{1}, t_{1}\right)=P\left(x_{n}, t_{n}\right)
$$

thus
$W_{n}\left(x_{n}, t_{n} ; \ldots ; x_{1}, t_{1}\right)=P\left(x_{n}, t_{n}\right) \ldots P\left(x_{1}, t_{1}\right)$
Hence, complete information of the process is contained in $P\left(x_{1}, t_{1}\right)=W_{1}\left(x_{1}, t_{1}\right)$. A purely random process cannot describe physical systems where the random variable is a continuous function of time.

Markov Process: If the conditional probability density depends only on the value $\xi\left(t_{n-1}\right)=x_{n-1}$ at $t_{n-1}$, but not on $\xi\left(t_{n-2}\right)=$ $x_{n-2}$ at $t_{n-2}$ and so on, then such a process is known as a Markov Process. This is given by

$$
\begin{align*}
& P\left(x_{n}, t_{n} \mid x_{n-1}, t_{n-1} ; \ldots ; x_{1}, t_{1}\right)= \\
& P\left(x_{n}, t_{n} \mid x_{n-1}, t_{n-1}\right) \tag{268}
\end{align*}
$$

Then it follows that

$$
\begin{align*}
& W_{n}\left(x_{n}, t_{n} ; \ldots ; x_{1}, t_{1}\right)= \\
& P\left(x_{n}, t_{n} \mid x_{n-1}, t_{n-1}\right) P\left(x_{n-1}, t_{n-1} \mid x_{n-2}, t_{n-2}\right) \\
& \ldots P\left(x_{2}, t_{2} \mid x_{1}, t_{1}\right) W_{1}\left(x_{1}, t_{1}\right) \tag{269}
\end{align*}
$$

For $n=2$

$$
P\left(x_{2}, t_{2} \mid x_{1}, t_{1}\right)=\frac{W_{2}\left(x_{2}, t_{2} ; x_{1} . t_{1}\right)}{W_{1}\left(x_{1}, t_{1}\right)}
$$

For a Markov process the complete information of the process is contained in $W_{2}\left(x_{2}, t_{2} ; x_{1}, t_{1}\right)$.

General processes: There can be other processes such that the complete information is contained in $W_{3}, W_{4}$ etc. However this classification is not suitable to describe nonMarkovian processes. For describing nonMarkovian processes, more random variables $\left(\right.$ other than $\left.\left.\xi(t)=\xi_{1}(t)\right), \xi_{2}(t), \ldots, \xi_{r}(t)\right)$ can be considered. Then by proper choice of these additional variables, one can have a Markov process for $r$ random variables.

## D. Markov Processes

The Markov ssumption is formulated in terms of conditional probabilities as follows :

$$
\begin{align*}
& P\left(x_{1}, t_{1} ; x_{2}, t_{2} ; \ldots \mid y_{1}, \tau_{1} ; y_{2}, \tau_{2} ; \ldots\right) \\
= & P\left(x_{1}, t_{1} ; x_{2}, t_{2} ; \ldots \mid y_{1}, \tau_{1}\right) \tag{270}
\end{align*}
$$

From the defintion of conditional probability

$$
\begin{align*}
& P\left(x_{1}, t_{1} ; x_{2}, t_{2} \mid y_{1}, \tau_{1}\right)= \\
& P\left(x_{1}, t_{1} \mid x_{2}, t_{2} ; y_{1}, \tau_{1}\right) P\left(x_{2}, t_{2} \mid y_{1}, \tau_{1}\right) \tag{271}
\end{align*}
$$

Using Markov property one can write

$$
\begin{align*}
& P\left(x_{1}, t_{1} ; x_{2}, t_{2} ; x_{3}, t_{3} ; \ldots ; x_{n}, t_{n}\right)= \\
& P\left(x_{1}, t_{1} \mid x_{2}, t_{2}\right) P\left(x_{2}, t_{2} \mid x_{3}, t_{3}\right) P\left(x_{3}, t_{3} \mid x_{4}, t_{4}\right) \\
& \ldots P\left(x_{n-1}, t_{n-1} \mid x_{n}, t_{n}\right) P\left(x_{n}, t_{n}\right) \tag{272}
\end{align*}
$$

provided $t_{1} \geq t_{2} \geq t_{3} \geq t_{4} \geq \ldots t_{n-1} \geq t_{n}$.

## 1. The Chapmann Kolmogorov Condition

The following condition holds for all stochastic processes :

$$
\begin{align*}
& P\left(x_{1}, t_{1}\right)=\int d x_{2} P\left(x_{1}, t_{1} ; x_{2}, t_{2}\right) \\
= & \int d x_{2} P\left(x_{1}, t_{1} \mid x_{2}, t_{2}\right) P\left(x_{2}, t_{2}\right) \tag{273}
\end{align*}
$$

Now

$$
\begin{aligned}
P\left(x_{1}, t_{1} \mid x_{3}, t_{3}\right)= & \int d x_{2} P\left(x_{1}, t_{1} ; x_{2}, t_{2} \mid x_{3}, t_{3}\right) \\
& \int d x_{2} P\left(x_{1}, t_{1} \mid x_{2}, t_{2} ; x_{3}, t_{3}\right) \\
& P\left(x_{2}, t_{2} \mid x_{3}, t_{3}\right)
\end{aligned}
$$

Introducing the Markov assumption, if $t_{1} \geq$ $t_{2} \geq t_{3}$, then we can drop dependence on $x_{3}, t_{3}$. Thus,

$$
\begin{align*}
& P\left(x_{1}, t_{1} \mid x_{3}, t_{3}\right)= \\
& \int d x_{2} P\left(x_{1}, t_{1} \mid x_{2}, t_{2}\right) P\left(x_{2}, t_{2} \mid x_{3}, t_{3}\right)(2 \tag{274}
\end{align*}
$$

This is the Chapmann Kolmogorov Equation. The differential form of this equation plays an important role in the description of stochastic processes that follows.

## 2. The Fokker-Planck equation

From the definition of transition probability one can write

$$
\begin{equation*}
W(x, t+\tau)=\int P\left(x, t+\tau \mid x^{\prime}, t\right) W\left(x^{\prime}, t\right) d x^{\prime} \tag{275}
\end{equation*}
$$

This connects $W(x, t+\tau)$ with $W\left(x^{\prime}, t\right)$. To obtain a differential equation for the above,
the following procedure can be carried out. Assume that all moments $M_{n}(x, t, \tau), n \geq 1$ are known

$$
\begin{aligned}
M_{n}\left(x^{\prime}, t, \tau\right) & =<[\xi(t+\tau)-\xi(t)]^{n}>\left.\right|_{\xi(t)=x^{\prime}} \\
& =\int(x-x)^{n} P\left(x, t+\tau \mid x^{\prime}, t\right) d x
\end{aligned}
$$

Introducing $\Delta=x-x^{\prime}$ in equation (275), the integrand can be expanded in a Taylor series, which after integration over $\delta$ can be put in the following form:
$W(x, t+\tau)-W(x, t)=\frac{\partial W(x, t)}{\partial t} \tau+O\left(\tau^{2}\right)=$ $\sum_{n=1}^{\infty}\left(-\frac{\partial}{\partial x}\right)^{n}\left(\frac{M_{n}(x, t, \tau)}{n!}\right) W(x, t)$
$M_{n}$ can be expanded into Taylor series with respect to $\tau$ (Risken, 1996) $(n \geq 1)$

$$
M_{n}(x, t, \tau) / n!=D^{(n)}(x, t) \tau+O\left(\tau^{2}\right)
$$

Considering only linear terms in $\tau$, Eqn. (276) can be written as

$$
\begin{align*}
\frac{\partial W(x, t)}{\partial t} & =\sum_{n=1}^{\infty}\left(-\frac{\partial}{\partial x}\right)^{n} D^{(n)}(x, t) W(x, t) \\
& =L_{K M} W(x, t) \\
L_{K M} & =\sum_{n=1}^{\infty}\left(-\frac{\partial}{\partial x}\right)^{n} D^{(n)}(x, t) \tag{277}
\end{align*}
$$

The above is the Kramers-Moyal Expansion. Pawula Theorem states that for a positive transition probability $P$ the Kramers-Moyal expansion stops after the second term; if not then it must contain an infinite number of terms.

If the K-M expansion stops after the second term, then it is called the Fokker -Planck equation, given by

$$
\begin{equation*}
\frac{\partial W(x, t)}{\partial t}=-\frac{\partial}{\partial x} D^{(1)}(x, t)+\frac{\partial^{2}}{\partial x^{2}} D^{(2)}(x, t) \tag{278}
\end{equation*}
$$

$D^{(1)}$ is called the drift coefficient, and $D^{(2)}$ is called the diffusion coefficient. The transition probability $P\left(x, t \mid x^{\prime}, t^{\prime}\right)$ is the distribution $W(x, t)$ for the initial condition $W\left(x, t^{\prime}\right)=$ $\delta\left(x-x^{\prime}\right)$. Therefore the transition probability must also satistfy 277). Hence,

$$
\begin{equation*}
\frac{\partial P\left(x, t \mid x^{\prime}, t^{\prime}\right)}{\partial t}=L_{K M}(x, t) P\left(x, t \mid x^{\prime}, t^{\prime}\right) \tag{279}
\end{equation*}
$$

where the initial condition is given by $P\left(x, t \mid x^{\prime}, t\right)=\delta\left(x-x^{\prime}\right)$.

The Fokker-Planck equation can also be written as

$$
\begin{equation*}
\frac{\partial W}{\partial t}+\frac{\partial S}{\partial x}=0 \tag{280}
\end{equation*}
$$

where

$$
S(x, t)=\left[D^{(1)}(x, t)-\frac{\partial}{\partial x} D^{(2)}(x, t)\right] W(x, t)
$$

Here $S$ can be interpreted as a probability current. We will discuss the one variable Fokker-Planck equation with timeindependent drift and diffusion coefficients given by

$$
\begin{equation*}
\frac{\partial W(x, t)}{\partial t}=-\frac{\partial}{\partial x} D^{(1)}(x)+\frac{\partial^{2}}{\partial x^{2}} D^{(2)}(x) \tag{281}
\end{equation*}
$$

For stationary processes the probability current $S=$ constant. Methods of solution
of such a F-P equation for stationary processes are discussed in (Risken, 1996). Nonstationary solutions of the F-P equation are in general difficult to obtain. A general expression for non-stationary solution can be found only for special drift and diffusion coefficients.

The Fokker-Planck equation can be taken as a starting point for introducing the concept of a stochastic differential equation. If the random variable $\xi(t)$ satisfies the initial condition

$$
\begin{equation*}
W(\xi(t), y)=\delta(\xi-y) \tag{282}
\end{equation*}
$$

that is, it is sharply peaked at the value $y$, it can be shown by solving the Fokker-Planck equation that a short time $\Delta t$ later, the solution is still sharply peaked, and is a Gaussian with mean $y+D^{(1)} \Delta t$ and variance $D^{(2)}$. The picture is that of a system moving with a systematic drift velocity $D^{(1)}$, and on this motion is superimposed a Gaussian fluctuation with variance $D^{(2)}$. Thus,

$$
\begin{equation*}
y(t+\Delta t)=y(t)+D^{(1)} \Delta t+\eta(t) \Delta t^{1 / 2} \tag{283}
\end{equation*}
$$

where $\langle\eta\rangle=0$ and $\left\langle\eta^{2}\right\rangle=D^{(2)}$. This picture gives sample paths which are always continuous but nowhere differentiable. As we will see shortly, this heuristic picture can be made much more precise and leads to the concept of the stochastic differential equation.

## 3. Wiener Process

A process which is described by Eqn. (281) with $D^{(1)}=0$ and $D^{(2)}(x)=D=$ constant, is called a Wiener process. Then the equation for transition probability $P=$ $P\left(x, t \mid x^{\prime}, t^{\prime}\right)$ is the diffusion equation, given by

$$
\begin{equation*}
\frac{\partial P}{\partial t}=D \frac{\partial^{2} P}{\partial x^{2}} \tag{284}
\end{equation*}
$$

with the initial condition $P\left(x, t^{\prime} \mid x^{\prime}, t^{\prime}\right)=$ $\delta\left(x-x^{\prime}\right)$. Then the solution for $t>t^{\prime}$ is given by the gaussian distribution

$$
\begin{align*}
& P\left(x, t \mid x^{\prime}, t^{\prime}\right)= \\
& \frac{1}{\sqrt{4 \pi D\left(t-t^{\prime}\right)}} \exp \left(-\frac{\left(x-x^{\prime}\right)^{2}}{4 D\left(t-t^{\prime}\right)}\right) \tag{285}
\end{align*}
$$

Thus the general solution for probaility density with initial distribution $W\left(x^{\prime}, t^{\prime}\right)$ is given by

$$
W(x, t)=\int P\left(x, t \mid x^{\prime}, t^{\prime}\right) W\left(x^{\prime}, t^{\prime}\right) d x^{\prime}
$$

An initially sharp distribution spreads in time. The one-variable Wiener process is often simply called Brownian motion, since it obeys the same differential equation of motion as Brownian motion.

## 4. Ornstein-Uhlenbeck Process

This process is described by Eqn. (281) when the drift coefficient is linear and diffusion coefficient is constant, i.e.

$$
D^{(1)}(x)=-\gamma x ; D^{(2)}(x)=D=\text { const }
$$

The Fokker-Planck equation then can be written as

$$
\begin{equation*}
\frac{\partial P}{\partial t}=\gamma \frac{\partial}{\partial x}(x P)+D \frac{\partial^{2}}{\partial x^{2}} P \tag{286}
\end{equation*}
$$

with initial condition $P\left(x, t^{\prime} \mid x^{\prime}, t^{\prime}\right)=\delta(x-$ $\left.x^{\prime}\right)$. The above equation can be solved by taking a Fourier transform w.r.t. $x$ i.e.

$$
P\left(x, t \mid x^{\prime}, t^{\prime}\right)=(2 \pi)^{-1} \int e^{i k x} \tilde{P}\left(k, t \mid x^{\prime}, t^{\prime}\right) d k
$$

Thus it results in the following equation

$$
\frac{\partial \tilde{P}}{\partial t}=-\gamma k \frac{\partial \tilde{P}}{\partial k}-D k^{2} \tilde{P}
$$

with initial condition $\tilde{P}\left(k, t^{\prime} \mid x^{\prime} t^{\prime}\right)=e^{i k x^{\prime}}$ for $t>t^{\prime}$. Then one gets
$\tilde{P}\left(k, t \mid x^{\prime}, t^{\prime}\right)=$
$\exp \left[-i k x^{\prime} e^{-\gamma\left(t-t^{\prime}\right)}-D k^{2}\left(1-e^{-2 \gamma\left(t-t^{\prime}\right)}\right) /(2 \gamma)\right]$

By applying the inverse Fourier transform one finally obtains the solution of the Fokker Planck equation describing the Ornstein Uhlenbeck process

$$
\begin{align*}
& P\left(x, t \mid x^{\prime}, t^{\prime}\right)=\sqrt{\frac{\gamma}{2 \pi D\left(1-e^{-2 \gamma\left(t-t^{\prime}\right)}\right)}} \times \\
& \exp \left[-\frac{\gamma\left(x-e^{-\gamma\left(t-t^{\prime}\right)} x^{\prime}\right)^{2}}{2 D\left(1-e^{-2 \gamma\left(t-t^{\prime}\right)}\right)}\right] \tag{288}
\end{align*}
$$

In the limit $\gamma \rightarrow 0$ we get the Gaussian distribution for a Wiener Process.

Eqn. (288) is valid for positive and negative values of $\gamma$. For positive $\gamma$ and large time
difference $\gamma\left(t-t^{\prime}\right) \gg 1$ the equation passes over to the stationary distribution given by

$$
\begin{equation*}
W_{s t}=\sqrt{\frac{\gamma}{2 \pi D}} \exp \left[-\frac{\gamma x^{2}}{2 D}\right] \tag{289}
\end{equation*}
$$

For $\gamma \leq 0$ no stationary solution exists.

## E. Langevin Equation

We introduce stochastic integration via the Langevin equation. In the presence of a viscous drag linearly proportional to velocity, the equation of motion for a particle of mass ' $m$ ' is given by

$$
\begin{equation*}
m \dot{v}+\alpha v=0 \tag{290}
\end{equation*}
$$

or

$$
\dot{v}+\gamma v=0
$$

where $\gamma=\alpha / m=1 / \tau, \tau$ being the relaxation time. The solution of the above equation is given by

$$
v(t)=v(0) e^{-t / \tau}=v(0) e^{-\gamma t}
$$

If the mass of the particle is small, so that the velocity due to thermal fluctuations is not negligible, then

$$
v_{t h}=\sqrt{\left\langle v^{2}\right\rangle}=\sqrt{k T / m}
$$

is observable and hence the velocity of the small paricle cannot be described by Eqn. (290). Thus this equation has to be modified as follows:

$$
\begin{equation*}
\dot{v}+\gamma v=\Gamma(t) \tag{291}
\end{equation*}
$$

where $\Gamma(t)=F_{f}(t) / m$ is the stochastic term, $F_{f}(t)$ is the fluctuating force acting on the particle. This is the Langevin equation.

## 1. Brownian Motion

The Langevin equation for Brownian motion is given by Eqn. (291) where $\Gamma$ describes the Langevin force with

$$
\langle\Gamma(t)\rangle=0, \quad\left\langle\Gamma(t) \Gamma\left(t^{\prime}\right)\right\rangle=q \delta\left(t-t^{\prime}\right)
$$

which after solving for the double integral gives

$$
\begin{align*}
& \left\langle v\left(t_{1}\right) v\left(t_{2}\right)\right\rangle=v_{0}^{2} e^{-\gamma\left(t_{1}+t_{2}\right)}+ \\
& \frac{q}{2 \gamma}\left(e^{-\gamma\left|t_{1}-t_{2}\right|}-e^{-\gamma\left(t_{1}+t_{2}\right)}\right) . \tag{294}
\end{align*}
$$

For large $t_{1}$ and $t_{2}$, i.e $\gamma t_{1}, \gamma t_{2} \gg 1$ the correlation is independent of $v_{0}$ and is a function of the time difference.

$$
\left\langle v\left(t_{1}\right) v\left(t_{2}\right)\right\rangle=\frac{q}{2 \gamma} e^{-\gamma\left|t_{1}-t_{2}\right|}
$$

Now in the stationary state

$$
\langle E\rangle=\frac{1}{2} m\left\langle v(t)^{2}\right\rangle=\frac{1}{2} m \frac{q}{2 \gamma}
$$

for a Brownian particle.
According to the law of equipartition

$$
\langle E\rangle=\frac{1}{2} k T
$$

and comparing with the earlier expression we get

$$
q=2 \gamma k T / m
$$

3. Overdamped Langevin equation

The overdamped Langevin Equation looks like

$$
v(t)=\frac{1}{\gamma} \Gamma(t)
$$

where the acceleration term is dropped because of the prominence of damping. In the large time limit, the two point correlation for velocity is given by

$$
\left\langle v\left(t_{1}\right) v\left(t_{2}\right)\right\rangle \approx \frac{1}{\gamma^{2}}\left\langle\Gamma\left(t_{1}\right) \Gamma\left(t_{2}\right)\right\rangle=\frac{q}{\gamma^{2}} \delta\left(t_{1}-t_{2}\right)
$$

## 4. Non-linear Langevin Equation

A nonlinear Langevin equation has the following form

$$
\begin{equation*}
\dot{\xi}=h(\xi, t)+g(\xi, t) \Gamma(t) \tag{295}
\end{equation*}
$$

Here $\Gamma(t)$ is assumed to be Gaussian random variable with zero mean and $\delta$ correlation function.

$$
\langle\Gamma(t)\rangle=0 ; \quad\left\langle\Gamma(t) \Gamma\left(t^{\prime}\right)\right\rangle=2 \delta\left(t-t^{\prime}\right)
$$

Integrating Eqn. (295)

$$
\begin{equation*}
\xi=\int_{0}^{t} h(\xi, t) d t+\int_{0}^{t} g(\xi, t) \Gamma(t) d t \tag{296}
\end{equation*}
$$

Here ' $g$ ' could be a constant or can depend on $\xi$. Constant $g$ gives additive noise, while if it depends on $\xi$ it is referred to as multiplicative noise. For $\xi$ dependent ' $g$ ' in the above equation, since $\Gamma(t)$ has no correlation time, it is not clear which value of $\xi$ one has to use in $g$ while evaluating the integral. Physicists use
an approximation of the $\delta$ function to ensure that one gets appropriate results. But from a purely mathematical point of view one cannot answer this question, unless some additional specification is given. Thus, this gives rise to the requirement of using the stochastic integrals, namely the Itô and Stratonovich integrals for solving such equations. In the following section, we give a brief introduction to these Stochastic Integrals. Prior to this we state an example of a particular form of noise and solve the nonlinear Langevin equation for this particular case.

Example For $g=a \xi$ where $a$ is a constant

$$
\dot{\xi}=a \xi \Gamma(t)
$$

The formal solution of the above equation is given by

$$
\xi(t)=x \exp \left[a \int_{0}^{t} \Gamma\left(t^{\prime}\right) d t^{\prime}\right]
$$

Assuming $\xi(0)=x$

$$
\begin{align*}
& \langle\xi(t)\rangle=\left\langle x \exp \left[a \int_{0}^{t} \Gamma\left(t^{\prime}\right) d t^{\prime}\right]\right\rangle \\
= & x\left[1+a \int_{0}^{t}\left\langle\Gamma\left(t_{1}\right)\right\rangle d t_{1}+\right. \\
& \left.\frac{1}{2!} a^{2} \int_{0}^{t} \int_{0}^{t}\left\langle\Gamma\left(t_{1}\right) \Gamma\left(t_{2}\right)\right\rangle d t_{1} d t_{2}+\ldots\right] \tag{297}
\end{align*}
$$

Since $\Gamma$ is delta correlated Gaussian white noise, all the higher correlations can be expressed in terms of two point correlations, therefore the above equation can be written
in the following form

$$
\begin{align*}
& =\frac{(2 n)!}{2^{n} n!}\left[\int_{0}^{t} \int_{0}^{t} \phi\left(t_{1}-t_{2}\right) d t_{1} d t_{2}\right]^{n}  \tag{298}\\
& =\exp \left[\frac{1}{2} a^{2} \int_{0}^{t} \int_{0}^{t}\left\langle\Gamma\left(t_{1}\right) \Gamma\left(t_{2}\right)\right\rangle d t_{1} d t_{2}\right] \tag{299}
\end{align*}
$$

For delta correlated Langevin force the double integral gives

$$
\langle\xi(t)\rangle=x \exp \left(a^{2} t\right)
$$

and

$$
\left.\frac{d}{d t}\langle\xi(t)\rangle\right|_{t=0}=a^{2} x
$$

is called spurious or noise induced drift.

## F. Stochastic integration, Itô calculus

 and stochastic differential equationsConsider a Langevin equation of the form

$$
\begin{equation*}
\frac{d x}{d t}=a(x, t)+b(x, t) \zeta(t) \tag{300}
\end{equation*}
$$

for a time-dependent variable $x$, where $a(x, t)$ and $b(x, t)$ are known functions, and $\zeta(t)$ is the rapidly fluctuating random term which induces stochasticity in the evolution. We want to examine the mathematical status of this equation as a differential equation. Since we expect it to be integrable, the integral

$$
\begin{equation*}
u(t)=\int_{0}^{t} d t^{\prime} \zeta\left(t^{\prime}\right) \tag{301}
\end{equation*}
$$

should exist. If we demand that $u$ is a continuous function of $t$, then it is a Markov process whose evolution can be described by a Fokker-Planck equation, which can be shown to have zero drift, and diffusion unity. Hence, $u(t)$ is a Wiener process, denoted say by $W(t)$; but we know that $W(t)$ is not differentiable. This would imply that in a mathematical sense the Langevin equation does not exist! However, the corresponding integral equation
$x(t)-x(0)=\int_{0}^{t} a[x(s), s] d s+\int_{0}^{t} b[x(s), s] \zeta(s) d s$
can be interpreted consistently. We make the replacement

$$
\begin{equation*}
d W(t) \equiv W(t+d t)-W(t)=\zeta(t) d t \tag{303}
\end{equation*}
$$

so that the second integral can be written as

$$
\begin{equation*}
\int_{0}^{t} b[x(s), s] d W(s) \tag{304}
\end{equation*}
$$

which is a stochastic Riemann-Stieltjes integral, which we now define.

Given that $G(t)$ is an arbitrary function of time and $W(t)$ is a stochastic process, the stochastic integral $\int_{0}^{t} G\left(t^{\prime}\right) d W\left(t^{\prime}\right)$ is defined by dividing the interval $(0, t)$ into $n$ subintervals $\left(t_{i-1}, t_{i}\right)$ such that

$$
\begin{equation*}
t_{0} \leq t_{1} \leq t_{2} \leq \cdots \leq t \tag{305}
\end{equation*}
$$

and defining intermediate points $\tau_{i}$ such that $t_{i-1} \leq \tau_{i} \leq t_{i}$. The stochastic integral is
defined as a limit of partial sums

$$
\begin{equation*}
S_{n}=\sum_{i=1}^{n} G\left(\tau_{i}\right)\left[W\left(t_{i}\right)-W\left(t_{i-1}\right)\right] . \tag{306}
\end{equation*}
$$

The challenge is that, $G(t)$ being function of a random variable, the limit of $S_{n}$ depends on the partcular choice of intermediate point $\tau_{i}$ ! Different choices of the intermediate point give different results for the integral.

Denoting $\Delta=\max \left(t_{i}-t_{i-1}\right)$, the It $\hat{o}$ stochastic integral is defined by taking $\tau_{i}=$ $t_{i-1}$ and taking the limit of the sum :

$$
\begin{align*}
& \int_{0}^{\tau} G\left(t^{\prime}\right) d W\left(t^{\prime}\right)= \\
& \lim _{\Delta \rightarrow 0} \sum_{i=1}^{n} G\left(t_{i-1}\right)\left[W\left(t_{i}\right)-W\left(t_{i-1}\right)\right] \tag{307}
\end{align*}
$$

As an example, it can be shown that

$$
\begin{align*}
& \int_{0}^{t} W\left(t^{\prime}\right) d W\left(t^{\prime}\right)= \\
& \frac{1}{2}\left[W(t)^{2}-W\left(t_{0}\right)^{2}-\left(t-t_{0}\right)\right] \tag{308}
\end{align*}
$$

Note that the result for the integration is no longer the same as the ordinary RiemannStieltjes integral, where the term $\left(t-t_{0}\right)$ would be absent; the reason being that the difference $W(t+\Delta t)-W(t)$ is almost always of the order $\sqrt{t}$, which implies that, unlike in ordinary integration, terms of second order in $\Delta W(t)$ do not vanish on taking the limit.

An alternative definition of the stochastic integral is the Stratanovich integral, denoted by $S$, and is such that the anomalous term above, $\left(t-t_{0}\right)$, does not occur. This happens if the intermediate point $\tau_{i}$ is taken as the
mid-point $\left.\tau_{i}=\left(t_{i}+t_{i+1}\right)\right) / 2$, and it can then be shown that

$$
\begin{equation*}
S \int_{0}^{t} W\left(t^{\prime}\right) d W\left(t^{\prime}\right)=\frac{1}{2}\left[W(t)^{2}-W\left(t_{0}\right)^{2}\right] . \tag{309}
\end{equation*}
$$

For arbitrary functions $G(t)$ there is no connection between the Itô integral and the Stratanovich integral. However, in cases where we can specify that $G(t)$ is related to some stochastic differential equation, a formula can be given relating the two differential equations.

## 1. Rules of Itô calculus

Mean square limit or the limit in the mean is defined as follows: Let $X_{n}(\omega)$ be a sequence of random variables $X_{n}$ on the probability space $\Omega$, where $\omega$ are the elements of the space which have probability density $p(\omega)$. Thus one can say that $X_{n}$ converges to $X$ in the mean square if

$$
\begin{align*}
& \lim _{n \rightarrow \infty} \int d \omega p(\omega)\left[X_{n}(\omega)-X(\omega)\right]^{2} \equiv \\
& \lim \left\langle\left(X_{n}-X\right)^{2}\right\rangle=0 \tag{310}
\end{align*}
$$

This is written as

$$
m s-\lim _{n \rightarrow \infty} X_{n}=X
$$

## Rules

1. $d W(t)^{2}=d t$
2. $d W^{2+N}(t)=0$

The above formulae mean the following

$$
\begin{align*}
& \int_{t_{0}}^{t}\left[d W\left(t^{\prime}\right)\right]^{2+N} G\left(t^{\prime}\right) \equiv \\
& m s-\lim n \rightarrow \infty \sum_{i} G_{i-1} \Delta W_{i}^{2+N} \\
= & \int_{t_{0}}^{t} d t^{\prime} G\left(t^{\prime}\right) \text { for } N=0 \\
= & 0 \text { for } N>0 \tag{311}
\end{align*}
$$

for an arbitrary non-anticipating function $G$.

## Proof:

For $N=0$ consider the following sum

$$
\begin{align*}
& \left.I=\lim _{n \rightarrow \infty}\left\langle\left[\sum_{i} G_{i-1}\left(\Delta W_{i}^{2}-\Delta t_{i}\right)\right]^{2}\right]\right\rangle  \tag{312}\\
& =\lim _{n \rightarrow \infty}\left\langle\sum_{i}\left(G_{i-1}\right)^{2}\left(\Delta W_{i}^{2}-\Delta t_{i}\right)^{2}+\right. \\
& \left.\sum_{i>j} 2 G_{i-1} G_{j-1}\left(\Delta W_{j}^{2}-\Delta t_{j}\right)\left(\Delta W_{i}^{2}-\Delta t_{i}\right)\right\rangle \tag{313}
\end{align*}
$$

Using the following results mentioned earlier

$$
\begin{gathered}
\left\langle\Delta W_{i}^{2}\right\rangle=\Delta t_{i} \\
\left(\Delta W_{i}^{2}-\Delta t_{i}\right)^{2}>=2 \Delta t_{i}^{2}
\end{gathered}
$$

one gets

$$
I=2 \lim _{n \rightarrow \infty}\left[\sum_{i} \Delta t_{i}^{2}\left\langle\left(G_{i-1}\right)^{2}\right\rangle\right]
$$

This can be written as
$m s-\lim _{n \rightarrow \infty}\left(\sum_{i} G_{i-1} \Delta W_{i}^{2}-\sum_{i} G_{i-1} \Delta t_{i}\right)=0$ since

$$
m s-\lim _{n \rightarrow \infty} \sum_{i} G_{i-1} \Delta t_{i}=\int_{t_{0}}^{t} d t^{\prime} G\left(t^{\prime}\right)
$$

we get

$$
\int_{t_{0}}^{t}\left[d W\left(t^{\prime}\right)\right]^{2} G\left(t^{\prime}\right)=\int_{t_{0}}^{t} d t^{\prime} G\left(t^{\prime}\right)
$$

from this we see that $d W\left(t^{\prime}\right)^{2}=d t^{\prime}$. Similarly one can show that $d W(t)^{2+N} \equiv 0(N>0)$.

Another important result that can be proved by the above method is

$$
\begin{align*}
& \int_{t_{0}}^{t} G\left(t^{\prime}\right) d t^{\prime} d W\left(t^{\prime}\right) \equiv \\
& m s-\lim _{n \rightarrow \infty} \sum G_{i-1} \Delta W_{i} \Delta t_{i}=0 \tag{314}
\end{align*}
$$

Rule for integration of polynomials :

$$
\begin{align*}
& \int_{t_{0}}^{t} W\left(t^{\prime}\right)^{n} d W\left(t^{\prime}\right) \\
= & \frac{1}{n+1}\left[W(t)^{(n+1)}-W\left(t_{0}\right)^{(n+1)}\right] \\
& \left.-\frac{n}{2} \int t_{0}^{t} W(t)^{( } n-1\right) d t \tag{315}
\end{align*}
$$

General rule for differentiation ;

$$
\begin{align*}
& d f[W(t), t]=\left(\frac{\partial f}{\partial t}+\frac{1}{2} \frac{\partial^{2} f}{\partial W^{2}}\right) d t+ \\
& \frac{\partial f}{\partial W} d W(t) . \tag{316}
\end{align*}
$$

## 2. Stochastic differential equations

The Itô integral is mathematically the most satisfactory, but not always the most natural physical choice. The Stratanovich integral is the natural choice for an interpretation where $\zeta(t)$ is a colored (not white) noise. Also, unlike in the Itô interpretation, the Stratanovich interpretation enables the use of ordinary calculus. From the mathematical point of view, it is more convenient to define the Itô SDE, develop its equvalence with the Stratanovich SDE, and use either form depending on circumstances.

A stochastic quantity $x(t)$ obeys an Itô differential equation

$$
\begin{equation*}
d x(t)=a[x(t), t] d t+b[x(t), t] d W(t) \tag{317}
\end{equation*}
$$

if for all $t$ and $t_{0}$

$$
\begin{align*}
& x(t)=x\left(t_{0}\right)+\int_{t_{0}}^{t} d t^{\prime} a\left[x\left(t^{\prime}\right), t^{\prime}\right]+ \\
& \int_{t_{0}}^{t} d W\left(t^{\prime}\right) b\left[x\left(t^{\prime}\right), t^{\prime}\right] . \tag{318}
\end{align*}
$$

If $f[x(t)]$ is an arbitrary function of $x(t)$ then Itô's formula gives the differential equation satisfied by f :

$$
\begin{align*}
& d f[x(t)]= \\
& \left(a[x(t), t] f^{\prime}[x(t)]+\frac{1}{2} b[x(t), t]^{2} f^{\prime \prime}[x(t)]\right) d t \\
& +b[x(t), t] f^{\prime}[x(t)] d W(t) . \tag{319}
\end{align*}
$$

Thus change of variables is not given by ordinary calculus unless $f[x(t)]$ is linear in $x(t)$.

Given the time development of an arbitrary $f[x(t)]$, the conditional probability density $p\left(x, t \mid x_{0}, t_{0}\right)$ for $x(t)$ can be shown to satisfy a Fokker-Planck equation with drift coefficient $a(x, t)$ and diffusion coefficient $b(x, t)^{2}$.

The stochastic differential equation studied in detail in the QMUPL model

$$
\begin{align*}
d \psi_{t}= & {\left[-\frac{i}{\hbar} H d t+\sqrt{\lambda}\left(q-\langle q\rangle_{t}\right) d W_{t}\right.} \\
& \left.-\frac{\lambda}{2}\left(q-\langle q\rangle_{t}\right)^{2} d t\right] \psi_{t} \tag{320}
\end{align*}
$$

is an Itô differential equation of the type (317). If we formally treat this as an equation for $\ln \psi$ then upon comparison we see
that $\ln \psi$ is $x$, and

$$
\begin{align*}
& a(x(t), t]=-\frac{i}{\hbar} H-\frac{\lambda}{2}\left(q-\langle q\rangle_{t}\right)^{2} \\
& b=\sqrt{\lambda}(q-\langle q\rangle)_{t} \tag{321}
\end{align*}
$$

The real part of the drift coefficient $a$ is related to the diffusion coefficient $b^{2}$ by

$$
\begin{equation*}
-2 a=b^{2} \tag{322}
\end{equation*}
$$

This non-trivial relation between diffusion and drift is what gives the equation its normpreserving martingale property which eventually leads to the Born rule. Why the drift and diffusion must be related this way is at present not understood in Trace Dynamics and there probably is some deep underlying reason for this relation.

Stratonovich's stochastic differential equation : Given the Itô differential equation (317) its solution $x(t)$ can also be expressed in terms of a Stratonovich integral

$$
\begin{align*}
& x(t)=x_{0}+\int_{t_{0}}^{t} d t^{\prime} \alpha\left[x\left(t^{\prime}\right), t^{\prime}\right]+ \\
& S \int_{t_{0}}^{t} d W\left(t^{\prime}\right) \beta\left[\left(x\left(t^{\prime}\right), t^{\prime}\right]\right. \tag{323}
\end{align*}
$$

where
$\alpha(x, t)=a(x, t)-\frac{1}{2} b(x, t) \partial_{x} b(x, t) ; \beta(x, t)=b(x, t)$

In other words, the Itô SDE (317) is the same as the Stratonovich SDE

$$
\begin{equation*}
d x=\left[a(x, t)-\frac{1}{2} b(x, t) \partial_{x} b(x, t)\right] d t+b d W(t) \tag{325}
\end{equation*}
$$

or conversely, the Stratonovich SDE

$$
\begin{equation*}
d x=\alpha d t+\beta d W(t) \tag{326}
\end{equation*}
$$

is the same as the Itô SDE
$d x=\left[\alpha(x, t)+\frac{1}{2} \beta(x, t) \partial_{x} \beta(x, t)\right] d t+\beta d W(t)$

It can be shown that in a Stratonovich SDE the rule for change of variable is the same as in ordinary calculus.

## G. Martingales

Martingales play a role in stochastic processes roughly similar to that played by conserved quantities in dynamical systems. Unlike a conserved quantity in dynamics, which remains constant in time, a martingale's value can change ; however, its expectation remains constant in time.

A martingale is defined as follows:
A discrete time martingale is a discrete time stochastic process, $X_{1}, X-2, \ldots$, that satisfies for any time $n$,

$$
\begin{aligned}
E\left(\left|X_{n}\right|\right) & <\infty \\
E\left(X_{n+1} \mid X_{1}, \ldots, X_{n}\right) & =X_{n}
\end{aligned}
$$

where $E(X)$ denotes the expectation of $X$.
Martingale Sequence with Respect to Another Sequence A sequence $Y_{1}, Y_{2}, \ldots$ is said to be a martingale w.r.t.
another sequence $X_{1}, X_{2}, X_{3} \ldots$ if for all $n$

$$
\begin{aligned}
E\left(\left|Y_{n}\right|\right) & <\infty \\
E\left(Y_{n+1} \mid X_{1}, \ldots, X_{n}\right) & =0
\end{aligned}
$$

In other words, a martingale is a model of a fair game, where no knowledge of past events can help to predict future winnings. It is a sequence of random variables for which at a particular time in a realized sequence, the expectation of the next value in the sequence is equal to the present observed value even given knowledge of all prior observed value at current time.

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[^1]:    T 125

[^2]:    ${ }^{1}$ The superscripts " R " and " I " denote, respectively, the real and imaginary parts of the corresponding quantities.

[^3]:    ${ }^{3}$ With an Hermitian coupling, one would have a standard quantum Hamiltonian with a random potential; the equation would be linear and no suppression of quantum superpositions would occur.

[^4]:    ${ }^{4} 6$ photons corresponds to the threshold of vision.

