Macroscopic Quantum Phenomena from the Correlation, Coupling and Criticality Perspectives

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Abstract.

In this sequel paper we explore how macroscopic quantum phenomena can be measured or understood from the behavior of quantum correlations which exist in a quantum system of many particles or components and how the interaction strengths change with energy or scale, under ordinary situations and when the system is near its critical point. We use the nPI (master) effective action related to the Boltzmann-BBGKY / Schwinger-Dyson hierarchy of equations as a tool for systemizing the contributions of higher order correlation functions to the dynamics of lower order correlation functions. Together with the large \( N \) expansion discussed in our first paper [1] we explore 1) the conditions whereby an H-theorem is obtained, which can be viewed as a signifier of the emergence of macroscopic behavior in the system. We give two more examples from past work: 2) the nonequilibrium dynamics of \( N \) atoms in an optical lattice under the large \( N \) (field components), \( 2PI \) and second order perturbative expansions, illustrating how \( N \) and \( \lambda \) enter in these three aspects of quantum correlations, coherence and coupling strength. 3) the behavior of an interacting quantum system near its critical point, the effects of quantum and thermal fluctuations and the conditions under which the system manifests infrared dimensional reduction. We also discuss how the effective field theory concept bears on macroscopic quantum phenomena: the running of the coupling parameters with energy or scale imparts a dynamical-dependent and an interaction-sensitive definition of ‘macroscopia’.

1. Key points: correlation, interaction strength, effective scales and dimensions
We continue our investigations begun in an earlier paper [1] (MQP1) into the key features of macroscopic quantum phenomena (MQP), seeking to identify and assemble the necessary ingredients toward the construction of a viable theoretical framework. Not only does this new field bear on the foundational issues of both quantum and statistical mechanics it also sees a widening range of manifestations and applications. (For background please refer to references in [1]). We want to examine the conditions or criteria whereby a macroscopic quantum system may take on classical attributes, and, more interestingly, that it keeps some of its quantum features. Our first paper focused on the large \( N \) approximation, highlighting the subtle yet important difference between results obtained from a leading-order large-\( N \) approximation, namely, the mean field theory which is quantum in nature, from a corresponding classical theory. Here we focus on the key features of correlations and couplings in a quantum system of many particles or components, especially its behavior near the critical point, to investigate which factors contribute
to, or determine a system as being macroscopic, and under what conditions would quantum features persist or dominate in that limit.

Before delving into these main subjects, it may be useful for the purpose of orientation to refresh what is already known in quantum statistical mechanics of a) how the thermodynamic properties of a quantum system depend on \( N \), the number of particles present, and b) how qualitative changes arise in a many-body quantum system, such as the formation of a Bose-Einstein condensate (BEC), one of the most commonly cited examples (the other being quantized fluxes in Josephson junctions superconductivity) of MQP. We want to point out the qualitative differences in the meanings of ‘quantumness’ in these familiar cases before we launch our investigations into the other quantum features we are more interested in, namely, quantum correlations, fluctuations, coherence and entanglement, and how they may show up or even persist at a macroscopic scale.

1.1. \( N \) dependence in quantum statistical mechanics

Note from the start that one of the two basic premises of quantum statistical mechanics (QSM) [2], namely, the assumption of random phase approximation, precludes quantum coherence considerations. All descriptions are in terms of probabilities, not amplitudes. The other basic premise, that the system has equal a priori probability to be found in any of its accessible states, refers to the equilibrium condition. Note, however, that decoherence and quantum to classical transition [3, 4] are fundamentally nonequilibrium processes.

The ‘quantum’ in QSM refers to effects due to spin-statistics, in the difference between bosons and fermions, and distinguishability: different combinatorics in distinguishable (classical feature) versus identical (quantum feature) particles, which underlie the Gibbs paradox [5]. Thus the answer to a) above is simple: the dependence of the partition function \( Z \) on \( N \) is explicit, and the thermodynamic limit is well known. With the correct counting of distinguishable versus indistinguishable particles included in the partition function, the difference between quantum and classical features in this restricted sense is also well accounted for.

QSM describes finite temperature equilibrium processes for a quantum (in the above restricted sense) system in or near equilibrium, such as in the ordinary (thermal) critical phenomena mediated by energy fluctuations in a canonical ensemble (temperature), or numbers / species fluctuations in a grand canonical ensemble (chemical potential) description, but cannot address issues related to quantum coherence and entanglement, the central issues in quantum information. To the extent entanglement is a useful marker for quantum phase transition QSM cannot address issues of quantum noise or fluctuation phenomena.

BEC is the well-known process where one can pinpoint the number \( N_c \) of a system of \( N \) bosonic atoms undergoing a transition where the salient features of the system undergo marked qualitative changes on the large scale, exemplifying MQP. There, the critical number \( N_c \) and temperature \( T_c \) which define the phase transition also depend on the dimensionality. (E.g., a two-dimensional Bose gas undergoes transition into vortex-antivortex pairs – see e.g., [6]). BEC is certainly a quantum entity, carrying macroscopic quantum coherence described by the \( N \)-atom wave function as many exquisite experiments carried out in the last decade have witnessed. Yet its pivotal feature, its ‘quantumness’, originates from the spin-statistics properties (bosonic vs fermionic condensate) and the critical phenomenon which describes the formation of a condensate is a classical, not a quantum phase transition. This addresses query b) above.

Making the distinction between ‘quantumness’ arising from particle spin-statistics as in quantum statistical mechanics and that of large scale quantum coherence as in BEC offers a way to discern what would be called MQP in the modern sense of the word: usually it is the latter, not the former, which is the deciding factor. To name two examples: 1) Can the behavior of a collection of quantum gas be regarded as MQP? In solid state physics one treats the conduction electrons in a metal as a degenerate Fermi gas because the room temperature is much lower
than the Fermi temperature of the metal. The electron’s de Broglie thermal wavelength is much larger than the atomic spacing. But one would not regard the thermal properties of metals as MQP. Similar situations occur in the white dwarfs and neutron stars where the degenerate pressure of electrons or neutrons respectively balances the gravitation attraction to keep the stars in relative stable configurations. One does not refer to the existence of white dwarfs or neutron stars as examples of MQP. 2) Can one view collective excitations as MQP? Phonons are the quantized long wavelength modes of lattice vibrations. Like waves all such collective variables involve a large number of atoms and can span a large spatial extent. But one would not call these collective phenomena as MQP. However for a large population of bosonic gas accumulating in the ground state at an extremely low temperature forming a Bose-Einstein condensate (BEC), we do regard it as a MQP. The crucial difference is the large scale quantum coherence established in such systems. Similarly, for superconductors: the Cooper pairs existing at large spatial separation (compared to the extent of electron wave functions) are quasiparticles with full quantum coherence.

1.2. Scaling and interaction, RG running and criticality

In parallel to explicating the (rather restrictive) meaning of ‘quantumness’ used widely in QSM mentioned above here we wish to explore the meaning of ‘macroscopia’ in like spirit. Macroscopia in the most direct way conjures large numbers: many particles or components and/or large sizes or scales. Paper I focused on the ‘many’ aspect, here we focus on the ‘scale’ aspect. If one goes by the scale of a quantum system alone without other considerations such as the interactions (or coupling strength) amongst its underlying constituents then the quantum behavior of a macro object such as the Universe would be similar to that of a micro one, such as the internal degrees of freedom of an atom. In fact, both obey a harmonic oscillator equation of motion, albeit the former (for the wave function of the universe) has a negative spring constant. The scale is usually measured by the inverse mass of the relevant process. The physically relevant scale (or mass) is different from the bare scale (or mass) by a ratio given by a renormalization constant which captures the relevant interactions involved. Interaction is measured by the coupling constants which vary (‘run’) with energy according to the renormalization group (RG) equations. The infrared behavior of an interacting quantum system at or near its critical point such as the universality class it belongs to, is of special significance. We will review briefly these familiar concepts below as the notion of ‘macroscopic’ for quantum systems near criticality is determined by its infrared behavior and the interaction strength of its constituents.

1.2.1. Scales, interaction strength and effective theories

Let us examine the calculation of the energy levels of a hydrogen atom as an example. In a standard textbook treatment one uses the Schrödinger equation for an electron moving in the static Coulomb field of the proton. To a good approximation, the only properties of a proton which are relevant to this problem are its mass and charge. The knowledge of the quark structure of the proton is not necessary to compute the energy levels of the hydrogen atom.

The required knowledge of the proton depends on how accurate one asks for the energy levels. A more detailed calculation, taking hyperfine splitting into account, for instance, requires the knowledge of the spin of the proton and the value of the magnetic dipole moment. An even more accurate calculation requires the knowledge of the proton charge radius and details of the proton structure.

Another more practical angle or functional criterion towards this demarkation is to ask whether one could use such a system to perform quantum information processing, where quantum coherence and entanglement are essential. Cold atoms in an optical lattice, atom assembly, as well as superconducting flux qubits are the well-known viable examples. Note, however, there are proposed QIP schemes such as the NMR based ones, which do not invoke quantum entanglement directly, yet could achieve speed-ups over classical computation.
The typical length scale characteristic of the hydrogen atom is the Bohr radius $r_0 = 1/(m_e \alpha)$ (in units where $\hbar = c = 1$). The typical momentum scale is $1/r_0 = m_e \alpha$, the typical energy scale characteristic of the hydrogen atom is $\sim m_e \alpha^2$, and the typical time scale is $1/(m_e \alpha^2)$. We can get a quantitative estimate of the errors caused by the neglected interactions.

In an effective theory description the relevant interactions also depend on the question being asked. For the hydrogen atom the energy spectrum can be computed to the accuracy $(m_e \alpha/M_w)^2$ while ignoring the weak interactions. But if one is interested in atomic parity violation, the weak interactions are the leading contributions because the electromagnetic and strong interactions conserve parity. The effect of atomic parity violation will be very small because the weak scale is much larger than the atomic scale.

As one approaches shorter and shorter distances (or higher and higher energies), one needs to invoke the relevant physical processes at that scale. The high energy processes affecting low energy physics appear in physical parameters measured at the low energy regime. This is captured nicely by effective field theories, the essence of which we will discuss in a later section.

1.2.2. RG running and criticality In an interacting quantum system the variation of interaction strength with energy is described by the RG equations which are derived from the renormalization of the coupling constants (for ultraviolet divergence). Assuming that the system scales homogeneously this enables one to obtain the infrared properties of a quantum system from its ultraviolet behavior. At the critical point the correlation functions of such a system usually diverge: the system ‘feels’, so to speak, all the way to infinity. Let us consider how this interesting physical situation bears on macroscopic quantum phenomena.

Critical phenomena involves many different length scales. Consider a physical system going through a continuous phase transition under some temperature change: the correlation length diverges as the temperature approaches the critical temperature, and fluctuations at all length scales are involved. When the temperature lowers across the critical point, the system’s order parameter will develop a nonzero value, where the relevant degrees of freedom act in a correlated manner. In this sense the system’s behavior near its critical point indeed manifests in a macroscopic scale.

Note traditional treatment of critical phenomena is classical in that the external parameters, temperature, magnetic field, etc, are classical. But it could also be quantum in nature, the system being driven by quantum noise or vacuum fluctuations. Quantum phase transition is brought about by the change of the coupling parameters when the system is at zero temperature. It results from the energy competition between different ground states as a function of the coupling parameter, and quantum entanglement in the system has been shown to provide a useful measure of such transitions. We hope to address this issue in a following paper when we turn our attention to the role of quantum entanglement in the description of MQP.

Naively one would expect that at the critical point as the correlation length goes to infinity, the system would not depend on the microscopic details such as the lattice spacing in the Ising model, or the interaction amongst the microscopic constituents. But it does, and that makes it more interesting. It is this particular aspect of MQP which fascinates us, namely, what and how known microscopic details determine a quantum system’s macroscopic attributes, or, even more challenging, how to decipher a system’s microscopic features from the macroscopic phenomena. An example is the dialectic relation between emergent and quantum gravity pertaining to the micro and macro structures of spacetime [7].

On this point it is perhaps worth making the following observation: systems in a critical state can exist for a long time to certain classes of observers with specially interesting physical consequences, and many qualitative features in the ensuing phase depends on what came out of the transition. An example is inflationary cosmology where it is believed that the universe underwent a first (old) or second (new) order transition at the GUT (10^{14} GeV) scale, with a
duration of 68 e-folding time to be able to account for the entropy content \((10^{80} \text{ photons})\) in the present universe. During the exponential expansion the universe appears to be static for a local observer, its scale factor described by a de-Sitter -Einstein solution. In fact, in such an epoch between the metastable false vacuum and reheating to the true vacuum, one can use the language of critical phenomena to describe its dynamics, replacing the evolution by scaling (see, e.g., [8]). The microscopic features of the inflaton field – the quantum fluctuations which existed in the early universe – are magnified during this phase transition into galaxies of today. The effect of inflation is like a giant zoom lens at work [9] and the universe we observe today is in this sense a truly macroscopic quantum phenomenon [10]. The exponential red-shifting of outgoing waves from a black hole which accounts for the thermal feature of Hawking radiation has a similar nature, acting like a microscope in a pictorial description [11].

1.3. Correlation in relation to fluctuations and coherence. Organization of this paper

Unlike the situations of large \(N\) where an expansion in inverse \(N\) yields meaningful physical results, e.g., the leading order corresponds to a semiclassical limit, quantum correlations are less direct or transparent in their physical meanings. Correlation is related to fluctuations (as one form of fluctuation-dissipation relation indicates) and at the quantum level, quantum correlation can be a measure of quantum coherence (as the correlation history approach to quantum decoherence shows [12]. It is also related to but distinct from quantum entanglement (see work of Cirac [13]) It would be very useful to identify the conditions whereby these three properties could offer a measure (no matter how tenuous) of the macroscopic features of a quantum system and how they signify the quantum (albeit residual) nature of a macroscopic system. For this we use three examples to explore these aspects. Part I of this paper consisting of Sections 2 and 3 discusses the relation between the loop, large \(N\) and nPI (n-particle irreducible) expansions, the first two signifying the appearance of quantum and macro features and how quantum correlations is placed in regard to them. In Sec. 2 we use the existence of a H theorem at the next-to-leading-order in the quantum mechanical \(O(N)\) model to demarcate which order of large \(N\) expansion will entropy generation first ensue and thus can be (approximately) regarded as acquiring some macroscopic attributes. In Sec. 3 we provide a concrete example of the above connections in the Bose-Hubbard model for \(N\) atoms in an optical lattice. The nonequilibrium dynamics of this system is described by the equations of motion obtained from the closed-time-path (CTP) two particle irreducible \((2PI)\) effective action. We compare results from a large \(\mathcal{N}\) expansion where \(\mathcal{N}\) is the number of fields (see Eq. (3.5) below) under second order coupling with the exact (numerical) solution to see how quantum correlation plays out against \(\mathcal{N}\), the number of components of the \(\mathcal{N}\) body wave function. Although \(\mathcal{N}\) or \(N\) are not exactly indicators of the macroscopic, they offer a probe into the interplay between these three aspects: Correlation, fluctuations and coupling strength. Part II of this essay discusses the effective field theory (Sec. 4) and effective infrared dimensional reduction (Sec. 5). Though we didn’t dwell on this point as we believe it is quite well known, we want to point out the fact that the critical behavior of interacting quantum systems can provide a sense of macroscopia quite different from the simplistic depictions (referring to the size of the system or how many components it possesses). Key to this is the fact that coupling strengths can vary (‘run’) with energy scale according to the RG equations. In Sec. 4 we describe effective field theory, where this key notion well adopted in critical phenomena studies is put to good use in that the low energy phenomenology (or long wavelength behavior) is effectively independent of or largely insensitive to the high energy processes. In Sec. 5 we discuss the conditions where an interacting quantum system near the critical point may take on a lower-dimensional appearance. This effective infrared dimensional reduction is shown by an eigenvalue analysis: wherever the eigen-spectrum of the invariant operator of the order parameter field possesses a band structure with a gap separating the lowest mode or band from the higher sector, the infrared behavior of this system behaves effectively as
that of a lower dimension. These examples serve to show the intricacy of even a simple notion such as ‘macroscopic’ for a quantum system, that it is invariably tethered with the interaction strength, enters into the correlation lengths and undergoes qualitative changes when the system is close to the condition of criticality. The characterization of quantum macroscopic phenomena by quantum entanglement is left for a later paper.

Part I: Correlation hierarchy, $nPI$, $NLO$-large $N$ and H-theorem

2. Correlation entropy and H theorem at $NLO$

To prepare for our discussions on how quantum correlations play out in the quantum - macro issues we first introduce the $nPI$ effective action with $n=2$ as example, which is a special case of the so-called master effective action [14] (MEA), with the specific model of a quantum mechanical $O(N)$ model (QMON) which we have used in our first paper (MQP1) to illustrate the large $N$ expansion. We then recount (see [15, 16]) how $nPI$ is related to the loop expansion on the one hand, where one customarily takes as a measure of quantum features, and the large $N$ expansion on the other, where one may regard as signifying macroscopic features. (These latter two relations were expounded in MQP1.) We then discuss the implications of the existence of a H theorem proven in [17] using the QMON model. In the next section we will use the example of the nonequilibrium dynamics of BEC atoms to illustrate how this threefold-relation amongst $1/N$, $2PI$ and expansion in the (second order) coupling strength pertain to quantum and macro issues.

We paraphrase the results from [17] where a correlation entropy for an interacting quantum field is proposed, and a proof of the existence of a H-theorem for the quantum mechanical $O(N)$ model [18, 19] is provided. For the former, Calzetta and Hu followed the paradigm of Boltzmann-BBGKY [20] and proposed a correlation entropy (of the nth order) for an interacting quantum field [21, 22] obtained by ‘slaving’ the higher (n+1 th) order correlation functions in the Schwinger-Dyson system of equations (See [15, 16] and below). They then derived the closed time path (CTP) [23] two particle irreducible (2PI) [24] effective action (EA) for this model up to the next-to-leading order ($NLO$) in $1/N$ [17] and used the $CTP$ $2PI$ EA to prove an H-theorem for the correlation entropy of a quantum mechanical $O(N)$ model at the $NLO$ level. We begin with an introduction to the correlation hierarchy.

2.1. Boltzmann-BBGKY hierarchy and Schwinger-Dyson equations

As is well known in nonequilibrium statistical mechanics, truncation of the BBGKY hierarchy at a finite nth order yields a closed system 2. The system of equations describing the n-particle distribution functions are time reversal invariant. When a causal factorization condition such as Boltzmann’s molecular chaos assumption is imposed – that is, assuming that the n+1 th correlation function can be factorized into a product of n th correlation functions initially but not finally (after the collisions) – time-irreversibility appears and an H-theorem is obtained. This type of coarse graining of the hierarchy i.e., truncation plus causal factorization into a set of coupled equations for the n-particle distribution functions is called ‘slaving’ in the language of [15, 16]. Slaving of the n+1 -particle distribution function renders an otherwise closed system (of the n+1 th order correlation functions) an effectively open system (of nth order correlation functions) and ushers in the appearance of dissipative dynamics [22]. Noise and fluctuations [25, 26] should also appear, as required by the fluctuation-dissipation relation (FDR) [27], now

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2 The truncation of the correlation hierarchy is not just an arbitrary mathematical procedure, it reflects the fact that in realistic conditions, measurement are of finite accuracy associated with limited resolution of the instruments. Thus the relevant physical degrees of freedom are often limited to the lower end of the hierarchy, namely, the mean field and two-point function.
manifesting not just for an open system near equilibrium (as depicted in linear response theory), but for an effectively open system [15, 16].

A familiar example is the Boltzmann equation for dilute gases. At sufficiently low density the description of a molecular gas by a truncated BBGKY hierarchy with only the one-particle and two-particle distribution functions may be justified. Because of the low density one may further assume that before the collisions the colliding partners are uncorrelated, i.e., the two-particle distribution function factorizes into a product of single-particle distribution functions. This is the physical basis for Boltzmann’s imposition of the molecular chaos hypothesis. Of course the colliding partners become correlated after interaction. This assumed distinction between initial and final conditions is the origin of the macroscopic arrow of time, the appearance of dissipation and foe entropy generation in the Boltzmann paradigm. FDR calls for a rightful place for fluctuations in the Boltzmann equation, as demonstrated in the derivation of a stochastic Boltzmann equation [28, 29].

There are other approximation schemes that allow for the derivation of the kinetic equations from the correlation hierarchy [20]. To see the relation between correlation and interaction in this context of kinetic theory consider a system in which particles interact via a potential of the form \( V(r_i, r_j) = \lambda v(r_i, r_j) \). For weakly coupled systems, i.e., \( \lambda \ll 1 \), the hierarchy of equations can be written for a certain order of \( \lambda \).

To \( O(\lambda) \) the equation for the one-particle distribution function, namely, the Vlasov equation, is closed, i.e., it does not involve any two-particle distribution function. We emphasize that in this case truncation alone not followed by slaving is sufficient to obtain a closed equation for the one-particle distribution function. There is no collision term for short-ranged interaction, but interactions between particles do exist represented by an average potential of all other particles present. Vlasov equation is useful for describing the kinetics of systems containing many particles interacting under long range forces. Recall we have seen it in our first paper as the equation obtained from the leading order large \( N \) expansion in the \( O(N) \) model. Vlasov equation is obtained as a mean field theory with reversible dynamics and no H-theorem.

To \( O(\lambda^2) \) the set of equations for the one-particle and two-particle distribution functions is also closed. By slaving the two-particle distribution function to the one-particle distribution function the Landau equation is obtained. Similar to the Boltzmann equation, Landau equation also has a collision term accounting for the collisions between individual particles at \( O(\lambda^2) \). Landau equation describes irreversible dynamics and its solutions satisfy the H-theorem. What is different from the Boltzmann equation is the form of the collision integral. This difference can be traced back to the absence of a term in the equation for the two-particle distribution function at \( O(\lambda^2) \) compared to the low density approximation.

The situation in quantum kinetic field theory is completely analogous. By quantum kinetic field theory (see, e.g., [21, 22]), we are referring to the hierarchy of coupled equations for the relativistic Wigner function and its higher-correlation analogs, obtainable from the variation of the master effective action [15, 16] whose variation yields the Schwinger-Dyson equations. This is a quantum analogue of the BBGKY hierarchy, expressed in a representation convenient for distinguishing between microscopic (quantum field-theoretic) and macroscopic (transport and relaxation) phenomena 3.

One may choose to work with a truncation of the hierarchy of the Wigner function and its higher correlation analogs, or one may instead perform a slaving of, for example, the Wigner-transformed four-point function, which leads (within the context of perturbation theory) directly to the relativistic Boltzmann equation [22] and the usual \( H \)-theorem. The truncation and subsequent slaving of the hierarchy within quantum kinetic field theory can be carried out at

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3 It should be pointed out that in order to identify the relativistic Wigner function with a distribution function for quasiparticles, one must show that the density matrix has decohered, and this is neither guaranteed nor required by the existence of a separation of macroscopic and microscopic time scales [30].
any desired order, as dictated by the initial conditions and relevant interactions.

2.2. Entropy from slaving the higher correlations: the CTP 2PI EA

A general procedure has been presented for obtaining coupled equations for correlation functions at any order \( l \) in the correlation hierarchy, which involves a truncation of the master effective action at a finite order in the loop expansion [15, 16]. By working with an \( l \) loop-order truncation of the master effective action, one obtains a closed, time-reversal invariant set of coupled equations for the first \( l + 1 \) correlation functions, \( \phi, G, C_3, \ldots, C_{l+1} \). In general, the equation of motion for the highest order correlation function will be linear, and thus can be formally solved using Green’s function methods. The existence of a unique solution depends on supplying causal boundary conditions. When the resulting solution for the highest correlation function is back-substituted into the evolution equations for the other lower-order correlation functions, the resulting dynamics is not time-reversal invariant, and generically dissipative. Thus, as was described before, with the slaving of the higher-order (Wigner-transformed) correlation function in quantum kinetic field theory, we have rendered a closed system (the truncated equations for correlation functions) into an effectively open system. In addition to dissipation, one expects that an effectively open system will manifest noise/fluctuations, as shown in [15, 16] for the case of the slaving of the four-point function to the two-point function in the symmetry-unbroken \( \lambda \Phi^4 \) field theory. Thus a framework exists for exploring irreversibility and fluctuations within the context of a unitary quantum field theory, using the truncation and slaving of the correlation hierarchy. The effectively open system framework is useful for precisely those situations, where a separation of macroscopic and microscopic time scales (which would permit an effective kinetic theory description) does not exist, such as is encountered in the thermalization issue.

As a particular coarse graining measure the slaving of higher correlation functions to lower-order correlation functions within a particular truncation of the correlation hierarchy has several important benefits. It can be implemented in a truly nonperturbative fashion. This necessitates a nonperturbative resummation of daisy graphs, which can be incorporated in the truncation/slaving of the correlation hierarchy in a natural way.

In [16] the authors are interested in the growth of entropy due to the coarse graining of the correlation hierarchy by slaving a higher correlation function. The simplest nonperturbative truncation of the Schwinger-Dyson equations for the \( \lambda \Phi^4 \) field theory which contains the time-dependent Hartree-Fock approximation is the two-loop truncation of the master effective action, in which only the mean field \( \phi \), the two-point function \( G \), the three-point function \( C_3 \) are dynamical. All higher order correlation functions obey algebraic constraints, and can thus be expressed in terms of the three dynamical correlation functions.

While this truncation of the Schwinger-Dyson equations is well-defined and could in principle be solved, it is disadvantageous because, as stated above, without some coarse graining, the system will not manifest irreversibility and will not equilibrate. Therefore we slave the three-point function to the mean field and two-point function, and thus arrive at an effectively open system. In principle, a systematic analysis of the coarse-grained dynamics of the mean field and two-point function should include stochasticity [15, 16].

The 2PI formalism is also suitable for addressing this question because, provided an auxiliary field is cleverly introduced, the 2PI CTP effective action can be found in closed form at each order in \( 1/N \) [17, 31]. We now use the familiar quantum mechanical \( O(N) \) model to illustrate how to derive the 2PI effective action and then explore the conditions for the existence of an H-theorem. This model was used in our first paper [1] for the discussion of large \( N \) and macroscopia.
2.3. $O(N)\lambda X^4$ theory

As we recall the system dynamics is described by the Hamiltonian with variables $X_A$ and their conjugate momenta $P_A$, where $A,B$ are the $O(N)$ group indices, with

$$H = \frac{1}{2} \left\{ P^B P^B + M^2 X_B X_B + \lambda \frac{4}{4N} (X_B X_B)^2 \right\}$$

(2.1)

The classical action

$$S = \int dt \frac{1}{2} \left\{ \dot{X}_B \dot{X}_B - M^2 X_B X_B - \frac{\lambda}{4N} (X_B X_B)^2 \right\}$$

(2.2)

where $M_0^2$ and $\lambda_0$ are the mass parameter and coupling constant. We rescale $X_B \equiv \sqrt{N} x_B$

$$S = N \int dt \frac{1}{2} \left\{ \dot{x}_B \dot{x}_B - \frac{M^2}{\sqrt{\lambda}} + \frac{\sqrt{\lambda}}{2} x_B x_B \right\}$$

(2.3)

Discarding a constant term, we may rewrite the classical action as

$$S = N \int dt \frac{1}{2} \left\{ \dot{x}_B \dot{x}_B - \left[ \frac{M^2}{\sqrt{\lambda}} + \frac{\lambda}{2} x_B x_B \right] \right\}$$

(2.4)

To set up the $1/N$ resummation scheme, it is customary to introduce the auxiliary field $\chi$, writing

$$S = \frac{N}{2} \int \left\{ \dot{x}_B \dot{x}_B - \left[ \frac{M^2}{\sqrt{\lambda}} + \frac{\lambda}{2} x_B x_B \right]^2 \right\}$$

whence

$$S = N \int dt \left\{ \frac{1}{2} \dot{x}_B \dot{x}_B - \chi \left[ \frac{M^2}{\lambda} + \frac{x_B x_B}{2} \right] + \frac{1}{2\lambda} \chi^2 \right\}$$

(2.5)

From now on, we consider $\chi$ and $x_B$ as fundamental variables on equal footing.

Because of the $O(N)$ symmetry, the symmetric point must be a solution of the equations of motion. For simplicity, we shall assume we are within this symmetric phase, and treat $x_B$ as a quantum fluctuation. We also split the auxiliary field $\chi = \bar{\chi} + \tilde{\chi}$ into a background field $\bar{\chi}$ and a fluctuation field $\tilde{\chi}$. The action becomes

$$S = S_0 + S_1 + S_2 + S_3$$

(2.7)

$S_0$ is just the classical action evaluated at $x_B = 0, \chi = \bar{\chi}$:

$$S_0 = \frac{N}{\lambda} \int dt \left\{ \frac{1}{2} \dot{\bar{\chi}}^2 - M^2 \bar{\chi} \right\}$$

(2.8)

$S_1$ contains terms linear in $\tilde{\chi}$ and can be set to zero by a suitable choice of the background field $\bar{\chi}$:

$$S_1 = \frac{N}{\lambda} \int dt \left\{ \tilde{\chi} - M^2 \bar{\chi} \right\}$$

(2.9)

$S_2$ contains the quadratic terms and yields the tree - level inverse propagators,

$$S_2 = N \int dt \left\{ \frac{1}{2} \dot{x}_B \dot{x}_B - \frac{\bar{\chi}}{2} x_B x_B + \frac{1}{2\lambda} \tilde{\chi}^2 \right\}$$

(2.10)
Finally $S_3$ contains the bare vertex

$$S_3 = \left( -\frac{N}{2} \right) \int d^dx \{ \bar{x} x_B x_B \}. \quad (2.11)$$

To write the $2PI$ CTP EA we double the degrees of freedom, incorporating a branch label $a = 1, 2$ (for simplicity, if not explicitly written, we assume that the label $a$ includes the time branch, i.e., $x^{Aa} \equiv x^{Aa}(t_a)$). We also introduce propagators $G^{Aa,Bb}$ for the path ordered expectation values

$$G^{Aa,Bb} = \langle x^{Aa} x^{Bb} \rangle \quad (2.12)$$

and $F^{ab}$ for

$$F^{ab} = \langle \bar{\chi}^a \bar{\chi}^b \rangle \quad (2.13)$$

Because of symmetry, it is not necessary to introduce a mixed propagator, for $\langle \bar{\chi}^a x^{Bb} \rangle \equiv 0$.

The $2PI$ CTP EA reads

$$\Gamma = \frac{N}{2} \left[ \bar{x}^1 - \bar{x}^2 \right] + \frac{1}{2} \int dudv \left\{ D_{Aa,Bb}(u,v)G^{Aa,Bb}(u,v) + \frac{N}{\lambda_0} c_{ab} \delta(u,v)F^{ab}(u,v) \right\} - \frac{i\hbar}{2} \left[ \text{Tr} \ln G + \text{Tr} \ln F \right] + \Gamma_Q \quad (2.14)$$

where, if the position variable is explicit, $c_{11} = -c_{22} = 1$, $c_{12} = c_{21} = 0$,

$$D_{Aa,Bb}(u,v) = N \delta_{AB} \left[ c_{ab} \partial_x^2 - c_{abc} \bar{\chi}^c \right] \delta(u,v), \quad (2.15)$$

and $c_{ab} = 1$ when all entries are 1, $c_{abc} = -1$ when all entries are 2, and $c_{abc} = 0$ otherwise. When we use the compressed notation, it is understood that $c_{ab} \equiv c_{ab}(t_a, t_b)$ and $c_{abc} \equiv c_{abc}(t_a, t_b, t_c)$. $\Gamma_Q$ is the sum of all $2PI$ vacuum bubbles with cubic vertices from $S_3$ and propagators $G^{Aa,Bb}$ and $F^{ab}$. Observe that $\Gamma_Q$ is independent of $\bar{\chi}^c$.

Taking variations of the $2PI$ CTP EA and identifying $\bar{\chi}^1 = \bar{\chi}^2 = \bar{\chi}$, we find the equations of motion

$$\frac{N}{2} \delta_{AB} D_{ab} - \frac{i\hbar}{2} \left[ G^{-1} \right]_{Aa,Bb} + \frac{1}{2} \Pi_{Aa,Bb} = 0 \quad (2.16)$$

$$\frac{N}{2\lambda} c_{ab} - \frac{i\hbar}{2} \left[ F^{-1} \right]_{ab} + \frac{1}{2} \Pi_{ab} = 0 \quad (2.17)$$

$$\frac{N}{\lambda} \left\{ \bar{x}(t) - M^2 \right\} - \frac{N}{2} \delta_{AB} G^{A1,B1}(t,t) = 0 \quad (2.18)$$

where $D_{ab}(u,v) = c_{ab} \left[ \partial_x^2 - \bar{x}(u) \right] \delta(u,v)$,

$$\Pi_{Aa,Bb} = 2 \frac{\delta \Gamma_Q}{\delta G^{Aa,Bb}}; \quad \Pi_{ab} = 2 \frac{\delta \Gamma_Q}{\delta F^{ab}} \quad (2.19)$$

We shall seek a solution with the structure

$$G^{Aa,Bb} = \frac{\hbar}{N} \delta^{AB} G^{ab}(u,v) \quad (2.20)$$
which is consistent with vanishing Noether charges. Then it is convenient to write

\[ F^{ab} = \frac{\hbar}{N} H^{ab}; \quad \Pi_{Aa,Bb} = \delta_{AB} P_{ab}; \quad \Pi_{ab}(x,y) = N Q_{ab}(x,y) \]  

(2.21)

The equations become

\[ D_{ab} - i \left[ G^{-1} \right]_{ab} + \frac{1}{N} P_{ab} = 0 \]  

(2.22)

\[ \frac{1}{\lambda} c_{ab} - i \left[ H^{-1} \right]_{ab} + Q_{ab} = 0 \]  

(2.23)

\[ \frac{1}{\lambda} \left\{ \bar{\chi} (t) - M^2 \right\} - \frac{\hbar}{2} G^{11}(t,t) = 0 \]  

(2.24)

Observe that

\[ P_{ab} = \frac{2}{\hbar} \frac{\delta \Gamma Q}{\delta G^{ab}}; \quad Q_{ab} = \frac{2}{\hbar} \frac{\delta \Gamma Q}{\delta H^{ab}} \]  

(2.25)

These are the exact equations we must solve. The successive $1/N$ approximations amount to different constitutive relations expressing $P_{ab}$ and $Q_{ab}$ in terms of the propagators.

For power counting in orders of $N$, observe that in any given Feynman graph each vertex contributes a power of $N$, each internal line a power of $N^{-1}$, and each trace over group indices another power of $N$. We have both $G$ and $H$ internal lines, but the $G$ lines only appear in closed loops. On each loop, the number of vertices equals the number of $G$ lines, so there only remains one power of $N$ from the single trace over group labels. Therefore the overall power of the graph is the number of $G$ loops minus the number of $H$ lines. Now, since we only consider $2PI$ graphs, there is a minimum number of $H$ lines for a given number of $G$ loops. For example, if there are two $G$ loops, they must be connected by no less than 3 $H$ lines, and so this graph cannot be higher than $NNLO$. A graph with 3 $G$ loops cannot have less than 5 $H$ lines, and so on.

We conclude that $\Gamma Q$ vanishes at $LO$, and therefore $P_{ab} = Q_{ab} = 0$. There is only one $NLO$ graph, consisting of a single $G$ loop and a single $H$ line. This graph leads to

\[ \Gamma^{NLO}_Q = \left( \frac{-i\hbar}{2} \right) \left( -\frac{N}{2\pi\hbar} \right)^2 2N \left( \frac{\hbar}{N} \right)^3 \int dudv H^{ad}(u,v) G^{be}(u,v) G^{cf}(u,v) \]  

(2.26)

Therefore, we get

\[ P_{ab} = i\hbar c_{acde} G^{ce} G^{df} \]  

(2.27)

\[ Q_{ab} = \frac{i\hbar}{2} c_{acde} G^{ce} G^{df} \]  

(2.28)

The $2PI$ EA yields equations of motion for the (arbitrary time) two-point functions of the theory. Given a solution of these equations, in principle we may find the expectation values of a large family of composite operators at any given time. Suppose we adopt a coarse-grained description where we choose a certain number of these expectation values as the relevant variables to describe the system. Then there will be a single density matrix which has maximum von Neumann entropy with respect to the class of states reproducing the preferred expectation values. This maximum entropy density matrix is the reduced density matrix for the system, and its entropy its correlation entropy. The $H$ theorem is the statement that the correlation entropy grows in time, when the correlations themselves are evolved using the equations derived from the $2PI$ EA truncated to some order in the $1/N$ expansion. For details in the proof refer to [17].
2.4. H theorem at NLOLN and its implications

It is clear that if we could solve the exact evolution, the $H$ theorem would be manifest: Given $\rho(t_0)$ at the initial time $t_0$, solve the exact Liouville equation up to a time $t_1$. Let $\tilde{\rho}(t_1)$ be the result. We extract the new expectation values from $\tilde{\rho}(t_1)$ and use them to construct the new maximum entropy density matrix $\rho(t_1)$. Then $S[\rho(t_1)] \geq S[\tilde{\rho}(t_1)]$, by definition, and $S[\rho(t_1)] = S[\rho(t_0)]$, because the exact evolution is unitary, thus the $H$ theorem.

What we need to determine is whether, given the approximate dynamics for the expectation values provided by the $1/N$ scheme, the $H$ theorem still holds. From our first paper (MQP1) in this series [1] we know that to $LO$ it does not because $LO$ large $N$ yields a mean field theory whose equation of motion, the Vlasov equation, is unitary. Thus there is no net entropy production and no $H$-theorem. In such a case, if one works with a Fock representation, for boson fields, the number of particles increases with time and can be used as a measure of field entropy, what Hu and Pavon [32] called an intrinsic measure of field entropy. (See also [33].) However, this should not be confused with the correlation entropy of the Boltzmann kind under study for which the $H$-theorem is defined.

Of course, we expect better approximations to be closer to the exact dynamics, therefore requiring less external control of (or rather, tampering with) the system. This reduces the entropy loss to the environment. Physically, whenever some higher order correlations are ignored, we expect to obtain an $H$ theorem because such a description of the system is incomplete. This is the physical origin of correlation entropy, as explained in the Introduction. In order to have the different correlations evolving according to the $2PI 1/N$ equations appropriate to the desired order, instead of the accurate description from the full Schwinger-Dyson hierarchy, we must keep nudging it to conform to this artificially created condition due to our inability to comprehend the complete picture. This is the source of a new kind of noise arising from ignoring the higher order correlation functions, called correlation noise, which, when incorporated in the Schwinger-Dyson hierarchy, yields the (nth-correlation order) stochastic SD equations, as explained in [16].

2.5. $2PI$ EA, $H$-theorem, quantum-classical and micro-macro

We can now ask the question: How does quantum correlations get registered in the $2PI$ EA and how does the existence of $H$-theorem at the $NLO$ expansion pertain to the quantum-Macro issue?

First, how do the quantum features show up in the $2PI$ effective action? Only equations of evolution for the first two moments can be obtained from the $2PI$ EA. It is known that [34] the equations of motion for the first two moments are identical for classical and quantum systems. Quantum corrections arise from the way the higher order moments are treated implicitly in the effective action approach. Because of this implicit dependence the $2PI$ EA approach is not easy to work with in addressing the differences of quantum and classical dynamics. One could consider going to higher orders, at least $3PI$, and see if the third moment equations possess the terms absent in classical dynamics at a given order in large $N$.

The micro-Macro (m-M) demarkation issue is as difficult to tackle, if not more, as the quantum-classical transition issue. Going from micro to macro requires coarse graining. QMON model is a “democratic” system, hence the coarse graining should be “democratic” as well. The $2PI$ approach provides us with a reduced description in terms of the two lowest order Green’s functions, which we shall refer to as the relevant variables. Higher order Green’s functions are referred to as irrelevant variables. If one keeps track only of the relevant variables information about irrelevant variables is lost, by choice. If these variables are coupled and one wishes to see

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4 Democratic’ is a figurative description of a system of identical particles and ‘autocratic’ refers to a system containing a distinct member from the rest. This corresponds to the Boltzmann versus Langevin paradigms describing an effectively open (by correlation order, which depends on the level of precision of observation) versus an open (by ab initio designation) system. See, [35].
how the relevant variables (our system) is affected by the irrelevant variables (its environment) one needs to take into account the back-action of the irrelevant variables. This back-action included reduced system (in terms of e.g., a reduced density matrix) would show dissipative dynamics and a H-theorem will obtain for the correlation entropy.

The question is whether the H-theorem holds for the dynamics obtained at a certain order of the $1/N$ expansion. There is no H-theorem at the leading order. This can be understood by realizing that the LO is equivalent to Gaussian approximation, and in this approximation correlations of order higher than two have no dynamics. As a result the mean field and the two point function contain all the available partial information because of the approximation imposed. The fact that H-theorem holds at NLO shows that the NLO approximation is capable of capturing the back-action from the irrelevant variables which engenders dissipation in the dynamics of the relevant variables. Very coarsely, in the large N perspective the NLO may be viewed as providing us a witness to the micro-macro transition. Note this does not mean that NLO large N is a good approximation in general. Actually we know from Mihaila et. al. [18] that it fails at long times although the H-theorem holds at long times, albeit only shown for a vacuum initial state [17]. The existence of an H-theorem, hence irreversibility, is a way of arguing that we have gone beyond the micro-domain.

In the QMON model $N$ plays a dual role. On the one hand $N$ is the number of degrees of freedom. On the other hand $N$ appears in the denominator of the interaction term in the system Hamiltonian 2.1. The large N limit is usually associated with the number of components $N \to \infty$ only, however for the QMON model it is the weak coupling limit as well. If we would like to make an analogy to the classical BBGKY hierarchy of section 2.1, large N approximation would have the same effect as the weak coupling approximation. The fact that there is no H-theorem at LO but there is an H-theorem at NLO is analogous to the fact that Vlasov equation doesn’t lead to entropy production, whereas the Landau equation does so.

Entropy increase is a signifier of irreversibility and irreversibility is usually associated with macroscopisa. LO approximation gives reversible dynamics, while at NLO irreversibility emerges. This fact appears to be in contradiction with the association of irreversibility with macroscopia, since LO theory is successful in describing systems with more components $N$ than NLO theory. The apparent paradox can be resolved by appreciating the dual role of $N$ in the QMON model. The above example shows the interplay between system-size and interaction strength in the emergence of macroscopic qualities in a system. The system-size alone is not enough to qualify the system as macroscopic.

The work of Mihaila et. al. [18] reveals a critical value of $N$ below which the NLO approximation does poorly ($N_c = 18.6$ for the parameters they chose for the simulations). As was discussed in MQP1 this is a way of quantifying when $N$ is actually large enough for a certain order to yield good results. This is a stronger condition than the validity of H-theorem (which holds at NLO for any $N$), since it requires the dynamics to follow the exact one, rather than just one condition of nondecreasing entropy. We may combine these two results into the following interpretation: The H-theorem is necessary for a satisfactory reduced description of a macroscopic system. It holds at NLO. NLO is a reasonable approximation for the system dynamics for $N > N_c$. Hence for $N > N_c$ the reduced description is consistent.

3. Model study: 2PI, NLOLN, loop expansions in nonequilibrium BEC dynamics

We begin with a description of the well-known Bose-Hubbard model and state our goal in this illustrated example. We introduce the 2PI generating functional to construct the 2PI effective action ($EA$) $\Gamma[\phi, G]$ and Green’s functions. We perform a perturbative expansions on $\Gamma_2$ and define the various approximation schemes. We then derive the equation of motion from the CTP 2PI $EA$ and discuss the results under each approximation scheme, starting with the Hartree-Fock-Bogoliubov (HFB) approximation followed by second order expansions which
includes the $1/N$ expansion and then the full second order expansion. The results from these approximations are compared with the exact numerical answers so as to exhibit the effects of quantum correlations, the coupling strength and how the results depend on increasing $N$.

The dynamics of an ultracold bosonic gas in an optical lattice can be described by a Bose-Hubbard model where the system parameters are controlled by laser light. For a one dimensional lattice the Hamiltonian is:

$$
\hat{H} = -J \sum_i (\hat{\Phi}_i^\dagger \hat{\Phi}_{i+1} + \hat{\Phi}_{i+1}^\dagger \hat{\Phi}_i) + \sum_i \epsilon_i \hat{\Phi}_i^\dagger \hat{\Phi}_i + \frac{1}{2} U \sum_i \hat{\Phi}_i^\dagger \hat{\Phi}_i^\dagger \hat{\Phi}_i \hat{\Phi}_i,
$$

where $\hat{\Phi}_i$ and $\hat{\Phi}_i^\dagger$ are the annihilation and creation operators at the site $i$ which obey the canonical commutation relations for bosons. Here, the parameter $U$ denotes the strength of the on-site repulsion of two atoms on the site $i$; the parameter $\epsilon_i$ denotes the energy offset of each lattice site due to an additional slow varying external potential such as a magnetic trap and $J$ denotes the hopping rate between adjacent sites. Because the next-to-nearest neighbor amplitudes are typically two orders of magnitude smaller, tunneling to them can be neglected. We denote the total number of atoms by $N$ and the number of lattice sites by $I$. Only a one-dimensional homogeneous lattice with periodic boundary conditions is considered.

A dimensionless parameter that is convenient to describe the different regimes of $H$ is the coupling strength $\lambda \equiv NU/IJ$. Different from a homogeneous system without a lattice where at zero temperature the superfluid fraction is always unity, the presence of the lattice changes the superfluid properties and even at zero temperature, the superfluid fraction decreases with the lattice depth. For strong-coupling strengths $\lambda > \lambda_{\text{crit}}$, it is known that the ground state undergoes a quantum phase transition from a superfluid to a Mott insulator. In the weakly interacting regime, $\lambda \ll 1$, where tunneling overwhelms the repulsion, to a good approximation quantum fluctuations can be neglected and the properties of the system can be described by replacing the operator on the lattice site $i$ by a classical $c$ number. It can be said that most of the atoms are in the zero quasimomentum state. In the intermediate regime $1 \ll \lambda \ll \lambda_{\text{crit}}/2$ the interactions between the bosons can be very strong but the ground state is nevertheless a superfluid. For these interaction parameters a self-consistent HFB-Popov theory gives a good description of the system. However, different from the weak interacting regime where the depletion of the zero quasimomentum state is very small and has little effect on the superfluid properties, in this intermediate regime, depleted atoms spread over the central part of the band and reduces the superfluid fraction. As interactions are further increased the depleted population completely fills the band and cancels the superfluid properties. The system reaches the Mott insulator regime, where atoms are localized at each lattice site and the eigenstates of the system are almost Fock states with vanishing number fluctuations per lattice site. The dynamics in the intermediate regime is the focus of study in [37], where the superfluid properties are important but quantum fluctuations cannot be ignored.

3.1. $2PI EA \Gamma[\phi, G]$

The first requirement for the study of nonequilibrium processes is a general initial-value formulation depicting the dynamics of interacting quantum fields. The $CTP$ or Schwinger-Keldysh effective action formalism [23] serves this purpose. The second requirement is to describe the evolution of the correlation functions and the mean field on an equal footing. The two particle irreducible ($2PI$) formalism [24] where the correlation functions appear also as independent variables serves this purpose. By requiring the generalized (master) $CTP$ effective...
In terms of these fields the classical action takes the form

$$G \text{ field by } \phi_{a,b} = (1/N) \sum_i \Phi_i^a(t) \Phi_i^b(t),$$

where, as before, $i$ denotes the lattice position, $J$ is the hopping rate and $U$ is the interaction strength. We limit the analysis to the case when no external potential is present and include only nearest neighbor hopping. To compactify our notation we introduce $\Phi^i$ by

$$\Phi_i = \Phi^1_i, \quad \Phi_i^* = \Phi^2_i.$$ (3.4)

In terms of these fields the classical action takes the form

$$S[\Phi_i, \Phi_i^*] = \int dt \sum_i \left[ i\hbar \Phi_i^*(t) \partial_t \Phi_i(t) + J (\Phi_i^*(t) \Phi_{i+1}^i(t) + \Phi_i(t) \Phi_{i+1}^*(t)) \right]$$

$$- \frac{U}{2} \Phi_i^*(t) \Phi_i^b(t) \Phi_i(t) \Phi_i(t),$$ (3.3)

where, as before, $i$ denotes the lattice position, $J$ is the hopping rate and $U$ is the interaction strength. We limit the analysis to the case when no external potential is present and include only nearest neighbor hopping. To compactify our notation we introduce $\Phi_i^a(a = 1, 2)$ defined by

$$\Phi_i = \Phi_i^1, \quad \Phi_i^* = \Phi_i^2.$$ (3.4)

In terms of these fields the classical action takes the form

$$S[\Phi_i, \Phi_i^*] = \int dt \sum_i \left[ \frac{1}{2} \hbar \Phi_i^a(t) \hbar \partial_t \Phi_i^b(t) + J \sigma_i \Phi_i^a(t) \Phi_{i+1}^b(t) - \frac{U}{4N} (\sigma_i \Phi_i^a(t) \Phi_i^b(t))^2 \right],$$ (3.5)

where $N$ is the number of fields (two in this case), and summation over repeated field indices $a, b = (1, 2)$ is implied. $\hbar$ and $\sigma_i$ are matrices defined as

$$\hbar = \begin{pmatrix} 0 & -1 \\ 1 & 0 \end{pmatrix}, \quad \sigma_i = \begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix}.$$ (3.6)

After second quantization the fields $\Phi_i^a$ are promoted to operators. We denote the expectation value of the field operator or mean field by $\phi_i^a(t)$ and the expectation value of the composite field by $G_{ij}^a(t, t')$. Physically, $|\phi_i^a(t)|^2$ is the condensate population and the composite fields determine the fluctuations around the mean field.

$$\phi_i^a(t) = \langle \Phi_i^a(t) \rangle,$$ (3.7)

$$\hbar G_{ij}^a(t, t') = \langle T_C \Phi_i^a(t) \Phi_j^a(t') \rangle - \langle \Phi_i^a(t) \rangle \langle \Phi_j^a(t') \rangle.$$ (3.8)

The brackets denote taking the expectation value with respect to the density matrix and $T_C$ denotes time ordering along a contour $C$ in the complex plane.

All correlation functions of the quantum theory can be obtained from the effective action $\Gamma[\phi, G]$, the two particle irreducible generating functional for Green’s functions parametrized by $\phi_i^a(t)$ and the composite field $G_{ij}^a(t, t')$. To get an expression for the effective action we first define the functional $Z[J, K]$ [24] as

$$Z[J, K] = e^{i/hW[J, K]} = \prod_i \int D\Phi_i^a \exp \left\{ \frac{i}{\hbar} \left[ S[\Phi] + \int dt \sum_i J_{ia}(t) \Phi_i^a(t) \right] \right.$$

$$\left. + \frac{1}{2} \int dt dt' \sum_{ij} \Phi_i^a(t) \Phi_j^a(t') K_{iab}(t, t') \right\},$$ (3.9)
where we have introduced the following index lowering convention

\[ X_a = \sigma_{ab} X^b. \]  

(3.10)

The functional integral (3.9) is a sum over classical histories of the field \( \Phi^a \) in the presence of the local source \( J_{ia} \) and the non local source \( K_{ijab} \). The coherent state measure is included in \( D\Phi \). The addition of the two-particle source term is what characterizes the 2PI formalism.

We define \( \Gamma[\phi, G] \) as the double Legendre transform of \( W[J, K] \) such that

\[
\frac{\delta W[J, K]}{\delta J_{ia}(t)} = \phi_i^a(t), \quad \frac{\delta W[J, K]}{\delta K_{ijab}(t,t')} = \frac{1}{2} [\phi_i^a(t)\phi_j^b(t') + \hbar G^{ab}_{ij}(t,t')].
\]

(3.11)

(3.12)

Expressing \( J \) and \( K \) in terms of \( \phi \) and \( G \) yields

\[
\Gamma[\phi, G] = W[J, K] - \int dt \sum_i J_{ia}(t)\phi_i^a(t) - \frac{1}{2} \int dt dt' \sum_{ij} \phi_i^a(t)\phi_j^b(t')K_{ijab}(t,t').
\]

(3.13)

From this equation the following identity can be derived:

\[
\frac{\delta \Gamma[\phi, G]}{\delta \phi_i^a(t)} = -\left( J_{ia}(t) + \int dt' \sum_j (K_{ijad}(t,t'))\phi_j^d(t') \right),
\]

(3.14)

\[
\frac{\delta \Gamma[\phi, G]}{\delta G^{ab}_{ij}(t,t')} = -\frac{\hbar}{2} K_{ijab}(t,t').
\]

(3.15)

In order to get an expression for \( \Gamma[\phi, G] \) notice that by using (3.9) for \( W[J, K] \) and placing it in (3.13) for \( \Gamma[\phi, G] \), it can be written as

\[
\exp\left(\frac{i}{\hbar} \Gamma[\phi, G] \right) = \prod_a \int D\Phi^a\exp \left\{ \frac{i}{\hbar} \left( S[\Phi] + \int dt_i J_{ia}(t) [\Phi_i^a(t) - \phi_i^a(t)] + \frac{1}{2} \int dt_i dt_j' \phi_i^a(t)K_{ijab}(t,t')\phi_j^b(t') - \phi_i^a(t)K_{ijab}(t,t')\phi_j^b(t') - \frac{\hbar}{2} \text{Tr} G K \right) \right\}
\]

(3.16)

\[
= \prod_a \int D\Phi^a\exp \left\{ \frac{i}{\hbar} \left( S[\Phi] - \int dt_i \frac{\delta \Gamma[\phi, G]}{\delta \phi_i^a(t)} [\Phi_i^a(t) - \phi_i^a(t)] - \frac{1}{\hbar} \int dt_i dt_j' [\Phi_i^a(t) - \phi_i^a(t)] \frac{\delta \Gamma[\phi, G]}{\delta G^{ab}_{ij}(t,t')} [\Phi_j^b(t') - \phi_j^b(t')] + \text{Tr} G \frac{\delta \Gamma[\phi, G]}{\delta G} \right) \right\},
\]

where \( \text{Tr} \) means taking the trace. For simplicity we have denoted \( \int dt \sum_i \) by \( \int dt_i \). Defining the fluctuation field, \( \varphi_i^a = \Phi_i^a - \phi_i^a \), we have

\[
\Gamma[\phi, G] = -\text{Tr} G \frac{\delta \Gamma[\phi, G]}{\delta G} = -i\hbar \ln \prod_a \int D\varphi^a \exp \left( \frac{i}{\hbar} S[\phi, G; \varphi] \right)
\]

(3.17)

\[
S[\phi, G; \varphi] = S[\phi + \varphi] - \int dt_i \frac{\delta \Gamma[\phi, G]}{\delta \phi_i^a(t)} \varphi_i^a(t) - \frac{1}{\hbar} \int dt_i dt_j' \varphi_i^a(t) \frac{\delta \Gamma[\phi, G]}{\delta G^{ab}_{ij}(t,t')} \varphi_j^b(t').
\]

(3.18)
By introducing the classical inverse propagator \( iD^{-1}(\phi) \) given by

\[
iD_{ijab}(t, t')^{-1} = \frac{\delta S[\phi]}{\delta \phi^a_i(t) \delta \phi^b_j(t')}
\]

and

\[
= (\delta_{ij} h_{ab} \partial_t + J(\delta_{i+1j} + \delta_{i-1j})\sigma_{ab} ) \delta(t - t')
- \frac{U}{\mathcal{N}} (2\phi_{ia}(t)\phi_{ib}(t) + \sigma_{ab}\phi_{ij}^c(t)\phi_{ic}(t)) \delta(t - t'),
\]

the solution of the functional integro-differential equation (3.17) can be expressed as

\[
\Gamma[\phi, G] = S[\phi] + \frac{i}{2} \text{Tr} \ln G^{-1} + \frac{i}{2} \text{Tr} D^{-1}(\phi)G + \Gamma_2[\phi, G] + \text{const.}
\]

The quantity \( \Gamma_2[\phi, G] \) is conveniently described in terms of the diagrams generated by the interaction terms in \( S[\phi + \varphi] \) which are of cubic and higher orders in \( \varphi \).

\[
S_{\text{int}}[\phi + \varphi] = -\frac{U}{4\mathcal{N}} \int dt_i (\varphi_{ib}(t)\varphi_i^a(t))^2 - \frac{U}{\mathcal{N}} \int dt_i \varphi_i^a(t)\phi_{ia}(t)\varphi_i^b(t)\varphi_{ib}(t).
\]

It consists of all two-particle irreducible vacuum graphs (the diagrams representing these interactions do not become disconnected by cutting two propagator lines) in the theory with propagators set equal to \( G \) and vertices determined by the interaction terms in \( S[\phi + \varphi] \).

Since physical processes correspond to vanishing sources \( J \) and \( K \), the dynamical equations of motion for the mean field and the propagators are found by using the expression (3.20) in equations (3.14) and (3.15), and setting the right hand side equal to zero. This procedure leads to the following equations:

\[
h_{ab} \partial_t \phi^b_i(t) = -J(\phi_{i+1a}(t) + \phi_{i-1a}(t)) + \frac{U}{\mathcal{N}} (\phi_{id}(t)\phi_{ia}(t) + G_{ic}^b(t, t))\phi_{ia}(t) + \frac{U}{\mathcal{N}}((G_{iia}(t, t) + G_{iiad}(t, t))\phi^d_i(t) - \frac{\delta \Gamma_2[\phi,G]}{\delta \phi^a_i(t)}),
\]

and

\[
G^{\text{ab}}_{ij}(t, t') = D_{ijab}(t, t')^{-1} - \Sigma_{ijab}(t, t'),
\]

\[
\Sigma_{ijab}(t, t') \equiv \frac{2i}{\mathcal{N}} \frac{\delta \Gamma_2[\phi,G]}{\delta G^{\text{ab}}_{ij}(t, t')}.\]

Equation (3.23) can be rewritten as a partial differential equation suitable for initial value problems by convolution with \( G \). This differential equation reads explicitly

\[
h_{ab} \partial_t G^{\text{ab}}_{ij}(t, t') = -J(G_{i+1j}^{\text{ab}}(t, t') + G_{i-1j}^{\text{ab}}(t, t')) + \frac{U}{\mathcal{N}}(\phi_{id}(t)\phi_{id}(t))G^{\text{ab}}_{ij}(t, t') + \frac{2U}{\mathcal{N}}\phi_{ia}^d(t)G^{\text{ab}}_{ij}(t, t')\phi_{ic}(t) + i \int dt'' \sum_a \phi^a_{kic}(t, t'')G^{\text{ab}}_{kj}(t'', t') + i\delta^{ab}\delta_{ij}\delta(t - t').
\]

The evolution of \( \phi^a \) and \( G^{ab} \) is determined by Eqs. (3.22) and (3.25) once \( \Gamma_2[\phi,G] \) is specified.
3.2. Perturbative expansion of $\Gamma_2(\phi, G)$ and approximation schemes

The diagrammatic expansion of $\Gamma_2$ is illustrated in the accompanying Figure where two and three-loop vacuum diagrams are shown.

The dots where four lines meet represent interaction vertices. The expression corresponding to each vacuum diagram should be multiplied by a factor $(-i)^l(i)^{s-2}$ where $l$ is the number of solid lines and $s$ the number of loops the diagram contains.

The action $\Gamma$ including the full diagrammatic series for $\Gamma_2$ gives the full dynamics. It is of course not feasible to obtain an exact expression for $\Gamma_2$ in a closed form. Various approximations for the full $2PI$ effective action can be obtained by truncating the diagrammatic expansion for $\Gamma_2$. Which approximation is most appropriate depends on the physical problem under consideration:

3.3. The standard approaches

(i) Bogoliubov (One-loop) Approximation:
   The simplest approximation consists of discarding $\Gamma_2$ altogether. This yields the so called Bogoliubov or one-loop approximation whose limitations have been extensively documented in the literature ([38, 39]).

(ii) Time-dependent Hartree-Fock-Bogoliubov (HFB) Approximation:
   A truncation of $\Gamma_2$ retaining only the first order diagram in $U$, i.e., keeping only the double-bubble diagram, Fig. a, yields equations of motion of $\phi$ and $G$ which correspond to the time dependent Hartree-Fock-Bogoliubov (HFB) approximation. This approximation violates Goldstone’s theorem, but conserves energy and particle number [40, 41, 42]). The HFB equations can also be obtained by using cumulant expansions up to the second order [43] in which all cumulants containing three or four field operators are neglected. The HFB approximation neglects multiple scattering. It can be interpreted as an expansion in terms of $Ut/J$, (where $t$ is the time of evolution) and is good for the description of short time dynamics or weak interaction strengths.

(i) Second Order expansion:
A truncation retaining diagrams of second order in $U$ containing besides the double-bubble, also the setting-sun and the basket-ball. By including the setting-sun and the Basket-ball in the approximations we are taking into account two particle scattering processes [44]. Second order terms lead to integro- differential equations which depend on the time history of the system.

(ii) Large-$N$ approximation

The $1/N$ expansion is a controlled non-perturbative approximation scheme which can be used to study non-equilibrium quantum field dynamics in the regime of strong interactions[31]. In the large $N$ approach the field is modeled by $N$ fields and the quantum field generating functional is expanded in powers of $1/N$. In this sense the method is a controlled expansion in a small parameter but unlike perturbation theory in the coupling constant, which corresponds to an expansion around the vacuum, the large $N$ expansion corresponds to an expansion of the theory about a strong quasiclassical field.

In [37] numerical solutions to the equations of motion for a moderate number of atoms and wells up to the second order in the coupling constant $U$ were given. These exact many body solutions are useful for determining the range of validity of the three types of approximations described above, namely, a) the HFB, b) the full second order and c) NLO large $N$ expansion up to second order in $U$ (the shorthands HFB, 2nd and $1/N$ respectively are used in the figure). For the effect of the total number of atoms on the dynamics Rey et al have given the numerical solutions for a double well system with fix ratio $UN/J = 4$ and three different values of $N$: $N=20, 40$ and $80$. The results are given for the evolution of the atomic population per well (Fig. 5), the condensate population per well and total condensate population (Fig. 6) and the quasi-momentum intensities (Fig. 7). (To make the comparisons easier the numerical results obtained for the three different values of $N$ are scaled by dividing them by the total number of atoms. In this way for all the cases we start with an atomic population of magnitude one in the initial populated well.)

In the exact dynamics we see that as the number of atoms is increased the damping effects occur at slower rates. This feature can be noticed in the quantum dynamics of all of the observables depicted in the Figures 5 to 7 of [37]. The decrease of the damping rates as the number of atoms is increased is not surprising because changing the number of atoms alters the quantum coherence properties of the system. As shown in reference [45], the collapse time of the condensate population is approximately given by $t_{\text{coll}} \sim \frac{t_{\text{rev}}}{\sigma}$ where $\sigma$ is the variance of the initial atomic distribution and $t_{\text{rev}}$ is the revival time which depends on the detailed spectrum for the Hamiltonian. In the kinetic energy dominated regime $t_{\text{rev}} \sim \hbar N/J$ (see [46]), thus $t_{\text{coll}} \sim \frac{N}{\sigma}$. For our initial conditions the variance is proportional to $\sigma = \sqrt{N}$ so $t_{\text{coll}} \sim \sqrt{N}$. Besides damping rates, the qualitative behavior of the exact quantum dynamics is not affected very much as the number of atoms is increased for a fixed $UN/J$.

The improvement of the $2PI$ approximations as $N$ is increased, as a result of the increase in the initial number of coherent atoms is in fact observed in the plots. Even though the problem of underdamping in the HFB approximation and the overdamping in the second order approaches are not cured, as the number of atoms is increased, we do observe a better matching with the full quantal solution. The $1/N$ expansion shows the fastest convergence, as can be more easily observed in the quasi-momentum distribution plots, Fig. 7. The better agreement of the $1/N$ expansion relies on the fact that even though the number of fields is only two in our calculations the $1/N$ expansion is an expansion about a strong quasiclassical field configuration, a feature we have explained in MQP1.
Part II: Effective Scale and Infrared Dimension

4. Effective field theory, coupling and the effective scale of a quantum system

Effective field theory is a very powerful tool in quantum field theory, and in particular it gives a new point of view about the meaning of renormalization.

Consider a quantum field theory with a characteristic energy scale $E_0$, and suppose that we are interested in physics at some lower energy scale $E$. The basic idea of effective field theory is that one can choose a cutoff energy scale $\Lambda$ at or slightly below $E_0$, and divide the fields in the path integral into low- and high- frequency parts, $\phi = \phi_L + \phi_H$, where $\phi_L(\phi_H)$ are the field modes with frequencies lower(higher) than $\Lambda$. Now integrate over the high frequency fields $\phi_H$

$$\int D\phi_H D\phi_L e^{iS(\phi_H,\phi_L)} = \int D\phi_L e^{iS(\phi_L)}$$

(4.1)

where $e^{iS_\Lambda(\phi_L)} = \int D\phi_H e^{iS(\phi_H,\phi_L)}$. The $S_\Lambda(\phi_L)$ is called the low-energy or Wilsonian effective action. It’s different from the 1PI effective action which are generated from integrating over all frequencies but keeping only 1PI graphs.

One can expand the low-energy effective action in terms of local operators which are consistent with the symmetries of the problem,

$$S_\Lambda(\phi_L) = \int d^D x \sum_i g_i \hat{Q}_i.$$  

(4.2)

In units of $\hbar = c = 1$, if an operator $\hat{O}_i$ has unit $E^{d_i}$ where $d_i$ is the dimension of operator $\hat{O}_i$ then the parameter $g_i$ has unit $E^{D-d_i}$ where $D$ is the spacetime dimension. In the case of a free scalar field in $D$ dimensions, with action

$$S = \int d^D x \left( \frac{1}{2} \partial_\mu \phi \partial^\mu \phi - \frac{1}{2} m^2 \phi^2 \right).$$

(4.3)

The action $S$ is dimensionless, so the dimension of $\phi(x)$ is determined from the kinetic term to be $[\phi] = D/2$ and the dimension of $m^2$ is $[m^2] = 2$. An operator $\hat{O}_i$ constructed from $M$ $\phi$’s and $N$ derivatives has dimension $d_i = M \frac{D-2}{2} + N$. Define dimensionless couplings $\lambda_i = \Lambda^{d_i-D} g_i$. Now for a process at energy scale $E$, one can dimensionally estimate the contribution from the $i$-th term of the effective action

$$\int d^D x g_i \hat{O}_i \sim \Lambda_i \left( \frac{E}{\Lambda} \right)^{d_i-D}.$$  

(4.4)

If $d_i > D$, this term will become less and less important at lower and lower energies, and is called an irrelevant operator. If $d_i < D$, the operator is more and more important at lower and lower energies and is called a relevant operator. If $d_i = D$ the operator is equally important at all scales and called marginal. In most cases there is only a finite number of relevant and marginal operators so the low energy physics depends only on a finite number of parameters.

The low energy physics depends on the short distance theory only through the relevant and marginal couplings, and possibly through some leading irrelevant couplings if one measures small enough effects.

It should be pointed out that the coupling strength measured will vary with energy(scale). This can easily be seen from the point of view of effective field theory. We could write the path integral as

$$\int_{k \leq \Lambda} D\phi_k e^{iS_\Lambda(\phi_k)} = \int_{k' \leq \Lambda'} D\phi_{k'} e^{iS'_\Lambda(\phi_{k'})}.$$  

(4.5)
The action $S_\Lambda(\phi_k)$ contains all momentum modes up to some maximum value $\Lambda$, whereas the new effective action $S'_\Lambda(\phi_{k'})$ contains momentum modes up to $\Lambda' < \Lambda$. If one takes $\Lambda'$ to be infinitesimally smaller than $\Lambda$, $S'_\Lambda$ will be infinitesimally different from $S_\Lambda$. This generates a differential equation, the Wilson equation

$$\frac{\partial S_\Lambda}{\partial \Lambda} = F(S_\Lambda), \quad (4.6)$$

where $F$ is some functional of the action. The Wilson equation is the renormalization group flow equation in an infinite dimensional space and it gives the change in action as a function of cutoff. Since

$$S_\Lambda(\phi) = \int d^D x \sum_i g_i \mathcal{Q}_i, \quad (4.7)$$

the Wilson equation then gives

$$\frac{\partial g_i}{\partial \Lambda} = F(\{g_i\}). \quad (4.8)$$

If one linearizes a solution of the equation, irrelevant operators will correspond to those directions with negative eigenvalues whereas the relevant operators correspond to those directions with positive eigenvalues.

5. Infrared behavior of quantum fields: finite-size / topology effects

In the above we pointed out that the infrared or large scale behavior of a system depends on many factors: a) Interaction strength: difference between a free theory (Gaussian) and an interacting theory, and for an interacting theory the Gaussian fixed point as different from non-Gaussian ones, relevant versus irrelevant fixed points; b) Coupling strength which runs with energy via the RG equation. All this show the intricacy of the notion of ‘large scale’ which we need to bear in mind in describing ‘macroscopic’ quantum behavior. In this section, we demonstrate how the infrared behavior of a quantum system can be altered qualitatively by the geometry and topology of space, or more precisely, the ‘finite size’ effect. This can be seen from analysis based on rather general arguments, e.g., on whether there exist a band structure in the spectrum of eigenvalues of the invariant operator of the order parameter field. If so there is a dimensional reduction effect where the IR behavior of the system becomes that of a lower dimension. What this conveys is that, for a quantum system near the critical point where its correlation length increases to (near) infinity its large scale or IR behavior or macroscopic features can become sensitive to the size and topology of the underlying spacetime. Cosmological phase transition is an extreme yet valid, even realistic, manifestation of this feature. A more rigorous and complete treatment of this effect is given in Hu and O’Connor [47] from which our present exposition is adapted.

Consider an $N$-component self-interacting scalar field $\Phi^a(a = 1, \cdots, N)$ on a manifold of dimension $D$, coupled to the background spacetime with curvature $R$ and coupling constant $\xi$ (conformal coupling for $\xi = 0$ and minimal coupling for $\xi = 1$) described by the action

$$S[\Phi] = \int d^D x \sqrt{g} \left[ \frac{1}{2} \Phi^a \nabla^2 \Phi^a + \frac{1}{2} M_\xi^2 \Phi^2 + \lambda \Phi^4 \right], \quad (5.1)$$

where $M_\xi^2 = m^2 + (1 - \xi) R/6$ and $\nabla^2 = -\sqrt{g} \partial_{\mu}(\sqrt{g} g^{\mu\nu} \partial_{\nu})$ is the Laplace-Beltrami operator for scalar fields. For convenience we develop the formalism here in Euclidean space version with signature $(+,+,+,+)$. We begin with a decomposition of the field $\Phi^a$ into a background field $\bar{\Phi}^a$
and a fluctuation field \( \varphi^a \), i.e., \( \Phi^a = \tilde{\varphi}^a + \varphi^a \). The background field \( \tilde{\varphi}^a \) is required to satisfy the classical equations of motion with an arbitrary external source. Such a shift eliminates the linear term in the fluctuation field, which is equivalent to performing a Legendre transform. The resultant action is

\[
S[\phi, \varphi] = S[\phi] + \int d^D x \sqrt{g} \times \left\{ \frac{1}{2} \varphi^a \left[ (\nabla^2 + M_2^2 + \frac{\lambda}{6} \tilde{\varphi}^2) \delta^{ab} + \frac{\lambda}{3} \tilde{\varphi}^a \tilde{\varphi}^b \right] \varphi^b + \frac{\lambda}{6} \tilde{\varphi}^a \varphi^a \varphi^2 + \frac{\lambda}{4!} \varphi^4 \right\}.
\] (5.2)

In the Feynman diagrammatics there are two kinds of vertices: a four point vertex proportional to \( \lambda \) and a three point vertex proportional to \( \lambda \tilde{\varphi}_{ab} (x) \).

The effective action \( \Gamma[\phi] \) is obtained by functionally integrating over the fluctuation fields:

\[
e^{-\Gamma[\phi]} = \int [d\varphi] e^{-S[\phi, \varphi]}.
\] (5.3)

The wave operator \( A^{ab} \) for the fluctuating field is given by

\[
A^{ab} = (\nabla^2 + M_1^2) \frac{\tilde{\varphi}^a \tilde{\varphi}^b}{\tilde{\varphi}^2} + (\nabla^2 + M_2^2) \left( \delta^{ab} - \frac{\tilde{\varphi}^a \tilde{\varphi}^b}{\tilde{\varphi}^2} \right)
\] (5.4)

where

\[
M_1^2 = M_2^2 + \frac{\lambda}{2} \tilde{\varphi}^2, \quad M_2^2 = M_2^2 + \frac{\lambda}{6} \tilde{\varphi}^2.
\] (5.5)

Here \( \tilde{\varphi}^a \tilde{\varphi}^b / \tilde{\varphi}^2 \) and \( \delta^{ab} - \tilde{\varphi}^a \tilde{\varphi}^b / \tilde{\varphi}^2 \) are orthogonal projectors, the former projects along the direction in the internal space picked out by \( \tilde{\varphi}^a \) and the latter projects into an \((N-1)\)-dimensional subspace orthogonal to the direction of \( \tilde{\varphi}^a \). Note that the operator \( \nabla^2 \) does not commute with the projectors unless \( \tilde{\varphi}^2 \) is a constant.

When the direction in group space picked out by \( \tilde{\varphi}^a \) does not vary from point to point around the manifold (this does not necessarily imply \( \tilde{\varphi}^2 \) is a constant) the Green’s function for \( A^{ab} \) is given by

\[
G^{ab} = G_1 \frac{\tilde{\varphi}^a \tilde{\varphi}^b}{\tilde{\varphi}^2} + G_2 \left( \delta^{ab} - \frac{\tilde{\varphi}^a \tilde{\varphi}^b}{\tilde{\varphi}^2} \right),
\] (5.6)

where the \( G_i \) \((i = 1, 2)\) are the Green’s functions for the operators \( \nabla^2 + M_i^2 \).

The one-loop correction is related to the sum of the logarithms of the determinants of the fluctuation operators. So in this case, by using the projection operators, the one-loop effective action is given by

\[
\Gamma[\phi] = S[\phi] - \frac{1}{2} \text{Tr} \ln G_1 - \frac{N - 1}{2} \text{Tr} \ln G_2
\]

\[
= S[\phi] + \frac{1}{2} \sum_i d_i \ln \lambda_{hi} + \frac{N - 1}{2} \sum_i d_i \ln \lambda_{2i},
\] (5.7)

where \( S[\phi] \) is the classical action, \( \lambda_{hi} \) and \( d_i \) are the eigenvalues and degeneracies of the operators \( \nabla^2 + M_i^2 \).

The determinant of \( A \) is formally divergent and needs to be regularized. There are a number of commonly used regularization methods. If the space is Riemannian and has sufficient symmetry so that the spectrum of the invariant operator is known explicitly, then the \( \zeta \)-function method is probably most convenient. The generalized \( \zeta \) function is defined by

\[
\zeta(\nu) = \sum_n (\mu^{-2} \lambda_n)^{-\nu},
\] (5.8)
where $\lambda_n$ are the eigenvalues of the operator $A$ on the Euclideanized metric obtained by a Wick rotation to imaginary time $\tau = it$. Here a constant mass scale $\mu$ is introduced to make the measure $d[\varphi]$ of the functional integral dimensionless. Using the regularization method of Dowker and Critchley, one can express the one-loop effective potential as

$$V^{(1)}(\bar{\phi}) = -\frac{1}{2} h(Vol)^{-1}[\zeta'(0) + \zeta(0)/\nu].$$

The ultraviolet divergence in $V^{(1)}$ can be canceled by the addition of counterterms which is not our concern here. The regularized effective action is useful for the analysis of the infrared behavior of quantum fields in a curved spacetime or spaces of nontrivial topology or finite extent, as shown below. Note, however that one-loop results are insufficient. Leading infrared contributions in higher loop terms need be included by using composite operator techniques.

We learn from the work of [47] that in spacetimes with some compact dimensions the lowest mode of the fluctuation operator has the strongest effect on the symmetry behavior of the system. When the lowest mode is massless it will give the dominant contribution to the effective action. The low energy behavior corresponds to a lower-dimensional system. In what follows we first give a formal derivation of infrared dimensional reduction by examining the result of the decoupling of the higher modes (or bands) in the functional integral for the effective action.

We will then give a physical explanation in terms of the correlation lengths and the notion of effective IR dimension (EIRD). An alternative way of seeing this problem of dimensional reduction is by spectral analysis. This is applied to direct product spaces with some compact dimensions and to spaces which can be reduced to product spaces.

5.1. Decoupling of the higher modes (or bands)

Let us examine systems where the eigenvalues of the fluctuation operator take on a band structure. By band structure we mean that the eigenvalues occur in continua with each continuum having a higher lowest eigenvalue than the previous one. This is true for fields on spacetimes with compact sections or for fields with discrete spectrum (e.g. the harmonic oscillator). The procedure is to expand the fields in terms of the band eigenfunctions and convert the functional integral over the fields to an integral over the amplitudes of the individual modes.

On a manifold with topology $R^d \times B^b$ where $B$ is compact, consider quantum fields where the fluctuation operator $A$ in (5.4) has the general form of a direct sum of operators $D$ and $B$

$$A^{ab}(x, y) = D^{ab}(x) + B^{ab}(y)$$

with coordinates $x$ on $R^d$ and $y$ on $B^b$. Assume that the eigenvalues $\omega_n$ associated with the eigenfunctions $\psi_n(y)$ of $B^{ab}$ are discrete:

$$B^{ab}\psi_n(y) = \omega^{ab}_n \psi_n(y)$$

Decomposing the field $\varphi^a(x, y)$ in terms of $\psi_n(y)$

$$\varphi^a(x, y) = \sum_n \varphi^a_n(x) \psi_n(y)$$

one obtains for the quadratic part of the action

$$\frac{1}{2} \int d^d x d^b y \varphi^a A^{ab} \varphi^b = \frac{1}{2} \int d^d x [\varphi^a_n f_{nm} D^{ab} \varphi^b_m + \omega^{ab}_n f_{nm} \varphi^a_n \varphi^b_m]$$

where $f_{nm} = \int d^b y \psi_n(y) \psi_m(y)$. When $\varphi_n$ are properly normalized $f_{nm} = \delta_{nm}$ (we will make such a choice here) the resulting theory in terms of the new fields $\varphi^a_n$ will involve massive fields.
with masses determined by the eigenvalue matrix $\lambda_n^{ab}$ of the operators $\nabla^2 + M_i$ [see (5.7)], even if the fields in terms of the old variables appeared massless. We will take the smallest eigenvalue to be given by $n = 0$ and assume that its only degeneracy is labeled by the indices $a$ and $b$. Assume also that the operator $D^{ab}$ is simply minus the Laplacian $\nabla_d^2$ on $R^d$ times $\delta^{ab}$, the $n = 0$ mode is then governed by the action whose quadratic term is

$$\frac{1}{2} \int d^d x [\varphi^a_0 (-\nabla^2_d) \delta^{ab} \varphi^b_0 + \omega^{ab}_0 \varphi^a_0 \varphi^b_0]$$

(5.14)

Thus this appears like a $d$-dimensional field with an apparent mass matrix $\omega^{ab}_0$. For the case of an $N$ component $\lambda \phi^4$ theory the action after this decomposition takes the form

$$S[\phi + \varphi] = \int d^d x \left[ \frac{1}{2} \varphi^a_n (-\nabla^2_d) \delta^{ab} + \omega^{ab}_n \right] \varphi^b_n + \frac{\lambda}{6} g_{n\ell m} \varphi^b_\ell \varphi^b_\ell \varphi^b_m + \frac{\lambda}{4!} f_{kn\ell m} \varphi^a_k \varphi^a_\ell \varphi^a_\ell \varphi^b_m,$$

(5.15)

where $g_{n\ell m} = \int d^d y \varphi^a_n \psi_\ell \psi_m$ and $f_{kn\ell m} = \int d^d y \psi_k \varphi^a_\ell \psi_m$. The effective action is now given by the functional integral

$$e^{-\Gamma[\phi]} = \int [d\varphi^a_n] e^{-S[\phi + \varphi]}.$$

(5.16)

The interesting case occurs when the lowest eigenvalue approaches zero. At low energy the Appelquist-Carazzone decoupling theorem assu...
The symmetry behavior of the system (described here by a $\lambda \phi^4$ scalar field as example) is determined by the correlation length $\Xi$ defined as the inverse of the effective mass $M_{\text{eff}}$ related to the effective potential $V_{\text{eff}}$ by (we use subscript eff to denote quantities including higher loop corrections)

$$\Xi^{-2} = \frac{\partial^2 V_{\text{eff}}}{\partial \phi^2} \bigg|_{\phi=0} = M_{\text{eff}}^2 = (\text{curvature-induced mass } M_{\text{eff}}^2 + \text{radiative corrections})$$

(5.17)

It measures the curvature of the effective potential at a minimum energy state ($\phi = 0$ for the symmetric state, or the false vacuum, $\phi = \phi_{\text{min}}$ for the broken-symmetry state or the true vacuum.) The effective mass is defined to include radiative corrections to the same order corresponding to the effective potential. (This quantity is called the generalized susceptibility function in condensed matter physics.) The critical point of a system is reached when $\Xi \to \infty$ or $M_{\text{eff}} \to 0$. In flat or open spaces or for bulk systems, the critical point can be reached without restriction from the geometry (note that in dynamical situations, exponential expansion can effectively introduce a finite size effect equivalent to event horizons, see [48]). However, in spaces with compact dimensions, the correlation length of fluctuations can only extend to infinity in the remaining non-compact dimensions, and thus the critical behavior becomes effectively equivalent to a lower $d$-dimensional system. One can also think of $\Xi$ as the Compton wavelength $\Lambda = 2\pi/M_{\text{eff}}$ of a system of quasi-particles with effective mass $M_{\text{eff}}$. Any fine structure of the background spacetime with scale $L$ is relevant only if $\Lambda \leq L$. Thus when $\Lambda$ is small or $\eta \ll 1$, (far away from critical point, at higher energy, with higher mode contributions) it sees the details of a spacetime of full dimensionality. At this wavelength, the apparent size of the universe is large in both compact and non-compact dimensions. When $\Lambda \to \infty$ or $\eta \gg 1$ (near critical point, IR limit, lowest mode dominant) finer features in the compact dimensions will not be important. The apparent ‘size’ of the universe will be measured by the non-compact directions and the EIRD is measured by the number of non-compact dimensions. The value of $\eta$ getting very large is an indication of when dimensional reduction can take place.

Notice that in flat space ($R^d$) critical phenomena the effective potential $V_{\text{eff}}$ (free energy density) depends on the coupling constants of fields which run with energy and temperature. In curved-space coupling parameters run also with curvature or the scale length of the space. This makes the concept of EIRD even more interesting, as there is now an interplay between $\Xi$ and $L$, and $\eta$ can either decrease or increase with curvature. For example, for $\lambda \phi^4$ fields in the Einstein universe near the symmetric state $\phi = 0$, the EIRD is equal to 1 but near the global minimum of broken symmetry state $\phi = \phi_{\text{min}}$ is equal to 4. Near the symmetric state, $\eta \gg 1$ signifies reduction of EIRD to one. This is consistent with the theorem of Hohenberg, Mermin and Wagner (for statistical mechanics on a lattice) and Coleman (for continuum field theory) which states that in dimensions less than or equal to two, the infrared divergence of the scalar field is so severe that there could be no possibility of spontaneous symmetry breaking: the only vacuum expectation value for $\phi$ allowed is zero. Away from the region $\phi \approx 0$ the one-dimensional behavior no longer prevails. Indeed a global minimum of the effective potential exists at $\phi_{\text{min}}$. Near $\phi_{\text{min}}$, $\eta \ll 1$ and decreases with curvature. Thus the apparent size of the universe near the global minimum actually increases with increasing curvature. There is therefore no dimensional reduction and the system has a full 4-dimensional IR behavior. A transition to the asymmetric ground state is not precluded as symmetry breaking via tunneling is in principle possible. The complete picture extending from $\phi = 0$ to $\phi = \phi_{\text{min}}$ is a combination of one dimensional and four dimensional infrared behavior. Similar arguments can be applied to other spacetimes or field theories. Using this notion one can understand, for example, why it is often said that at high temperatures (small radius limit of $S^1$) the finite temperature theory becomes an effective three-dimensional theory.
5.3. Dimensional reduction: an eigenvalue analysis

In the above we have introduced the notion of effective infrared -dimension and suggested the ratio of the correlation length $\Xi$ to the geometric scale of spacetime $L$ as a measure of the conditions for the system to behave effectively as in a reduced dimension in the infrared regime. We suggested that for product spaces $R^d \times B^e$ with some noncompact dimension $d$, the effective IR dimension is usually just $d$. We will now verify this assertion by analyzing the spectrum of the fluctuation operators in these spacetimes directly.

Consider for simplicity direct product spaces. Examples are a) physical cosmological spacetimes with topology $R^1 (time) \times S^3 (or \ R^3, H^3, T^3)$, b) Kaluza-Klein cosmology with $M^4 \times S^7$ (or other internal space), c) finite-temperature (imaginary-time) theory has $R^3 \times S^4$. Take a simple example $S^2 \times S^1$ for illustration. (This could be the spatial geometry of a “handled” Gowdy universe.) Similar reasoning can be extended to a wide range of product spaces.

From (5.7) the wave operator $A$ governing the fluctuation fields in the large $N$ limit is $A \equiv \nabla^2 + M_2^2$, where $\nabla^2$ is the Laplace-Beltrami operator on a general curved spacetime. For $S^2 \times S^1$ with radii $a_2$ and $a_1$ respectively, $\nabla^2$ is a sum of the total angular momentum operator $L$ on $S^2$, and $L_z$ on $S^1$:

$$\nabla^2 = \frac{L^2}{a_2^2} + \frac{L_z^2}{a_1^2}$$

(5.18)

The eigenfunction is a product of $Y_{\ell m}(\theta, \phi)e^{imz}$ belonging to the eigenvalues

$$\kappa_n^2 = \frac{\ell(\ell + 1)}{a_2^2} + \frac{n^2}{a_1^2},$$

(5.19)

where $n = (\ell, m, n), \ell = 0, \cdots, \infty, m = -\ell$ to $\ell, n = 0 \pm 1, \cdots$. The eigenvalues of $A$ are then $\lambda_n = \kappa_n^2 + M_2^2$. In the infrared region, we are interested in the contribution of the lowest eigenvalue (zero mode) to the effective potential. We will consider the two limiting cases of 1) $S^2 \times R^1$ and 2) $R^2 \times S^1$ obtained when $a_1$ and $a_2 \to \infty$ respectively, and show that the EIRD is equal to 1 and 2 respectively.

The effective potential $V_{eff}$ can be constructed from the $\zeta$ function $\zeta(0)$ and its derivative $\zeta'(0)$ by (5.8).

$$\zeta(\nu) = \sum_n (\mu^{-2} \lambda_n)^{-\nu} = \mu^{2\nu} \sum_{\ell, n} (2\ell + 1)\frac{\ell(\ell + 1)}{a_2^2} + \frac{n^2}{a_1^2} + M_2^2)^{-\nu}$$

(5.20)

where $2\ell + 1$ is the degeneracy of $m$. In the limiting cases the summation over discrete quantum numbers will be replaced by integrals of the form $\int k^{D-1}dk$. We will derive the dimensionality of the reduced system by finding $D$. Thus in case a) $a_1 \to \infty$, the lowest eigenvalues belong to the band $\ell = 0$ as $a_1 \to \infty$, $k = n/a_1$ assumes continuous value and $\zeta(\nu)$ becomes

$$\zeta(\nu) \sim a_1 \int_0^\infty dk (k^2 + M_2^2)^{-\nu}, a_1 \to \infty, \ M_2a_1 << 1$$

(5.21)

This is a one-dimensional integral. In case (b) $a_2 \to \infty$, the lowest eigenvalues come from the lowest band $n=0$ given by the first term in (5.19), where $\ell$ assumes continuous values as $a_2 \to \infty$. Now call $k = \ell/a_2$, $\zeta(\nu)$ becomes ($n = 0$)

$$\zeta(\nu) \sim 2a_2 \int_0^\infty dkk (k^2 + M_2^2)^{-\nu}, a_2 \to \infty, \ M_2a_1 << 1$$

(5.22)
The extra factor of $k$ comes from the degeneracy of $m$ for nonzero $\ell$. The integral is thus 2-dimensional, as expected.

Physically, the two limiting cases may represent two different nonzero $\ell$. The integral is thus 2-dimensional, as expected.

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We then explore the issue of how MQP can be measured or understood from the behavior of quantum correlations which exist in a quantum system and how the interaction strengths change with energy or scale, under ordinary situations and when the system is near its critical point. Noting that quantum correlation is further related to quantum fluctuations and quantum coherence, we used three examples from known results to illustrate these aspects: a) the existence of an H-theorem at the next-to-leading order large \( N \) expansion for an \( O(N) \) quantum mechanical model, with entropy generation signifying the emergence of macroscopic (thermodynamic) properties. b) the nonequilibrium dynamics of \( N \) atoms in an optical lattice under the large \( N \) (field components), \( 2PI \) and second order perturbative expansions, illustrating how \( N \) and \( N \) enter in these three aspects of quantum correlations, fluctuations and coupling strength. c) the infrared behavior of an interacting quantum system, we discuss the conditions where dimensional reduction shows up. The effective IR dimension is determined by the spectrum of the Laplacian of an interacting quantum system, in particular, whether a lowest eigenmode exists. On the same theme, we also discuss how the effective field theory concept bears on MQP: the running of the coupling parameters with energy or scale imparts a dynamical-dependent and an interaction-sensitive definition of ‘macroscopia’.

Quantum entanglement will be the theme of our third essay where we intend to focus on how this uniquely quantum feature shows up in quantum many-body systems and inquire to what extent it quantifies a quantum system as micro, meso versus macro. In this enquiry we will necessarily also touch on related subjects such as ‘small systems’ and emergent concepts in the new field known as quantum thermodynamics (ostensibly this is not the thermodynamics of quantum systems, treated in QSM textbooks, where quantum pertains to spin-statistics of particles). We can see why this is relevant to MQP in a simplified way: thermodynamics is a macroscopic theory valid when the volume \( V \) and number of constituents \( N \) both go to infinity while their ratio \( N/V \) remains constant. Quantum dynamics is used for the description of micro-objects or small systems. As one increases the ‘size’ (number and volume, say) of a quantum system, in what parameter ranges will a thermodynamic description of such a many body quantum system become useful, at least begin to make sense? These inquires into the meaning of MQP will unavoidably lead us to confront some foundational issues of quantum and statistical mechanics.

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