Quantum disentangled liquids

Tarun Grover\textsuperscript{1} and Matthew P A Fisher\textsuperscript{2}

\textsuperscript{1} Kavli Institute for Theoretical Physics, University of California, Santa Barbara, CA 93106, USA
\textsuperscript{2} Department of Physics, University of California, Santa Barbara, CA 93106, USA

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Abstract. We propose and explore a new finite temperature phase of translationally invariant multi-component liquids which we call a ‘Quantum Disentangled Liquid’ (QDL) phase. We contemplate the possibility that in fluids consisting of two (or more) species of indistinguishable quantum particles with a large mass ratio, the light particles might ‘localize’ on the heavy particles. We give a precise, formal definition of this QDL phase in terms of the finite energy density many-particle wavefunctions. While the heavy particles are fully thermalized, for a typical fixed configuration of the heavy particles, the entanglement entropy of the light particles satisfies an area law; this implies that the light particles have not thermalized. Equivalently, but more intuitively, if the positions of all the heavy particles are measured, the projected wavefunction for the unmeasured light particles has as an area law entanglement entropy. Thus, in a QDL phase, thermal equilibration is incomplete, and the canonical assumptions of statistical mechanics are not fully operative.

The definition of the QDL phase for heavy/light particles can be readily generalized to other cases with two (or more) conserved currents, such as spin/charge in a system of spin-1/2 fermions (as in a Hubbard model). Indeed, we argue that the finite energy-density eigenstates of the $t$–$J$ model will generically be in such a spin/charge QDL, although the fate of the QDL in the large $U$ Hubbard model is uncertain. We explore the possibility of QDL in water, with the light proton degrees of freedom becoming ‘localized’ on the oxygen ions. While we do not presently know whether a local, generic Hamiltonian can have eigenstates of the QDL form, if not, then the non-thermal behavior discussed here will exist as an interesting crossover phenomena at a time scale that diverges as the ratio of the mass of the heavy to the light particles also diverges.
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1. Introduction

Conventional wisdom holds that in ordinary liquids at finite temperature, local thermal equilibration is established on scales long compared to inelastic scattering lengths and times, and classical hydrodynamics is operative. Quantum mechanics, which plays a role on short scales and in setting the magnitudes of the crossover scales, can be safely ignored in describing qualitative behavior in the hydrodynamic regime. For example, classical statistical mechanics should be sufficient in capturing the equilibrium and linear response behavior of ordinary room temperature liquids, e.g. water, on long length and time scales.

In this paper we question these assumptions. In particular, we propose and explore a qualitative role that quantum effects might play in determining the macroscopic behavior of finite temperature liquids. We focus on two-component (or more general multi-component) liquids in which there are two species of indistinguishable quantum particles with a large mass ratio, for example the ions and electrons that constitute the atoms in an atomic or molecular liquid (such as liquid helium), or two different ions with large mass ratio in a molecular fluid, such as oxygen and hydrogen ions in water.

To illustrate the general class of questions, consider the properties of helium-3 fluid at temperatures above the superfluid transition (of several millikelvin). For temperatures below the Fermi energy (roughly 5 K) He-3 is well described by Fermi liquid theory, while at higher temperatures a description as a non-degenerate—essentially classical—fluid would be appropriate. At much higher temperatures, on the electron volt scale, as electrons are highly excited, significant ionization is expected, and a picture in terms of a plasma of electrons and He-3 nucleons would be appropriate. At much higher temperatures still, the nucleons would start breaking into protons and neutrons, and with continued heating one should eventually enter into a regime described as a quark–gluon and electron plasma. The canonical wisdom maintains that all such regimes are in the same phase, separated by crossovers rather than phase transitions\(^3\). But might this conventional wisdom be incomplete or even incorrect?

In this paper we propose and explore a potentially new phase of finite temperature two-component quantum fluids in which the constituents have a large mass ratio. A description of this new phase, which we will call a ‘Quantum Disentangled Liquid’ (QDL), requires treating both species of particles quantum mechanically. In the QDL phase the light particles are, in a sense to be made more precise below, ‘localized’ on the heavy particles, and due to quantum effects, do not thermalize. Due to the lack of full thermalization, a correct description requires a ‘microcanonical’ formulation wherein the fluid, confined to a very large box, is presumed to be in a pure quantum state, perhaps an eigenstate, with a finite energy density above the ground state.

In section 2 we revisit the entanglement properties of single-component finite ‘temperature’ fluids of indistinguishable quantum particles. We focus on the nature of highly excited pure quantum states—with a finite energy density above the ground state. In a fully thermalized phase, for such states the entanglement entropy, for a spatial bi-partitioning, will satisfy a volume law. As we will argue, this high degree of spatial entanglement is encoded in the ‘sign-structure’ of these finite energy-density states. Using this insight,

\(^3\) There could, in principle, always exist first order transitions that separate various regimes that are otherwise smoothly connected (for example, liquid-to-gas transition in a conventional fluid).
we construct a volume law wavefunction for bosons, which can be shown to possess a maximal entanglement entropy density corresponding to an infinite ‘temperature’.

Section 3 is devoted to a discussion of finite ‘temperature’ two-component quantum fluids, with one species much more massive than the other. We give a precise mathematical definition of the QDL phase which involves a diagnostic to be performed on the many-particle eigenstate. Roughly, when the positions of the heavy particles are fixed, the light particles are in a ‘disentangled’ area law phase. Equivalently, if the positions of the heavy particles are measured, the wavefunction of the unmeasured light particles will have an area law entanglement entropy. For a lattice gas model of two boson species, we construct an explicit QDL wavefunction which can be shown to satisfy all of the defining criterion of the QDL state.

The essence of the QDL phase can be conveyed by considering a wavefunction for a two-component fluid of quantum particles which takes a Born–Oppenheimer [1] type form. To be explicit, consider a system of heavy, mass $M$, bosons and light, mass $m$, fermions, with $m \ll M$. A Born–Oppenheimer wavefunction, expressed in terms of the coordinates of the heavy and light particles, $R_\alpha$ and $r_\beta$, respectively, takes the form,

$$\Psi(R_\alpha, r_\beta) = \psi(R_\alpha) \det[\Phi_R(r'_\beta)],$$

where the second term is a Slater determinant for the light fermions constructed from single fermion orbitals, denoted, $\Phi_R(r'_\beta)$, which are eigenstates of the effective Born–Oppenheimer Hamiltonian,

$$H = \sum_\beta \left[ \frac{p^2_\beta}{2m} + V_R(r_\beta) \right],$$

with effective potential,

$$V_R(x) = \sum_\alpha U^{Mm}(x - R_\alpha).$$

Here $U^{Mm}$ is the interaction potential between the light and heavy particles and the heavy particle coordinates $\{R_\alpha\}$ are c-numbers.

A caricature wavefunction for the conjectured QDL phase can be constructed by taking the first piece of the wavefunction in equation (1), $\psi(\{R_\alpha\})$, to correspond to a finite-energy density fluid phase. For the heavy bosons, a typical finite energy density state for an isotropic liquid can be constructed from a permanent of plane waves, $\psi \sim \text{per}[e^{i\hat{k}_\alpha \cdot R_\alpha'}]$, with wavevectors chosen with random directions, $\hat{k}_\alpha$, and with random amplitudes taken from some appropriate distribution, for example a Boltzmann distribution, $p_k \sim \exp(-\epsilon_k/\tau)$ with $\epsilon_k = \hbar^2 k^2/2M$ and $\tau$ corresponding to an ‘effective temperature’. A simple wavefunction which describes the proposed QDL phase of two-component quantum particles should thus takes the form,

$$\Psi_{\text{QDL}}(\{R_\alpha\}, r_\beta) = S(\text{per}[e^{i\hat{k}_\alpha \cdot R_\alpha'}] \det[\Phi_R(r'_\beta)]),$$

where $S$ denotes symmetrization over the set of coordinates $\{R_\alpha\}$. Due to the random wavevectors, the typical positions of the heavy particles will themselves be in an essentially random configuration, appropriate to a finite energy-density fluid phase. As a result, the
effective potential $V_R(x)$ in the Born–Oppenheimer Hamiltonian for the light fermions will be an essentially random potential. And one expects that for strong inter-species interaction energy $U^{Mm}$ the low energy single particle eigenstates will be spatially localized [2]. The Slater determinant wavefunction thus describes a localized phase of fermions. One can imagine generalizing the above Born–Oppenheimer wavefunction to incorporate (weak) inter-fermion interactions, replacing the Slater determinant by an interacting wavefunction for a ‘many-body-localized’ (MBL) phase of fermions, as introduced in [3–5]. In either case, under a spatial bi-partitioning, into regions $A$ and $B$ with linear dimension $L$, this localized eigenstates of fermions will have an area law entanglement entropy, $S_A \sim L^{d-1}$.

In contrast, the permanent wavefunction for the heavy particles $\psi \sim \text{per}[e^{ik_{\alpha} \cdot R_{\alpha}^\prime}]$, being built from (random) plane waves is expected to have a volume law entanglement entropy, $S \sim L^d$, due to the essentially random sign structure upon varying the positions of the particles (as we shall discuss in considerable detail in section 2). For a given fixed typical configuration of the coordinates of the light fermions, say $\{\tilde{r}_\beta\}$, the Slater determinant will have a magnitude and sign that varies rapidly when the positions of the heavy particles are changed. But since the Slater determinant is in any event multiplied by a heavy particle wavefunction, $\psi(\{R_{\alpha}\})$, with an essentially random sign structure, the full QDL wavefunction $\Psi_{QDL}(\{R_{\alpha}\}, \{\tilde{r}_\beta\})$, is still expected to have a volume law entanglement entropy. These are the two primary properties of a QDL wavefunction—when the heavy particles are fixed in a typical configuration the light particles are in an area law phase, while, with the light particle coordinates fixed, the heavy particles have a volume law entanglement entropy. In essence, the heavy particles have thermalized, but the light particles, ‘localized’ by the heavy particles, have not.

Going beyond wavefunctions, the generic stability of the QDL phase in local Hamiltonians of interacting particles, requires, at the very least, the generic stability of the MBL [3–5] state. While an Anderson insulator of non-interacting fermions in a random potential can exist at finite energy densities—provided that all the single particle states entering the Slater determinant are localized—establishing the stability of the MBL, a finite-energy density state of interacting electrons, is comparatively much more non-trivial and recent numerical progress [6–24] has corroborated the original picture presented in [4,5]. Returning to QDL, even the stability of the interacting MBL to arbitrary perturbations does not imply the stability of the QDL phase, which is much more tricky. As illustrated in the Born–Oppenheimer discussion, if the heavy particle mass is taken to infinity, their positions will act as a quenched random potential for the light quantum particles which, if in an area law MBL state at finite energy density, will not thermalize. For very large but finite mass ratio, $M/m$, when the heavy particles become quantum dynamical and are assumed to thermalize, there are two possibilities for the typical finite-energy density eigenstates of the two-component quantum system: (i) The light particles also thermalize, among themselves, at the same ‘temperature’ as the heavy particles—this is the conventional implicit assumption underlying the hypothesis of quantum statistical mechanics, (ii) The light particles do not thermalize—this is the QDL phase. While the wavefunction of the QDL phase has a well defined and precise definition, as detailed in section 3, it might be that it simply is not possible for a (generic) local Hamiltonian to have eigenstates of the QDL form. If this is the case, the $M/m \to \infty$ limit would be singular, and the non-thermal behavior of the light particles could only occur at strictly $M = \infty$. Presumably, in this case, one could define a typical thermal equilibration time for the light particles, say starting from the MBL...
state, and this time would diverge as $M \to \infty$. On times scales short compared to this crossover time the QDL physics would be operative, but would break down at longer times. Whether or not this equilibration time can remain infinite at finite $M$—i.e. whether the QDL phase can exist as a ‘true phase’—is not something that we can presently establish.

Section 4.1 will be devoted to a more general discussion of the physics of the QDL phase, drawing further analogies with the MBL phase of quantum particles in a random potential. In particular, the (linear response) conductivity in an MBL phase with a conserved particle number is expected to vanish, even at finite ‘temperature’—the particles cannot propagate. Analogously, in a QDL phase where the two species carry opposite ‘electric’ charge, one expects that the (linear response) electrical conductivity should vanish identically. We will also discuss hypothetical experiments in QDL which can detect lack of thermalization of the light particles.

In section 5.1 we will briefly discuss the prospects of establishing whether a QDL phase can exist in a local Hamiltonian for a two-fluid system, and the feasibility of verifying a QDL phase in numerics. In section 5.2 we will define a natural generalization of the QDL phase, appropriate in systems with a conserved spin and charge, such as a Hubbard model. Here, the role of the heavy/light particles is replaced by the spin/charge. Section 5.3 will be devoted to a brief discussion of the prospects that a large $U$ Hubbard model might be in such a spin/charge QDL phase, informed by the generic existence of a QDL in the $t$–$J$ model limit. In section 5.4 we will explore the possibility that pure water might be in a QDL phase, where the protons are ‘localized’ by the heavier oxygen ions. This possibility is informed by the remarkable phenomenology of pure water—its incredibly high resistivity, and the intricate hydrogen bonding which strongly correlates the motion of the oxygen ions with the protons. Finally, in section 5.5 we briefly remark on the implications of the QDL physics in the broader context of (multi-species) ‘finite-temperature’ quantum field theories with both heavy (semi-classical) and light (strongly quantum) fields present.

2. Quantum entanglement in finite ‘temperature’ fluids

Consider a single component translationally invariant fluid of indistinguishable quantum particles (fermions or bosons). For now we will ignore any internal ‘spin’ quantum numbers of the particles. We imagine that the dynamics of the fluid is controlled by an underlying quantum Hamiltonian. The precise nature of the Hamiltonian will not be critical, but we can think of non-relativistic particles moving in the continuum interacting with a translationally invariant (say, two-body) interaction, see below. The quantum particles are assumed to be isolated from any thermal bath, for example confined in a box. Our primary focus will be on the nature of eigenstates, with a finite energy-density above the ground state. The eigenstate thermalization hypothesis [25–28] posits that a large sub-region (the ‘system’) will behave thermally—as if at finite temperature—if one traces out the degrees of freedom outside this region (the ‘environment’). We will review and discuss the entanglement properties of pure states in such a system.

2.1. Operator and wavefunction representations

Since we will be discussing such finite temperature quantum fluids from several different angles, it will prove convenient to introduce and define several different
Quantum disentangled liquids—employing first and second quantized operators and states/wavefunctions in the continuum, as well as a lattice gas model.

2.1.1. Second quantization in the continuum. We will denote general particle field operators by, \( \hat{d}_x, \hat{d}^\dagger_x \) which satisfy canonical (anti-)commutation relations, \( [\hat{d}_x, \hat{d}^\dagger_{x'}]_\pm = \delta(x - x') \), with \( \pm \) sign for fermions/bosons respectively. When we wish to specialize to fermions or bosons we will replace the operators, \( \hat{d}_x \) by the more familiar notations, \( \hat{c}_x \) for fermion operators and \( \hat{b}_x \) for boson operators. In terms of the field operators a representative fluid Hamiltonian is given by,

\[
\hat{H} = \int_x \hat{d}_x^\dagger (-\nabla_x^2/2m) \hat{d}_x + \int_{x,x'} U(|x - x'|) \hat{n}_x \hat{n}_{x'},
\]

where we have defined a particle density operator, \( \hat{n}_x = \hat{d}_x^\dagger \hat{d}_x \). Here the particles have mass \( m \), and interact via a two-body interaction \( U(|x|) \). While we are primarily interested in three-dimensional fluids, we do not need to specify the dimensionality here.

For bosons the ground state of this model will likely be either a boson superfluid or a boson crystal. And the fermion ground state could be a Fermi liquid, a paired superfluid or a fermion crystal. But we are not interested in ground states. We will focus on highly excited states at energy densities well above any condensation or crystallization temperatures.

As mentioned above, we employ a microcanonical ensemble in which the fluid is confined within a box of volume \( V \), possibly with periodic boundary conditions, and isolated from any thermal bath. We are interested in eigenstates, with a finite energy density above the ground state. Since our model is time reversal invariant we can take the exact eigenstates to be real.

2.1.2. Particle wavefunctions in the continuum. To discuss first quantized wavefunctions expressed in terms of particle coordinates, it is convenient to define a complete set of orthogonal position kets, \( |\{r_\alpha\}\rangle = \prod_{i=1}^{N_m} \hat{d}_i^\dagger \text{vac} \rangle \) where \( \alpha = 1, 2, \ldots, N_m \) labels the particles, with \( N_m \) denoting the total number of (mass \( m \)) particles. For fermions, these kets are normalized, while the normalization for the boson states depends on how many bosons occupy the same spatial point. To dispense with these unimportant yet distracting subtleties, we assume the bosons have a hard core, with radius \( a \) setting the size of the particles. Associated with a general state vector \( |\Psi\rangle \), is a position space wavefunction \( \psi(\{r_\alpha\}) = \langle \{r_\alpha\}|\Psi\rangle \) which is symmetric/antisymmetric under exchange of any two particles, for bosons/fermions.

2.1.3. Quantum lattice gas. At times below it will be convenient to place the particles on a lattice. We define a lattice gas model in terms of creation/annihilation operators \( \hat{d}_j^\dagger, \hat{d}_j \), where the subscript \( j = 1, 2, \ldots, N \) labels the lattice sites. We assume the lattice Hamiltonian has a 'canonical hopping term and local interactions,

\[
\hat{H} = -\sum_{ij} t_{ij}(\hat{d}_i^\dagger \hat{d}_j + h.c.) + \sum_{ij} U_{ij} \hat{n}_i \hat{n}_j,
\]

where both \( t_{ij} \geq 0 \) and \( U_{ij} \) are local. Here, \( \hat{n}_j = \hat{d}_j^\dagger \hat{d}_j \) is the on-site density operator—the hard core condition implies \( \hat{n}_j \) has eigenvalues, \( n_j = 0, 1 \). The model is taken to respect the point and space group symmetries of the lattice, especially translations.

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It is helpful to introduce position-space occupation number basis states, \( \{|n_j\rangle = \prod_{j=1}^{N} [\hat{d}_j^\dagger]^{n_j} |\text{vac}\rangle \) where \( n_j = 0, 1 \) denotes the absence/presence of a particle on site \( j \). For fermions one must choose some ordering for labeling the sites of the lattice—we do not specify this unless it is necessary. Associated with a general state vector \( |\Psi\rangle \) we can define a wavefunction in the occupation number basis, \( \psi(\{n_j\}) = \langle \{n_j\} | \Psi \rangle \).

### 2.2. Entanglement entropy for spatial bi-partitioning

Throughout we will assume the system is in a pure quantum state with density matrix, \( \hat{\rho} = |\Psi\rangle \langle \Psi| \). The density matrix can be either expressed in terms of the particle-position basis wavefunctions,

\[
\rho(\{r_\alpha\}, \{r_\alpha'\}) = \psi(\{r_\alpha\})\psi^*(\{r_\alpha'\}),
\]

(7)

or the particle-occupation basis wavefunctions,

\[
\rho(\{n_j\}, \{n'_j\}) = \psi(\{n_j\})\psi^*(\{n'_j\}).
\]

(8)

Consider a simple bi-partitioning into two spatial regions, denoted \( A \) and \( B \) respectively. In the continuum we denote the volume of region \( A \) as \( V_A = L_A^d \), the ‘system’, and of region \( B \) as \( V_B \), the “environment”, with \( V_A + V_B = V \). On the lattice \( N_A \) and \( N_B \) denote the number of sites in the two regions, with \( N_A + N_B = N \).

The reduced density matrix for sub-region \( A \) is obtained as usual by tracing out the particles in region \( B \),

\[
\hat{\rho}_A = \text{Tr}_B[\hat{\rho}].
\]

(9)

And the (Renyi) entanglement entropies follow,

\[
S_{A,\alpha} = \frac{1}{1-\alpha} \ln \text{Tr}_A[(\hat{\rho}_A)^\alpha],
\]

(10)

with \( S_{A}^{\text{VN}} = \lim_{\alpha \to 1} S_{A,\alpha} \) the von Neumann entropy.

### 2.3. Area and volume laws

Numerous studies have shown that the ground state of boson systems described by the representative Hamiltonians above will have area law entanglement entropy [29–31], \( S_A^{(0)} \sim L_A^{d-1} \). Indeed, both the boson superfluid and boson crystal ground states exhibit an area law, and this is believed to be a general property of (unfrustrated) boson ground states. A paired fermion superfluid or fermion crystal ground state will also have an area law, while the entanglement entropy of a Fermi liquid ground state has a multiplicative log correction to the area law [32,33], \( S_{A}^{\text{FL}} \sim L_A^{d-1} \ln L_A \).

The entanglement entropy for eigenstates with a finite energy density above the ground state are expected to be qualitatively different. These eigenstates, which offer a microcanonical description of a finite ‘temperature’ fluid, are generally assumed to thermalize (although see section 3). Indeed, the eigenstate thermalization hypothesis [25–28] maintains that when \( V_A \ll V_B \), the reduced density matrix \( \hat{\rho}_A \) is the same as a thermal density matrix for the sub-system \( A \) at a temperature for which the thermodynamic internal energy equals the energy per unit volume of the pure state.

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That is, the large sub-system $B$ can be viewed as a thermal bath which effectively causes thermalization of the much smaller ‘system in volume $A$. In particular, equilibrium statistical mechanics employing the reduced density matrix, $\rho_A$ can be used to compute the various thermodynamic properties of the sub-system $A$.

A consequence of the eigenstate thermalization hypothesis is that the von Neumann entanglement entropy satisfies a volume law, $S_{V}^{VN} = sV_A$, and that $s$ can be equated with the thermal entropy density in the sub-system $A$. Similarly, the Renyi entropies also satisfy a volume law scaling, $S_{A,\alpha} \sim V_A$.

The eigenstate thermalization hypothesis is tantamount to the assumption that statistical mechanics is valid in a large quantum system; a pure quantum state has sub-systems which are correctly described using conventional statistical mechanics in the canonical ensemble wherein the sub-system has thermalized with an appropriate heat bath at some temperature. (In section 3 we will introduce a new finite temperature phase of multi-component quantum particles which only partially thermalizes.)

While the assumed volume law behavior for finite-energy density states is most plausible, it would be desirable to have explicit volume law wavefunctions in order to develop a physical picture for (quantum) thermalization. In the next subsection we construct simple wavefunctions that exhibit area and volume laws.

2.4. Wavefunctions for area and volume law entanglement

For simplicity we focus now on boson wavefunctions. For the lattice gas model, the most general state vector can be expressed as,

$$|\Psi\rangle = \sum_{\{n_j\}} \psi(\{n_j\})|\{n_j\}\rangle. \quad (11)$$

Since the wavefunction amplitudes can be taken as real, they can be decomposed as,

$$\psi(\{n_j\}) = |\psi(\{n_j\})| \text{sgn}(\{n_j\}), \quad (12)$$

where $\text{sgn}(\{n_j\}) = \pm 1$.

2.4.1. Boson ground state wavefunctions: area law. The simplicity of boson ground states, $\psi_0$, (of the representative Hamiltonians above) is that their wavefunctions are nodeless, with $\text{sgn}_0(\{n_j\}) = +1$ for all $2^N$ configurations. The wavefunctions in terms of particle coordinates are likewise nodeless, $\psi_0(\{r_\alpha\}) \geq 0$ for any particle configuration.

The entanglement entropy of such non-negative boson ground states will generically follow the aforementioned area law. To illustrate this, consider a state that is expressed an equal weight superposition over all occupation-number basis states,

$$\psi_0(\{n_j\}) = \frac{1}{\sqrt{2^N}}. \quad (13)$$

This state encodes large density fluctuations of the particles and should describe a boson superfluid. While it has been established that boson superfluid states have an area rather than volume law entanglement [34, 35], this can be explicitly verified for the above equal
amplitude wavefunction, which can be re-expressed as a direct product state,

$$\psi_0(\{n_j\}) = \prod_{j=1}^{\mathcal{N}} \frac{1}{\sqrt{2}} [\delta_{n_j,0} + \delta_{n_j,1}].$$

For this product state the entanglement entropy actually vanishes identically, $S_{A,\alpha} = 0$. Slight modifications of this superfluid wavefunction would lead to small corrections giving an area law [34,35], $S_{A,\alpha} \sim L_A^{d-1}$.

### 2.4.2. Volume law wavefunctions.

We now wish to construct wavefunctions for bosons which exhibit a volume law entanglement entropy, which are hopefully representative of typical finite energy density eigenstates in such systems. While the ground state of the boson system is nodeless, excited states necessarily have nodes, with positive and negative wavefunction amplitudes. And very highly excited eigenstates—with a finite energy density above the ground state—should be riddled with nodes, likely exhibiting an intricate and rich sign structure.

Intuition for such sign structure intricacies can be gained by considering particle-coordinate eigen wavefunctions for the ideal Bose gas, which are expressed as permanents of plane wave states,

$$\psi_{\text{IBG}}(\{r_\alpha\}) = \text{per}[M],$$

with matrix,

$$M_{\alpha\beta} = \frac{1}{\sqrt{V}} e^{ik_\alpha \cdot r_\beta},$$

with $k_\alpha$ a set of wavevectors with $\alpha = 1, 2, \ldots, \mathcal{N}$. While the ground state Bose condensate has all $k_\alpha = 0$, one expects that a typical finite energy density eigenstate would correspond to choosing the wavevectors $k_\alpha$ to be essentially random, with directions $\hat{k}$ equally weighted over the unit sphere, and with amplitudes chosen from an appropriate distribution function, such as $p_k \sim \exp(-\epsilon_k/T)$ with kinetic energy $\epsilon_k = \hbar^2 k^2/2m$, and $T$ an average kinetic energy. Such a finite energy wavefunction should correspond to an effective temperature of order $k_B T_{\text{eff}} \sim \tau$. We note that the wavefunction $\psi_{\text{IBG}}$ can be made real by taking the wavevectors to come in $\pm k$ pairs.

It is most plausible that permanent wavefunctions, $\psi_{\text{IBG}}(\{r_\alpha\})$, constructed from plane waves with random wavevectors, should exhibit a very complicated, and essentially random sign structure, as the particle positions are moved about.

Based on this reasoning, one would anticipate that the finite energy density eigenstates for the Bose lattice gas, have an occupation-number wavefunction representation with essentially random signs. To illustrate this we will explore a wavefunction for lattice bosons which consists of an equal amplitude superposition of occupation basis states, but with randomly chosen signs:

$$\psi(\{n_j\}) = \frac{1}{\sqrt{2^\mathcal{N}}} \text{sgn}(\{n_j\}),$$

with the $\text{sgn}$ function chosen randomly $\text{sgn}(\{n_j\}) = \pm 1$, independently for each of the $2^\mathcal{N}$ configurations $\{n_j = 0, 1\}$. 

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A short calculation (see appendix A), shows that the second Renyi entropy $S_{A,2}$ corresponding to the wavefunction in equation (17) for a spatial bi-partitioning of the system into regions $A$ and $B$ is given by

$$S_{A,2} = - \log \left( 2^{-N_A} + 2^{-N_B} - 2^{-(N_A + N_B)} \right),$$

$$\approx \ln(2) N_A$$

where in the second line, we have assumed that $N_A \ll N_B$ (‘system’ much smaller than ‘environment’). This confirms that the maximally random-sign wavefunction exhibits a volume law entanglement entropy for the von-Neumann entropy $S^\text{vN}_A$ as well (recall the inequality $S^\text{vN}_A > S_{A,2}$). Moreover, the amplitude, $\ln(2)$, is the maximum possible value. When interpreted as a thermal entropy this corresponds to ‘infinite temperature’, with all $2^{N_A}$ states in region $A$ summed over with equal weight in the canonical partition function. It is natural that this maximally entropic state, corresponds to the wavefunction with completely random and uncorrelated signs. If one were to modify the wavefunction by introducing some (spatial) correlations into the sign structure (still with half the signs positive and half the signs negative), one would expect that the coefficient of the volume law would be reduced$^4$.

Before turning to the two-component quantum fluid, in section 3, where we propose, define and discuss the QDL phase, it will be helpful to briefly review the entanglement entropy in localized states of quantum particles moving in a spatially dependent quenched random potential.

### 2.4.3. Area law with disorder: MBL states.

While we are interested in translationally invariant fluids, it will be instructive to remind the reader of the properties of wavefunctions for bosons (or fermions) moving through a spatially random quenched potential, described, for example, by adding a term to the Hamiltonian of the form,

$$\hat{H}_{\text{dis}} = \sum_j V_j \hat{d}_j^\dagger \hat{d}_j.$$  

(20)

Here, for example, one might take $V_j$ to be a site-uncorrelated random potential. When the disorder strength greatly exceeds the particles kinetic and interaction energies, the ground state wavefunction is expected to be localized, either in a fermion insulator [2] or a ‘Bose glass’ phase [37]. A caricature wavefunction for the localized fermion insulator ground state might be constructed in terms of a determinant built from the lowest energy localized single particle orbitals, $\phi_\alpha(r)$,

$$\psi_{\text{MBL}}(\{r_\alpha\}) = \det[M],$$

(21)

with the matrix $M_{\alpha\beta} = \phi_\alpha(r_\beta)$. While this wavefunction is an eigenstate for non-interacting fermions, it is generally believed that the system will still retain a localized insulating ground state with weak enough interactions.

While the ground state should be localized with strong enough disorder, recent work [4–24] has suggested that such strongly disordered systems can have highly excited finite energy density states that are likewise localized. These so-called MBL phases [4,5] while having a finite energy density above the ground state, as for a thermal state, are not

$^4$ Note also that if one were to ‘bias’ the signs so that the positive signs occurred with some probability $p > 1/2$ then one would find that $S_{A,2} \sim O(1)$.  

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thermalized. Indeed, the entanglement entropy for the eigenstates in an MBL state is believed to satisfy an area law, $S_{\text{MBL}}^A \sim L_A^{d-1}$. Loosely speaking, the MBL states can be thought of as real-space direct product states of single particle localized orbitals.

As we shall see in the next section, for certain finite ‘temperature’ translationally invariant multi-component fluids, a new kind of entanglement entropy can be defined, which does have an area law, despite the absence of any quenched disorder.

3. QDL: a possible new phase in two-component fluids?

We next consider a translationally invariant two-component fluid of indistinguishable quantum particles, a set of heavy particles with large mass $M$ and a set of light particles with mass $m \ll M$. Initially we will not specify the quantum statistics of the two sets of particles. The Hamiltonian we have in mind is a generalization of equation (5) consisting of kinetic energies for the two particle types, along with short-range two-body interactions,

$$H = H_0 + H_U$$

with

$$H_0 = \int_x \hat{D}_x^\dagger (-\nabla_x^2/2M) \hat{D}_x + \int_x \hat{d}_x^\dagger (-\nabla_x^2/2m) \hat{d}_x,$$

$$H_U = \int_{x,x'} [U_{xx'}^m \hat{n}_x \hat{n}_{x'} + U_{xx'}^M \hat{N}_x \hat{N}_{x'} + U_{xx'}^{mM} \hat{n}_x \hat{N}_{x'}],$$

where $\hat{n}_x \equiv \hat{d}_x^\dagger \hat{d}_x$ and $\hat{N}_x \equiv \hat{D}_x^\dagger \hat{D}_x$.

We will again focus on the microcanonical ensemble in which the fluids are confined in a box with volume $V$, possibly with periodic boundary conditions, and isolated from any thermal bath. We will primarily be interested in the situation where the system is placed in an exact many-body eigenstate of the Hamiltonian with a finite energy density above the ground state. Since our model is time reversal invariant we can take the exact eigenstates to be real.

It will again be convenient to consider first quantized particle wavefunctions, which we can define via the position kets,

$$|\{R_{\alpha}\}, \{r_{\beta}\}\rangle = N_M^{R} \prod_{\alpha=1}^{N_M} \hat{D}_R^\dagger \prod_{\beta=1}^{N_m} \hat{d}_{r_{\beta}}^\dagger |\text{vac}\rangle,$$

where $N_M$ and $N_m$ denote the number of heavy and light particles, respectively. We make no assumptions about $N_M, N_m$ except that the ratio $N_M/N_m$ remains finite in the thermodynamic limit. Denoting the state vectors as $|\Psi\rangle$ we have position space wavefunctions, $\Psi(\{R_{\alpha}\}, \{r_{\beta}\}) = \langle \{R_{\alpha}\}, \{r_{\beta}\} |\Psi\rangle$. For bosons/fermions these wavefunctions will be appropriately symmetrized/antisymmetrized under exchange of any two particles.

As for the single component case it will prove convenient to also define a quantum lattice gas model, in terms of operators $\hat{d}_j, \hat{D}_j$ with $j = 1, 2, \ldots, N$ labeling the lattice sites. The analogous lattice tight binding Hamiltonian for the two-component fluid takes

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the form $\hat{H} = \hat{H}_0 + \hat{H}_U$ with kinetic energy,
\[
\hat{H} = -\sum_{ij} (t^M_{ij} \hat{D}_i^\dagger \hat{D}_j + t^m_{ij} \hat{d}_i^\dagger \hat{d}_j + \text{h.c.})
\] (26)

where the positive hopping amplitudes satisfy the inequality, $t^m_{ij} \gg t^M_{ij} > 0$, and an interaction energy,
\[
\hat{H}_U = \sum_{ij} \left[ U^m_{ij} \hat{n}_i \hat{n}_j + U^M_{ij} \hat{N}_i \hat{N}_j + U^{mM}_{ij} \hat{n}_i \hat{N}_j \right],
\] (27)

with the definitions $\hat{n}_j = \hat{d}_j^\dagger \hat{d}_j$ and $\hat{N}_j = \hat{D}_j^\dagger \hat{D}_j$.

We again introduce position-space occupation number basis states,
\[
|\{N_j\},\{n_j\}\rangle = \prod_{j=1}^N [\hat{D}_j]^{N_j} \prod_{j=1}^N [\hat{d}_j^\dagger]^{n_j} |\text{vac}\rangle,
\] (28)

where $N_j = 0, 1$ and $n_j = 0, 1$ denotes the absence/presence of a heavy particle and light particle on site $j$, respectively. Associated with a general state vector $|\Psi\rangle$ we define a wavefunction in the occupation number basis, $\Psi(\{N_j\},\{n_j\}) = \langle \{N_j\},\{n_j\} | \Psi \rangle$.

3.1. Definition of QDL

We will assume that the system is in a translationally invariant pure state $\Psi(\{\mathbf{R}_a\},\{\mathbf{r}_a\})$, a finite energy density eigenstate of the interacting Hamiltonian in equation (22). We will assume that the wavefunction is real and has the exchange properties appropriate for the particles’ statistics.

We make the following assumptions about the wavefunction. Under a spatial bipartitioning into region $A$ and $B$ with $V_A \ll V_B$, we will assume that the entanglement entropy exhibits a volume law, that is for $\hat{\rho}_A = \text{Tr}_B |\Psi\rangle \langle \Psi|$, the reduced density matrix varies as,
\[
S_A = -\text{Tr}_A [\hat{\rho}_A \ln \hat{\rho}_A] \sim V_A.
\] (30)

We will also assume that the light and heavy particles have a ‘mutual’ volume law entanglement, under a partitioning between the two components. Specifically, define a reduced density matrix by integrating out only the light particles throughout the whole volume, $V = V_A + V_B$,
\[
\hat{\rho}_M = \text{Tr}_m |\Psi\rangle \langle \Psi|.
\] (31)

Then define an entropy by tracing over the heavy particles, also throughout the whole volume of the system,
\[
S_{Mm} = -\text{Tr}_M \hat{\rho}_M \ln \hat{\rho}_M.
\] (32)

This ‘inter-component’ mutual entropy is assumed to vary with the whole volume of the system, $S_{Mm} \sim V$.
3.1.1. Spatial and inter-component bi-partitioning diagnostic. We now introduce the following diagnostic of such two-component wavefunctions. It consists of several steps. In step one, we construct the probability distribution for the positions of the heavy particles were they to be measured (with the light particles unmeasured),

$$\mathcal{P}(\{R_\alpha\}) = \int \prod_{\beta=1}^{N_m} dr_\beta |\Psi(\{R_\alpha\}, \{r_\beta\})|^2. \quad (33)$$

Next, we draw sets of coordinates, \(\{R_\alpha^{(1)}\}, \{R_\alpha^{(2)}\}, \{R_\alpha^{(3)}\}, \ldots\), from this probability distribution \(\mathcal{P}(\{R_\alpha\})\). And for a given member of this ensemble we return to the original wavefunction and now imagine it as a wavefunction solely of the light particles, with the position of the heavy particles \(\{\tilde{R}_\alpha\}\) playing the role of a fixed parameter. This requires renormalizing the wavefunction as

$$|\Psi\rangle_{\tilde{R}} \equiv \frac{1}{\sqrt{Z_{\tilde{R}}}} |\Psi\rangle,$$  

(34)

with

$$Z_{\tilde{R}} = \int \prod_{\beta=1}^{N_m} dr_\beta |\Psi(\{\tilde{R}_\alpha^{(n)}\}, \{r_\beta\})|^2. \quad (35)$$

so that \(\int \prod_{\beta=1}^{N_m} dr_\beta |\Psi_{\tilde{R}}(\{r_\beta\})|^2 = 1\).

We then define a density matrix,

$$\hat{\rho}_{\tilde{R}} = |\Psi\rangle_{\tilde{R}} \langle \Psi\rangle,$$  

(36)

and with the coordinates of the heavy particles fixed, construct a reduced matrix from \(\hat{\rho}_{\tilde{R}}\) by tracing out the light particles in region \(B\),

$$\hat{\rho}_{A} = Tr_{r_B} \hat{\rho}_{\tilde{R}}. \quad (37)$$

Next, we use the above reduced density matrix to compute the von Neumann entanglement entropy,

$$S_{A}^{\tilde{R}} = -Tr_{r_A}[\hat{\rho}_{A} \ln \hat{\rho}_{A}] - Tr_{r_B}[\hat{\rho}_{B} \ln \hat{\rho}_{B}]. \quad (38)$$

Note that here we have performed a trace solely over the light particles within region \(A\). The von Neumann entropy thus defined depends on the coordinates of all the heavy particles, both in regions \(A\) and \(B\), which we are denoting \(\tilde{R} = \{\tilde{R}_\alpha\}\).

Finally, we ensemble average \(S_{A}^{\tilde{R}}\) over the probability distribution function \(\mathcal{P}(\{R_\alpha\})\) of the heavy particles (equation (33)):

$$S_{A}^{m/M} = \int \prod_{\alpha=1}^{N_M} d\tilde{R}_\alpha \mathcal{P}(\tilde{R}) S_{A}^{\tilde{R}}. \quad (39)$$

Here the superscript \(m/M\) is to indicate that this entropy refers to the entanglement entropy obtained by spatially bi-partitioning the light particles, with mass \(m\) and coordinates \(r_\alpha\), given a fixed (ie measured) position of the heavy particles with mass \(M\) and coordinates \(R_\alpha\), only afterwards averaging over the distribution of the heavy particle positions.

By simply switching the coordinates, \(r \leftrightarrow \tilde{R}\), we can carry through this procedure to obtain the entropy of spatial entanglement of the heavy particles given we fixed the positions of the light particles, which we will denote \(S_{A}^{M/m}\).
Quantum disentangled liquids

In the ‘conventional’ description of a translationally invariant two-component liquid phase, one would typically assume that the light and heavy particles have both thermally equilibrated, implying a volume law for both entropies,

\[ S_A^{m/M} \sim L_A^d; \quad S_A^{M/m} \sim L_A^d; \quad \text{(thermal)} \] (40)

These volume law behaviors imply that each species of particle contributes a piece to the entropy per unit volume in region \( A \).

We are finally in a position to give a precise definition of the QDL. The QDL phase is described by finite-energy density wavefunctions each of which has a volume law under a full spatial bi-partitioning, \( S_A \sim V_A \), an inter-component entanglement varying as the volume of the whole system, \( S_{Mm} \sim V \), while for each wavefunction the two spatial/inter-component bi-partitioning entropies behave differently—fixing (measuring) the light particles leaves the heavy particles in a spatial volume law state, but with the heavy particles fixed (measured) the light particles are in a spatial area law,

\[ S_A^{m/M} \sim L_A^{-1}; \quad S_A^{M/m} \sim L_A^d; \quad \text{(QDL)} \] (41)

3.2. Illustrative wavefunction for the QDL

In order to construct an explicit wavefunction that we can prove is in a QDL phase, it will be convenient to focus on the lattice model with both heavy and light boson particles. Consider the boson wavefunction in the occupation number basis,

\[ \Psi_{QDL}(\{N_j\}, \{n_j\}) = \psi(\{N_j\}) \prod_{j=1}^N \frac{1}{\sqrt{2}} \left[ \delta_{n_j,0} + e^{i\pi N_j} \delta_{n_j,1} \right]. \] (42)

Here, the first term on the right hand sign, \( \psi(\{N_j\}) \), is a wavefunction for the heavy particles alone, and we assume that this is in the ‘random-sign’ volume law state from section 2, \( \psi(\{N_j\}) = 2^{-N/2} \text{sgn}(\{N_j\}) \) with random signs, \( \text{sgn}(\{N_j\}) = \pm 1 \), chosen independently for each of the \( 2^N \) configurations \( \{N_j = 0, 1\} \).

We must now check each of the conditions for the QDL state. Below, when convenient, we will employ the simplified notation, \( \Psi_{QDL}(N, n) \).

We first consider tracing out the light degrees of freedom throughout the whole system, \( \hat{\rho}_M = Tr_m |\Psi_{QDL} \rangle \langle \Psi_{QDL}| \), writing

\[ \rho_M(N, N') = \sum_{\{n_j\}} \Psi_{QDL}(N, n) \Psi_{QDL}^*(N', n), \] (43)

and explicitly perform the summation over \( \{n_j\} \) to get,

\[ \rho_M(N, N') = \psi(N) \psi^*(N') \prod_j \frac{1}{2} (1 + e^{i\pi (N_j + N_j')}). \] (44)

To assess the inter-species entanglement entropy we consider,

\[ Tr_M(\rho_M^2) = \sum_{\{N_j\}\{N'_j\}} \rho_M(N, N') \rho_M(N', N). \] (45)
and find,
\[ Tr_M(\hat{\rho}_M)^2 = \sum_{\{N\}} |\psi(N)|^4 = 2^N 2^{-2N} = 2^{-N}, \] (46)

giving the required volume law for the inter-species (second Renyi) entanglement entropy, \( S_{Mm,2} = -\ln Tr_M(\hat{\rho}_M)^2 = (\ln 2)N. \)

Next we consider a spatial bi-partition to check that the wavefunction \( \Psi_{QDL} \) has a volume law spatial entanglement, when both the heavy and light particles are traced out in a region \( B \). It is convenient to introduce the notation, \( N_A = \{N_{j\in A}\} \) and \( N_B = \{N_{j\in B}\} \), and similarly for \( N \rightarrow n \), so that we can write the reduced density matrix, \( \hat{\rho}_A = Tr_B[\hat{\rho}] \), in a shorthand notation as, \( \rho_A \equiv \rho_A(N_A, n_A, N'_A, n'_A) \) with,
\[ \rho_A = \sum_{N_B, n_B} \Psi_{QDL}(N_A, N_B, n_A, n_B) \Psi_{QDL}^\ast(N'_A, N_B, n'_A, n_B). \] (47)

The summation over \( \{n_B\} \) can be performed explicitly to give,
\[ \rho_A = f(N_A, n_A)f(N'_A, n'_A) \sum_{N_B} \psi(N_A, N_B)\psi^\ast(N'_A, N_B), \] (48)

where we have defined,
\[ f(N_A, n_A) = \frac{1}{\sqrt{2}} \prod_j (\delta_{n_A,0} + e^{i\pi N_A \delta_{n_A,1}}). \] (49)

We now examine \( Tr_A[\hat{\rho}_A^2] \),
\[ \sum_{N_A, n_A, N'_A, n'_A} \rho_A(N_A, n_A, N'_A, n'_A)\rho_A(N'_A, n'_A, N_A, n_A). \] (50)

We can use the identity,
\[ \sum_{n_A, n'_A} [f(N_A, n_A)f(N'_A, n'_A)]^2 = 2^{2N_A} 4^{-N_A} = 1, \] (51)

to obtain the expression,
\[ Tr_A[\hat{\rho}_A^2] = \sum_{N_B, N'_B} |\sum_{N_A} \psi(N_A, N_B)\psi^\ast(N_A, N'_B)|^2, \] (52)

With the presumed form of the heavy particle wavefunction, as a sum over all real-space occupation number configurations with a random sign, we obtain,
\[ Tr_A[\hat{\rho}_A^2] = 2^{-N_A} + 2^{-N_B} - 2^{-(N_A+N_B)}. \] (53)

This gives the required volume law for the spatial bi-partition Renyi entanglement entropy, \( S_{A2} = -\ln Tr_A[\hat{\rho}_A^2] = (\ln 2)N_A \) for \( N_A \ll N_B \). Again, this implies that von Neumann entropy \( S_A \) also follows the volume law.

We next turn to the wavefunction diagnostic which involves freezing the coordinates of one species of particle, and performing a spatial bi-partitioning into regions \( A \) and \( B \) and integration over the other particle type in region \( B \). Consider first fixing the positions of the heavy particles, choosing a configuration of the occupation numbers, \( \{N_j\} \). Once the
heavy particles are fixed in $\Psi_{QDL}(\hat{N}, n)$, the wavefunction in terms of the light particles is a product law in space. Thus, the spatial entanglement entropy for the light particles will actually vanish, so that $S_{A}^{3/m} = 0$. This is morally an area law, $S_{A}^{3/m} \sim L_{A}^{d-1}$.

Next we must fix the positions of the light particles, choosing some configuration for $\{\tilde{n}_j\} = 0, 1$. Let $\tilde{j}$ denote the subset of sites with $\tilde{n}_j = 1$. The QDL wavefunction can then be written as,

$$\Psi_{QDL}(\{N_j\}, \{\tilde{n}_j\}) = C(\tilde{n})\psi(\{N_j\}) \prod_{j} e^{i\pi n_j},$$

(54)

where the coefficient $C(\tilde{n}) = \prod_{j}(1/\sqrt{2})$. Following section 3.1.1, this requires normalizing this wavefunction over the heavy particles, which simply sets $C(\tilde{n}) = 1$ for all configurations of the light particles, giving $\Psi_{QDL}(N_A, N_B, \tilde{n}) = \psi(N_A, N_B) \prod_{j} \exp(i\pi N_j)$.

We must now spatially bi-partition, and integrate out the heavy particles in region $B$, $\hat{\rho}_A = Tr_{NB} |\Psi_{QDL}\rangle \langle \Psi_{QDL}|$, 

$$\rho_{A}^{\tilde{n}}(N_A, N_A) = \prod_{\tilde{j} \in A} e^{i\pi (N_{A}^{\tilde{j}} + N_{A}^{\tilde{j}}')} \times \sum_{N_B} \psi(N_A, N_B)\psi^*(N_A', N_B).$$  

(55)

Finally, we can compute the trace over the density matrix squared, giving,

$$Tr_{N_A}[\rho_{A}^{\tilde{n}}]^2 = 2^{-N_A} + 2^{-N_B} - 2^{-N_A + N_B},$$

(56)

or a volume law for the second Renyi entropy,

$$S_{A,2}^{m/m} = -\sum_{\{\tilde{n}\}} \mathcal{P}(\{\tilde{n}_j\}) \ln Tr_{N_A}[\rho_{A}^{\tilde{n}}]^2 = (\ln 2)N_A.$$  

(57)

Here $\mathcal{P}(\{\tilde{n}_j\})$ is the probability of finding the light particles in the configuration $\{\tilde{n}_j\}$,

$$\mathcal{P}(\{\tilde{n}_j\}) = \sum_{\{N_j\}} |\Psi_{QDL}(\{N_j\}, \{\tilde{n}_j\})|^2.$$  

(58)

Again, this implies volume law for the von Neumann entropy as well$^5$. Therefore, our illustrative wavefunction, equation (42), indeed satisfies all the properties postulated for the QDL.

3.3. QDL wavefunction of the Born–Oppenheimer form

We now consider a wavefunction for a QDL phase of two-component quantum particles moving in the continuum. This wavefunction will be constructed by using a Born–Oppenheimer type approach [1]. To be concrete we take the heavy particles to be bosons and the light particles to be fermions. We assume that the number of heavy and light particles, $N_M, N_m$, are comparable, as are their mean interparticle spacings, $\ell^{-d}_m = N_M/V$ and $\ell^{-d}_m = N_m/V$. We are interested in a wavefunction with average energy per particle above the ground state, $\tau$, which satisfies,

$$\frac{\hbar^2}{M\ell^{2}_m} \ll \tau \ll \frac{\hbar^2}{m\ell^{2}_m}.$$  

(59)

$^5$ The von Neumann entropy $S$ satisfies $S > S_2$ [36]. Therefore, $\sum_i p_i S_i > \sum_i p_i S_{i,2}$, for a set of positive numbers $\{p_i\}$.

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This corresponds to an ‘effective temperature’ where the heavy particles are in a ‘classical’ regime while the light particles are in a ‘quantum’ regime—although, as throughout this paper, we are always treating both heavy and light particles as quantum particles. We will also assume that the heavy and light particle are interacting with one another via a strong potential, $U_{MM}(x)$ with a characteristic energy scale large compared to the ‘degeneracy’ energy scale for the light particles, $U_{Mm}(0) \gg \hbar^2/m\ell_m^2$.

Consider a wavefunction of the form,

$$
\Psi_{QDL}^{BO}(\{R_\alpha\}, \{r_\beta\}) = S\left(\text{per}[e^{ik_\alpha \cdot R_\alpha'}] \det[\Phi^R_\beta(r_\beta')]\right),
$$

where the indices $\alpha, \alpha'$ run from 1, 2, $\ldots$, $N_M$, the indices $\beta, \beta'$ run from 1, 2, $\ldots$, $N_m$, and the symmetrization $S$ acts on the set of coordinates $\{R_\alpha\}$. The first term in this wavefunction denotes a permanent for the heavy bosons with wavevectors $k_\alpha$ that are chosen with a random direction and an amplitude pulled from an appropriate Boltzmann distribution, $p_k \sim \exp(-\epsilon_k/\tau)$ with $\epsilon_k = \hbar^2 k^2/2M$.

The second term is a determinant wavefunction for the light fermion particles. Here, the single particle orbitals, $\Phi^R_\beta(r)$, denote the lowest energy single particle eigenstates for the light fermions in the presence of a static configuration of the heavy particles, $\{R_\alpha\}$, described by the single particle ‘Born–Oppenheimer’ Hamiltonian,

$$
\hat{H}^{BO}_c = \int_x \hat{c}_x^\dagger (-\nabla_x^2/2m) \hat{c}_x + \int_x V_x \hat{c}_x^\dagger \hat{c}_x,
$$

where we have defined a potential energy,

$$
V_x = \sum_{\alpha=1}^{N_M} U_{MM}(x - R_\alpha).
$$

It is important to emphasize that the coordinates of the heavy particles $\{R_j\}$ that enter into the effective potential are c-numbers, not operators.

Since the heavy particles have been placed into a wavefunction appropriate for a fluid, constructed from a permanent of randomly selected single boson plane waves, a typical configuration of heavy particles that carries appreciable weight in the full wavefunction will correspond to essentially ‘random’ locations of the heavy particles. Thus, in the effective Born–Oppenheimer Hamiltonian for the light fermions, the effective potential $V_x$ will vary randomly in space. Under the assumption that the inter-component interaction energy scale is large compared to the degeneracy energy of the light particles, $U_{Mm}(0) \gg \hbar^2/m\ell_m^2$, all of the low energy single fermion eigenstates in $\hat{H}^{BO}_c$ should be localized.

Due to this localization, for given fixed positions of the heavy particles, the Slater determinant of the light fermions describes a localized phase. While we have assumed that the light fermions are occupying the lowest energy localized eigenstates, which is reasonable in the extreme limit, $\tau \ll \hbar^2/m\ell_m^2$, this can be relaxed. Provided the higher energy occupied orbitals in the fermion Slater determinant are all spatially localized one expects that the properties of the QDL wavefunction should not change qualitatively. Even upon inclusion of interactions between the light fermions, the many fermion wave function can still describe a localized phase—a MBL phase [4, 5].
Since the MBL wavefunction of the light fermions has an area law entanglement entropy under spatial bi-partitioning into regions $A$ and $B$, the entropy $S_{A}^{m/M}$ defined in section 3.1 obtained by fixing the positions of the heavy particles should satisfy an area law, $S_{A}^{m/M} \sim L_{A}^{d-1}$, one of the two important defining properties of the QDL phase.

On the other hand, consider fixing the positions of the light particles in the QDL wavefunction, with some typical configuration, denoted $\{\tilde{r}_\beta\}$. While the magnitude, and more importantly the sign, of the Slater determinant $\det[\Phi^{R}_{\beta}(\tilde{r}_\beta')]$, for given fixed $\{\tilde{r}_\beta\}$, will vary appreciably as the coordinates of the heavy particles are moved about, since the sign of the permanent term in equation (60) has, in any case, an essentially random sign structure already, one expects that the entanglement entropy for the heavy particles under spatial bi-partitioning should be a volume law, $S_{A}^{M/m} \sim L_{A}^{d}$. Thus, the Born–Oppenheimer type wavefunction defined above is indeed expected to describe a QDL phase.

4. Physics of the QDL

The key property of the QDL phase as defined above is that it is not fully thermalized. Physically, if one is given the positions of the heavy particles, the light particles appear as if they were localized due to the effective potential provided by the heavy particles. Therefore, the dynamics and the mutual entanglement of the light particles is largely dictated by the instantaneous positions of the heavy particles.

4.1. Analogy with MBL physics

Consider the limit where the mass $M$ of the heavy particles is taken to infinity. In this limit, the heavy particles can be taken as fixed ‘impurities’, and the problem reduces to solving for the quantum dynamics of the light particles moving in the random background potential of the infinitely heavy particles. Let’s assume that this background static random potential is strong enough that the light particles are in a MBL phase [4, 5]—that is, generally, a finite energy density state within which the particles cannot move, and have zero conductivity (even at finite energy density). In the MBL phase, the light particles are in area law. In this phase equilibrium statistical mechanics essentially breaks down—the entropy per unit volume is zero, even though the system is at a finite energy density. The microcanonical ensemble, when one can take the quantum system in a pure state, is not equivalent to the canonical ensemble for a system in thermal equilibrium with a bath. Essentially, in the presence of the random potential the quantum spectrum of the light particles is locally discrete. Due to this discreteness, it is essentially impossible for energy to move about, and one part of the system cannot act as a thermal reservoir for the remaining part of the system.

Due to the area law, one can think of the particles in the MBL phase as essentially being in a direct product state. This state is a very low entangled state having an area law—indeed a generic finite energy density state will have a volume law entanglement entropy. It is important to emphasize that such ‘disentangling’ requires quantum mechanics, and there is no classical analog to the MBL phase.
Quantum disentangled liquids

One can think of the QDL as a phase within which the light particles fail to thermalize due to the presence of the heavy particles. While the heavy particles can move around once their masses are finite, \( M^{-1} \neq 0 \), the light particles can move much more quickly, and can become ‘localized’ by the (nearly) static potential of the heavy particles. While this picture of localization is no longer strictly valid once the heavy particles can move, the definition in the preceding section of the QDL phase in terms of the wavefunction is still precise. Effectively, with the positions of the heavy particles ‘fixed’ (i.e. measured) in the many-particle wavefunction, the light particles are in an area law state.

To expose the partial non-thermalization of the QDL, consider a hypothetical scenario where one suddenly turns off all interactions between the particles (intra-, as well as, inter-species). This allows one to assign a contribution to the entropy solely from the light particles, \( S_{\text{light}} \). At the same time, since the heavy particles in the QDL are thermalized, one can also associate a well-defined temperature \( T_{\text{QDL}} \) with the QDL, which can be measured prior to the moment interactions are turned off. This can be measured, for example, by connecting the QDL to a non-QDL ideal gas (‘thermometer’) and measuring the temperature of this ideal gas when there is no transfer of energy between the two systems. Repeating the above process at different temperatures, one can obtain a \( S_{\text{light}}(T_{\text{QDL}}) \) curve. The main point is that since the light particles in the QDL are not thermalized, the function \( S_{\text{light}}(T_{\text{QDL}}) \) will, in general, be different for different ensembles of the system, and for a given ensemble, may even be independent of \( T_{\text{QDL}} \), signaling a breakdown of the thermodynamics in QDL. Such a process might be realizable in a two-fluid cold atomic system with large mass ratio for the two components, for example, a \(^{133}\text{Cs}-^{6}\text{Li} \) mixture [38].

In such systems, the interactions can be tuned by sweeping the system through Feshbach resonances which changes the intra- and inter-species scattering length. Subsequently the entropy \( S_{\text{QDL}} \) can be measured by adiabatically transforming the light particles into an ideal gas.

It is also interesting to contemplate the propagation of information in QDL and MBL phases. To illustrate this, consider an MBL eigenstate and compute the entanglement entropy, \( S_A(L) \), for some region \( A \), with linear dimension \( L \), which is taken to be much smaller than the total system size. In the MBL phase one expects an area law for this quantity, \( S_A(L) \sim L^{d-1} \). Now imagine perturbing the Hamiltonian with a time dependent perturbation at time \( t = 0 \) in the middle of region \( A \), and then integrating the time dependent Schrodinger equation. After some fixed amount of time, turn off the local perturbation, but let the system evolve for infinitely long times. Denote as \( S_A^f(L) \) the ‘final’ entanglement entropy, and examine the change in entropy, \( \delta S(L) = S_A(L, t = 0) - S_A^f(L, t = \infty) \). In the MBL phase we expect that this should behave as \( \delta S(L) \sim \exp(-L/\xi_{\text{ent}}) \), falling off exponentially with some ‘entanglement length’, \( \xi_{\text{ent}} \). The point here is that in the MBL phase, ‘entanglement does not propagate’, or more correctly quantum information does not propagate.

However, in the context of QDL, under a time dependent local perturbation \( \delta S_A^{\text{MD}}(L) = S_A^{\text{MD}}(L, t = 0) - S_A^{\text{MD}, f}(L, t = \infty) \) is not expected to fall off exponentially, since the system is partially thermalized. However, even then, \( S_A^{\text{MD}}(L, t) \) will follow an area law for all times \( t \). Focusing on \( S_A^{\text{MD}}(L, t) \) allows one to distill the partial localization in QDL. In essence, the only way to transfer quantum information between light ‘particles’ (excitations of the light quantum field) is by having the heavy ‘particles’ convect along this information between two spatially separated points.
5. Extensions and open issues

5.1. Stability of the QDL for heavy and light particles?

Perhaps the most important question pertinent to our exploration is whether or not the QDL phase can exist as typical finite energy-density eigenstates for generic local Hamiltonians which possess the appropriate particle content—heavy and light particles. For a given Hamiltonian straightforward (full) exact diagonalization is in principle possible—with the full set of exact eigenstates in hand the QDL wavefunction diagnostics can be implemented. This will clearly be challenging, since ED will only be possible in very small systems with very few particles—perhaps up to 12 sites in a 1d tight binding model of heavy and light fermions. It is unclear how effective DMRG will be in exploring QDL physics in even simple toy models. The DMRG is best suited in accessing low entanglement states, usually ground states, while the QDL state has a volume law entanglement entropy. Perhaps some variant of dynamical DMRG could be helpful in using short-time dynamics to explore the expected small area law entanglement of the light particles, once the heavy particles are fixed. It appears that numerical progress on the QDL might require modifications of existing algorithms/approaches, at the very least, and perhaps the development of new algorithms tailored to access and evolve (typical) volume law wavefunctions.

5.2. Spin/charge QDL phases

The definition of the QDL phase is not restricted to fluid systems with two particles types (i.e. heavy and light) but can be readily generalized to any quantum system which possesses more than one global symmetry and associated conserved currents. Of particular physical relevance are quantum systems with a conserved $U(1)$ charge and a conserved $SU(2)$ spin, as in a Hubbard-like model, for example. Here, we briefly discuss two different QDL phases which can be naturally defined in quantum systems with both a conserved charge and a conserved spin, which we will refer to as a ‘Quantum Disentangled-charge Liquid’ and a ‘Quantum Disentangled-spin Liquid’, respectively. We will denote these two phases as QD$^c$L and QD$^s$L, where the subscript $c$ ($s$) on the ‘D’ indicates that the charge (spin) degrees of freedom are disentangled.

To define these phases consider spin-1/2 fermions moving on a tight-binding lattice with conserved charge and spin. We are interested in the entanglement properties of a (typical) finite energy-density eigenstate, which we denote as, $\Psi(\{n\},\{s_z\})$, where $n_j = 0, 1, 2$ and $s_{z,j} = 0, \pm 1/2$ label the charge and $z$-component of spin quantum numbers on site $j$.

The diagnostics for these spin/charge QDL phases proceeds just as in section 3.1. In brief, first draw a charge configuration, denoted $\{\tilde{n}_j\}$, from the probability distribution that would be found if the charge were measured on each site,

$$P(\{n\}) = \sum_{\{s_z\}} |\Psi(\{n\},\{s_z\})|^2.$$  \hspace{1cm} (63)

Then, for a given measured charge configuration construct a spin wavefunction, $\Psi(\{\tilde{n}\},\{s_z\}) = \Psi(\{\tilde{n}\},\{s_z\})$. After normalizing this spin wavefunction (over the spin numerical results.
configurations), for given fixed charge configuration \( \{ \tilde{n} \} \), compute the entanglement entropy of the spins under a spatial bi-partitioning, between a (small) sub-region \( A \), of linear dimension \( L_A \), and the rest of the sample, and denote this entropy as \( S_A^{s/c} \). Finally, average this entanglement entropy over the distribution of charge configurations from \( P(\{n\}) \) to obtain \( S_A^{s/c} \) where the superscript refers to the spin-entanglement for given configuration of the charge.

Switching charge and spin, one can carry through the same procedure, fixing the spin configuration (effectively by measuring the spin on each site) to obtain a charge wavefunction, and then computing the spatial entanglement of the charge degrees of freedom. Averaging the resulting charge entanglement entropy over the spin configurations gives the entropy \( S_A^{c/s} \).

In a Quantum Disentangled-charge Liquid (QD\(_c\)L) the finite energy-density eigenstates have spin degrees of freedom which are in a volume law state for given fixed (measured) charge, while the charge degrees of freedom are in an area law state when the spins are fixed (measured);

\[
S_A^{s/c} \sim L_A^d; \quad S_A^{c/s} \sim L_A^{-1}; QDcL.
\]

The area law for the charge degrees of freedom, is indicative that the charges are disentangled for a given fixed spin configuration.

Conversely, in a Quantum Disentangled-spin Liquid (QD\(_s\)L) one has,

\[
S_A^{s/c} \sim L_A^{-1}; \quad S_A^{c/s} \sim L_A^d; QD_sL,
\]

indicating that the spins are spatially disentangled once the charge is measured.

### 5.3. Stability of the QDL in Hubbard-type models?

To address the energetic stability of the spin/charge QDL phases for Hubbard-type models of spin-1/2 fermions, we shall first consider the large \( U \) limit where the model reduces to the \( t-J \) model.

#### 5.3.1. The \( t-J \) model is in a QD\(_c\)L phase.

Remarkably, one can argue that the finite energy-density eigenstates of a \( t-J \) model should always be in a Quantum Disentangled-charge Liquid phase. As we shall see, this follows from the no-double occupancy constraint.

Consider first the case with no holes, where the model reduces to the Heisenberg model. A typical finite energy-density eigenstate of the Heisenberg model will have a volume law entanglement entropy.

Next dope in some holes, and implement the QDL diagnostics. Consider first measuring the charge in the \( t-J \) model wavefunction, which is tantamount to fixing the positions of the electrons and the holes (empty sites). The remaining spin wavefunction is expected to have a volume law entanglement entropy, as it has with no holes present.

Next measure the spin in the \( t-J \) model wavefunction. The sites with \( s_z = \pm 1/2 \) are occupied by an electron (charge one), and due to the no double-occupancy constraints those sites with \( s_z = 0 \) necessarily have charge zero. Thus, in the \( t-J \) model Hilbert space, measuring the spins on each site fixes the charges into a single configuration. Thus, with measured spins the charge wavefunction is a trivial direct product of the corresponding occupied and empty sites, and is fully disentangled!
A volume law spin wavefunction (with fixed/measured charge) and a disentangled (or area law) charge wavefunction (with fixed/measured spin) is precisely the conditions for a Quantum Disentangled-charge Liquid phase. We thereby conclude that the finite energy-density eigenstates of a $t$–$J$ model should be in a QDcL phase.

### 5.3.2. Finite $U$ Hubbard models.

In contrast to the $t$–$J$ model, in the Hubbard model with finite $U$, measuring (ie fixing) the spin configuration on each site does not uniquely specify the charge configuration, since sites with $s_z = 0$ can either have zero charge (hole) or two electrons in a singlet (a doublon) with charge 2. Thus, in the QDL diagnostics, while the spin wavefunction for fixed charge is still expected to be in a volume law, the charge wavefunction, for fixed spin configuration, is now a non-trivial wavefunction for the doublons—and it is unclear whether this doublon wavefunction will have an area law or a volume law entanglement entropy.

When $U$ is very large the typical spatial-density of doublons will be very small. And when the hole doping is much smaller than one, the number of sites that the doublons can occupy is very low. Plausibly, in this limit, the doublon wavefunction might be ‘localized, with an area law entanglement entropy. If so, the Hubbard model in this regime would be in a QDcL phase. It might be possible to establish whether or not this is the case by using exact diagonalization studies to extract the eigenstates of the large $U$ Hubbard model, and to implement the QDL diagnostics to extract the doublon entanglement entropy.

In the absence of interactions ($U = 0$) the exact many-body eigenstates are Slater determinants. A typical finite energy-density state can be constructed by building the determinant from a set of plane waves with randomly chosen wavevectors. Such wavefunctions will be riddled with doublons, and it seems most likely that both the spin-wavefunction (for fixed charge) and the charge-wavefunction (for fixed spin) will have a volume law entanglement entropy, that is $S_{A}^{s/c} \sim S_{A}^{c/s} \sim L_{A}^{d}$—this could perhaps be checked numerically. Provided this is so, the free fermion model would not be in a QDL phase.

In view of the above discussion, the important question to address (perhaps numerically) in the Hubbard model is whether the transition between the eigenvalue thermalized phase for small $U$ and the QDcL phase in the $U \rightarrow \infty$ ($t$–$J$) limit, occurs at finite or infinite $U$. If the latter, then the QDcL would only be a consequence of the strict no-double occupancy constraint of the Hilbert space in the $t$–$J$ model, and should not be considered a generic phase. On the other hand, a finite $U$ transition would imply the generic stability of the QDcL for large enough $U$.

### 5.4. Exploring the possibility of QDL physics in water

It seems worthwhile to ask if a QDL phase existed in an actual fluid, what would be its experimental manifestations? To put our ideas into concrete form, below we ask could pure water be in a QDL phase, with the light proton degrees of freedom becoming ‘localized’ on the oxygen ions? We then discuss the consequences if such was indeed the case, in light of some of the transport experiments on water.

The oxygen to proton mass ratio is 16, so water offers, at the very least, a natural setting to explore possible QDL physics. Moreover, while 16 is not so tremendously large, the covalent oxygen–hydrogen bonds (together with the hydrogen bonds themselves)
will strongly correlate the relative motion of the protons and the oxygen ions. Strong correlations between the heavy and light particles are central to the QDL. In addition, the proton must be treated quantum mechanically with a thermal de Broglie wavelength comparable to the inter oxygen–hydrogen separations. One might be tempted to treat the heavier oxygen ions semiclassically, using a Born–Oppenheimer approximation. As discussed in the introduction, the Born–Oppenheimer potential produced by a random (fluid) configuration of the oxygen ions, might ‘localize’ the protons on the oxygen ions. The corresponding localized Born–Oppenheimer wavefunction, would then take the QDL form in equation (4).

Let’s then imagine that water at standard temperature and pressure (STP) is in a QDL phase. Consider perfectly pure water, with exactly twice as many hydrogen as oxygen ions, and with no other ‘impurity’ ions present. The protons are the light, charge \( e \) fermions, and the oxygen ions are the heavy, charge \( -2e \) bosons. If in a QDL phase, water should be a ‘perfect insulator’ with a strictly zero (linear response) electrical conductivity, despite being at finite temperature. An applied electrical field which pushes the protons and oxygen ions in opposite directions, could not set up an electrical current (linear in the field) since the protons are ‘enslaved’ by the oxygen ions.

Experimentally, salt water is a fairly good conductor [39], with a conductivity of order \( 10^{-1} \) S/cm (a 10 \( \Omega \) cm resistivity). The conduction is dominated by solvated ions (salts) and tap water has a much smaller conductivity, typically \( 10^{-4} \) S/cm. Data on ‘pure’ water, of interest to us here, is hard to come by. The often quoted ‘theoretical’ value for the conductivity of pure water [40] is tiny, \( 5.5 \times 10^{-8} \) S/cm, and measured conductivities in the \( 10^{-7} \) S/cm ballpark have been mentioned in the literature [41]. This conductivity is very, very small, some 13 orders of magnitude smaller than the conductivity of a metal.

It will clearly be challenging experimentally to ascertain the difference between \( 10^{-7} \) S/cm and strictly zero conductivity, as in a QDL phase, especially since even nominally ‘pure’ water contains unwanted solvated ions which will conduct. Some progress can be made by measuring the electrical conductivity of ‘supercritical water’ using high pressures [42, 43], up to 500 kbar using ‘shock’ methods. Upon pressurizing (nominally) pure water the conductivity is found to rise. Remarkably [42, 43], for very high pressures (under ‘shock’) the conductivity increases dramatically by some 8 orders of magnitude, from \( 10^{-7} \) S/cm at STP to \( 10^{-4} \) S/cm at 50 kbar before saturating at 20 S/cm above 200 kbar (estimated temperature of 5000 K). At these temperatures/pressures, pure water conducts over 100 times better than salt water! If pure water at STP were in fact in an insulating QDL phase with zero conductivity, at such pressures a highly conducting and presumably fully thermalized phase has been reached. Another direction to explore is the effect of deuteration on the electrical conductivity of water. How well does heavy water conduct? Further experiments on the electrical properties of water could be most interesting.

One can also ask about the physics of the proton nuclear spins in water. Since the spin-exchange interaction between two protons on a single water molecule, of order 30K, is more than 6 orders of magnitude larger than the inter-proton nuclear spin dipole–dipole interaction, it is a good approximation to assume a global \( SU(2) \) nuclear spin symmetry in water. As such, we can contemplate the possibility of a spin/charge QDL phase, in particular a Quantum Disentangled-spin Liquid. Imagine first measuring (fixing) the electrical charge in water, which fixes the locations of the proton and oxygen ions. In the absence of any spin-exchange between protons on neighboring water molecules, the
proton spins will all be ‘paired-up on each water molecule, and the spin wavefunction, for fixed charge, will be in a direct product state (area law). If the spin-exchange rate between protons on neighboring water molecules is small enough compared to the intra-molecule proton spin exchange rate, it is plausible that an area law spin-wavefunction persists. On the other hand, measuring (fixing) the spins in the water wavefunction, while determining the locations of the protons, will leave a wavefunction for the oxygen ions, likely to be in a volume law state. If water were actually in such a QD,\textsubscript{L} phase, the disentangling (localization) of the proton spins might have implications for nuclear spin-dephasing of ions/molecules solvated in water.

5.5. Relevance to finite ‘temperature’ quantum field theory?

Viewed more broadly, a fluid of heavy and light quantum particles at finite ‘temperatures’ is a special case of a two-component finite ‘temperature’ quantum field theory. Of relevance here is a QFT Lagrangian with two quantum fields, a ‘light’ field with strong quantum fluctuations interacting strongly with a ‘heavy’ field which is in a ‘semi-classical’ regime. The eigenstate thermalization hypothesis (ETH) [25–28] posits that the reduced density matrix in a large sub-volume $A$ (the ‘system’) obtained by tracing out the degrees of freedom in the even-larger sub-volume $B$ (the ‘environment’), equals the thermal-density matrix in region $A$:

$$\hat{\rho}_A \equiv \text{Tr}_B |\psi\rangle\langle \psi| = Z_A^{-1} e^{-\beta \hat{H}_{\text{QFT}}^A}.$$ (66)

Here $\hat{H}_{\text{QFT}}^A$ is the QFT Hamiltonian projected into region $A$, $Z_A = \text{Tr}_A e^{-\beta \hat{H}_{\text{QFT}}^A}$ and $\beta$ is determined by the condition that the von Neumann entanglement entropy, $S_{\text{vN}} = -A T_A \hat{\rho}_A \ln(\hat{\rho}_A)$ satisfies a volume law, $S_{\text{vN}} = sV_A$ with entropy density $s$ equal to the thermal entropy density: $s = \partial_T (T \ln Z_A) / V_A$.

Imagine a two-component QFT, with a strongly fluctuating quantum field coupled to a ‘semi-classical’ quantum field, with finite energy-density eigenstates that are in a QDL phase. Since the QDL wavefunction is not fully thermalized, it cannot be accessed using the canonical thermal-ensemble and the ETH is not fully operative. For a QFT in the QDL phase, if one fixes the background ‘semi-classical field’, the entanglement entropy of the strongly fluctuating quantum field satisfies an area law. It might be especially interesting to explore such ideas further in the context of an elementary quantum particle interacting with a background semi-classical gravitational field.

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Appendix A. Volume law entanglement entropy for random-sign wavefunction

The ‘random-sign’ wavefunction is given by

$$\psi(\{n\}_j) = \frac{1}{\sqrt{2^N}} sgn(\{n\}_j),$$  \hspace{1cm} (A.1)

with the \textit{sgn} function chosen randomly \textit{sgn}(\{n\}_j) = \pm 1, independently for each of the \(2^N\) configurations \(\{n_j = 0, 1\}\).

To demonstrate that this maximally random-sign wavefunction has a volume law entanglement entropy, it will be convenient to introduce a shorthand notation,

$$\psi(n_A, n_B) \equiv \psi(\{n_{j \in A}\}, \{n_{j \in B}\}).$$  \hspace{1cm} (A.2)

With this notation the density matrix is simply,

$$\rho(n_A, n_B, n'_A, n'_B) = \psi(n_A, n_B)\psi^*(n'_A, n'_B).$$  \hspace{1cm} (A.3)

The reduced density matrix can be obtained by tracing over the sites in region \(B\):

$$\rho_A(n_A, n'_A) = \sum_{n_B} \rho(n_A, n_B, n'_A, n_B),$$  \hspace{1cm} (A.4)

where \(\sum_{n_B}\) denotes a summation over \(n_j = 0, 1\) for \(j \in B\).

The volume law is easiest to establish from the second Renyi entropy, which requires computing,

$$Tr_A[\hat{\rho}_A^2] = \sum_{n_A, n'_A} \rho_A(n_A, n'_A)\rho_A(n'_A, n_A).$$  \hspace{1cm} (A.5)

Re-expressing in terms of the wavefunctions gives,

$$Tr_A[\hat{\rho}_A^2] = \sum_{n_A, n'_A} |\mathcal{M}_A(n_A, n'_A)|^2 = \sum_{n_B, n'_B} |\mathcal{M}_B(n_B, n'_B)|^2,$$  \hspace{1cm} (A.6)

with the definitions,

$$\mathcal{M}_A(n_A, n'_A) = \sum_{n_B} \psi(n_A, n_B)\psi^*(n'_A, n_B),$$  \hspace{1cm} (A.7)

$$\mathcal{M}_B(n_B, n'_B) = \sum_{n_A} \psi(n_A, n_B)\psi^*(n_A, n'_B).$$  \hspace{1cm} (A.8)

For the random-sign wavefunction, \(\mathcal{M}_B\) can be expressed as,

$$\mathcal{M}_B(n_B, n'_B) = \frac{1}{2^N} \sum_{n_A} sgn(n_A, n_B)sgn(n_A, n'_B).$$  \hspace{1cm} (A.9)

Consider two cases, \(n_B = n'_B\) and \(n_B \neq n'_B\). In the former case we have, \(\mathcal{M}_B(n_B, n_B) = 2^{N_A}/2^N\). For the latter case we can use the central limit theorem,

$$\langle |\mathcal{M}_B(n_B \neq n'_B)|^2 \rangle = 2^{N_A}/2^{2N}. \hspace{1cm} (A.10)$$

Here, the brackets \(\langle \cdot \cdot \cdot \rangle\) indicate an average over an ensemble of random-sign wavefunctions, but for a given typical random-sign wavefunction this form should hold in the large \(N_A\) limit.

Combining the above expressions gives,

$$Tr_A[\hat{\rho}_A^2] = 2^{-N_A} + 2^{-N_B} - 2^{-(N_A+N_B)}.$$  \hspace{1cm} (A.11)
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