Many-body Localization

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Submitted for the Degree

Master of Science

University of Oslo
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May 2017
Abstract

We study a random-field Heisenberg model numerically and argue that as we vary the strength of the disorder $W$ there is a phase transition between a thermal phase and a many-body localized phase at a critical disorder strength $W_C = 3.5 \pm 1.0$. We show some properties of the many-body localized phase and contrast them to the properties of the thermal phase. We have considered transport properties, scaling of entanglement entropy, level statistics, participation ratios and whether or not the system satisfies the ETH to distinguish between the phases. We also study the dynamics in the thermal phase near the transition and argue that the phase transition is governed by an infinite randomness fixed point. Furthermore we review much of what is known and conjectured about the physics of generic closed quantum systems.
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Chapter 1

Introduction

When we are concerned with closed many-body quantum systems a fundamental question to ask is which state does the unitary time evolution bring the system to, after an arbitrarily long time? There appear to be two generic answers which are robust under small, local perturbations of the Hamiltonian, namely thermalization and many-body localization (MBL). MBL systems remember local details of the initial state at all times and thus do not thermally equilibrate, and it is the only known generic exception to thermalization in closed strongly interacting quantum systems. Other known non-thermalizing systems are in some way fine-tuned. Whether the system is MBL or thermal depends on the nature of the system and on the the initial state, and the system can have a quantum phase transition between the two phases. MBL generally occurs in systems with disorder and the transition is driven by the disorder strength.

The transition is not captured by the statistical mechanics ensembles and there are no singularities in thermodynamic quantities as the transition is crossed. It is a dynamic phase transition and it can be observed in the eigenstates of the Hamiltonian. We call it an “eigenstate phase transition”, which is marked by a singular change in the properties of the many-body energy eigenstates. The MBL transition is a quantum phase transition with no classical counterpart, but in contrast to most quantum phase transitions it can occur at energy densities corresponding to finite, or even infinite, temperatures.

It has been known for a long time that in the presence of disorder, quantum systems can host a variety of interesting phenomena. In 1958 Anderson [1] showed that the quantum mechanical wavefunction of a non-interacting particle in a sufficiently disordered landscape will be exponentially localized. This has profound consequences for the system’s transport properties as it entails that these states cannot carry current over macroscopic distances. He also conjectured that this effect could in some way occur also in an interacting system, and that it would lead to non-thermalization. The question of the possibility of Anderson localization in an interacting system went mostly unanswered until roughly ten years ago
when first Mirlin, Gornyi and Polyakov [3] and shortly thereafter Basko, Aleiner and Altshuler [2] showed perturbatively that Anderson localization can persist when we turn on interactions between the particles. During the last decade there has been a lot of theoretical research on MBL, both numerical [4, 5, 53] and analytical [49, 50], and recently it has also been reported to be experimentally observed [6, 7, 8].

Statistical mechanics and thermodynamics are some of the most successful physical theories and can be used to explain a large number of phenomena, both in many of the natural sciences and in our daily lives. Statistical mechanics allows us to treat systems of $N \sim 10^{23}$ particles, for which an exact solution is unfeasible. We do this by considering different microstates labeling the properties (momenta, positions etc.) of every particle of the system. Statistical mechanics assumes that all microstates of the system are equally likely and that the system dynamically explores all the microstates. As a result of this exploration the system eventually reaches a thermal equilibrium and forgets the details of its initial state. The system in equilibrium can thus be described by just a few macroscopic variable. This allows us to not care about the details of the microscopic dynamics and just consider the much simpler statistical average over possible macroscopic states.

It is not a priori obvious that this procedure should work. In classical systems the justification comes from the connection between chaos and thermalization, and chaotic ergodicity seems to be a requirement for classical statistical mechanics to apply. We do however know that our world is ultimately quantum mechanical in nature and we are therefore compelled to consider quantum statistical mechanics. Strict dynamical chaos however is not present in closed quantum systems and it is not completely understood which mechanism justifies the ensemble approach to quantum statistical mechanics. Despite its success we still need to put the understanding of quantum statistical mechanics on a more solid foundation. Seminal steps in this direction were made by Deutsch [11] and Srednicki [10], culminating in the “eigenstate thermalization hypothesis” (ETH) which should determine which closed quantum systems thermalize.

The purpose of this thesis will be to define, study and discuss these two phases and to review much of what is known and conjectured about them. We do this by studying an isotropic random-field Heisenberg Hamiltonian numerically and investigating its properties as we vary the disorder strength. The random-field Heisenberg Hamiltonian is a paradigmatic model in the context of MBL, and it has been studied extensively in recent years. Alongside with original considerations, many of our numeric results are inspired by these previous studies. In particular we have reproduced the results of Pal and Huse [4] and some from the more recent paper by Serbyn, Papic and Abanin [54]. Our aim will thus be twofold; first we wish to numerically show the existence of the two phases and consider some properties of the phase transition between them. Second this thesis should serve as a more or less
This thesis is structured as follows: We start in Chap. 2 by heuristically discussing localization, with and without interactions. We will then introduce the mathematical framework of Random Matrix Theory in Chap. 3, which lies at the core of quantum thermalization, and in Chap. 4 we discuss some of the issues of justifying the foundations of quantum statistical mechanics and we define and discuss the ETH. We then turn our attention to MBL and introduce the model with which we will be concerned and the numerical methods used to study it in Chap. 5. In this chapter we also introduce the formalism of quasi-Local Integrals of Motion (LIOM's) which are expected to emerge in the MBL phase. In Chap. 6 we will argue through our numerical results that a MBL phase does indeed occur in the random-field Heisenberg model for sufficiently strong disorder. We discuss some of the main differences between the thermal and the MBL phase and investigate at which disorder strength the phase transition occurs. In Chap. 7 we venture to investigate the phase transition itself, and we will consider if it has universality and in particular argue for the possibility of it belonging to an infinite-randomness universality class. We also briefly discuss the dynamics in the thermal phase near the phase transition.
Chapter 2

From Anderson Localization to Many-body Localization

It is about 60 years since Anderson published his seminal paper [1] “Absence of Diffusion in Certain Random Lattices”. His purpose was to “lay the foundation for a quantum-mechanical theory of transport” and he showed that single-particle wavefunctions in a disordered landscape can be exponentially localized in real space. This is due to quantum interference which gives the wavefunction an envelope function and constrains it to a region of space. Anderson basically considered a quantum random walk and showed that this random walker under certain circumstances will be localized. This is a remarkable result since it was showed by Einstein [12] that all random and memory-less walks will lead to diffusion. Such processes are called Markovian, and during which the random walker should obey

\[ \langle r^2(t) \rangle = Dt \]  

(2.1)

Where \( \langle r^2(t) \rangle \) is the net movement of the walker at time \( t \), and \( D \) the diffusion constant. Anderson showed that for a quantum particle randomly propagating on a lattice, it can be the case that \( \langle r^2(t) \rangle \rightarrow \text{const.} \). This implies that quantum propagation in some sense has memory, and that information about the initial state can be contained in the system for arbitrarily long times.

Anderson touched upon two very interesting subjects within condensed matter physics in this paper. First the study of quantum transport in solids and second the question of whether ubiquitously present disorder can cause a closed system to fail to reach thermal equilibrium. He hypothesized that the system would not thermalize in the presence of large disorder, since its subsystems would not be able act as a thermal reservoirs for each other. He did indeed make an immense contribution to the making of a quantum-theory of transport, for which he was awarded the Nobel prize in 1977. He was in many ways ahead of his time in posing the second question, because at his time the mathematical and numerical tools to investigate it properly were not available. He could only analyze the non-interacting problem, and this system is known to be non-thermal, with or without of disorder. To say something
interesting about the thermalization of such disordered systems we would have to properly take the interactions between the particles into account, and this is why it took about 50 years before an answer to question of non-thermalization began to emerge.

Anderson’s work was inspired by the experiments of Fletcher’s group at Bell laboratories [13, 14, 15], which had observed anomalously slow relaxation times for electrons in Silicon-doped semiconductors with impurities. They observed relaxation times of spins of the order of minutes as opposed to milliseconds which was predicted by Fermi’s Golden Rule. The impurities in such semiconductors are frozen into the material in some random way during production. They can be a variety of different ions which enter the semiconductor lattice in several ways; either as ions which are located at interstitial sites, as vacancies where an ion should have been or as impurity ions which have replaced the lattice ions. Such impurities usually are quenched since they are frozen into the material too quickly to thermally equilibrate. We distinguish this from annealed disorder which is frozen into the system very slowly. For annealed disorder we can thus treat averages over disorder and thermal averages on the same footing. This is not possible for quenched disorder, making it much harder to handle. Thus we basically have a lattice with random impurities frozen into random sites, and it was this scenario which triggered Anderson’s interest.

2.1 The Anderson Model

Anderson simplified the situation and completely disregarded interactions; he considered a tight-binding Hamiltonian in three dimensions on a lattice with random on-site disorder and short-range hopping matrix elements

\[
H_A = \sum_i h_i c_i^\dagger c_i + \sum_{i,j} (J_{ij} c_i^\dagger c_j - J_{ji} c_i c_j^\dagger) \tag{2.2}
\]

where \(h_i\) are random static disorder potentials drawn from some distribution, which we for simplicity take to be uniform on \([-W/2, W/2]\), \(J\) is the hopping strength which falls off at least as \(1/r^3\) and \(c_i^\dagger\) and \(c_i\) are fermionic creation and annihilation operators acting on site \(i\). The Anderson Hamiltonian describes a substantially simplified physical model, but we will see that this model does indeed contain rich and interesting physics. Here we have described a fermionic model, but it can be mapped onto a spin model through a Jordan-Wigner transformation as we have shown in Chap. 5. So we will take Eq. (2.2) to essentially describe both spin systems and fermionic systems.

In a clean system there is no disorder and \(H_A\) reduces to a standard tight-binding Hamiltonian which is diagonalized by Bloch waves

\[
|\psi_k\rangle = \frac{1}{\sqrt{N}} \sum_i e^{i\vec{k} \cdot \vec{r}_i} |i\rangle \tag{2.3}
\]
Figure 2.1: We see in b) that in the tight binding model all wavefunctions are extended in space since the particle in a) is free to move all over the lattice. When we introduce quenched disorder at the lattice sites as in c) we get localized wavefunctions as in d). The wavefunctions in b) and d) are just examples of how extended and localized wavefunctions can look like.
Where $\vec{k}$ is the wavenumber and $\vec{r}_i$ is the location of a particle at site $i$ with wavefunction $|i\rangle$. Such a system is known to generally be conductive. In the opposite limit of finite $W$ and $J = 0$, the eigenstates are the $|i\rangle$'s with eigenenergies $h_i$'s, i.e. all the eigenstates are localized on the individual lattice sites and the system is thus insulating. After considering the two limits of $J/W$ it is interesting to look at where the transition between the localized system at $J/W = \infty$ and the metallic system at $J/W = 0$ is. More concretely we wish to know if there is a transition at a finite value of $J/W$. Anderson showed perturbatively that in three dimensions the transition will indeed occur at finite $J/W$, i.e. the conductivity will remain zero for small but finite $J/W$. He performed the perturbation theory in the localized limit, treating the hopping as the perturbation. This is usually referred to as the locator expansion.

To the lowest order in $J/W$ the perturbed eigenstates are

$$|\psi_A\rangle = |i\rangle + \sum_j \frac{J}{h_i - h_j} |j\rangle$$

(2.4)

Since $h_i$'s are random we can only make probabilistic statements about whether or not the second term is small, which we need in order for the perturbation theory to be valid. The typical value of $h_i - h_j$ is $W/2$, which means that the typical smallest value of $h_i - h_j$ for any given $i$ is $W/2z$, where $z$ is the coordination number of the lattice, i.e. the number of nearest neighbors. Therefore we naively expect the perturbation theory to be valid if $2Jz/W < 1$.

This conclusion turns out to be correct, although the argument above is clearly far from foolproof. There is always a possibility for $h_i - h_j$ being small, and to higher order in perturbation theory we will eventually with certainty encounter sites where $h_i$ is very close to $h_j$. We would need to perform a careful probabilistic analysis of the disorder in order to make any conclusions regarding whether or not such resonances will hurt the perturbation theory. Anderson set up the perturbation theory and showed that localization persists with probability one in the thermodynamic limit. However, he was also not completely rigorous in proving that the states nearby in energy do not harm the perturbation theory, and it was not until later that this was rigorously proven by Fröhlich and Spencer [37].

Thus for small $J/W$ the states of three dimensional system are localized and its wavefunction $\psi_i(\vec{r})$ has an envelope which goes as

$$\psi_i(\vec{r}) \sim e^{-|\vec{r} - \vec{r}_i|/\xi}$$

(2.5)

where $\xi$ is the localization length and $\vec{r}_i$ the localization center. Conversely a non-localized state has an extended wavefunction which is spread out over the entirety of space like $\psi_i(\vec{r}) \sim 1/\sqrt{V}$, where $V$ is the volume of space. It is widely believed that extended and localized wavefunctions cannot co-exist in the same energy range. Therefore they split into bands which are separated by so-called mobility edges $E_m$. As we have argued above, in three dimensions there is a transition from extended states to exponentially localized states. This
transition can happen through special “critical states” at the mobility edge which displays power-law localization.

2.2 Scaling Theory to Anderson Localization

Apart from Anderson’s perturbative study, there are many other ways to approach the problem of localization. The so-called “Gang of Four” consisting of Abrahams, Anderson, Licciardello, and Ramakrishnan [17] were some of the first to consider a scaling theory of Anderson localization. We review some of their results in the following.

Thouless and Edwards [16] defined the essential quantity for the scaling theory, namely the Thouless energy $E_T$. It is a measure of the sensitivity of the energy levels in a finite system to a change in boundary conditions. It quantifies how much the energy levels are affected when we twist the boundary conditions and measures the correlations between wavefunctions at different energies. Intuitively we expect that a state exponentially localized in the bulk of the system, with a wavefunction described by Eq. (2.5), would not be significantly affected by changes in boundary conditions and that the Thouless energy should therefore be exponentially small in system size. We thus expect that

$$E_T \propto e^{-\frac{L}{\xi}}$$

(2.6)

for a localized system, where $L$ is the system’s linear size and $\xi$ the localization length. Conversely an ergodic system could get a considerable change in eigenenergies due to a change in boundary conditions and it can be shown that the Thouless energy then is proportional to the inverse of the diffusion time $\tau_T$ across the sample. The Thouless energy in the ergodic system is

$$E_T = \frac{\hbar}{\tau_T} = \frac{\hbar D}{L^2}$$

(2.7)

where $D$ is the diffusion constant. Thouless and Edwards considered the conductance $G$ of a macroscopically homogeneous material. The conductance always has units of Siemens, so they define the dimensionless conductance as

$$g(L) = \frac{G(L)}{e^2/\hbar}$$

(2.8)

They showed that $g(L)$ is the ratio of $E_T$ to the mean level spacing $\Delta$. The mean level spacing is inversely proportional to the Heisenberg time $\tau_H = 2\pi\hbar/\Delta$, which is the longest time a particle can travel in the sample without visiting the same region twice. Since the Thouless energy is inversely proportional to the diffusion time $\tau_D$, which is the time it takes a conducting particle to arrive from the bulk of the material to the boundary of the sample, we get the conductance in an ergodic system as

$$g(L) \propto \frac{\tau_H}{\tau_T}$$

(2.9)
This leads us to an intuitive assumption for when the system fails to be ergodic in terms of $g(L)$. When the Thouless time is larger than the Heisenberg time, particles will not reach the boundary from the bulk. Thus, we expect that for $g < 1$, i.e. when the Thouless energy is much smaller than the level spacing, the system is not ergodic. Conversely when $g > 1$ the particles will eventually travel all around the system, and the system is ergodic. The transition between the Anderson localized and non-localized phases is thus expected to occur when $E_T$ and $\Delta$ are of the same order of magnitude.

The great insight of the “Gang of Four” which lead to the scaling theory of Anderson localization was that $g(L)$ should be the only relevant parameter for determining the conductive properties of the system, and that it depends on $L$ in an universal manner. Having recognized $g(L)$ as the universal variable, we consider putting $n^d$ identical blocks of length $L$ together into a hypercube of linear dimension $nL$. Then the conductance of the hypercube $g(nL)$ should only depend on the conductance of the smaller system $g(L)$. That is, we have

$$g(nL) = h(n, g(L))$$

(2.10)

That this equation should hold was an educated guess known as one-parameter scaling, and it is essentially an application of the renormalization group to the Anderson localization problem. We wish to have the scaling equation Eq. (2.10) on a continuous form, so we consider the case where $n = 1 + dn$, which gives us

$$g((1 + dn)L) = h((1 + dn), g(L))
= g(L) + g(L)\beta(g(L))dn
\Rightarrow \beta(g(L)) = \frac{L \cdot g(L + Ldn) - g(L)}{Ldn}$$

(2.11)

Where $\beta(g(L))$ is a scaling function. We take the limit $dn \to 0$ and get

$$\frac{d\log(g)}{d\log(L)} = \beta(g(L))$$

(2.12)

Which is the renormalization group equation governing $g(L)$. The physical significance of the scaling function $\beta(g)$ is that if we start out with a system of linear size $L$ and conductance $g(L)$ for which $\beta(g) < 0$, then the conductance will decrease upon enlarging the system, and conversely the conductance will increase if $\beta(g) > 0$. That is, the $\beta$-function encodes the transport properties of the system in the thermodynamic limit. We do not know exactly how this $\beta$-function looks like, but we can easily find its asymptotic behavior in the limits of very large and very small conductance.

We now consider the limit of weak disorder, which we expect leads to large conductance $g \gg 1$. In this regime we expect the $G(L)$ to be given by Ohm’s law

$$G(L) = \sigma \frac{L}{A} \Rightarrow g(L) = \sigma_0 L^{d-2}$$

(2.13)
where $\sigma_0$ proportional to the conductivity. Inserting this into the renormalization group equation Eq. (2.12) to obtain

$$\lim_{g \to \infty} \beta(g) = d - 2$$

(2.14)

We have seen that for strong disorder all wavefunctions are exponentially localized, therefore the conductance should also be exponentially small in system size. We therefore assume the scaling $g(L) = \exp(-L/\xi)$, which means that

$$\lim_{g \to 0} \beta(g) = \log \left( \frac{g}{g_c} \right)$$

(2.15)

Where $g_C$ is the critical value of conductance. Knowing the asymptotic form of the $\beta(g)$ for very strong and very weak disorder, we can use the simplest interpolation between the two limits to arrive at the famous scaling plot from the “Gang of Four” paper in Fig. 2.2. We see here that in one and two dimensions all states are localized for arbitrarily small disorder, but in three dimensions we have a mobility edge. In three dimensions there is a transition for at the critical value of conductance $g_C$, which we above argued should be $g_C \approx 1$.

It is not obvious that Eq. (2.10) should hold, but the same results as we have presented in this section have been verified through the use of a renormalization group approach to a version of the non-linear $\sigma$-model. This is a field theory which was first proposed by Wegner [18] and later Efetov [21] pioneered a supersymmetric version of the $\sigma$-model, which was used.

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We note that in two dimensions the picture is actually a bit more nuanced, and there can be delocalized states depending on the symmetry class of the Hamiltonian.
to verify some of the scaling theory’s results for Anderson localization \cite{19, 20}.

2.3 Localization with Interactions

After understanding single-particle Anderson localization reasonably well, we wish to consider what happens to an Anderson localization when we turn on interactions.

The Anderson model is a rather unrealistic model for any conceivable physical system, as there will generally be some coupling to the external world and particles will interact with each other. We will here consider what happens when we try to make the Anderson model a bit more general. We first let the system interact with phonons, i.e. couple it to a heat bath, and then we look at a closed system with inter-particle interactions.

2.3.1 Coupling to Heat Bath

It is believed that if we turn on, even a very week, coupling to a heat bath with a continuous spectrum the conductivity will become finite.

We provide a crude “derivation” of the conductivity in the the variable-range hopping model, as first discussed by Mott \cite{22}. The model describes conduction in a $d$-dimensional system where all charge carriers are localized, and which is weakly coupled to a heat bath. We consider a system of fermions at low temperatures with strong quenched disorder and a well-defined Fermi-level, in which all the states near the Fermi-level is occupied. Two adjacent states in energy are generally localized far apart in space. If the system is coupled to a heat bath there are delocalized phonons with energies arbitrarily close to zero, and the fermions can then exchange energy with the phonons of the heat bath and hop over long distances. This happens because there will always be a phonon of the correct energy to match the energy difference between two single-particle states, and the phonons can thus excite fermions above the Fermi-level and induce conduction.

We consider tunneling between states with localization centers separated by $R$ and with energies $E_1$ and $E_2$ lying above and below the Fermi-level respectively. The probability for tunneling decays as $\exp(-2R/\xi)$ where $\xi$ is the localization length. The probability to produce excitations of order $E_1 - E_2$ in the heat bath goes as $\exp((E_1 - E_2)/\beta)$. Which leads us to assume that the conductivity to leading order is

$$\sigma(\beta) \sim \exp\left(-\frac{2R}{\xi} - (E_1 - E_2)/\beta\right)$$

(2.16)
Mott suggested to optimize the competition of the overlap term $e^{\exp(-2R/\xi)}$ which favors short hops and the energy activation $e^{\exp((E_1 - E_2)\beta)}$ which generally favors long hops. He did this because he claimed the conductivity should be dominated by states where the activation and tunneling is optimal. In the tunneling term we approximate $R$ with its typical value $R_{\text{typ}} \sim \left(\frac{dN(E_F)}{dE}\right)^{-\frac{1}{4}}$

\begin{equation}
R_{\text{typ}} \sim \left(\frac{(E_1 - E_2)}{dN(E_F)}\right)^{-\frac{1}{4}}
\end{equation}

Where $dN(E_F)/dE$ is the energy level spacing at the Fermi-level. We now optimize and find that the “optimal value”, which we call $\epsilon^*$, of $E_1 - E_2$ is

\begin{equation}
\epsilon^* \sim \beta^{\frac{1}{4+4d}}
\end{equation}

Which yields the conductivity of the system

\begin{equation}
\sigma(\beta) = \sigma_0(\beta)e^{-(\frac{T_0\beta}{1+\beta})^{\frac{1}{1+4d}}}
\end{equation}

where $\sigma_0(\beta)$ has a power-law dependence on $\beta$, and $\sigma_0(\beta)$ and $T_0$ depend on the details of the system. This result is valid for small temperatures. If we consider larger temperatures then conduction will be dominated by activation across the mobility edge of the sample and we get an other exponent for our conductivity. In both cases, with fermion-phonon interaction, the conductivity is finite (although it can be very small) at all finite temperatures even when all one-particle states are localized.

### 2.3.2 In a Closed System

Having seen already in the sixties that an Anderson localized system interacting with a heat bath will thermalize, the question of whether inter-particle interactions alone can make the
system thermal was timely. However it took almost 50 years from Anderson’s original paper until Basko et al. were able to answer this question rigorously. Their calculations are rather lengthy so we will just mention their results and which sorts of systems it applies to.

We assume we have a highly disordered system, in which all the single-particle states are Anderson localized, and then let the particles interact with each other with an interaction strength $J_{int}$. We start by considering the two limits of $J_{int}$. If $J_{int} = 0$ we retain Anderson’s model and the system is obviously localized and conversely if $J_{int}$ is very large we intuitively expect the model to be non-localized since the disorder in this case is insignificant in comparison with the strong interaction. Therefore there should be some crossover from the localized to the ergodic phase, and the natural question to ask is if this crossover happens at finite $J_{int}$. It was this question Basko et al. tried to answer as they considered a closed system at energy densities corresponding to low, but finite, temperatures. They considered the Hamiltonian

$$H = \sum_i \epsilon_i c_i^+ c_i + \sum_{ijkl} J_{ijkl} c_i^+ c_j^+ c_k c_l$$

(2.20)

where $c_i^+$ creates a single particle state which is Anderson localized with localization center $\vec{r}_i$, localization length $\xi$ and energy $\epsilon_i$. They performed perturbation theory in the low temperature-limit of weakly interacting fermions, in a similar manner to Anderson’s locator expansion. The perturbation theory is performed in the basis of occupied single-particle eigenstates

$$|\phi_\alpha\rangle = |n_{\alpha_0} n_{\alpha_1} \ldots n_{\alpha_i}\rangle$$

(2.21)

Where $n_{\alpha_i}$ are the occupation numbers of the localized eigenstates. The occupation numbers completely determines $|\phi_\alpha\rangle$, which is a state in fermionic Fock space. The interactions are short range and the matrix elements $J_{ijkl}$ are constrained in both in space and energy, and the interaction term $J_{ijkl}$ is thus treated as a perturbation. It plays a similar role as the hopping between sites did in Anderson localization, and the full MBL problem looks like the Anderson problem on a hypercubic lattice in $N$ dimensions, where each site is a basis state in Fock space and $J_{ijkl}$ gives rise to “hopping” in Fock space. The interaction mixes the single-particle states that are close in Fock space, but assuming $J_{ijkl} \ll \hbar$ the mixing is suppressed with probability $O(J/\hbar)$.

Basko et al. showed to all powers in perturbation theory that, for small enough $J_{ijkl}$, localization in Fock space persists up to a finite energy density which is extensive in the system size, and conductivity can be zero at finite temperatures. This was proof that Anderson localization can occur in an interacting system and answers the question posed by Anderson; disorder can indeed prohibit a closed system from thermalizing.

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†Actually Mirlin et al. showed this about a year earlier. However this paper has not received as much attention as the paper by Basko et al. [3]
Chapter 3

Random Matrix Theory

We will in this chapter review “random matrix theory” (RMT), which will be crucial when discussing quantum thermalization later. RMT has had tremendous success in many areas of physics and it was first brought to the fore by Wigner [23, 24] and Dyson [25], who developed the theory in order to explain the spectra of complex nuclei. Wigner realized that it would be hopeless to try to calculate the exact energy eigenvalues of huge quantum systems, such as heavy nuclei, instead he considered focusing on their statistical properties. Wigner postulated that in an energy range far from the ground state, the Hamiltonian of the nuclei should not, from a statistical point of view, differ significantly from an ensemble of random matrices. He demanded that the ensemble supports unitary quantum evolution, i.e. its matrices must be hermitian, that all symmetries must be taken into account and that no additional information is encoded into the ensemble, in particular there should be no privileged direction of Hilbert space. When these constrains were met, Wigner claimed that the exact details of the distributions do not matter much.

This idea might indeed seem quite counter-intuitive, but Wigner was able to rather accurately predict the spacing between the lines in the spectra of heavy nuclei. If we look at a small energy-window where the density of states is constant, the Hamiltonian of many large, complex systems will, in a non fine-tuned basis, appear much like a random matrix. Therefore it does indeed make sense that we may gain insights into complex physical systems by studying random matrices subject to the same symmetries as those of the systems’ Hamiltonian. Wigner and Dyson considered approximating the Hamiltonian by a Gaussian ensemble of finite large $N \times N$-matrices $H$ which have the probability density of its independent elements as

$$P(H_{nm}) \propto e^{-\frac{\zeta tr(H^2)}{2a^2}} \quad (3.1)$$

Where the factor $\zeta$ depends on the ensemble and $a$ sets the overall energy scale. In other words the matrix elements are essentially independent Gaussian random variables, but they have to comply with the symmetries of the Hamiltonian. The matrix elements are real in the orthogonal, complex in the unitary or real quaternions in the symplectic ensemble. The
Gaussian orthogonal ensemble (GOE) contains real, symmetric matrices and corresponds to systems with time-reversal symmetry and it is this case which will interest us in the following. This is mainly because the model we will be studying later is time-reversal invariant. However, it is also of particular interest to see that equilibrium statistical mechanics, with its arrow of time, can emerge in time-reversal invariant systems. That is, how a system which is microscopically time-reversal invariant can break the the invariance on a macroscopic level.

### 3.1 Wigner Surmise

We will now derive a probability distribution function (PDF) for the spacings of the energy levels of GOE systems. We will use the $2 \times 2$ case to deduce the so-called Wigner Surmise. We consider the general real, symmetric matrix

$$
H_{2 \times 2} = \begin{pmatrix}
\epsilon_1 & \frac{V}{\sqrt{2}} \\
\frac{V}{\sqrt{2}} & \epsilon_2
\end{pmatrix}
$$

with eigenvalues

$$
\lambda_{1/2} = \frac{\epsilon_1 + \epsilon_2}{2} \pm \frac{1}{2} \sqrt{(\epsilon_1 - \epsilon_2)^2 + 2V^2}
$$

Where the factor of $1/\sqrt{2}$ is inserted since it leaves the Hamiltonian invariant under basis rotations. Since $\epsilon_1$, $\epsilon_2$, and $V$ are independent variables drawn from a Gaussian distribution, which we for simplicity assume to have zero mean and unit variance, we can easily find the statistics of the separation between the energy levels

$$
P_1(\lambda_1 - \lambda_2 = \omega) = \frac{1}{(2\pi)^{3/2}} \int d\epsilon_1 d\epsilon_2 dV \delta(\omega - \sqrt{(\epsilon_1 - \epsilon_2)^2 + 2V^2}) e^{-\frac{\epsilon_1^2 + \epsilon_2^2}{2}}
$$

We here change variables to $\epsilon_2 = \epsilon_1 + \sqrt{2} \eta$, which gives us a Gaussian integral in $E_1$, which is trivial to integrate. We then get

$$
P(\omega) = \frac{1}{2\pi} \int d\eta \ dV \ \delta(\sqrt{2\eta^2 + 2V^2} - \omega)e^{\frac{-\eta^2 + V^2}{2}}
$$

$$
= \frac{1}{2\pi} \int_0^{2\pi} d\theta \int_0^\infty dr \ r \ \delta(\sqrt{2r} - \omega)e^{-\frac{r^2}{2}}
$$

$$
= \frac{\omega}{2} e^{-\frac{\omega^2}{4}}
$$

Where we changed to spherical coordinates $\eta = r \cos(\theta)$ and $V = r \sin(\theta)$ before integrating. We have deduced the distribution function for level separation for the Gaussian Orthogonal Ensemble in two dimensions. This is the celebrated Wigner Surmise with which Wigner was able to explain statistical properties of the spectra of complex nuclei.

Of course such nuclei contains much more than two degrees of freedom, but Wigner made a leap of faith and assumed the two-dimensional result also to be approximately valid.
for systems with more degrees of freedom. In fact, the two-point level spacing function
has, some time after Wigner, been solved exactly in \( N \) dimensions and it turns out that
Wigner’s Surmise is indeed a very good approximation for the probability distribution of
level spacings as \( N \to \infty \). We can easily verify this numerically by generating many large
GOE matrices, which we then diagonalize and estimate the level spacing and average over
many realizations of such matrices to estimate the PDF numerically.

### 3.2 Probability Distribution for the Eigenvalues

Having derived the PDF for the two energy levels of two-dimensional GOE-matrices, we
generalize this to \( N \) dimensions in the following. To make predictions about the spectra
\( \{E_k\} \) of systems of random matrices belonging to GOE, we need to deduce the statistics
of the eigenvalues of \( H \). Whereas the matrix elements are roughly uncorrelated random
numbers, we will see that the eigenvalues are highly correlated. We now wish to find the joint
probability distribution \( P(\{E_k\}) \) for the \( N \) eigenvalues. We write the eigenvalue equation as

\[
H = VEV^T \iff H_{nm} = \sum_i E_i v_{ni} v_{mi} \tag{3.6}
\]

Where \( E \) is a \( N \times N \) diagonal matrix of eigenvalues and \( V \) a \( N \times N \) orthogonal matrix
whose columns consist of the eigenvectors of \( H \), which is a real, symmetric matrix and
has \( N(N + 1)/2 \) independent variables. The matrix of eigenvectors \( V \) must satisfy \( N(N - 1)/2 \)
orthogonality constraint and can therefore be determined by \( N(N - 1)/2 \) independent
parameters which we denote \( \beta_1, \beta_2 \ldots \beta_{N(N-1)/2} \). Since we wish to obtain the probability
distribution function of the eigenvalues, we make a change of variables from \( H_{nm} \) to \( E_k \) and
\( \beta_j \), which we substitute into \( P(H) \) defined in Eq. (3.1). We can use the orthogonality of the
eigenvectors to see that

\[
\text{tr}\{H^2\} = \sum_n \sum_{kl} E_k E_l v_{nk} v_{mk} v_{ol} v_{nl} = \sum_k E_k^2. \tag{3.7}
\]

Due to the fact that the probability is conserved and using Eq. (3.7) we have

\[
P(\mathcal{E})d\mathcal{E} = P(H)dH = Ce^{-\zeta \sum_k E_k^2} dH \tag{3.8}
\]

where

\[
dH = dH_{11}dH_{12}\ldots dH_{N}\quad \wedge \quad d\mathcal{E} = dE_1\ldots dE_N d\beta_1\ldots d\beta_{N(N-1)/2} \tag{3.9}
\]

We will now make the coordinate transformation \( dH = Jd\mathcal{E} \), which leaves us with

\[
P(\mathcal{E})d\mathcal{E} \propto J e^{-\zeta \sum_k E_k^2} d\mathcal{E} \tag{3.10}
\]
Then all that remains is to calculate the Jacobian \( J(E_1, \ldots, E_N; \beta_1, \ldots, \beta_{N(N-1)/2}) \) for this transformation. The Jacobian is

\[
J = \left| \frac{\partial(H_{11}, H_{12} \ldots H_{NN})}{\partial(E_1 \ldots E_N, \beta_1 \ldots \beta_{N(N-1)/2})} \right|
\tag{3.11}
\]

Where tensor notation is implied. We see from Eq. (3.6) that \( H_{nm} \) is a linear function of the eigenvalues which implies that \( \partial H_{nm}/\partial \beta_i \) is linear in eigenvalues as well, and that \( \partial H_{nm}/\partial E_i \) is independent of the eigenvalues. We thus see that \( J \) must be a polynomial of degree \( N(N-1)/2 \) in each of the eigenvalues.

If two eigenvalues are equal their corresponding eigenvectors are not uniquely determined and the inverse transformation of Eq. (3.6) is not properly defined, so \( J \) must therefore vanish when \( E_n = E_m \) for all \( n \) and \( m \). This may also be seen from the fact that determinants with duplicate columns always evaluates to zero. \( J \) thus contains every possible distinct combinations of \( |E_n - E_m| \) as a factor. There exists \( N(N-1)/2 \) such combinations and since \( J \) is a \( N(N-1)/2 \)'th degree polynomial in eigenvalues, all of \( J \)'s dependence on the eigenvalues is accounted for and we have

\[
J = \Pi_{n<m}|E_n - E_m|f(\beta_1, \ldots \beta_{N(N-1)/2})
\tag{3.12}
\]

We do not care about the dependence on \( \beta_i \)'s since we only wish to know the statistics of the eigenvalues \( \{E_k\} \), therefore we simply assume all of the \( \beta_j \)-dependence to be contained in some function \( f(\beta_1, \ldots \beta_{N(N-1)/2}) \). Now we can use this expression for the Jacobian and we get the following PDF

\[
P(\mathcal{E}) \propto e^{-\zeta \sum_k E_k^2} \Pi_{n<m}|E_n - E_m|f(\beta_1, \ldots \beta_{N(N-1)/2})
\tag{3.13}
\]

We integrate out the dependence on the \( \beta \)'s, this yields some normalizing constant which we disregard at this stage. We can always normalize our PDF at some later time. We then arrive at the joint probability distribution for the eigenvalues

\[
P_{\text{GOE}}(\{E_k\}) = C \Pi_{n<m}|E_n - E_m|e^{-\zeta \sum_k E_k^2}
\tag{3.14}
\]

We see clearly from Eq. (3.14), that the probability for having degeneracies in the spectrum is zero and that the probability for finding energy levels very close to each other is small. This effect is called level repulsion and it is one of of the main characteristics of RMT.

The probability distribution function in Eq. (3.14) should be used to obtain results for expectation values of correlation functions of the energy levels within the GOE. However it might get extremely complicated to obtain exact result even for a two-level correlation function when we are dealing with large systems. We will see an example of the PDF in use later; when we calculate a three-point correlation function in Appendix B.
3.3 Matrix Elements in RMT

The eigenvectors $\phi^n = (\phi^n_1, \ldots, \phi^n_N)$ of large GOE matrices are given by the following probability distribution of its components [40, 44]

$$P(\phi_1, \ldots, \phi_N) \propto \delta \left(1 - \sum_k \phi^2_k\right)$$

(3.15)

Where $\phi^n_k$ is the $k$'th component of the $n$'th eigenvector. This form follows from the fact that the orthogonal invariance of the GOE implies that the PDF only depends on the norm $\sqrt{\sum_k \phi^2_k}$ and should thus be proportional to the $\delta$-functions. We see from Eq. (3.3) that the eigenvectors are basically random and uncorrelated unit vectors. Due to the orthogonality restriction this cannot be completely accurate, but since two uncorrelated vectors in a large-dimensional space are usually nearly orthogonal we assume that we can disregard of the orthogonality restrictions between the eigenvectors.

We note that the matrices of GOE can be diagonalized and the eigenvectors form a basis, in which the matrix is diagonal and GOE gives the statistics of the eigenvalues. The statistical properties of the eigenvectors given by the PDF in Eq. (3.3) are specified in a fixed basis for an entire ensemble of random matrices. It thus holds for $N \to \infty$, that the projections of GOE eigenvectors onto a fixed vector in Hilbert space have a Gaussian distribution with zero mean and unit variance [45].

We now move on to study matrix elements of hermitian operators within the framework of RMT. We consider the some given local operator

$$A = \sum_k A_k |k\rangle\langle k|$$

(3.16)

where $\{|k\rangle\}$ are the eigenvectors of some given GOE-matrix. We have the matrix elements

$$A_{nm} = \langle n | A | m \rangle = \sum_k A_k \langle n | k \rangle \langle k | m \rangle = \sum_k A_k (\phi^n_k)^* (\phi^m_k)$$

(3.17)

Using that the eigenvectors are essentially random orthogonal unit vectors we see that to leading order in $1/N$, where $N$ is the dimension of the matrices, we have

$$\langle \phi^n_k \phi^m_l \rangle \approx \frac{1}{N} \delta_{kl} \delta_{nm}$$

(3.18)

where $\langle \phi^n_k \phi^m_l \rangle$ is an average over $|n\rangle$ and $|m\rangle$. We can now utilize Eq. (3.18) to give us the expectation values of matrix elements

$$\langle A_{nm} \rangle \approx \frac{\delta_{nm}}{N} \sum_k A_k = \delta_{nm} \bar{A}$$

(3.19)

Where we see that the expectation values of the off-diagonal elements are zero and all of the diagonal elements have the average value $\bar{A} \equiv \sum_k A_k/N$ as their expectation value. We move on to consider the fluctuations. We have

$$\langle A_{nm}^2 \rangle - \langle A_{nm} \rangle^2 = \sum_{i,j} A_i A_j \langle \phi^n_i \phi^m_i \phi^n_j \phi^m_j \rangle - \sum_{i,j} A_i A_j \langle \phi^n_i \phi^m_i \rangle \langle \phi^n_j \phi^m_j \rangle$$

(3.20)
To evaluate the average of the fluctuations we will need to use Isserli’s or Wick’s Therem [27], which in our notation and in four dimensions states that

\[
\langle \phi_i^n \phi_j^m \phi_j^m \phi_i^n \rangle = \langle \phi_i^n \phi_i^m \rangle \langle \phi_j^m \phi_j^n \rangle + \langle \phi_i^n \phi_j^m \rangle \langle \phi_j^m \phi_i^n \rangle + \langle \phi_i^n \phi_j^m \rangle \langle \phi_j^m \phi_i^n \rangle \tag{3.21}
\]

We will consider the diagonal and off-diagonal elements separately. Using Eq. (3.21) we get the following for the fluctuations of the diagonal elements

\[
(A_{mm}^2) - (A_{mm})^2 = \sum_i A_i^2 (\langle \phi_i^m \rangle^4 - (\langle \phi_i^m \rangle^2)^2)
= 2 \sum_i A_i^2 (\langle \phi_i^m \rangle^2)^2
= \frac{2}{N} A^2
\]

(3.22)

Whereas for the off-diagonal elements we have

\[
(A_{nm}^2) - (A_{nm})^2 = \sum_k A_k^2 (\langle \phi_k^m \rangle^2 (\phi_n^2))^2 = \frac{1}{N} A^2
\]

(3.23)

We can now approximate the matrix elements to leading order in \(1/N\) as

\[
A_{nm} \approx A \delta_{nm} + \sqrt{A \frac{A^2}{N} R_{nm}}
\]

(3.24)

where \(R_{nm}\) is a random variable with zero mean and unit variance on the off-diagonal elements and the diagonal elements have variance two. It is simple to see that the ansatz in Eq. (3.24) gives the correct fluctuations and average of matrix elements within the GOE. We averaged over the random Hamiltonian ensemble to arrive at the above expression but when \(N\) is large the fluctuations should be very small, and therefore the expression should be applicable also to single Hamiltonians.

The question concerning which systems RMT generally applies to still remains. RMT has found applications in nuclear physics, quantum gravity, quantum chromodynamics, the fractional quantum Hall effect and many other places.

In the same way as classical thermodynamics can describe a huge variety of different microscopical systems at a macroscopic level, also RMT can arise from many different microscopical systems, and RMT is insensitive to the interactions at a microscopical level. There exists many other random matrix ensembles and one of them we will see in Chap. 6, namely the Wishart matrix. This is rather similar to the GOE, but it has a slightly different PDF for its matrix elements.

As an aside, it is worth mentioning that there exists an interesting connection between the level statistics in RMT and the statistics of non-trivial zeros of the Riemann zeta function. The Riemann zeta function can be defined as

\[
\zeta(s) = \sum_{k=1}^{\infty} \frac{1}{k^s} = \prod_{p \in \mathcal{P}} \frac{1}{1 - p^{-s}}
\]

(3.25)
Where $\mathcal{P}$ contains all primes and $\zeta(s)$ is defined by Eq. (3.25) for $\Re\{s\} > 1$ and elsewhere through analytic continuation. It has been conjectured that the Riemann zeta function should have all its non-trivial zeros lying on the line $s = 1/2 + iE_k$ with $E_k \in \mathbb{R}$, this is the infamous Riemann hypothesis. It has also been shown numerically for about a billion zeros that the $E_k$’s fluctuate like the eigenvalues of a unitary Gaussian matrix. This hints toward a profound connection between prime numbers and RMT, which as we will see in Chap. 4, lies at the core of quantum chaos and thermalization.
Chapter 4

Chaos & Thermalization

In this section we will consider the statistical mechanics of closed quantum systems. If such systems thermalize, they should approach an equilibrium which is described by the microcanonical ensemble. It is however not always the case that a closed system is properly described by the microcanonical distribution, and we want to consider which conditions need to be satisfied in order for closed quantum systems to be thermal.

There are two common approaches to justify why the rules of statistical mechanics work. The first is to couple the system to a heat bath with certain properties, with which the system can exchange energy, particles etc, and whose heat capacity is unaffected upon coming in contact with the system. This is not the approach we are going to take since it seems in some way to be rather heuristic and only pushes the problem back one level and would force us to justify why the heat bath has its thermal properties. Furthermore it does not seem sound to justify the inherent properties of a closed system by coupling it to an external reservoir.

The other approach is to consider the system in isolation and make some assumption of ergodicity and mixing, i.e. assuming that the dynamics of the system is in some way chaotic. It is then possible to derive the thermodynamic distributions. Although completely isolated quantum systems are not really feasible experimentally, there has recently been made progress in approximating isolated many-body quantum systems [28], and it is this approach we will follow.

We thus have to look at what it means for a system to be ergodic and chaotic. We therefore start by considering chaos, first for classical systems and then for quantum systems, and thereafter look at how this is related to quantum thermalization.

4.1 Classical Chaos & Integrability

Classically chaotic systems are generally governed by non-linear equations and chaos is usually defined as an exponential sensitivity of the phase-space trajectories to arbitrary small perturbations of the initial conditions. Chaos is often quantified through Lyapunov
exponents \( \lambda \), which for chaotic systems show how the difference in phase space trajectories \( \delta Z(t) \) evolves in time from the initial separation \( \delta Z(0) \).

\[
|\delta Z(t)| \approx e^{\lambda t} |\delta Z(0)|
\] (4.1)

There are classes of systems which are not chaotic and have phase-space trajectories whose separation cannot be described by Lyapunov exponents, and these systems are called integrable. Textbook examples of classical chaotic and integrable systems are systems of one particle moving in a Bunimovich and a circular stadium respectively, as seen in Fig. 4.1. In the circular cavity the particle would not change its phase space trajectory drastically if we changed its initial conditions slightly. That is, two trajectories which at one time are close to each other will after an arbitrarily long time still lie close to each other in the non-chaotic circular stadium. In the chaotic Bunimovich stadium, two trajectories which initially lie close together, would lie arbitrarily far apart from each other after a long time. Chaotic systems are generally ergodic, which means that they with time will cover their entire available phase space \(^\dagger\). A particle in the chaotic Bunimovich stadium will after a while have had explored all of its available phase-space, whereas in the circular billiards this is clearly not the case.

Figure 4.1: Trajectories of particle in a cavity. a) Circular stadium with integrable dynamics. b) Bunimovich stadium with chaotic dynamics. Figure from Scholarpedia [32]

We will now to formalize the notion of integrability. Assume we have an Hamiltonian \( H(p,q) \) with the canonical coordinates \( q = \{q_1, ..., q_N\} \) and \( p = \{p_1, ..., p_N\} \). We further assume that the system have some functionally independent conserved quantities \( K = \{K_1, ..., K_j\} \) in involution (meaning they have vanishing Poisson brackets between each other). The conserved quantities are represented by constants of motion with vanishing Poisson brackets with the Hamiltonian. That is, we have

\[
\frac{d}{dt} K_j = \{K_j, H\} = 0 \quad \land \quad \{K_j, K_i\} = 0
\] (4.2)

\(^\dagger\)There are some nuanced differences between chaos and ergodicity, depending on how we define them. We will simply assume that chaotic systems are ergodic, which is indeed usually the case.
For all the $K_j$’s, where the standard Poisson brackets are defined as

$$\{f, g\} = \sum_{j=1}^{N} \frac{\partial f}{\partial p_j} \frac{\partial g}{\partial q_j} - \frac{\partial g}{\partial p_j} \frac{\partial f}{\partial q_j}$$

(4.3)

A system which is Liouville integrable is defined as a system governed by Hamiltonian $H(p, q)$ defined on $\mathbb{R}^{2N}$ with $N$ or more independent constants of motion in involution.

For Liouville integrable systems there thus exists a canonical transformation $(p, q) \to (K, \Theta)$ from the momentum and position variables to the so-called action-angle variables which yields $H(p, q) = H(K)$. The equations of motion are now trivial to solve

$$K_j(t) = K_j(0) \quad \land \quad \Theta_j(t) = \Omega_j t + \Theta_j(0)$$

(4.4)

We know that the trajectories in phase space lie on surfaces corresponding to the constants of motion. For a Liouville integrable system the trajectories are thus restricted to lie on a $N$-dimensional tori, and the dynamics is not chaotic.

When we generalize the above to many-particle systems we usually consider systems with an extensive number of conserved quantities integrable. As an example, a high dimensional system of many non-interacting particles would not be considered chaotic in this sense, even if each particle is chaotic in the part of phase-space associated with its own degrees of freedom, since the energy of each particle is separately conserved.

### 4.2 Quantum Chaology

Knowing that chaotic systems are governed by non-linear equations and have exponential sensitivity to initial conditions, quantum chaos is something of a misnomer. Closed quantum systems do not have exponential sensitivity to initial conditions\footnote{Furthermore it is not completely clear how to consider the divergence of initially small phase space separation, since the uncertainty principle does not allow us to determine positions in phase space sharply within regions smaller than $\hbar$} and are governed by the linear Schrödinger equation, in other words, closed quantum systems are never actually chaotic.

For these reasons Micheal Berry suggested the term “quantum chaology”, as a better word to use to describe unpredictable quantum behavior. Berry defined quantum chaology\textsuperscript{[33]} as

“\textit{The study of semi-classical, but non-classical, phenomena characteristic of systems whose classical counterparts exhibit chaos.}”

\textsuperscript{[33]}
Although chaos theory and quantum mechanics are two of the most successful theories of the 20th century, the interplay between these theories is generally not known. Due to Niels Bohr’s correspondence principle we know that classical mechanics is reached in the limit of large quantum numbers, and at large enough energies the predictions from classical and quantum mechanics does indeed coincide. The Kolmogorov–Arnold–Moser theorem provides a link between classical mechanics and classical chaos, as it describes how integrable systems behave under small non-linear perturbation.

Whereas the connections between classical mechanics and quantum mechanics and classical mechanics and classical chaos are both fairly well understood, it is not generally known how to reach the non-linear dynamics of classical chaos as a limit of quantum mechanics. It is not straightforward to see how the extreme irregularity of classical chaos can arise from the smooth and wavelike nature of phenomenon in the quantum world. It appears we need novel way of telling which quantum systems are “chaotic” and conversely which are not. The main objective of quantum chaology is to identify characteristic properties of quantum systems which, in the semi-classical limit, reflect the integrable or chaotic features of the corresponding classical dynamics.

One of the hallmarks of quantum mechanics is its quantized energy levels, as opposed to classical mechanics where energy is continuous. It seems that the first place to look for a way to distinguish quantum chaotic from non-chaotic systems, is in the spectra of the systems. However it has also been shown to be a remarkable connection between the wavefunctions of chaotic systems in the semi-classical limit and classical chaos. A lot has been conjectured about systems within quantum chaology. We will discuss some of these conjectures in the following.

4.2.1 Level Spacings of Chaotic and Integrable System

It appears that quantum chaos does not make itself felt at any particular energy level, however its presence can be seen in the spectrum of energy levels. It has been hypothesized that we can distinguish between chaotic and non-chaotic quantum systems by looking at the distributions of energy levels. We present two conjectures which say something about how to recognize quantum chaotic and non-chaotic behavior.

BGS conjecture

Bohigas, Gionnoni and Schmit [30] conjectured that quantum chaotic systems should have energy levels which follow RMT statistics. The types of systems for which the conjecture should apply are simple quantum systems with a well defined classical limit.

"Spectra of time reversal-invariant systems whose classical analogs are chaotic show the same fluctuation properties as predicted by GOE."

26
For system without time-reversal symmetry, GUE replaces GOE. There has been a few attempts to prove the BGS conjecture, but they all have fallen short. One's confidence in its applicability stems from that it has been verified numerically on a vast number of simple systems. Its general validity has now been largely accepted, and it is standard to use it to statistically address quantum systems which have chaotic behavior. The BGS conjecture reveals a strong connection between classical chaos and RMT. We have already seen in Chap. 3 can be used as a tool to describe spectra of quantum systems which are too complex to solve exactly. Wigner's work on heavy nuclei could thus be viewed as a special case of the BGS conjecture.

**Berry-Tabor conjecture**

The question of quantum integrability was considered by Berry and Tabor [31]. A simple example of a non-chaotic system with many degrees of freedom is an array of independent harmonic oscillators with incommensurable† frequencies. Here the energies will not be correlated with each other and can be viewed as random numbers. Their distribution should then be described by Poisson statistics

\[
p(s) = \frac{1}{\delta} e^{-\frac{s}{\delta}}
\]

Where \(s\) is the spacing between two adjacent energy levels and \(\delta\) a normalization constant. Berry and Tabor postulated that this should be a general feature of quantum integrable systems. The Barry-Tabor conjecture states that for quantum systems whose classical counterpart is integrable, the energy eigenvalues generically behave like a sequence of random numbers, i.e. the spectra obey Poisson statistics. There are known examples where this fails, but then usually as a result of emergent symmetries in the Hamiltonian which lead to extra degeneracies, resulting in commensurability of the spectra.

Poisson and RMT statistics are very different, in that in the former there is no level-repulsion. Poisson spectra tends to cluster, as can be readily seen from Fig. 4.2.

---

† \(a, b \in \mathbb{R}\) are said to be commensurable if \(a/b \in \mathbb{Z}\)
Figure 4.2: Level spacing of some known systems. We note that the zeros of the Riemann zeta function $\zeta(s)$ are believed to be identical to the spectrum of the GUE. We see in e) that the levels in the GUE spectrum are evenly distributed, whereas in the Poisson spectrum in a) the levels tend to cluster. Figure from [34].

4.2.2 Berry’s Conjecture

Berry made a remarkable connection between the structure of energy eigenstates in the semi-classical limit and classical chaos, which is known as Berry’s conjecture. We saw that it is expected that we can apply RMT to the energy level statistics of quantum systems whose classical analog is chaotic, Berry’s conjecture claims that RMT can be used also to describe the wavefunctions of such systems. We will in the following formulate the conjecture and then, following Srednicki [9], outline how Berry’s conjecture imply thermalization when applied to dilute gas of hard spheres. To formulate the hypothesis we need to introduce the Wigner function, which is the Wigner-Weyl transform of the density operator $\rho$

$$W(\vec{x}, \vec{p}) = \frac{1}{(2\pi\hbar)^3} \int d^{3N} \xi \left\langle \vec{x} + \frac{\xi}{2} | \rho | \vec{x} - \frac{\xi}{2} \right\rangle e^{-i\frac{\xi}{\hbar} \vec{p}}$$  \hspace{1cm} (4.6)

Where $\vec{x}$ and $\vec{p}$ are the coordinates and momenta of $N$ particles in three dimensions, spanning a $6N$ dimensional phase space [2]. We could also defined the Wigner function in terms of coherent state variables, angular momentum variables or any other set of canonically conjugate variables. We will in the following restrict ourselves to pure states $\rho = |\psi\rangle\langle\psi|$, but the generalization to mixed states is simple. The Wigner function was introduced by Wigner in [35], when he wanted it link the quantum wavefunction to a probability distribution on phase space and thereby study quantum corrections to statistical mechanics. In the same way as classical probability distributions on phase space, also $W(\vec{x}, \vec{p})$ can be integrated over
momentum to give a probability distribution over $\vec{x}$ as

$$
\int d^{3N} p \, W(\vec{x}, \vec{p}) = |\psi(\vec{x})|^2 
$$

(4.7)

It is however not a true probability distribution since it can become negative, and these regions of negative $W(\vec{x}, \vec{p})$ can be seen as signatures of the presence quantum effects. The Wigner function is uniquely defined for any state and it plays the role of a quasi-probability distribution on phase space. The Wigner function can be used to calculate expectation values of any standard operator $A$ as a phase-space average

$$
\langle A \rangle = \int d^{3N} x \, d^{3N} p \, A_W(\vec{x}, \vec{p}) W(\vec{x}, \vec{p})
$$

(4.8)

Where $A_W(\vec{x}, \vec{p})$ is the Wigner-Weyl transform of the operator $A$ as defined in Eq. (4.6). The Wigner-Weyl transform maps the operator onto phase-space, in a way which allows us to take the phase-space average by integrating over $\vec{x}$ and $\vec{p}$. Berry’s conjecture states that in the semi-classical limit of quantum systems whose classical counterpart is chaotic, the Wigner function of energy eigenstates averaged over a vanishingly small phase space reduces to the microcanonical distribution. To specify, we consider the following locally averaged Wigner function

$$
W_n(\vec{X}, \vec{P}) = \frac{1}{(2\pi\hbar)^N} \int_{\Delta \Gamma_1} dp_1 dx_1 \cdots \int_{\Delta \Gamma_N} dp_N dx_N W_n(\vec{x}, \vec{p})
$$

(4.9)

where $\Delta \Gamma_i$ is a small phase space volume around $X_i$ and $P_i$ and $W_n$ is the Wigner-Weyl transformation of a pure energy eigenstate $\rho_n = |n\rangle \langle n|$. This volume is chosen such that when we take the classical limit of Planck’s constant going to zero, we have both $\Delta \Gamma_i \to 0$ and $\hbar/\Delta \Gamma_i \to 0$ hold. Mathematically we have Berry’s conjecture as

$$
\lim_{\hbar \to 0} W_n(\vec{X}, \vec{P}) = \frac{\delta(E - H(\vec{X}, \vec{P}))}{\int d^{3N} X \, d^{3N} P \, \delta(E - H(\vec{X}, \vec{P}))}
$$

(4.10)

where $H(\vec{X}, \vec{P})$ is the systems classical Hamiltonian. In Berry’s words “$\psi_n(\vec{x})$ are Gaussian random functions in $\vec{x}$ whose spectrum at $\vec{x}$ are the local averages of their Wigner functions $W_n$”. Berry also postulated that the energy eigenfunctions of systems whose classical counterpart is integrable would have a very different structure.

**Application of Berry’s Conjecture**

We now follow Srednicki [9] and consider a system of a dilute gas consisting of identical hard spheres and we will show that the validity of Berry’s conjecture implies that the system will be thermal. It is well known that the classical distribution of momenta for the individual particles of such a system will follow the Maxwell-Boltzmann distribution at a temperature $T$

$$
f_{MB}(p_i, T) = \frac{e^{-\frac{p_i^2}{2mk_BT}}}{(2\pi mk_BT)^{\frac{3}{2}}}
$$

(4.11)
where \( m \) is the mass of the particles and \( k_B \) the Boltzmann constant. This is a quantum system which has a chaotic classical counterpart, since there is no extensive amount of constants of motion. Thus it is reasonable to assume the validity of Berry’s conjecture and we will show that this assumption is enough to “re-derive” the distribution of momenta \( f_{MB}(\vec{p}_i, T) \) for the quantum system. Srednicki used that an energy eigenstate corresponding to a high-energy eigenvalue \( E_n \) can always be chosen to be real and can generally be written as a superposition of plane waves with momentum \( \vec{p}^2 = 2mE_n. \)

\[
\psi_n(\vec{x}) = N_n \int d^3N \ A_n(\vec{p}) \delta(\vec{p}^2 - 2mE_n) e^{i\vec{p} \cdot \vec{x}} \tag{4.12}
\]

where \( N_n \) is a normalization constant and \( A^*_n(\vec{p}) = A_n(-\vec{p}) \). Srednicki introduced the fictitious “eigenstate ensemble” (EE) of energy eigenstates of the system to average over instead of Berry’s average over a small phase-space volume. The eigenstate ensemble should describe the properties of typical energy eigenfunctions and individual eigenfunctions behave as if they were chosen randomly from the eigenstate ensemble. Berry’s conjecture implies that in the eigenstate ensemble \( A(\vec{p}) \) should be a Gaussian random variable and the two-point correlation function is given by

\[
\langle A_m(\vec{p})A_n(\vec{k}) \rangle_{EE} = \frac{\delta^{3N}(\vec{p} + \vec{k})}{\delta(|\vec{p}|^2 - |\vec{k}|^2)} \delta_{nm} \tag{4.13}
\]

Where the denominator is needed for proper normalization. We Fourier transform our wave function

\[
\phi_n(\vec{p}) = (2\pi \hbar)^{-\frac{3N}{2}} \int d^3N x \ \psi_n(\vec{x}) e^{-i\vec{p} \cdot \vec{x}} \\
= (2\pi \hbar)^{-\frac{3N}{2}} \ N_n \int d^3N k \ A_n(\vec{k}) \delta(\vec{k}^2 - 2mE_n) \ \delta^{3N}(\vec{k} - \vec{p}) \tag{4.14}
\]

Where we have introduced \( \delta^{3N}_V(\vec{k}) = (2\pi \hbar) \int_V d^3\vec{x} \ \exp(i\vec{p} \cdot \vec{x}/\hbar) \). We consider the thermal de Broglie wavelength \( \lambda_{th} = (2\pi \hbar^2/(m k_B T))^{1/2} \), which is roughly the average de Broglie wavelength of the particles of an ideal gas. We assume that the \( \lambda_{th} < a \), where \( a \) is the radius of the particles. We also have that \( L^3 \gg a^3N \) since we assumed a low density gas. This gives

\[
\langle \phi^*_m(\vec{p})\phi_n(\vec{k}) \rangle_{EE} = N^2_n (2\pi \hbar)^{3N} \delta_{mn} \delta(\vec{p}^2 - 2mE_n) \ \delta^{3N}_V(\vec{k} - \vec{p}) \tag{4.15}
\]

From this we can calculate the momentum distribution of particles in the eigenstate ensemble. We integrate out the dependence of all but one particle to get the probability distribution of momentum of the single particles

\[
\langle \phi_{nm}(\vec{k_1}) \rangle_{EE} \equiv \int d^3p_2 \ldots d^3p_N \ \langle \phi^*_n(\vec{p})\phi_m(\vec{p}) \rangle = N^2_n L^{3N} \int \int d^3p_2 \ldots d^3p_N \ \delta(\vec{p}^2 - 2mE_n) \tag{4.16}
\]

Where
This should equal the Maxwell-Boltzmann distribution at approximately the temperature given by the ideal gas law \( T = \frac{2U}{3k_B N} \). To show this we to make use of

\[
I_N(q) = \int d^N p \, \delta(p^2 - q) = \frac{(\pi q)^{\frac{N}{2}}}{q^{\Gamma\left(\frac{N}{2}\right)}} \quad (4.17)
\]

Where \( \Gamma(z) \) is the gamma-function\(^†\). Using this we get that the normalization of \( \phi_n(p) \) demands that \( N_n^{-2} = I^{3N}_3 I_{3N}(2mE_n) \). We use this to rewrite Eq. (4.16) and we arrive at

\[
\langle \phi_{nm}(\vec{p}_1) \rangle_{EE} = \frac{I_{3N}(2mE_n - \vec{p}^2_1)}{I_{3N}(2mE_n)} = \frac{\Gamma\left(\frac{3N}{2}\right)}{\Gamma\left(\frac{3(N-1)}{2}\right)} \left( \frac{1}{2\pi mE_n} \right)^{\frac{3}{2}} \left( 1 - \frac{\vec{p}_1^2}{2mE_n} \right)^{\frac{3N-3}{2}} \quad (4.18)
\]

We now substitute the energy \( E_n \) for the temperature corresponding to the energy given by the ideal gas law \( E_n = \frac{3Nk_B T_n}{2} \). We use that \( \Gamma(3N/2)/\Gamma(3(N-1)/2) \) quickly approaches one for large values of \( N \). We are interested in the thermodynamic limit, so we now get

\[
\lim_{N \to \infty} \langle \phi_{nm}(\vec{p}_1) \rangle_{EE} = \left( \frac{1}{2\pi m k_B T_n} \right)^{\frac{3}{2}} e^{-\frac{\vec{p}_1^2}{2m k_B T_n}} \quad (4.19)
\]

Which is indeed the Maxwell-Boltzmann distribution of momentum for thermal particles. Srednicki \cite{9} also showed that if we assume the wavefunction to be completely anti-symmetric or completely symmetric, we instead would have gotten the Fermi-Dirac or Bose-Einstein distribution respectively.

### 4.3 Eigenstate Thermalization Hypothesis

In classical mechanics we know that chaotic systems which cover their phase spaces homogeneously will generally thermalize, and have macroscopic properties which are accurately described by the appropriate thermal ensembles. However there can be no dynamical chaos in closed quantum system, and we therefore need some other way of explaining why and when closed quantum systems can be described by equilibrium statistical mechanics.

The solution seems to be the “eigenstate thermalization hypothesis” (ETH), which we already saw signs of in Berry’s conjecture and Srednicki’s work \cite{9}. In this section we will motivate, define and discuss the ETH.

#### 4.3.1 Background

We start by outlining what sort of systems we will be considering, and more thoroughly define our notation and what we mean by thermalization.

\(^†\)Which for \( \Re\{z\} > 0 \) is defined as \( \Gamma(z) = \int_0^\infty x^{-z} e^x \, dx \)
The System

We assume we have a closed system of many degrees of freedom $N$ described by the Hamiltonian $H$ such that $H|n⟩ = E_n|n⟩$ determines the eigenenergies and energy eigenstates. The system is bounded, ensuring discrete eigenenergies, and closed, which means that the time-evolution is governed by the Schrödinger equation. A general state’s time-evolution can be written as $|ψ(t)⟩ = \sum_n C_n e^{-iE_nt}|n⟩$, where we assume $⟨ψ(t)|ψ(t)⟩ = 1$ and from here on set $\hbar = 1$. We assume that the expectation value of energy in the state we are considering is extensive

$$⟨E⟩ = \sum_n |C_n|^2 E_n \sim N \quad (4.20)$$

Which should be roughly satisfied when interactions are short-range, and we also assume the state to be excited well above the ground state. The uncertainty in energy is assumed to be much smaller than the total energy and to be decreasing inversely with the system size

$$\Delta = \sqrt{⟨H^2⟩ - ⟨H⟩^2} \ll ⟨E⟩ \sim N^{-\nu}⟨E⟩ \quad (0 < \nu \leq 1) \quad (4.21)$$

This seems to be a reasonable assumption since many physically interesting systems scale like this when $N$ is large. Furthermore we assume the spectrum of the Hamiltonian be non-degenerate, which is the case for most chaotic systems. This is consistent with the BGS conjecture which assumes RMT to be valid for quantum chaotic systems and non-degeneracy is indeed a property of RMT-spectra.

We let $A$ be some local operator and $A_{nm} = ⟨n|A|m⟩$ its matrix elements in the energy eigenstate basis. We have the expectation value of $A$ in a general state

$$⟨A(t)⟩ = ⟨ψ(t)|A|ψ(t)⟩ = \sum_n |C_n|^2 A_{nn} + \sum_{n\neq m} C_n^* C_m e^{-i(E_m - E_n)t} A_{nm} \quad (4.22)$$

We define the infinite-time average as

$$\overline{⟨A⟩} = \lim_{t \to \infty} \frac{1}{t} \int_0^t d\tau ⟨A(\tau)⟩ \quad (4.23)$$

We note that the long-time average might be insufficient when considering equilibration, since for large systems the energy levels tend to lie exponentially close, and the equilibration time might thus be unfeasibly long; possibly even longer than that of the universe’s age. We circumvent this problem and use the purely mathematical infinite time average. We have the infinite time average of $A$ in a general state as

$$\overline{⟨A⟩} = \sum_n |C_n|^2 A_{nn} + i\hbar \lim_{\tau \to \infty} \left[ \sum_{n\neq m} C_n^* C_m A_{nm} e^{-i(E_m - E_n)\tau} - 1 \right] \quad (4.24)$$

Where we have used that the spectrum is non-degenerate.
Thermalization

We will consider thermalization in terms of the local operator $A$ and we know that if the system comes to thermal equilibrium the $\langle A \rangle$ should be close to $\langle A \rangle_{th}$ most of the time. We thus define a thermal system as one which fulfills

$$|\langle A \rangle - \langle A \rangle_{th}| \to 0 \land (\langle A \rangle - \langle A \rangle)^2 \to 0$$

(4.25)

For most initial states, where $\langle \cdots \rangle_{th}$ is the thermal average in the appropriate ensemble and the arrow means “in the thermodynamic limit”. The thermodynamic limit is taken by letting number of degrees of freedom go to infinity. More precisely we have

$$N \to \infty \land V \to \infty \land \frac{N}{V} = K$$

(4.26)

Where $V$ the system’s volume and $K$ some constant. In other words, we call a system thermal if, in the thermodynamic limit, the infinite-time average approaches the ensemble average and the average fluctuations approach zero. The thermal average might be given by the micro-canonical ensemble at a given energy $E$ with the quantum statistical average

$$\langle A \rangle_{m.c.}(E) = \frac{1}{N_{\Delta}} \sum_{n: E_n \in \mathbb{X}} A_{nn}$$

(4.27)

Where $\mathbb{X} = [E - \Delta_{m.c.}, E + \Delta_{m.c.}]$, $\Delta_{m.c.}$ the energy width of the microcanonical distribution and $N_{\Delta}$ is the number of states in $\mathbb{X}$. This is the appropriate ensemble for closed systems with given energy, volume and particle number. For systems at inverse temperature $\beta$ we might use the canonical ensemble

$$\langle A \rangle_{can}(\beta) = \frac{1}{Z} \sum_n A_{nn} e^{-\beta E_n}$$

(4.28)

Where $Z = tr\{e^{-\beta H}\}$ is the standard partition function. The canonical ensemble should be used when we have given temperature, particle number and volume, i.e. for systems in contact with a heat bath. We will in the following switch between the ensembles rather uncritically. The assumption underlying this is that for a system of very many degrees of freedom and for most non-pathological operators $A$, the difference between the prediction of the canonical ensemble at given $\beta$ and the microcanonical ensemble at given $E$ is small. The different ensembles do indeed yield exactly the same averages in the thermodynamic limit.

4.3.2 Eigenstate Thermalization Hypothesis

We will consider what it takes for a closed quantum system to thermalize and put forth the ETH which should tell us which systems are thermal. For a closed system to be thermal we need the the following to be satisfied in the thermodynamic limit

$$\langle A \rangle = \sum_n |C_n|^2 A_{nn} = \frac{1}{N_{\Delta}} \sum_{n: E_n \in \mathbb{X}} A_{nn} = \langle A \rangle_{m.c.}(\langle E \rangle)$$

(4.29)
We also need the fluctuations to be small
\[
(\langle A \rangle - \langle A \rangle)^2 = \sum_{n \neq m} |C_n|^2 |C_m|^2 |A_{nm}|^2 \leq 0
\] (4.30)

We note that whereas the time averaged expectation value is determined by the initial state through the \(C_n\)'s, the microcanonical prediction makes no reference to the initial state as it is determined completely by \(\langle E \rangle\), which can be the same for many different initial states.

If the eigenstate expectation values \(A_{nn}\) practically do not fluctuate at all between eigenstates that are close in energy, Eq. (4.29) is satisfied for literally all states which are sufficiently narrow in energy, i.e. for which \(\Delta\) is small. We thus need \(A_{nn}\) to be smoothly varying with \(n\) and effectively constant over the relevant small energy window. We know that both \(\Delta_{m.c.}\) and \(\Delta\) are decreasing with increasing system size, and thus for large systems we can use that \(A_{nn}\) varies very slowly with \(n\) to draw it outside the sum. Furthermore this implies that also energy eigenstates themselves are thermal, i.e. we have \(A_{nn} = \langle A \rangle_{m.c.}(E_n)\). This is what we call eigenstates thermalization.

We consider the fluctuations and immediately see that for Eq. (4.30) to be satisfied for all initial states we need the off-diagonal matrix elements \(A_{nm}\) to be small. That is, the system thermalizes if
\[
|A_{n+1,n+1} - A_{n,n}| \to 0 \quad \land \quad \forall n \neq m \quad |A_{nm}| \to 0
\] (4.31)

We notice a similarity between the form of the eigenstate matrix elements imposed on \(A\) by Eq. (4.31) and the expectation values of operators in the GOE. In the GOE the expectation values of local operators on matrix form\(^1\) has the average value of the operator on the diagonal and the off-diagonal fluctuations are decreasing inversely with the system size. By analogy with Eq. (4.31) and Eq. (3.24) we now arrive at the ansatz of the ETH as defined by Srednicki \([10]\).

The ETH Ansatz

The ETH ansatz states that local observables should have matrix elements in the eigenstate basis whose statistics should be described by
\[
A_{nm} = \mathcal{A}(\bar{E}) \delta_{nm} + e^{-\frac{s(E)}{2}} g(\bar{E}, \omega) R_{nm}
\] (4.32)

We first define all the quantities of the equation; we let \(\bar{E} = (E_n + E_m)/2\), \(\omega = (E_n - E_m)\) and \(R_{nm}\) be a factor which varies erratically in \(n\) and \(m\)\(^1\). Furthermore we demand \(g(\bar{E}, \omega)\) and \(\mathcal{A}(\bar{E})\) to be smooth functions of their arguments, where the former is an envelope function on top of the fluctuations and the latter can be viewed as a spectral function of \(A\). Without loss

\(^1\)I.e. the statistical expectation value over an ensemble expressed in a fixed basis.

\(^1\)We assume \(R_{nm}\) to essentially be i.i.d. Gaussian numbers.
of generality we assume \( g(\bar{E}, \omega) \) to be positive, real and an even function of \( \omega \). We demand that \( R_{nm} \) has zero mean and unit variance, and the hermiticity of \( A \) implies that \( R_{nm} = R_{nm}^\ast \).

The thermodynamic entropy \( S(\bar{E}) \) at energy \( \bar{E} \) is defined by the density of states as

\[
e^{S(\bar{E})} = E \sum_n \delta(\bar{E} - E_n)
\]

where the Dirac delta function has been smeared out enough to ensure monotonicity of \( S(\bar{E}) \). The regularized Dirac delta function could be defined e.g. as

\[
\delta_\eta(x) = \frac{1}{\pi \eta^2 + x^2}
\]

(4.33)

For some small \( \eta \). The ETH ansatz differs from the RMT prediction only in that the diagonal elements are not all the same and that the small Gaussian fluctuations of the off-diagonal elements have an envelope function \( g(\bar{E}, \omega) \). The ETH ansatz reduces to the RMT form if we focus on a very narrow energy range where \( g(\bar{E}, \omega) \) is constant. The scale at which ETH and RMT are equal seems to be given by the Thouless energy \( E_T \) [39]. That is, if we only consider energies such that \( \omega < E_T \) it holds that \( g(\bar{E}, \omega) \) is approximately constant and that the ETH ansatz is identical to RMT. The Thouless energy goes to zero in the thermodynamic limit, which implies that RMT has a limited range of applicability. However the level spacing decreases faster in \( L \) than the Thouless energy and thus there are still a large number of energy levels in the region where RMT applies. Conversely the ETH ansatz applies to arbitrary energies, with the possible exception of the edges of the spectrum.

A striking feature of the ETH is that knowledge of a single energy eigenstate is sufficient to compute thermal averages and any eigenstate in the microcanonical window will do, since they all give the same average.

We show explicitly in Appendix A that when the matrix elements are well described by the ETH ansatz, the system is indeed thermal. Although the ETH implies that the system thermalizes, we do not know if the ETH necessarily must hold for all thermal systems. There has been some numeric evidence for that thermal quantum systems obey the ETH [36]. It is however hard to study this thoroughly because the only method of investigation is numerically costly exact diagonalization. We can therefore only test the ETH for very small systems.

4.4 Quantum Thermalization

We now address some issues of the thermal equilibration process in closed quantum systems.

In classical, thermal systems, most initial out-of-equilibrium states approach thermal equilibrium as the states are evolved in time. This thermal equilibrium is characterized by a small amount of parameters corresponding to the extensive conserved quantities such as temperature, chemical potential etc. Most information about the initial state of the system gets lost during the time evolution.

We do however know that closed quantum systems are governed by the linear Schrödinger equation. Thus all information about the initial state must be contained in the system at all times, since unitary time evolution cannot erase information.
In order to take a closer look at this we consider the density operator of the system \( \rho(t) \), which we use to find expectation values as \( \langle A(t) \rangle = \text{tr}\{\rho(t)A\} \). Naively we might expect that the system coming to equilibrium, means that the initial states evolve into the Maxwell-Boltzmann distribution \( \rho(t) \to \exp(-\beta H)/Z \), but this is not the case. To see why, we consider \( \rho(t) \) in terms of its matrix elements in the eigenstate basis

\[
\rho(t) = \sum_n \rho_{nn}(0)|n\rangle\langle n| + \sum_{n \neq m} \rho_{nm}(0)e^{-i(E_m - E_n)t}|m\rangle\langle n|
\] (4.34)

We see that the dynamics is simple and that the time evolution merely makes the off-diagonal matrix elements precess in the complex plane at a constant rate given by the energy difference between the eigenstates involved. Generally the full system described by \( \rho(t) \) does therefore not approach \( \exp(-\beta H)/Z \) at long times; i.e. the distributions of the thermodynamic ensembles are not attractors of the dynamics.

We have in Eq. (4.25) considered quantum thermalization in terms of local observables, and thermalization thus requires that the contributions from the off-diagonal matrix elements to local observables must vanish at long times. Due to the ETH, all energy eigenstates implicitly contain thermal states, i.e.

\[
\text{tr}\{|n\rangle\langle n|A\} = \text{tr}\left\{\frac{e^{-\beta H}}{Z}A\right\}
\] (4.35)

However we can prepare the system in out-of-equilibrium states by giving the phases in \( \rho(0) \) some special structure.

Thermalization of out-of-equilibrium states thus happens because at long times the off-diagonal terms in \( \rho(t) \) come with effectively random phases and their contribution to expectation values of local operators cancel. Thus states for which \( \langle A(0) \rangle \neq \langle A \rangle_{\text{th}} \) will after a long time have dephasing between the off-diagonal contributions and therefore \( \langle A(t) \rangle = \langle A \rangle_{\text{th}} \) at long times. In other words the thermal properties of out-of-equilibrium states are hidden by the coherence between the eigenstates, but the time evolution reveals it through dephasing.

Thus we see a distinction from classical mechanics where the time evolution "creates" a thermal state, which has forgotten everything about its initial state. A quantum system does not loose any information about the local properties of its initial state, the information is merely "hidden" somewhere else in the system. What we mean by "hiding" the information, is that it is made inaccessible, in the sense that recovering it means measuring global operators.

We started this section with stating that some formulations of quantum statistical mechanics postulate that the system in question is in contact with an external heat bath. A viewpoint which we dismissed, largely because we would rather consider truly isolated systems. However we can think about thermalization of closed quantum systems as the

\[\text{Or some other equilibrium distribution.}\]
different parts of the system acting as heat baths for each other. We consider a small subsystem $S$ with density operator $\rho_S(t)$. The density operator on $S$ does not have simple dynamics since it is an open system with non-unitary time evolution. We thus define thermalization as:

$$\lim_{t \to \infty} \rho_S(t) = \text{tr}_B \left\{ e^{-\beta H_Z} \right\}$$

(4.36)

Where $B$ contains all the degrees of freedom which are not in $S$. Even though $\rho(t)$ does not approach the Maxwell-Boltzmann distribution, the density matrices of small subsystems appear as if it does. The system can equilibrate under its own dynamics if its different parts are able to interact in a way which mimics the interaction to a thermal bath. Information is moved around in the system, and this makes all the subsystems come to a thermal equilibrium described by Eq. (4.36).

---

$^1$Some care need to be taken when taking the thermodynamic limit; it should be taken simultaneously with the infinite time limit, since finite size systems cannot equilibrate due to quasi-periodic dynamics and infinite systems need an infinite amount of time to equilibrate.

$^{††}$We do however note that the $S$ and $B$ are strongly interacting, so contrary to systems which are weakly coupled to a heat bath, the Hamiltonian appearing in $\rho_S$ is the full system+bath-Hamiltonian, not just $H_S$.

$^1$We note that the equilibration times could presumably differ between different parts of the system.
Chapter 5

Model and Methods

5.1 The Model

We will in the following be working with the isotropic Heisenberg Hamiltonian with random on-site magnetic fields along the $z$-direction

$$H = \sum_{i=1}^{N} S_i^z h_i + J \sum_{\langle i,j \rangle} (\vec{S}_i \cdot \vec{S}_j)$$

(5.1)

Where $J$ is the exchange interaction and the $h_i$'s are random fields. The distribution of $h_i$ have some width $W$ and we choose its probability distribution to be uniform and centered at zero. We have the standard spin operators $S_z = \sigma_z / 2$ in terms of Pauli operators and all operators are filled out with identity operators to a tensor product of $N$ operators. E.g. for a four-particle system: $S_2^z S_3^z = I \otimes S^z \otimes S^z \otimes I$. We can rewrite the Hamiltonian in a more useful form

$$H = \sum_{i=1}^{N} S_i^z h_i + J \sum_{\langle i,j \rangle} (S_i^z S_j^z + S_i^y S_j^y + S_i^x S_j^x)$$

(5.2)

$$= \sum_{i=1}^{N} S_i^z h_i + J \sum_{\langle i,j \rangle} S_i^z S_j^z + \frac{1}{2} (S_i^- S_j^+ + S_i^+ S_j^-)$$

Where we have used the spin raising and lowering operators $S_i^\pm = S_i^x \pm i S_i^y$. We will in the following assume periodic boundary conditions and work in one dimension, in which the Hamiltonian looks like

$$H = \sum_{i=1}^{N} \frac{h_i}{2} \sigma_i^z + \frac{J}{4} \sigma_i^z \sigma_{i+1}^z + \frac{J}{2} (\sigma_i^+ \sigma_{i+1}^- + \sigma_i^- \sigma_{i+1}^+)$$

(5.3)

We will now map this onto a fermionic problem and show how this is related to the model Andersen discussed in his paper [1]. The fermionic creation and annihilation operators $c_j^\dagger$ and $c_j$ have the following anti-commutation relations

$$\{c_j^\dagger, c_i\} = \delta_{ij} \land \{c_j, c_i\} = \{c_j^\dagger, c_i^\dagger\} = 0$$

(5.4)
The spin raising and lowering operators have similar commutation relations and we wish to exploit this fact to make a change of variables. The same site anti-commutation relations for raising and lowering Pauli operators are \( \{ \sigma_j^+, \sigma_j^- \} = 1 \) just like the fermionic operators. However on different sites the Pauli operators commute \( [\sigma_j^+, \sigma_i^-] = 0 \), whereas the fermionic operators anti-commute. We thus need to rectify that the different sites commute differently for the two sets of operators. We must use the Jordan-Wigner transformation, which is defined as

\[
\sigma_j^+ = e^{+i\pi \sum_{k<j} n_k} c_j^+ \\
\sigma_j^- = e^{-i\pi \sum_{k<j} n_k} c_j \\
\sigma_j^z = 2n_j - 1
\] (5.5)

Where \( n_j = c_j^+ c_j \) are the number operators. The spin operators and fermionic operators commute identically after the Jordan-Wigner transformation. Before we insert these operators into our Hamiltonian, we need the some in-between results. First we note that we can Taylor expand and rewrite the exponential operator, and show that it commutes with the fermionic operators

\[
e^{+i\pi \sum_{k<j} n_k} c_j^+ = \prod_{k<j} \left[ 1 + c_k^+ c_k \sum_{l=1}^{\infty} \frac{(i\pi)^l}{l!} \right] c_j^+ = c_j^+ e^{+i\pi \sum_{k<j} n_k}
\] (5.6)

Where we have exploited the anti-commutation relations of \( c_j^+ \)'s and that the \( n_k \)'s are fermionic with spectrum \{0, 1\}. Using this we get

\[
\sigma_j^+ \sigma_{j+1}^- = e^{+i\pi \sum_{k<j} n_k} c_j^+ e^{-i\pi \sum_{k<j+1} n_k} c_{j+1} \\
= c_j^+ e^{-i\pi n_j} c_{j+1} = c_j^+ c_{j+1} \\
= c_j^+ \left[ 1 + c_j^+ c_j \sum_{l=1}^{\infty} \frac{(i\pi)^l}{l!} \right] c_{j+1} = c_j^+ c_{j+1}
\]

\[
\sigma_j^- \sigma_{j+1}^+ = c_j e^{i\pi n_j} c_{j+1}^+ \\
= c_j \left[ 1 + c_j^+ c_j \sum_{l=1}^{\infty} \frac{(i\pi)^l}{l!} \right] c_{j+1}^+ \\
= \left[ 1 + \sum_{l=1}^{\infty} \frac{(i\pi)^l}{l!} \right] c_j c_{j+1}^+ = -c_j c_{j+1}^+
\] (5.7)
We can now readily map our spin-1/2 Hamiltonian onto a fermionic model. We get

\[ H = \sum_{i=1}^{N} \frac{h_i}{2}(2n_i - 1) + \frac{J}{4}(2n_i - 1)(2n_{i+1} - 1) + \frac{J}{4}(c_i^\dagger c_{i+1} - c_i c_{i+1}^\dagger) \]

\[ = \sum_{i=1}^{N} (h_i - J)n_i + Jn_i n_{i+1} + \frac{J}{2}(c_i^\dagger c_{i+1} - c_i c_{i+1}^\dagger) + \frac{J}{4}N - \sum_{i=1}^{N} h_i \]  

(5.8)

\[ = \sum_{i=1}^{N} \tilde{h}_i n_i + Jn_i n_{i+1} + \tilde{J}(c_i^\dagger c_{i+1} - c_i c_{i+1}^\dagger) \]

Where we have rescaled the energy by a constant and redefined the quenched disorder \( \tilde{h}_i \) and the interaction strength \( \tilde{J} \). Anderson considered the non-interacting case by linearizing the Hamiltonian in number operators, which allowed him to consider one particle at the time. This corresponds to disregarding the coupling between the z-components of spin in the Hamiltonian of spin-1/2 Hamiltonian. We have thus showed that if there is only one fermion in the system, which corresponds to haveing all but one spin pointing in the same direction along the z-axis in the spin model, we get

\[ H = \sum_{i=1}^{N} \tilde{h}_i n_i + \tilde{J}(c_i^\dagger c_{i+1} - c_i c_{i+1}^\dagger) \]

(5.9)

Because the terms with \( c_i^\dagger c_{i+1}^\dagger c_{i+1} c_i \) obviously vanish in the one-particle setting. This is exactly the Anderson model in one dimension, which means that when we study the numerics of our MBL Hamiltonian we can consider the Anderson Hamiltonian by simply setting the coupling between spin-z’s equal zero or by only considering initial states where all but one spin point in the same direction along the z-axis.

### 5.2 Numerical Methods

We will study the Hamiltonian defined in Eq. (5.3) numerically. Out method of investigation will be exact diagonalization of many matrices with different disorder realizations. The matrices are sparse since the interactions are short range, but we generally need all the eigenvalues and eigenvectors, or at least all of the states in the middle of the spectrum. This excludes methods such Lanczos algorithm, since it only yields the extreme eigenvalues and eigenstates, which means that we cannot really benefit from the sparsity of the Hamiltonian matrix. Monte Carlo methods are also not applicable since the MBL phase is not thermal. There have indeed been some attempts at studying MBL numerically through density matrix renormalization group schemes [46,47], but exact diagonalization is still the only reliable way study MBL systems numerically.

Exact diagonalization is numerically very costly and the computational requirements scale as \( O(N^3) \), where \( N \) is the size of our Hilbert space. The Hilbert space scales as
$2^L$ and seeing that we can only diagonalize matrices of the order $\mathcal{O}(10000 \times 10000)$ with exact diagonalization, we are restricted to only consider short spin chains. We use a binary representation of the spin-$z$ product state basis states to find the matrix elements of the Hamiltonian and to calculate the eigenstate expectation values, and we use the LAPACK library to perform the diagonalization. Some elements of the C++ program which was used are found in Appendix E.

The Hamiltonian has conserved total spin along the $z$-direction. We can utilize this by choosing our basis in a way that makes the Hamiltonian block diagonal. This allows us to diagonalize the different spin-sectors separately, and thereby consider larger systems. We will mostly only consider the sector where total spin along the $z$-axis equals zero. This means that half the spins point up and the other half down, and there are $\binom{L}{L/2}$ ways to place the spins pointing up. Our Hilbert space thus shrinks from $2^L$ to $L!/\left(\frac{L}{2}\right)!^2$. This is the largest spin sector and we presume that it will mimic the behavior of the full system well. When we restrict ourselves to states with zero total spin, we can investigate spin chains of length up to $L = 16$.

The exact diagonalization yields a huge amount of data. This means that although we can only work with rather small systems we still have the possibility of obtaining interesting information about the system, since we can average over a vast number of eigenstates and disorder realizations. So if we ask the system clever questions we can indeed investigate the different behavior in the two phases. We have used from roughly 10000 realizations of disorder for $L = 8$ and about 50 realizations for $L = 16$. However we have on some occasions needed to consider more disorder realizations in the critical region near the phase transition. In our figures we show one-standard deviation error bars.

Whereas the perturbative results on MBL are valid for systems at low temperature and low energy densities, we work with states with high energy densities and at infinite temperature. We will only use the states with energies in the middle of the spectrum; unless otherwise specified we have used the states from the middle one-third of the energy spectrum. We work in the limit infinite temperatures to remove one parameter from our problem, and this implies that all the Boltzmann weights are equal. That is, we weight every eigenstate from every disorder realization equally when averaging over eigenstates.

At infinite temperatures we cannot distinguish between the paramagnetic and the diamagnetic case so the sign of $J$ is irrelevant, and we will set $J = 1$ in the following. Furthermore the distinction between fermions and bosons vanish in this limit since both the Fermi-Dirac distribution and the Bose-Einstein distribution goes to the classical distribution function.
5.3 Local Integrals of Motion

It is a common belief that the failure of ergodicity in the MBL phase is related to some sort of mechanism which is similar to integrability. By analogy to classical integrable systems it is thus assumed that in the MBL phase it is generally satisfied that

(i) There exist as many integrals of motion in involution as degrees of freedom

(ii) The constants of motion forms a complete set of quasi-local operators

It has not been proven that this must always be the case for MBL systems, but the phenomenology of MBL can be deduced from these two assumptions. So a system which supports local integrals of motion (LIOM’s) should be MBL. From assumption (i) there exists a complete set of $N$ local integrals of motion $\tau_i$ which satisfies

$$[\tau_i, \tau_j] = [\tau_i, H] = 0 \quad (5.10)$$

We can exploit that $H$ and $\tau_j$ have vanishing commutators, and therefore simultaneous eigenvalues, to write the Hamiltonian as some linear combination of $\{\tau_j\}$. That is, the Hamiltonian is a non-linear functional of a complete set of integrals of motion as

$$H = \sum_{n=1}^{N} J_n \tau_n + \sum_{n,m=1}^{N} J_{nm} \tau_n \tau_m^z + \sum_{n,m,k=1}^{N} J_{nmk} \tau_n \tau_k \tau_m + \sum_{n,m,k,l=1}^{N} J_{nmkl} \tau_n \tau_k \tau_l \tau_m + \ldots \quad (5.11)$$

This expansion is however completely generic, insofar that all Hamiltonians can be written on the form of Eq. (5.11), if we for instance choose the $\tau_i$’s to be the projection operators onto the eigenstates of the Hamiltonian $P_n = |n\rangle\langle n|$. However from assumption (ii) these operators are quasi-local, meaning that they mainly act on a small region of space. More precisely we say that the operator norm of $\tau_j$ decay exponentially away from a region of size $\xi$ around the localization center $R_j$. We expand the quasi-local operators $\tau_i$’s in terms of strictly local operators $O_\alpha$’s as

$$\tau_j = \sum_{\alpha} K^j_\alpha O_\alpha \quad (5.12)$$

If we let $S(O_\alpha)$ be the support of the operator $O_\alpha$, i.e. the set of local degrees of freedom on which the operator act non-trivially, then quasi-locality is expressed as

$$|K^j_\alpha| \lesssim A_j e^{\max(R_j-S(O_\alpha))\xi} \quad (5.13)$$

In other words the contribution to the expansion by the local operators is suppressed by the element in their support which is the furthest away from $R_j$. This entails that $\tau_j$ is itself localized around $R_j$ and can be viewed as a weak deformation of the local, physical degrees of freedom $O_\alpha$, which can be e.g. the spin operators $S_i$ or the number operators $c_i^\dagger c_i$. The
set of LIOM’s is complete in the sense that we can label all of the $2^N$ eigenstate in terms of the eigenvalues of the $\tau_j$’s as $|\Psi_{\tau_1\tau_2...\tau_N}\rangle$. The quasi-locality of $\tau_j$’s entails that the $J$’s in Eq. (5.11) decay exponentially with the distance between the localization centers of the $\tau_j$’s. Due to the quasi-locality of the $\tau_j$’s MBL is, unlike integrability, robust to small, local perturbations of the Hamiltonian.

The the set of conserved and mutually commuting quantities is by no means unique. It is usually assumed that we can choose the LIOM’s such that the full LIOM algebra can be constructed through raising and lowering operators and that they have a binary spectrum. This was brought in as an additional assumption in e.g. Ref. [48], but followed straightforwardly from the constructions in Refs. [49, 50]. We will now very briefly review two examples of systems which support LIOM’s, one very simple and one rather involved.

**Single-Particle Localization**

It is not very hard to realize that the Anderson model supports LIOM’s and we will here briefly discuss how to make sense of Anderson localization in terms of LIOM’s. In the non-interacting case we have the Hamiltonian expressed in fermionic operators as in Eq. (2.2). In this case the single-particle level occupation number operators $n_\beta$ are our LIOM’s. That these operators are quasi-local follows immediately from their expansion in the basis of lattice operators

$$n_\beta = \sum_{i,j} \phi_\beta^*(i)\phi_\beta(j) c_i^\dagger c_j \tag{5.14}$$

where $\phi_\beta$ are the single-particle localized eigenfunctions with localization centers at $\vec{r}_\beta$. Due to the localization of the wave functions the operators $c_i^\dagger c_j$ contributes to $n_\beta$ with a weight which vanishes exponentially in the distance between its support and the localization center $R_\beta$. We can thus write the Anderson Hamiltonian as

$$H_A = \sum_{\beta=1}^N \epsilon_\beta n_\beta \tag{5.15}$$

which is on the same form as Eq. (5.11), but with the $J$’s of the higher order terms equal to zero.

**Ising Model**

John Imbrie [49] proved that MBL does indeed occur in an one-dimensional Ising model on an interval $\Lambda = [-K,K'] \subset \mathbb{Z}$, with random fields, random transverse fields and random exchanges. The Hamiltonian for the system is

$$H_I = \sum_{i=-K}^{K'} h_i S_i^z + \sum_{i=-K}^{K'} \gamma_i S_i^x + \sum_{i=-K-1}^{K'} J_i S_i^z S_{i+1}^z \tag{5.16}$$
Where $\gamma_i = \gamma \Gamma_i$ with $\gamma$ small and $\gamma_i$, $h_i$ and $J_i$ are random bounded and independent variables. The way he proved MBL for such a system was to show that this particular model supports LIOM’s. This is done by performing a set of quasi-local unitary rotations $\Omega$. These rotations transform the Hamiltonian into the form of Eq. (5.11)

$$\Omega^\dagger H_I \Omega = -\sum_{n=1}^{N} J_n S_n^z - \sum_{n,m=1}^{N} J_{nm} S_n^z S_m^z + \sum_{n,m,k=1}^{N} J_{nmk} S_n^z S_k^z S_m^z + \ldots$$ (5.17)

These rotations also show how the LIOM’s are related to the physical spin operators, namely $\tau_k = \Omega S_j^z \Omega^\dagger$ and $\tau_k^\pm = \Omega S_j^\pm \Omega^\dagger$. For completeness it should be mentioned that Imbrie could only prove MBL under a certain assumption, namely limited level attraction. It is a rather mild physical assumption, saying that with high probability the smallest spacing between two energy levels should be no smaller than some exponential in volume. There are no known generic systems whose spectrum violate the assumption of limited level attraction.
Chapter 6

Many-body Localization

We will in this section show numerically that there are two distinct phases in our model, namely a thermal phase for small disorder strengths and a MBL phase for sufficiently large disorder. We discuss and show different properties of the MBL phase and contrast them to the properties of thermal systems. We also want to find an estimate for the critical value of disorder $W_C$. We also discuss the phenomenology of MBL in terms of the LIOM’s.

We start by considering in which way the MBL states are localized.

6.1 Localization in Configuration Space

In much the same way as Anderson localized states are localized in real space, we can think of MBL states as being localized in configuration space. This was shown to be true for the fermionic system considered by Basko et al. [2]. They showed that the MBL eigenstates are localized in the fermionic Fock space of Anderson localized states, which is a standard configuration space basis for systems of identical particles.

For our spin-Hamiltonian the configuration space basis consists of the non-entangled product states on the form $|\alpha\rangle = |+\ldots\rangle$, where $|\pm\rangle$ are the eigenstates of $S_z$. We will in the following use greek indices to represent the configuration space basis states and latin indices to represent the energy eigenstates. Since our spin-Hamiltonian is linked to the fermionic one through a Jordan-Wigner transformation and the non-entangled product states and the Fock space states both are the basis states for the respective non-interacting systems, these two configuration space bases are closely linked.

We will show that in the MBL phase the energy eigenstates are localized in configuration space, in the sense that they are linear combinations of only a few of the non-entangled product states. Thus sharply localized eigenstates are eigenstates for which $|n\rangle \approx |\alpha\rangle$.

When we assume that our system supports LIOM’s, the exponential decay of the amplitudes in Eq. (5.13) around the localization centers $R_j$ can be put in an one-to-one correspondence with states localized in the fermionic Fock space of single-particle localized
states or with the non-entangled product states in the spin case [50] [51].

To show localization numerically, we start by investigating the expectation value of the magnetization 
\( m_{i,n} \equiv \langle n | S^z_i | n \rangle \) at site \( i \) in energy eigenstate \( |n\rangle \). If the system is thermal we expect Berry’s conjecture to hold and that the energy eigenstates should have weights which are as random as possible when expressed in a fixed basis. Thus for small disorder we expect the eigenstates to be linear combinations of many product states where the weights are Gaussian random numbers, which implies that 
\( m_{i,n} \approx 0 \) since we are in the \( S^z_{\text{tot}} = 0 \) -sector. In the MBL phase the eigenstates should be localized around only a few of the product states and we therefore expect 
\( m_{i,n} \approx \pm 1/2 \).

\[
\sum_{\alpha} = 1 |\langle \alpha | \psi \rangle|^2q
\]

\[
PR_q = \frac{1}{\sum_{\alpha=1}^N |\langle \alpha | \psi \rangle|^{2q}}
\]

Figure 6.1: Distribution of the magnetization \( m_{i,n} \) for the system in an eigenstate in the thermal phase (a) and in the MBL phase (c) and roughly at the phase transition (b). The systems consists of 12 spins and the disorder distribution have a width of \( W \)

We see from Fig. 6.1a that for small disorder the distribution of \( m_{i,n} \) is Gaussian and centered around zero, which is consistent with the eigenstates satisfying Berry’s conjecture in the thermal phase. In the MBL phase we see from Fig. 6.1c that the distribution of \( m_{i,n} \) is sharply peaked around \( \pm 1/2 \), which is consistent with the eigenstates being approximate non-entangled product states.

We also note that from Fig. 6.1b we see that in between the MBL and the thermal phase the distribution have peaks at both \( m_{i,n} = 0 \) and at \( m_{i,n} = \pm 1/2 \). After these first indications of localization in configuration space, we move on to discuss more quantitative measures of localization, starting with the participation ratio.

### 6.1.1 Participation Ratios

The participation ratio is a measure of how many of the basis states \( \{|\alpha\rangle\} \) are contributing to a general state \( |\psi\rangle \), which we assume to be normalized. We define it as

\[
PR_q = \frac{1}{\sum_{\alpha=1}^N |\langle \alpha | \psi \rangle|^{2q}}
\]

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In the following we will mainly be interested in $PR_2$. We see that if only one basis state contributes to $|\psi\rangle$ then we have $PR_2 = 1$ and if all $N$ states contribute equally then $PR_2 = N$. We note that in the literature the participation ratio as defined in Eq. (6.1) is on some occasions called the inverse participation ratio. Higher order $PR_q$’s can be useful for describing finer details of the system. For Anderson localization, it is well known that the wavefunctions will fluctuate strongly in the vicinity the transition, and these fluctuations can be described by the set of $PR_q$’s.

We start by briefly looking at $PR_2$ in the context of Anderson localization. It is known that we can distinguish the single-particle Anderson localized phase from the delocalized phase, by its eigenstates satisfying $PR_2 = O(1)$ [38], and the $PR_2$ as a measure of localization was discussed already by Thouless and Edwards almost 50 years ago [16].

![Figure 6.2: Participation ratio of the energy eigenstates as a function of disorder strength. The average is over eigenstates and disorder realizations. We have used the eigenstates of our Hamiltonian in the $S_{\text{tot}}^z = (L/2 - 1)$-sector. System sizes in legend.](image)

We will show numerically that Anderson localized states have $PR_2 = O(1)$. To do this we consider our Hamiltonian under the restriction that all spins except one are pointing in

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$^1$A multifractal structure emerges. At criticality $PR_q \propto L^{-D_q(q-1)}$, where the scaling is characterized by an infinite set of critical exponents $D_q$. This should be contrasted with $PR_q \propto L^{-d(q-1)}$ and $PR_q \propto L^0$ in the thermal and MBL phase respectively [38]. Multifractality is characteristic of a variety of complex systems; e.g. turbulence and strange attractors. The scale invariance and self-similarity of fractals suggests that there might possibly be a more general connection between phase transitions and fractal behavior.
the same direction, i.e. we are looking at the \( S^z_{\text{tot}} = (L/2 - 1) \) -sector. We let \(|\psi\rangle\) be an energy eigenstate and \(|{\alpha}\rangle\) the non-entangled product states in the \( S^z_{\text{tot}} = (L/2 - 1) \) -sector. This corresponds to only having one fermion in the fermionic system, and is equivalent to the Anderson model.

We see from Fig. 6.2 that the participation ratio of the Anderson localized eigenstates does not appear to have any systematic increase or decrease with system size, which indicates that we have \( PR_2 = \mathcal{O}(1) \). We also see that the participation ratio decreases with increasing \( W \). That is, for increasing disorder the eigenstates are localized on fewer and fewer sites, corresponding to shorter localization lengths. At very large \( W \) the localization length is comparable to the lattice spacing.

When discussing the scaling theory to Anderson localization we saw that in one dimension, the system should be localized for all disorder strengths in the thermodynamic limit, and this does indeed appear to hold for our model. We have however only considered disorder strengths down to \( W = 0.5 \), because for very small systems the localization lengths can be very large, even comparable to the size of the system. Thus for very small disorder finite-size effects might make such small systems appear delocalized.

We now turn our attention back to the MBL case, and we wish to consider if we can distinguish the MBL from the thermal phase using the \( PR_2 \). We therefore look at our model in the \( S^z_{\text{tot}} = 0 \) -sector and calculate the participation ratios of the MBL eigenstates. We start by looking at how the probability distribution of \( PR_2 \) in the MBL phase changes when we increase \( L \). We see from Fig. 6.3 that the participation ratio does not scale as \( \mathcal{O}(1) \), on the contrary it gets larger for increasing system sizes. Since \( PR_2 \) is increasing with increasing \( L \) it is problematic to use it to distinguish the thermal from the MBL phase.

The reason for the participation ratio’s increase with increasing \( L \) in the MBL phase is that, even though we have exponentially decaying amplitudes in configuration space, we also have a rapidly growing number of states which can lie inside the localization radius of a MBL state. Thus since the many-body Hilbert space is growing exponentially with the number of spins, the states will have to lie exponentially close to each other in configuration space and thus the number of states which lie inside of the localization radius of any given MBL state will increase with the system size.
Figure 6.3: Distribution of participation ratios for MBL energy eigenstates for increasingly large systems, which all have quenched disorder from a distribution with a width $W = 5$. System sizes in legend.

We will therefore instead consider the following related function

$$P_H = \frac{PR_2}{N}$$

(6.2)

Where $N$ is the size of the Hilbert space. Whereas there is an obvious physical interpretation of $PR_2$ ready at hand, it is not equally obvious which physical significance $P_H$ has, other than a rescaled participation ratio. Therefore we will show that we can relate $PR_2$ and $P_H$ to the quantum dynamics and argue that $P_H$ can be used to pinpoint when the system enters the MBL phase.

We consider the survival probability encoded in $G(t) = \langle \psi(0)| e^{-iHt} |\psi(0)\rangle$, where $|\langle \psi(0)| e^{-iHt} |\psi(0)\rangle|^2$ quantifies the probability for a state to remain the same after some time $t$ has passed. We start by showing how the survival probability of a basis state $|\alpha\rangle$ in the configuration space is linked to $PR_2$. We calculate the infinite-time average of $|G(t)|^2$, where the infinite-time average is taken in order to remove finite-size effects like quantum recurrence.
$$\overline{S} = \lim_{\tau \to \infty} \frac{1}{\tau} \int_{0}^{\tau} dt \left| G(t) \right|^2$$

$$= \lim_{\tau \to \infty} \frac{1}{\tau} \int_{0}^{\tau} dt \left| \langle \alpha | e^{-iHt} | \alpha \rangle \right|^2$$

$$= \sum_{nm} |\langle \alpha | m \rangle|^2 |\langle n | \alpha \rangle|^2 \lim_{\tau \to \infty} \frac{1}{\tau} \int_{0}^{\tau} dt \ e^{-i(E_n - E_m)t}$$

$$= \sum_{n} |\langle n | \alpha \rangle|^4 = PR_2^{-1} \quad (6.3)$$

Where we have inserted complete sets of energy eigenstates. We assumed that the system at time \( t = 0 \) is in one of the basis states \( | \alpha \rangle \), which is an eigenstate of the disorder part of the Hamiltonian. Starting the dynamics therefore coincides with turning on the interaction part of the Hamiltonian, and the survival probability can thus serve as an indicator of whether or not the system is localized.

If the system is thermal we expect that the Hilbert space will be rather uniformly explored during the quantum dynamics, and that the survival probability is approximately inversely proportional to the size of the Hilbert space. Therefore we expect to have \( \overline{S} = O(1/N) \) for small disorder.

Conversely if the system is MBL it must remain close to its initial state at all times because the quantum dynamics only lets the system explore a small amount of the configuration space. Therefore \( \overline{S} \) should certainly be much larger in the MBL phase than in the thermal phase. We define \( \overline{S}_{\text{erg}} = 1/N \) as the survival probability for an ideal ergodic system. We then have \( P_H = \overline{S}_{\text{erg}} / \overline{S} \), and we can use \( P_H \) to measure whether the system is localized or not. When we increase the system size, \( P_H \) must approach zero in the MBL phase, whereas it remains finite in the thermal phase.

We can also make a more detailed prediction for \( P_H \) in the thermal phase, since we expect that Berry’s conjecture holds and that GOE should describe the statistical behavior of the eigenvectors’ weights \( |\langle n | \alpha \rangle| \). Then \( |\langle n | \alpha \rangle| \) are Gaussian random numbers for which Isserlis’ therem holds, as in Eq. (3.21), and \( P_H \) should approach \( 1/3 \) in the ergodic phase for \( N \to \infty \).

We can use \( \langle P_H \rangle \) in Fig. 6.4 to distinguish between the MBL and the thermal phase. We see that \( \langle P_H \rangle \) is clearly tending towards zero in the thermodynamic limit for disorder strengths greater than \( W \approx 3 \). For \( W \lesssim 2.5 \), it appears that \( \langle P_H \rangle \) approaches finite values when we increase the system size. That is, for small disorder the survival probability does indeed scale as \( 1/N \) and the system is thermal.
To make a more detailed analysis of the dependence on system size we plot $\langle P_H \rangle$ against $1/L$. This probes the behavior of $\langle P_H \rangle$ in the thermodynamic limit. We see from Fig. 6.5 that deep in the MBL phase $\langle P_H \rangle$ does indeed tend to zero and in the thermal phase it appears to remain finite in the thermodynamic limit. We have performed similar finite size analysis to probe the behavior in the thermodynamic limit for many of the other quantities which we have calculated. From this we get a first estimate of the critical value of disorder, namely $W_C = 3$.

We note that the $\langle P_H \rangle$ does not seem to approach exactly $1/3$ in the thermal phase. That is, the GOE prediction is qualitatively correct but not quantitatively exact for our model. This could be due to some many-body structure of the eigenstates or finite size effects which we have not taken in to account.
6.1.2 Information Entropy

The MBL eigenstates are localized in a small portion of the basis states in configuration space whereas the thermal eigenstates generally are spread out over most of the configuration space basis.

In this sense, when expressed in the non-entangled product state basis, the energy eigenstates are more complex in the thermal phase than in the MBL phase. We will in the following quantify this by using the information entropy as a measure of complexity, and we consider the Shannon entropy defined as

$$I_n = - \sum_{\alpha} |\langle n|\alpha \rangle|^2 \log (|\langle n|\alpha \rangle|^2)$$  \hspace{1cm} (6.4)

The information entropy says something about the interrelation between the energy eigenstates basis and the product space basis and it quantifies the complexity of their relationship. We thus use $I_n$ to probe how localized the energy eigenstates are in configuration space. If the eigenstates are completely localized in configuration space we get $C_n^\alpha = \delta_{n\alpha}$ and thus zero complexity, and if the eigenstates are uniformly spread out over the non-entangled product states we have maximal complexity $\log(N)$. We thus expect the complexity to be almost maximal deep in the thermal phase and diminish as we increase the disorder.

We expect that the MBL eigenstates are not completely localized but rather spread out over a small portion of the configuration space, and that an increasing number of basis states...
contributes to the eigenstates for increasing system sizes. Thus the number of non-zero $C^\alpha_n$’s should increase with $N$. We will naively assume that the MBL eigenstates are uniformly spread out over $N^\nu$ basis states, where $\nu \in [0, 1] \subset \mathbb{R}$. This is indeed an oversimplification since we have expect to have exponentially localized states, and therefore the non-zero $C^\alpha_n$’s are not going to be equal. However, it could still serve as a crude measure of how many of the product states contribute to the MBL states. Under this assumption we get

$$I_n^{(loc)} = -\sum_{\alpha=1}^{N^\nu} \left| \frac{1}{\sqrt{N^\nu}} \right|^2 \log\left( \left| \frac{1}{\sqrt{N^\nu}} \right|^2 \right)$$

$$= \sum_{\alpha=1}^{N^\nu} N^{-\nu} \log(N^\nu)$$

$$= \nu \log(N) \quad (6.5)$$

We thus expect the complexity of the system to increase with increasing system size, and that the complexity differs from the maximal complexity by a factor $\nu$, which is some admittedly naive measure of how many of the product states contribute to an MBL state. Since the MBL eigenstates are localized only on a small number of states, there should not be any significant finite size effects to this approximation, and it should be equally good for all system sizes. We expect $\nu$ to be approximately independent of system system size, since the number of states inside the localization radius of a MBL state should increase at the same rate as the size of the Hilbert space. This is because the localization length is only weakly dependent on $L$ whereas the Hilbert space grows exponentially in $L$.

In the thermal phase we expect that GOE will describe the statistics of the system asymptotically for increasing $N$’s. As discussed in Chap. 3 the eigenvectors of the GOE in $N$ dimensions projected onto a fixed basis have a Gaussian distribution of its components

$$P(C^\alpha_n) \propto e^{-\frac{N}{2} |C^\alpha_n|^2} \quad (6.6)$$

Where $C^\alpha_n = \langle n|\alpha \rangle$. For such states the information entropy has been shown to be $I_n^{(GOE)} = \log(0.482N) + \mathcal{O}(1/N)$ in Ref. [52], which we see rapidly approaches maximal entropy. For realistic systems this presumably serves as an upper bound on the information entropy. Both since the GOE only accurately describes the eigenvectors statistics for large $N$ and since the GOE prediction entropy only asymptotically approaches $\log(N)$ with increasing $N$, the information entropy in the thermal phase should initially be much smaller than $\log(N)$ but quickly approaching $\log(N)$ as we increase $N$.

We calculate the information entropy divided by maximal entropy numerically in Fig. 6.6. We see that when we take the thermodynamic limit, $I_N/\log(N)$ appears to be approaching unity for small $W$ and for larger $W$ we see that $I_N/\log(N)$ is only weakly dependent on $N$ and decreasing with increasing disorder. This means that when we approach the thermodynamic limit, we tend toward maximally complex eigenstates in the thermal phase and in the MBL
phase the complexity is $O(\log(N))$. We note that although $I_n \sim O(\log(N))$ holds in the MBL phase we see that, apart from for $L = 8$, $I_N/\log(N)$ is slightly larger for larger $N$ and this is consistent with a weak dependence of the localization length on the system size. We can then approximate the critical value of disorder by the value for which $I_n/\log(N)$ gets roughly independent on the system size. We thus see that $W_C = 3.4$ appears to be the critical value of disorder strength.

![Figure 6.6: Information entropy divided by maximal entropy as a function of disorder strength. The average over eigenstates and disorder realizations. System sizes in legend](image)

**6.2 Absence of Level Repulsion**

We will now consider the spectral statistics of our many-body Hamiltonian. The BGS conjecture states that the statistics of thermal systems should be like that of the appropriate RMT ensemble. Since the MBL phase in some respect is similar to an integrable system, we expect the statistics in the MBL phase to have Poisson statistics, which according to the Berry-Tabor conjecture is the case for integrable systems. We know that in GOE statistics, the levels repel each other, whereas Poisson statistics has no such level repulsion. This is believed to be a distinguishing characteristic of the MBL phase.

If we assume our model to support LIOM’s we can easily argue that the levels do not repel each other. This is because adjacent energy levels in the spectrum typically differ by an extensive amount of eigenvalues of the $\tau_j$’s. Therefore the levels cannot hybridize and do not repel on the scale of the mean level spacing.
We calculate the energy difference between adjacent states in the spectrum and show histograms of the level spacings in the thermal and MBL phase in Fig. 6.8. We see that they fit very nicely with the Wigner Surmise and Poisson distribution respectively, and we clearly see the level repulsion in the weak disorder regime, and the absence thereof in the strong disorder regime.

![Histograms of level spacings in thermal and MBL phase](image)

Figure 6.7: The distribution of energy level spacings in the thermal and MBL phase. We found the level spacing $\Delta$ for different eigenstates and different realizations of disorder for systems of size $L = 14$.

We would like to study the crossover between GOE and Poisson statistics in our model and pinpoint at which disorder strength the MBL transition is.

The choice of quantity to use for such a finite-size scaling analysis is to some degree arbitrary, and there are many different quantities which we could have used. We need a dimensionless measure of the statistics of spectral properties, which in the thermodynamic limit takes different finite values in the ergodic and MBL phase. Generally when we want to compare energy level spacings we first need to perform an unfolding procedure, to ensure that the mean level spacing is set to unity. For that reason we will follow Refs. [4, 53] and consider the following function which requires no unfolding

$$r^{(n)} = \frac{\min\{\Delta_n, \Delta_{n-1}\}}{\max\{\Delta_n, \Delta_{n-1}\}}$$ (6.7)

Where $\Delta_n = |E_n - E_{n-1}|$ and we assume that the energy levels $\{E_n\}$ are listed in ascending order. This function requires no unfolding since it only compares ratios of gaps between energy levels. We will calculate expectations values of $r^{(n)}$ for the energy levels of our Hamiltonian and compare this to GOE and Poisson statistics. The expectation value of $r^{(n)}$ is calculated in Appendix B to be $\langle r^{(n)} \rangle_{Poisson} \approx 0.39$ and $\langle r^{(n)} \rangle_{GOE} \approx 0.53$ for spectra following Poisson and GOE statistics respectively.

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Figure 6.8: The ratio of adjacent energy gaps as a function of disorder strength. The average is over disorder realization and eigenstates. System sizes in legend.

We see from Fig. 6.8 that $\langle r^{(n)} \rangle$ is rapidly approaching the GOE and the Poisson expectation values with increasing $L$, for small and large $W$ respectively.

We note that our model is integrable at zero disorder, thus $\langle r^{(n)} \rangle$ might not show GOE statistics in this limit. We presumably see signs of this for $L = 8$ and small $W$. It was argued by Deutsch [11] that integrable many-body systems become thermal upon an arbitrary small perturbation of the Hamiltonian in the form of a random matrix. However his argument relied upon having systems with many degrees of freedom. This could explain why $\langle r^{(n)} \rangle$ appear to not approach $\langle r^{(n)} \rangle_{GOE}$ for small values of $L$ and $W$.

We estimate the location of the transition by considering the crossing of the curves for different values of $L$. These crossings are drifting towards the localized phase, however the drift is only very slight and is decreasing with increasing $L$. In Ref. [53] they were not able to say something conclusive about whether this drift will diminish or continue indefinitely, and they argued that this drift could stem from that behavior of the system in the critical region in some sense is more like the MBL phase. Although any proper finite-size scaling is difficult, we argue as in Ref. [4] that the crossings appear to converge to a finite $W_C$. Thus we conclude that the plot indicates that there will be a phase transition at a finite disorder strength, which we from Fig. 6.8 thus estimate to be at $W_C = 3.1$. 

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6.3 Area Law Entanglement

We will in this section consider the entanglement between two subsystems. Thermal systems are known to have entanglement which follows a volume law $O(L^d)$ whereas MBL systems have area-law entanglement $O(L^{d-1})$. Area-law entanglement is known to occur in certain quantum ground states of gapped Hamiltonians, but we will see that also highly excited MBL states have entanglement which scales as $O(L^{d-1})$.

We can see that MBL systems have area-law entanglement by assuming that our model supports LIOM’s. The eigenstates of our Hamiltonian is then product states of the $\tau_j$’s and when we split the system in two and consider the entanglement between the subsystems, we only get contributions to the entanglement from the integrals of motion which are localized within a distance $\xi$ away from the cut. Thus the entanglement scales as the area separating the subsystems and not the volume of the subsystems.

We will start by discussing and defining entanglement entropy in more detail and then we consider the entanglement in a thermal system more thoroughly, before we discuss our numeric results.

6.3.1 Bipartite Entanglement Entropy

A quantum system can have correlations which cannot be described by classical mechanics. This is called entanglement and is one of the fundamental features of quantum mechanics. In the following we describe bipartite entanglement mathematically. Bipartite means we have entanglement between exactly two subsystems. We consider a system with Hilbert space $\mathcal{H}$, which we divide into the subsystems $S$ and $B$ such that $\mathcal{H} = \mathcal{H}_S \otimes \mathcal{H}_B$. Then a general state in $\mathcal{H}$ can be expressed as

$$|\psi\rangle = \sum_{n=1}^{N_S} \sum_{m=1}^{N_B} C_{nm} |n\rangle_S \otimes |m\rangle_B$$

(6.8)

We have the density operator defined as

$$\rho = \sum_n p_n |\psi_n\rangle \langle \psi_n|$$

(6.9)

where $\{|\psi_n\rangle\}$ is some ensemble of states which are on the form of Eq. (6.8), and $p_n$ is the “classical” probability for having the system in state $|\psi_n\rangle$. If we assume $|\psi_n\rangle$ to be normalized to unity the density operator need to satisfy the following conditions

$$\text{tr}\{\rho\} = 1$$

$$\rho^\dagger = \rho$$

$$\langle \psi|\rho|\psi\rangle \geq 0 \ \forall |\psi\rangle$$

(6.10)

\[\text{\textsuperscript{\dagger}}\text{There is no unambiguously way to define multipartite entanglement}\]
If one of the $p_n$’s is one and all the others are zero we call it a pure state. From now on we restrict our discussion to pure states, since we will later measure entanglement of systems which we know are in their energy eigenstate, and as such we have maximal “classical information” about the system. A pure state is said to be quantum mechanically entangled if it cannot be expressed as a product state, i.e. if

$$|\psi\rangle \neq |\phi\rangle_S \otimes |\xi\rangle_B \quad (6.11)$$

Which is the case if it is not possible to express the weights in Eq. (6.8) as $C_n^S C_m^B$. Since we have maximal classical information about the full system, these correlations between the two subsystems are purely quantum in nature.

The reduced density matrix is obtained by tracing out the degrees of freedom of one of the systems, e.g. $\rho_S = \text{tr}_B{\rho}$, and this density matrix also satisfies the constraints of Eq. (6.10). The standard way of studying the entanglement is to look at the spectrum $\sigma(\rho_S) = \{\lambda_n\}$. The normalization and positivity restrictions leads us to think of $\{\lambda_n\}$ as a probability distribution. A distribution dominated by one value of $\lambda_n$ close to one and the rest roughly zero is less entangled than a more uniform distribution. In analogy with statistical physics we assign an entropy to this probability distribution, namely the von Neumann entropy defined as

$$S_S = - \text{tr}(\rho_S \log \rho_S) = - \sum \lambda_i \log \lambda_i \quad (6.12)$$

Here we see that a minimally entangled state has $S_S = 0$, while a maximally entangled state has $S_S = \log(N)$. We interpret the von Neumann entropy of the $\rho_S$ as a measure of how many degrees of freedom in $S$ are entangled with degrees of freedom in $B$. We assumed our system to be in a pure state, which means that the von Neumann entropy $S_F$ of the full system is zero. It holds for quantum systems that

$$S_F \leq S_S + S_B \quad \land \quad S_F \geq |S_S - S_B| \quad (6.13)$$

Where this should be contrasted with the classical constraints on entropy $S_F \geq \text{max}\{S_S, S_B\}$. In the classical regime the full system cannot be less ordered than its parts, whereas for quantum system the full system’s entropy can indeed be less than that of its parts. We thus have that in the case where the full system $F$ is in a pure state the von Neumann entropy of the two subsystems are equal and we call this the entanglement entropy $S_E$.

**Entanglement in a Thermal System**

We will assume a thermal system and consider what the entanglement entropy between the $B$ and $S$ of a full system in an energy eigenstate should be. We assume that thermal
systems satisfy Berry’s conjecture, and that we therefore can write a general eigenstate as

$$|n\rangle = \sum_{k=1}^{N} R_k |\alpha\rangle_S \otimes |\beta\rangle_B$$

(6.14)

Where we have changed indices on the random variable, which we can do since $N = N_B N_S$. This gives us the following form of the density matrix for the full system

$$\rho^{(n)} = \sum_{\alpha, \gamma = 1}^{N_S} \sum_{\beta, \delta = 1}^{N_B} R_{\alpha \gamma} R_{\beta \delta}^* |\alpha\rangle_S \langle \gamma |_S \otimes |\beta\rangle_B \langle \delta |_B$$

(6.15)

We wish to consider the reduced density operator, so we trace out the degrees of freedom in $B$ which we assume to be the larger of the two subsystems and we get

$$\rho_S^{(n)} = \text{tr}_B \rho = \sum_{\alpha, \gamma = 1}^{N_S} \sum_{\beta = 1}^{N_B} R_{\alpha \beta} R_{\gamma \beta}^* |\alpha\rangle_S \langle \gamma |_S$$

(6.16)

$$= \sum_{\alpha, \gamma = 1}^{N_S} X_{\alpha \gamma} |\alpha\rangle_S \langle \gamma |_S$$

Where we have $X = RR^\dagger$ as a $N_S \times N_S$-matrix and $R$ is a $N_S \times N_B$-matrix. The matrix elements of $R$ are by assumption i.i.d. Gaussian random numbers and for the sake of generality we assume $X$ to be unitary. Such matrices are called Wishart matrices and they have the following probability distribution

$$P(X) = \frac{1}{2^{N_B N_S} \Gamma_p\left(\frac{N_B}{2}\right) |\Sigma|^{N_S/2}} |X|^{N_B - N_S - 1} e^{-\frac{1}{2} \text{tr}(\Sigma^{-1} X)}$$

(6.17)

Where $\Sigma$ is the covariance matrix and $\Gamma_p(x)$ the multivariate gamma function. These types of matrices have been thoroughly studied for many years. If we assume the normalization constraint $\sum_n \lambda_n = 1$ on the spectrum $\{\lambda_n\}$, the joint probability distribution of the eigenvalues of $X$ is given by

$$P(\{\lambda_n\}) \propto \delta \left( \sum_{n=1}^{N_S} \lambda_n - 1 \right) \prod_{n=1}^{N_S} \lambda_n^{N_S - N_B} \prod_{k<l} (\lambda_k - \lambda_l)^2$$

(6.18)

Wishart matrices have also been studied in the context of entanglement entropy in random pure states and the average entropy of a subsystem $S$ where $1 << N_S \leq N_B$ has been considered in [42, 43]. They assumed that the spectrum of the reduced density matrix was described by Eq. (6.18), and then calculated the average entanglement entropy to be

$$\langle S_E \rangle = \ln(N_S) - \frac{N_S}{2 N_B}$$

(6.19)
Where we see that a random pure state has bipartite entanglement entropy which is dependent on the volume of the subsystem and that the states are almost maximally entangled.

We will calculate the entropy of entanglement between two halves of our spin chain numerically, and we will use the full Hamiltonian with all of its $S^z$-sectors. Since $N_B = N_S = 2^{L/2}$ we expect that the average entropy to be $\langle S_E \rangle = L \ln(2)/2 - 1/2$ in the thermal phase.

![Figure 6.9: The entropy of entanglement between two subsystems of equal size for a full system in its energy eigenstate as a function of disorder. The average is over eigenstates and disorder realizations. We have considered the full Hamiltonian, not just the $S^z = 0$-sector. The dotted line is the thermal entanglement entropy. Disorder strength in legend.](image)

We see from Fig. 6.9 that deep in the thermal phase the average entropy scales roughly as Eq. (6.19) and in the localized phase the entanglement entropy in our one-dimensional system does not change significantly when we increase the system size. That is, the entanglement scales as a volume-law in the thermal phase and an area-law in the MBL phase.

### 6.4 Failure to Thermalize

An essential characteristic of MBL systems is that they are unable to thermalize under their own dynamics. We can see this from the assumption that MBL systems supports LIOM’s. We then get that the quasi-locality of the $\tau_j$’s imply that information about the initial conditions is stored in the system at all times. This means that no information can be exchanged on the scale of the system size and thus the system fail to reach thermal equilibrium.
6.4.1 Violation of the ETH

We would now like to test the ETH, which claims that single energy eigenstates should reproduce the microcanonical ensemble average at given energy. For this to happen we need that the difference of the expectation values of a local operator in two adjacent energy eigenstates should be vanishingly small when approaching the thermodynamic limit. We let $S_i^z$ be our local operator and we consider the magnetization in energy eigenstates $m_{i,n}$, and if the ETH is satisfied it must hold that

\[ m_{i,n+1} - m_{i,n} \to 0 \quad (6.20) \]

That is, the difference in eigenstate expectation values of $S_i^z$ between two adjacent eigenstates should decrease exponentially when we increase $L$.

Assuming that in the MBL phase there exist LIOM’s which are weak deformation of the local degrees of freedom, namely the spin-operators, we typically get $m_{i,n} \approx \pm 1/2$. Since the adjacent energy eigenstates generally differ in an extensive amount of eigenvalues of the $\tau_j$’s, $m_{i,n+1} - m_{i,n}$ is thus equally likely to evaluate to approximately zero and to approximately one. We therefore expect that in the MBL phase $\langle m_{i,n+1} - m_{i,n} \rangle \approx 1/2$.

\[ \text{Figure 6.10: The difference of expectation values of } S_i^z \text{ in adjacent energy eigenstates as a function of disorder. The average is over sites, disorder realizations and eigenstates. System sizes in legend.} \]

We see clearly from Fig. 6.10 that for small disorder the difference in magnetization becomes exponentially small when we increase the system size. That $\langle m_{i,n+1} - m_{i,n} \rangle$ tends exponentially fast to zero with increasing $L$ pertains up to disorder strength $W \approx 4$, but
the the dependence on $L$ is weaker for for larger $W$, which we interpret as the system thermalizing slower close to the transition. In the MBL phase we see that the difference between the magnetization in adjacent eigenstates apparently tends to $1/2$. For $W = 4$ we have that $\langle m_{i,n+1} - m_{i,n} \rangle \approx 0.43$ for all system sizes, and it is increasing slightly with $W$. We thus get another estimate for the critical value of disorder at the point where the graphs gets independent of $L$, namely $W_C = 3.8$.

6.4.2 Spatial Correlations

We move on to consider spatial correlations on the scale of the system size, and we will show that the correlations behave differently in the two phases. Our object of investigation will be the same as in Ref. [4], namely the following connected correlation function

$$ C(k, l) = \langle n | S_k^z S_l^z | n \rangle - \langle n | S_k^z | n \rangle \langle n | S_l^z | n \rangle $$

Due to periodic boundary conditions, the correlations for a given eigenstate $|n\rangle$ for $|k - l| > L/2$ is identically equal to the correlations for $L - |k - l|$. We can use that we have restricted ourselves to the $S_z=0$-sector to evaluate the correlation function summed over $l$ for any fixed $k$. That we are in the $S_z=0$-sector means that when we let $S_z^\alpha |\alpha\rangle = s^\alpha |\alpha\rangle$ we have

$$ \sum_j s^\alpha_j = 0. $$

This gives

$$ \sum_l C(k, l) = \sum_l \sum_{\alpha, \beta} C_\alpha^\alpha C_\beta^\beta \langle \alpha | S_k^z S_l^z | \beta \rangle - \sum_{\alpha, \beta} C_\alpha^\alpha C_\beta^\beta \langle \alpha | S_k^z | \beta \rangle \sum_{\alpha', \beta'} C_\alpha'^\alpha' C_\beta'^\beta' \sum_l \langle \alpha' | S_l^z | \beta' \rangle $$

$$ = \sum_\alpha |C_\alpha|^2 s_k^\alpha \sum_l s_l^\alpha - \sum_\alpha |C_\alpha|^2 s_k^\alpha \sum_{\alpha'} |C_{\alpha'}|^2 \sum_l s_l^{\alpha'} $$

$$ = 0 $$

This summation rule holds for all eigenstates and at all disorders; simply due to spin conservation. We now consider how this correlation function differs in the two phases. Since we consider states with zero total $S_z$ we should have that $\langle n | S_k^z | n \rangle \approx 0$ for the thermal eigenstates. We already saw that this approximately holds in Fig. 6.1. This implies that the same-site correlation function must be $C(k, k) \approx 1/4$, from which it follows

$$ \sum_{k \neq l} C(k, l) = -\frac{1}{4} $$

Since we are at infinite temperatures the Boltzmann weights are all one and the thermal behavior of $C(j, l)$ at large distances is determined entirely by the sum rule in Eq. (6.23). We therefore assume that $C(k, l) \approx -1/(4(L - 1))$ for well-separated spins in the thermal phase. Two well-separated spins $k$ and $l$ are thus entangled and anti-correlated, and for growing $L$ the anti-correlations will become smaller.

In the MBL phase however the eigenstates are not thermal and as we saw in Fig. 6.1c the expectation value of $\langle n | S_k^z | n \rangle$ is non-zero. We thus expect the correlation function to have
an amplitude which falls off exponentially with distance due to the exponential localization of spins in configuration space. That is, we expect

\[ C(k, l) \sim e^{-|k-l|/\xi} \]  

(6.24)

Where \( \xi \) is the localization length. If we assume that our model supports LIOM’s this follows naturally, since we can think of the LIOM’s as dressed spins. Therefore the exponential localizations of the \( \tau_j \)’s imply that the spins far away from each other will not influence each other significantly.

Whereas we expect mostly negative correlations in the thermal phase, we have no reason to expect any particular sign of the correlations in the MBL phase.

![Figure 6.11](image)

Figure 6.11: The logarithm of the correlation function \( C(i, i + d) \) as a function of \( d \). The average is over eigenstates, disorder realizations and \( i \), and the disorder distribution have a width of \( W \). System sizes in legend

We see from Fig. 6.11 that the correlations in the MBL phase do indeed fall off exponentially with increasing \( d \), except for \( d \) close to \( L/2 \), due to periodic boundary conditions. In the thermal phase the correlations are roughly constant for large \( d \) and smaller for larger systems, as we expected. We have used the absolute value of the correlations in Fig. 6.11 but we wish to also consider if the system tends to have mostly correlations or anti-correlations. We therefore check that in the MBL phase the correlations do indeed have random signs, whereas the correlations are mostly negative in the thermal phase. We quantify this by considering the function

\[ N_C = \frac{N_- - N_+}{N_- + N_+} \]  

(6.25)

Where \( N_+ \) and \( N_- \) is the number of eigenstates for which the correlation function \( C(i, i+L/2) \) is positive and negative respectively.
6.5 Absence of DC Transport

We will in this section show that there is no transport of conserved quantities in the MBL phase. For an Anderson localized system this is rather obvious since its wavefunctions are localized in a finite region of space. However, MBL system have eigenfunctions which are extended in real space so it is not equally obvious that conserved quantities cannot be carried over large distances.

It was shown by Basko et al. [2] that the MBL states they considered have zero conductivity. It does not necessarily need to be any absence of transport for MBL to occur, since MBL has been shown to exist also in Floquet systems, which generally have
no conserved quantities and thus no transport \[26, 41\]. In the following we will argue that LIOM’s suppress transport and we will show numerically that there is no transport of spin in the random-field Heisenberg model.

### 6.5.1 Absence of DC Transport in a Model with LIOM’s

We argue that a complete set of constants of motions suppress transport, following the arguments of Ros, Mueller and Scardicchio \[50\]. We first show that the DC conductivity is zero when our model has strictly local integrals of motion. We then also argue that the argument should hold also for quasi-local integrals of motion, but this can only be seen rigorously from the convergence of the perturbative construction of the LIOM’s in Ref. \[50\]. In Appendix \[C\] we use the Kubo formula for the conductivity tensor to show that the DC conductivity is governed by

$$
\text{Re}\{\sigma(\omega \to 0)\} = \frac{\pi \beta}{8\Omega} \sum_{r} \sum_{m} \langle m | J_{r} | n \rangle \langle n | J_{r} | m \rangle e^{-\beta E_{n}} \delta_{\eta}(E_{m} - E_{n})
$$

(6.26)

Where \( \Omega \) is the volume of the system and \( J_{r} \) the local current density. We first assume that the integrals of motion are strictly local, i.e. their support is restricted to a small region of diameter \( \xi \). The energy eigenstates can be completely determined by the eigenvalues of the LIOM’s, due to the completeness of the set of integrals of motion. For two different eigenstates, there will therefore always exist integrals of motion which have different eigenvalues for the two states. We assume \( \tau \) to be such an integral of motion for the two states \( |m\rangle \) and \( |n\rangle \). We then have

$$
\tau |m\rangle = \tau_{m} |m\rangle \quad \land \quad \tau |n\rangle = \tau_{n} |n\rangle
$$

(6.27)

Where \( \tau_{m} \neq \tau_{n} \), where \( \tau_{m} \) and \( \tau_{n} \) are the eigenvalues of \( \tau \). This allows us to write the all of the matrix elements of the current operators as

$$
\langle m | J_{r} | n \rangle = \frac{\langle m | [J_{r}, \tau] | n \rangle}{\tau_{n} - \tau_{m}},
\langle n | J_{r}^{+} | m \rangle = \frac{\langle n | [J_{r}^{+}, \tau] | m \rangle}{\tau_{m} - \tau_{n}}
$$

(6.28)

Where the \( \tau \) is different for the different matrix elements. Since we assumed strictly local integrals of motion and the current operators are local, one of these terms has to vanish when \( r' \) is larger than \( \xi \), since one of the commutators then evaluates to zero. That is, the sum over \( r \) in Eq. (6.26) is restricted to \( r \lesssim \xi \).

Furthermore the locality of the \( \tau_{j} \)'s and \( J_{r} \), implies that \( J_{r} |n\rangle \) is different from \( |n\rangle \) only for a finite number of \( \tau_{j} \)'s. This is because we have LIOM’s whose eigenvalues unequivocally determine the eigenstates of the Hamiltonian, and we therefore have

$$
J_{r} |n\rangle = J_{r} |\tau_{n_{1}}, \tau_{n_{2}}, \ldots \tau_{n_{N}}\rangle
\propto |\tau_{n_{1}}, \ldots J_{r} \tau_{n_{1}}, J_{r} \tau_{n_{1}+1}, J_{r} \tau_{n_{1}+2}, \ldots \tau_{n_{N}}\rangle
$$

(6.29)
That is, the current operator only affect the $\tau_j$’s whose support overlap with that of $J_r$’s. For a given $r$ and $|n\rangle$ the matrix element $\langle m|J_r|n\rangle$ can thus be non-zero only for the $m$’s which have the same eigenvalue as $|n\rangle$ for all the LIOM’s whose support do not overlap with $J_r$’s support. This means that for every $m$ the sum over $n$ is restricted to a finite set. The eigenstates for which $\langle m|J_r|n\rangle$ is non-zero do not have to be close in energy, because adjacent energy levels in the spectrum typically differ in an extensive number of eigenvalues of the LIOM’s. Therefore the contribution to the delta function goes to zero when we take the thermodynamic limit and send $\eta \to 0$.

Thus we have shown that $\Re\{\sigma(\omega \to 0)\} = 0$ in a system with strictly local integrals of motion. However the LIOM’s in MBL systems are be quasi-local. Quasi-locality implies that the matrix elements $\langle m|J_r|n\rangle$ for eigenstates $|m\rangle$ and $|n\rangle$ who are different only in integrals of motion which are centered up to a distance $x\xi$ away from $r$, do no longer need to be zero. They will be exponentially small in $x$, but there are exponentially many states $m$ and $n$ that satisfies this criteria. So some energy differences $E_m - E_n$ in Eq. (6.28) might become exponentially small. The energy denominators are however dominated by the exponential decay of the matrix elements with probability one, and this is the key statement that guarantees the stability of the MBL. To see explicitly why this is the case we refer to Ref. [50], and their perturbative construction if the LIOM’s. The convergence of the construction procedure ensures that the exponential smallness of the energy denominators is dominated by the decay of the matrix elements and that the contribution to the $\delta$-function is zero also for a system with quasi-local integrals of motion.

### 6.5.2 Absence of Spin Transport in our Model

There are two conserved quantities in our model, namely total $S^z$ and energy. We have showed that the system fails to thermalize in the MBL phase, and this implies that there are no transport of energy. We will now show numerically that there is also no transport of spin in our random-field Heisenberg model. We will largely follow Ref. [4] and consider the longest wavelength Fourier mode of the spin density

$$M = \sum_{j=1}^{L} S^z_j e^{2\pi i \frac{j}{L}}$$

(6.30)

We wish to study transport over long length scales and therefore look at the relaxation of an initial inhomogeneous spin density given by $M$. We assume to have prepared our system with a tiny modulation of order $\epsilon$ of the spin density in this mode and that the initial density operator for the system is

$$\rho(0) = \frac{e^{\epsilon M}}{Z} \approx \frac{1 + \epsilon M^\dagger}{Z}$$

(6.31)
Where we have used that we have $\beta = 0$ and that $\epsilon$ is very small. The initial spin polarization of this mode is then
\[
\langle M \rangle_0 = \text{tr}\{\rho(0)M\} = \frac{\epsilon}{Z} \sum_n \langle n|M^{\dagger}M|n\rangle
\]  
(6.32)

We now wish to consider how this initial spin polarization evolves in time. We are in the Schrödinger-picture so the density matrix is evolved in time by the unitary $\exp(-iHt)$ and get the following expectation value for $M$ at an arbitrary time $t$
\[
\langle M \rangle_t = \text{tr}\{\rho(t)M\} = \text{tr}\{e^{-iHt}\rho(0)e^{iHt}M\} = \frac{\epsilon}{Z} \sum_{nm} |\langle m|M^{\dagger}|n\rangle|^2 e^{i(E_n - E_m)t}
\]  
(6.33)

We take the infinite-time average of $\langle M \rangle_t$ and get that it is diagonal in the energy eigenstates basis. We have
\[
\langle M \rangle_\infty = \lim_{\tau \to \infty} \frac{1}{\tau} \int_0^\tau dt \text{tr}\{\rho(t)M\} = \frac{\epsilon}{Z} \sum_n \langle n|M|n\rangle \langle n|M|n\rangle
\]  
(6.34)

If there is transport over long distances the initial spin polarization should decay away with time, and therefore the initial spin polarization should not equal the infinite-time averaged polarization. We thus expect that in the MBL phase $\langle M \rangle_0 \approx \langle M \rangle_\infty$ should hold, whereas in the thermal phase we expect that $\langle M \rangle_0/\langle M \rangle_\infty \to 0$. To test this we define the following function as a measure of how much of the contribution to $\langle M \rangle$ from an energy eigenstate $|n\rangle$ is dynamic and therefore decays away
\[
f_n = 1 - \frac{\langle n|M^{\dagger}|n\rangle \langle n|M|n\rangle}{\langle n|M^{\dagger}M|n\rangle}
\]  
(6.35)

This function should approach zero in the MBL phase since the spin polarization is not transported away, and conversely in the thermal phase the spin polarization should decay and $f_n$ should approach one.

We do indeed see from Fig. 6.13 that $f_n$ appears to approach zero as we increase $L$ in the MBL phase, i.e. that there is no long distance spin transport. In the thermal phase $f_n$ quickly approaches one for large $L$, which means that the polarization decays under the dynamics. We use the points of crossing of the lines for different system sizes as an estimate for $W_C$. In contrast to Ref. [4] we find a somewhat slighter drift of the crossings, and it appears that the critical point is at $W_C = 3.1$. 

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Figure 6.13: Transport of spin on the scale of the system size as a function of disorder strength. We have measured spin transport through the function $f_n$ as defined in Eq. (6.35). The average is over eigenstates and realizations of disorder. System sizes in legend.
Chapter 7

The Phase Transition & Universality

We have in Chap. 6 shown that our model has two distinct phases with very different properties, and we have argued through finite size scaling that the phase transition is at finite values of $W$. We will in the following consider the MBL transition itself in more detail. Unlike most quantum phase transitions the MBL transition occurs for non-zero, and even infinite, temperatures. Or more precisely at energy densities, which would correspond to finite temperatures if the system was thermal. We will assume that the MBL transition is a continuous phase transition with a vanishing characteristic energy scale $\delta$, and a diverging characteristic length scale $l_C$.

We start by discussing which role disorder plays in determining the critical behavior and how we can use the renormalization group on systems with disorder. We will then use some results from the strong disorder renormalization group and argue that the MBL transition is governed by a fixed point with infinite randomness. We shall also consider the long-time dynamics near the transition and argue that there is a broad critical region near the phase transition and show that the dynamics can be sub-diffusive in the thermal phase.

7.1 Critical Phenomena & the Renormalization Group

We will in the following briefly discuss some ideas of the renormalization group in the context of phase transitions in general and the strong disorder renormalization group in particular. Renormalization was first consider in particle physics and was used in quantum electrodynamics by Gell-Mann and Low [60]. In the context of critical phenomena such ideas were considered in 1966 by Kadanoff [61], when he considered “block-spin renormalization”. He blocked the components of his model together to define the components at a large scale in terms of the components at shorter distances, and by this coarse graining he was able to extract essential features of the macroscopic system near criticality. The renormalization group was subsequently properly developed into a powerful calculational tool by Wilson [62].
in 1971, as he formalized and generalized the ideas of Kadanoff.

### 7.1.1 The Renormalization Group

A hallmark of critical phenomena is that fluctuations appear simultaneously at all length scales and this causes non-analytic behavior of physical quantities. These singularities make perturbative approaches inappropriate. The renormalization group is a mathematical tool which we can use to systematically look at how a physical system appears when viewed at different length scales. The basic concept is to follow the change of physical quantities as we increase the length scale by coarse graining and rescaling, and these operations constitute the essence of the renormalization group transformation. We assume a Hamiltonian $H$ and denote the renormalization group transformation

$$H' = R_b(H) \quad (7.1)$$

Where $R_b$ generally is a complicated non-linear transformation, and the system described by $H'$ is now identical to that described by $H$ except that the spacing between the degrees of freedom has increased by $b$. $R_b$ reduces the total number of degrees of freedom by a factor of $b^{-d}$ and length scales by $b^{-1}$.

The renormalization transformation is a map in Hamiltonian space that mathematically defines a semi-group. It is a semi-group because, although it has an identity operation $H = R_1(H)$ and is associative $R_{b_1b_2}(H) = R_{b_1}R_{b_2}(H)$, it has no inverse mapping. The reason for the nonexistence of $R_b^{-1}$ is that when we perform the coarse graining we trace out many of the degrees of freedom and information is irretrievably lost. Exactly how $R_b$ looks is dependent upon the system in question.

The features of the system near criticality should only depend on a small number of variables such as dimensionality, symmetry and range of interactions. Thus when the renormalization group transformation removes the short-range details of the system, we should see that many properties are independent of its microscopical details. This is called universality and systems belonging to the same universality class have equal critical exponents.

It is generally the case that after applying the renormalization transformation many times the system asymptotically approaches a fixed point

$$H^* = \lim_{n \to \infty} R_b^n(H) \quad (7.2)$$

Where the Hamiltonian at a fixed point is defined through $H^* = R_b(H^*)$. Lengths scales are reduced by a factor of $1/b$ after renormalization, and in particular the correlation length renormalizes as $\xi = b^{-1}\xi$. Since the system is invariant under the renormalization transformation at fixed points the correlation length must be either infinite or zero. We
call fixed points with infinite correlation length critical fixed points, whereas “trivial” fixed points have correlation length equal to zero.

To make calculations with the renormalization group we write the Hamiltonian in terms of the parameters $\vec{u}$ and the operators $\vec{O}$ as $H = \vec{u} \cdot \vec{O}$, and consider how the parameters change under renormalization. We have the renormalization group equation $\vec{u}' = \tilde{R}\vec{u}$, or on continuous form $d\vec{u}/d\log(b) = \beta(\vec{u})$. The beta function $\beta(\vec{u})$ encodes the parameters’ dependence on the scale $b$ and its zeros define the fixed points. The renormalization group equation determines the flow towards and away from the fixed points in parameter space. Critical fixed points repel the renormalization flow and describe the singular critical behavior, whereas the “trivial” fixed points attract the renormalization flow and describe the phases of the system.

7.1.2 Strong Disorder Renormalization Group

We will now discuss how we can use the renormalization group on strongly disordered systems, a subject which was originally developed by Dasgupta and Ma [63], and eventually given a firm theoretical footing by Fisher [65, 66]. Systems with disorder may be classified according to which effect the disorder has at large length scales and we divide into three cases. First we have systems controlled by pure fixed points, for which the disorder tends to average out on large length scales and does not play any role in determining critical behavior. Second, we have systems for which the effective disorder converges to a finite level. Such systems are said to be controlled by a finite disorder fixed point. Third we have systems governed by infinite disorder fixed points, and such systems have effective disorder which grows without bound when we coarse grain them and the disorder dominate with respect to thermal or quantum fluctuations near criticality. It is the latter case which will interest us in the following.

Whereas renormalization in systems without disorder involves a finite number of parameters, the strong disorder renormalization group involves probability distributions $\rho_0(h)dh$, describing the disorder. Therefore the analysis of the fixed points and how the systems evolve under renormalization transformations generally becomes much harder. The various strong disorder renormalization groups have different implementations of the renormalization transformation but there are some properties which are general. First the renormalization should concern the degree of freedom with the highest energy, and this high energy grain constitutes the scale of renormalization. Second the renormalization should be local in real space and only be concerned with the immediate neighbors of the largest valued

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The “Gang of Four” scaling theory was essentially an application of the renormalization group, where they assumed a one-parameter scaling [17].
degree of freedom, and we pertubatively eliminate the highest energy grain. We discuss one example of strong disorder RG rules in Appendix D.

After having defined such RG rules, we apply them systematically to eliminate high energy modes. We can safely eliminate these high energy modes without changing the low energy physics because the strong randomness ensures that there will be a local separation of scales. A grain of the system with atypically high energy is likely to be surrounded by grains of lower energy. The broader the distributions of random variables the more the extreme-valued grain sticks out from its surroundings, the more accurate the perturbative elimination of the high energy grain is. Along the way we must keep track of how the distributions $\rho(h)dh$ changes with the renormalization transformations. If the width of these distributions grows indefinitely after performing the renormalization procedures, the disorder will with certainty dominate the quantum and thermal fluctuations, and the obtained results are asymptotically exact.

Fisher [64] performed a renormalization group analysis of an Ising model with quenched disorder and showed that it is governed by an infinite disorder fixed point. The RG rules he used are briefly discussed in Appendix D. He found that the system have a strong dynamical anisotropy, i.e. it formally has an infinite dynamical critical exponent because the characteristic length scale diverges as the logarithm of the characteristic time scale at the critical point. Furthermore, he found that these systems have very broad and uneven distributions of physical quantities near criticality. This stems from that the typical values are generally very different from the mean values, and that the average values of physical quantities are generally dominated by rare values (or rare regions of the system).

7.2 Infinite Disorder Fixed Point

There has been some suggestions for how to describe MBL by phenomenological RG in various models [67, 68], but we will not perform a RG-analysis ourselves. We rather exploit the results of Fisher [64] to argue that the MBL transition might be governed by an infinite disorder fixed point. We will first show that near criticality the distributions of spin correlation functions are broad and uneven and then we argue that the transition has dynamical anisotropy, since the Thouless energy is of order the level spacing near criticality.

7.2.1 Distribution of Correlation Functions

We will follow Ref. [3] and consider the distribution for the correlation function $C(k,l)$ defined in Eq. (6.21). For quantum-critical ground states governed by infinite randomness fixed points Fisher [64] showed that the distributions of correlation function should be very
broad and asymmetric, since typical values and mean values differ significantly. So if the MBL transition is in the infinite disorder universality class, this must also be the case for our model. We consider

$$
\Phi_k = \log|C(k, k + \frac{L}{2})| \quad (7.3)
$$

We see from Fig. 7.1 that the distribution of $\Phi_k$ is broad in the MBL phase and narrow in the thermal phase. In the critical region near the phase transition the distribution is asymmetric, with a fat tail.

Figure 7.1: Distribution of the spin correlation function $\Phi_k$ for systems of size $L = 14$. For even smaller disorder strengths the $\Phi$ is even more narrowly distributed. Disorder strengths in legend.

To quantify the width of the distribution and to be able to study how it changes with system size we need a dimensionless measure of the distribution width. We define $\eta = \Phi_k / \overline{\Phi_k}$ and we consider the standard deviation of $\eta$

$$
\sigma_L = \sqrt{\eta^2 - 1} = \sqrt{\frac{\Phi_k^2 - \overline{\Phi_k}^2}{\overline{\Phi_k}}} \quad (7.4)
$$

In the non-localized phase $\overline{\Phi_k}$ is roughly constant and the standard deviation of $\Phi_k$ decreases with increasing system sizes so we expect $\sigma_L$ to vanish as $L \to \infty$. In the localized regime Fisher \[64\] showed that $\overline{\Phi}$ should grow linearly in $L$ but the standard deviation of $\Phi_k$ only grows like $\sqrt{L}$, so $\sigma_L$ should vanish for $L \to \infty$. At the critical point the typical values of the correlation function should go as $-\sqrt{L}$ whereas the mean value should be $\overline{\Phi_k} \sim \log(L^{\phi-2})$, 73
where $\phi = (\sqrt{5} + 1)/2$ is the golden ratio. The standard deviation of $\Phi_k$ should be increasing fast since the discrepancies between typical values and the mean value is increasing quickly with $L$. Thus $\sigma_L$ should be finite, and presumably increase with $L$, at the critical point if the MBL transition is governed by an infinite disorder fixed point.

Figure 7.2: The width of the scaled distribution of the long-distance correlation function $\eta$ as a function of disorder strength. We estimated $\sigma_L$ by averaging over sites and eigenstates, and then we find the statistical errors by using the variations of $\sigma_L$ between the different disorder realizations. System sizes in legend.

We calculate $\sigma_L$ numerically and see from Fig. 7.2 that it does indeed appear to go towards zero in the ergodic and in the MBL phase, and that it approaches zero faster for larger $L$. We also see that at the critical point $\sigma_L$ remains finite, and it even increases slightly with $L$. This is exactly what we would expect from a transition in an infinite-randomness universality class. The plot also yields another estimate for the critical disorder strength, namely $W_C = 4$.

### 7.2.2 Thouless Energy

The Thouless energy is inversely proportional to the relaxation time on the scale of the system size. There are two conserved quantities in our model, namely energy and total spin. Their relaxation times, and thus Thouless energies, might behave differently close to the critical point. We will in the following consider the spin transport time, and we cannot exclude that the energy transport time will scale differently close to the transition. We follow Ref. [4] and argue that the level spacing is comparable to the Thouless energy at criticality.
We assume the Thouless energy to be given roughly by the relaxation rate of the spin density modulation of $M$ as given in Eq. (6.30), and we can think of the Thouless energy as a measure of how much different eigenstates affect each other. In the thermal phase $E_T$ is large and many states affect each other, whereas in the MBL phase $E_T$ is exponentially small, so different states do not influence each another significantly. We will in the following argue that the Thouless energy is of the same order of magnitude as the level spacing at the transition, and this implies that only the states which are adjacent in the energy spectrum will affect each other near criticality.

Thus a non-zero fraction of the dynamic part of $\langle M \rangle$ should be from its matrix elements between adjacent energy levels, also in the thermodynamic limit.

Continuing our discussion in Sect. 6.5.2, the contribution to the dynamical part of $\langle M \rangle$ from a given energy eigenstate should be given by

$$
\Delta M_n = \langle n|M^\dagger M|n \rangle - |\langle n|M|n \rangle|^2
$$

(7.5)

We showed that this appears to approach zero in the MBL phase and remains non-zero in the thermal phase. In the thermal phase $\Delta M_n$ has contributions from matrix elements with many other eigenstates and $E_T$ is a measure of the energy range over which these contributions occur. We look at contributions to the dynamic part of $\Delta M_n$ from the matrix elements between state $n$ and $n \pm i$

$$Q_i = |\langle n - i|M|n \rangle|^2 - |\langle n|M|n + i \rangle|^2
$$

(7.6)

We insert a complete set of eigenstates to see that we have

$$
\Delta M_n = \sum_m \langle n|M^\dagger m \rangle \langle m|M|n \rangle - |\langle n|M|n \rangle|^2
= \sum_{i \neq 0} |\langle n - i|M|n \rangle|^2 - |\langle n|M|n + i \rangle|^2 = \sum_{i \neq 0} Q_i
$$

(7.7)

We define

$$P_n = \frac{Q_1}{\Delta M_n}
$$

(7.8)

Where $P_n$ is the fraction of the dynamics which is due to interference between adjacent energy levels. In the thermal phase we expect matrix elements between many different states to contribute to the dynamic part of $\Delta M_n$, so many of the $Q_i$’s should contribute significantly. However in the MBL phase the Thouless energy is exponentially small and therefore none of the $Q_i$’s give significant contributions to the dynamic part of $\Delta M_n$. In the MBL phase we have both $Q_1$ and $\Delta M_n$ going to zero. Thus both in the MBL and in the thermal phase we expect $P_n$ to approach zero for increasing system size. If $E_T$ is of the same order of magnitude as $\delta$ at the transition, we expect $Q_i$ to contribute to $\Delta M_n$ only for $i = 1$ close to
the critical point. Thus $P_n$ should be non-zero at the critical point, also when we take the thermodynamic limit.

![Figure 7.3: Fraction of dynamics which comes from adjacent eigenstates as a function of disorder strength. Average is over eigenstates and disorder realizations. System sizes in legend.](image)

We see from Fig. 7.3 that both in the MBL and thermal phase $\langle P_n \rangle$ appears to be approaching zero as we increase $L$. Furthermore we do indeed see a strong peak in $\langle P_n \rangle$ around the critical point, indicating that the maximal contribution from adjacent states is near the phase transition. The peaks decrease a little for increasing $L$ but not as much as the system size is increased, so the results seem to comply with $E_T \sim \delta$ scaling at the transition. If we disregard $L = 8$, the peaks seems to drift very slightly towards larger $W$ and the peak appears at $W \approx 2.8$ for $L = 16$, suggesting that the critical disorder strength is $W_C = 3$.

**Dynamical Anisotropy**

Continuous quantum phase transition generally have some characteristic time scale $\tau_C$ and characteristic length scale $l_C$ which diverge at the critical point. The dynamical critical exponent $z$ is given by the scaling of $\tau_C$ with $l_C$ near the critical point as $\tau_C \sim l_C^z$. In other words the characteristic time diverges as the $z$’th power of the characteristic length when we approach the phase transition.

We know that Anderson localization transition occurs when $E_T \sim \delta$. The single-particle level spacing for a $d$-dimensional system of linear size $L$ behaves roughly as $\delta \sim L^{-d}$ and we therefore have $E_T \sim L^{-d}$. We have a diverging correlation length $\xi$ and therefore the
characteristic length scale of the system equals $L$. The characteristic time scale is the Thouless time, which is the inverse of $E_T$. We therefore have $\tau_T \sim L^d$ scaling at criticality and the system thus has a dynamic critical exponent which is $z = d$.

We now assume that the MBL transition occurs when the Thouless energy and the level spacing are of the same order of magnitude, as we argued above. The level spacing goes roughly as $\log(\delta) \sim -L^d$. This gives us the scaling $\log(\tau_T) \sim L^d$ which is characteristic for infinite critical exponents. We here have the typical length scale related to the logarithm of the typical time scale and thus a formally infinite dynamical exponent $z = \infty$. We note that even for our model with $d = 1$ this is a stronger divergence of the critical time scale than what is known from infinite randomness critical points in ground states [64], where $\log(\tau_C) \sim \psi_C^\infty$ with $\psi \leq 1/2$ [64].

### 7.3 Dynamics & Spectral Functions

In this section we will investigate the long-time dynamics in the thermal phase when we approach the phase transition. We will show that there can be anomalous diffusion on the thermal side of the transition and argue that there is a broad critical region. We also further argue that $E_T \sim \delta$ at criticality.

Some recent studies have reported sub-diffusive transport close to the critical point of the MBL transition [55, 56, 57]. This is believed to be due to so-called “Griffiths effects”. This is when the low-frequency response is dominated by contributions from rare regions. In our case it is regions which locally behave as the MBL phase on the thermal side of the transition. These insulating regions impede the transport but do not prevent thermalization. Particularly in one dimension this can severely affect the dynamics because the locally insulating regions can act as bottlenecks [59]. Griffiths effects arise due to regions with atypical configurations of the quenched randomness.

Griffiths effects were first brought to the fore by Griffiths [58] when considering thermodynamic and dynamic singularities due to rare regions in disordered systems near phase transitions.

We assume that the ETH ansatz in Eq. (4.32) describes the matrix elements of some local observable $A$, and we show that we can relate $g(E, \omega)$ to the dynamics of the system. If we neglect the variations of the entropy near $E=0$, it can be shown that $g^2(\omega)$ is the average spectral function of $A$ for the states with energy close to zero. We will therefore denote $g^2(\omega)$ the spectral function in the following. Our discussion will be similar to the recent work of
Abanin, Papic and Serbyn [54]. We consider the following connected correlation function

\[ F_n(t) = \langle n|A(t)A(0)|n\rangle - \langle n|A(t)|n\rangle\langle n|A(0)|n\rangle \]

\[ = \sum_{m \neq n} |\langle n|A|m\rangle|^2 e^{i(E_n - E_m)t} \quad (7.9) \]

Since we assumed that the ETH ansatz holds for the matrix elements of \( A \) we can readily evaluate the correlation function

\[ F_n(t) = \sum_{m \neq n} g^2(E_n, \omega) R^2_{mn} e^{-S(E)} e^{i\omega t} \quad (7.10) \]

We use that \( g(E, \omega) \) is a smooth function to argue that \( R^2_{mn} \) should level out in the sum, and switch from the summation over the eigenstates to an integral over the density of states as \( \sum_{m \neq n} R^2_{mn} \rightarrow \int dE_n \rho(E_n) = \int d\omega \ e^{(E_n + \omega)} \), where we also switch integration variable to \( \omega \). This gives

\[ F_n(t) = \int d\omega \ g^2(E_n + \frac{\omega}{2}, \omega) e^{S(E_n + \omega)} e^{i\omega t} \quad (7.11) \]

We are interested in the long-time dynamics of the system so we need to consider small frequencies, and we therefore Taylor expand the entropy and the spectral function to the first order about \( \omega = 0 \) as

\[ S(E_n + \omega) - S(E_n + \frac{\omega}{2}) = \beta \omega + \mathcal{O}(\omega^2) \]

\[ g^2(E_n + \frac{\omega}{2}, \omega) = g^2(E_n, \omega) + \frac{\omega}{2} \frac{dg^2(E_n, \omega)}{d\omega} \mathcal{O}(\omega^2) \quad (7.12) \]

Using this we arrive at

\[ F_n(t) \approx \int d\omega \ e^{-i\omega t} e^{\frac{\omega}{2}} [g^2(E_n, \omega) + \frac{\omega}{2} \frac{dg^2(E_n, \omega)}{d\omega}] \quad (7.13) \]

We also Taylor expand \( \exp(\beta \omega/2) = 1 + \beta^2 \omega^2 + \mathcal{O}(\omega^2) \). The long-time correlation function is thus the Fourier transform of \( g^2(E_n, \omega) \) at small frequencies

\[ F_n(t) \approx \int d\omega \ e^{-i\omega t} g^2(E_n, \omega) \quad (7.14) \]

Since we are in the thermal phase where we have assumed ETH to hold, the dynamics should be diffusive or super/sub-diffusive. We consider how the correlation function should behave under diffusive transport. We have the one-dimensional diffusion equation

\[ \frac{\partial \phi}{\partial t} = D \frac{\partial^2 \phi}{\partial x^2} \quad (7.15) \]

which is solved by the Green’s function

\[ G(x_0, t_0|x, t) = \frac{1}{\sqrt{4\pi D(t-t_0)}} e^{-\frac{(x-x_0)^2}{2(D(t-t_0))}} \quad (7.16) \]
Where $D$ is the diffusion constant. We associate our correlation function $F_n(t)$ to the Green’s function for the diffusion equation $G(x_0,0|x_0,t)$. This leads us to expect $F_n(t) \sim |t|^{-\gamma}$, where $\gamma = 1/2$. However if we assume anomalous sub-diffusion† we get $\gamma < 1/2$.

We thus assume that $F_n(t)$ describes the diffusive or sub-diffusive transport and Fourier transform Eq. (7.14) to estimate $g^2(\omega)$

$$g^2(\omega) = \int_{-\infty}^{\infty} dt \ |t|^{-\gamma} e^{i\omega t} \propto \omega^{\gamma-1} \quad (7.17)$$

Furthermore we expect the diffusive transport to saturate at a time approximately equal to the Thouless time $\tau_T$, since for times longer than $\tau_T$ local excitations explore the entire system, and the diffusion saturates. We get saturation $g^2(\omega) \sim K$ for some constant $K$ for frequencies $\omega < E_T$; in this region the ETH ansatz reduces to RMT prediction for the matrix elements. We therefore conjecture the approximate functional form of the spectral function

$$g^2(\omega) \approx \frac{K}{1 + \left(\frac{\omega}{E_T}\right)^{1-\gamma}} \quad (7.18)$$

Which encapsulates the diffusive behavior and saturates at frequencies smaller than the Thouless energy. We have thus showed that we can obtain information about the dynamics as encoded in $F_n(t)$ by studying the $g^2(\omega)$, and found an expression for the behavior of $g^2(\omega)$ under diffusive transport. We rewrite and modify the ETH ansatz slightly to get an alternative expression for the spectral function

$$g^2(\omega) = e^{S(E)} \langle |A_{nm}^2| \delta(\omega - (E_n - E_m)) \rangle \quad (7.19)$$

If we broaden the $\delta$-function to be much larger than the level spacing and assume that the fluctuations $R_{nm}^2$ average out, we see that Eq. (7.19) is equivalent to the ETH ansatz in Eq. (4.32).

†We note that anomalous diffusion is not described by the standard diffusion equation. One of the easiest way to model anomalous diffusion is to assume a time-dependent diffusivity $D(t) = \alpha t^{\alpha-1} D$ in Eq. (7.15), where $\alpha = \gamma/2$. 

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Figure 7.4: \( g^2(\omega) \) as a function of \( \omega/\Delta \), where \( g^2(\omega) \) is the spectral function of \( \sigma_i^z \). We found \( g^2(\omega) \) numerically through Eq. (7.19) and used only the center two percent of the eigenstates. System sizes in legend.

We found \( g^2(\omega) \) as a function of \( \omega/\Delta \) numerically for several disorder strengths and system sizes, where \( \Delta = W \sqrt{L/N} \) is the typical many body level spacing. We see from Fig. 7.4 that \( g^2(\omega) \) appears to go as \( \omega^{\gamma-1} \) for \( E_T \ll \omega \ll 1 \) and saturate for \( \omega \lesssim E_T \), which we expected. We extract \( \gamma \) by a fitting \( \omega^{\gamma-1} \) to \( g^2(\omega) \) in the region before it saturates. For \( W = 1 \) we find that \( \gamma \approx 1/2 \), which means that deep in the thermal phase the transport is indeed diffusive. However when we increase disorder we find that \( \gamma \) decreases and at \( W \approx 2 \) we find that \( \gamma \approx 0 \).

For the disorder strengths showed in Fig. 7.4 the saturation occurs at larger frequencies for larger systems, meaning that the Thouless energy is increasing faster than the level spacing with \( L \). However we see that the Thouless energy decreases rapidly with increasing disorder.
Already for $W = 2.5$ we see that $g^2(\omega)$ does not fully saturate, which signifies that $E_T \sim \delta$. However $g^2(\omega)$ retains a slight upward curvature through the MBL transition, and is is not before $W > 4$ that we see a power law behavior for $g^2(\omega)$ even for energies smaller than the level spacing.

We fit the curves in Fig. 7.4 with the functional form of $g^2(\omega)$ in Eq. (7.18) and use $K$, $\gamma$ and $E_T$ as parameters of the fit, and thereby extract the Thouless energy.

We see from Fig. 7.5 that the Thouless energy is comparable to the level spacing in a broad region before the MBL transition. We have plotted the Thouless energy of a diffusive system $E_T \propto L^{-2}$ and see that this seems to be roughly consistent with the data for $W = 1$. When we increase the disorder we see that $E_T/\Delta$ increases slower with $L$. For disorder larger than $W = 2$, the Thouless energy increases only a little faster than the level spacing with increasing $L$. This could possibly mean that the Thouless energy in goes as $E_T \propto L^{\frac{1}{2}}$ with a very small $\gamma$. It seems however more natural to interpret this as $E_T \propto \exp(-\kappa L)$, for some sufficiently small $\kappa$, similar to in a MBL system. The many-body level spacing scales roughly as $2^{-L}$ so we need to have $\kappa \lesssim \log(2)$, since $E_T/\Delta$ is indeed increasing slightly. We interpret this exponential scaling of $E_T$ as a sign of our system entering a critical region which starts at disorder strength $W \approx 2$.

We assume that there is correlation a length $\xi(W)$ which diverges at MBL transition,
since the MBL transition is a continuous phase transition. Thus close to $W_C$, it could be the case that $\xi(W) \geq L$, and when this is the case the system will appear to be at the transition, since it is too small to capture the delocalized behavior. That is, if we consider a system at scales smaller than the correlation length, it will appear critical. We see that for disorder $W \approx 2$ the correlation length appears to be larger than $L = 14$, and therefore $g^2(\omega)$ can no longer capture the delocalized behavior of the system.
Chapter 8

Summary and Outlook

We have shown the existence of two distinct phases in the random-field Heisenberg model, namely the thermal phase and the MBL phase. We transition from the thermal to the MBL phase by increasing the disorder strength $W$ and we found the critical value to be $W_C = 3.5 \pm 1.0$. We have distinguished between the phases through the scaling of entanglement entropy, level statistics, transport properties, participation ratios and whether or not the system satisfies the ETH.

We can view this transition as an eigenstate phase transition, where the eigenstates have very different properties in the two phases. Very crudely we say that in the thermal phase all eigenstates appear the same and in the MBL phase all eigenstates appear different. What we mean by this is that the all thermal eigenstates within some energy window should, according to the ETH, reproduce the thermodynamic ensembles at given energy density, whereas the MBL states are exponentially localized around different basis states in configuration space. This eigenstate transition is invisible to equilibrium statistical mechanics, because statistical mechanics averages over many eigenstates and thus washes out the sharp change in terms of single eigenstates.

The fact that there can exist two distinct phases in closed, strongly interacting, disordered quantum systems has been well established over the course of the last 10 years. However not everything about the MBL transition is known. We have argued that the the Thouless energy is of the order the level spacing at the transition, which implies a formally infinite dynamical critical exponent. We have also seen that the dynamics can be sub-diffusive in the thermal phase and signs of a critical region near the phase transition. Furthermore we discussed the possibility of the MBL transition belonging to the infinite disorder universality class.

There are some sides of the MBL phase and the MBL transition which we have not touched upon in this thesis. One of these is localization protected quantum order. It has
been shown that individual energy eigenstates of MBL systems can break symmetries or display topological order in the thermodynamic limit. This can happen at energy densities where the corresponding thermally equilibrated systems are disordered, and MBL systems can move between ordered and disordered phases via non-thermodynamic transitions in the properties of the many-body eigenstates [69, 70].

We have also not properly touched upon multifractality, which is believed to occur near the phase transition. There have been some studies interpreting the matrix elements of an observable as the wavefunction of a local excitation and reporting on a strong multifractal spectrum at the critical point [54, 71].

It would have also been interesting to consider the spreading of entanglement. MBL systems initially prepared in states with low entanglement have a fast growth of entanglement, and after some time the entanglement has a slow, logarithmic growth [72]. For an infinite system the entanglement is expected to grow indefinitely, whereas for a finite system it saturates to some value, which depends on the system size and should be smaller than in the thermal phase. This unbounded growth is compatible with the absence of transport, and it stems from the interaction-induced dephasing involved in the decomposition of the initial state. This is different from Anderson localization, where the entanglement saturates to a finite level, also in thermodynamically large systems.

Most of the MBL theory is formulated in terms of exact eigenstates of closed systems. This is a useful idealization which sheds light on the phenomenology of MBL, but to make contact with experiments we should presumably consider other approaches and it would therefore be interesting to consider a MBL system weakly coupled to a heat bath. Such a system can be studied using spectral functions of local observables, similar to what we did in Sect. 7.3 since these retain signatures of MBL even when coupled to a heat bath [74]. Furthermore the picture of the MBL transition as an “eigenstate phase transition” has been critiqued in Ref. [73], where they argue for the possibility that there is an “eigenstate phase transition” within the MBL phase. That is, they argued through approximately conserved quasi-local operators that the MBL phase can have eigenstates which satisfies the ETH.

The MBL phase has recently been experimentally observed in a couple of different systems. In Ref. [6] MBL was observed for interacting fermions in an one-dimensional optical lattice, where they considered the relaxation dynamics of an initially prepared charge density wave to identify MBL. Similarly the relaxation dynamics of bosons in two dimensions has been used to observe the MBL transition in Ref. [8]. They prepared an out-of-equilibrium density pattern and found evidence for a diverging length scale when approaching the transition. MBL has also been reported to occur in linear arrays of cold, trapped atomic ions [7].

There are still many open questions concerning MBL. One interesting issue is if it is
possible to have MBL without disorder. There have been some studies arguing for the existence of translation invariant MBL. In Ref. \[75\] they found numerical evidence for MBL in a disorder free Josephson-junction array. Effects similar to MBL have been reported to occur in a translation invariant systems of hard-core bosons with infinite range and periodic interactions in Ref. \[77\], in a system which contains two species of hard-core particles with very different masses in Ref. \[76\] and in an exactly soluble model with an extensive number of conserved quantities in Ref. \[78\]. In these disorder-free models the systems seem to somehow generate their own randomness dynamically. Which role disorder plays in MBL is still not fully understood, and further study of localization of a purely dynamic origin would be interesting.

It is also not completely understood if it is possible to have mobility edges in MBL system, i.e. MBL transitions as functions if energy. It was generally believed that MBL systems can indeed have mobility edges, but in Ref. \[79\] it was argued that MBL can only occur for lattice model which are localized at all energies and that previously reported mobility edges cannot be distinguished from finite-size effects. They argued that local hot thermal spots, which they denotes “bubbles”, constitute a mechanism for global delocalization.

An effective field theory approach of MBL has also recently been considered in Ref. \[80\] where XXZ-chains in the presence of local disorder were considered. It would be interesting to consider this model (or a similar one) further and possibly us it to identify the universality class of the MBL transition.

It has recently been shown that spontaneous breaking of time-translation symmetry can occur in periodically driven MBL systems with discrete time-translation symmetries \[83\] \[82\] \[84\]. Wilczek \[81\] considered the possibility of time-crystals, which intriguingly has a spontaneous breaking of the continuous time-translation symmetry. This was subsequently proven to be disallowed in equilibrium, but in oscillating quantum systems spontaneous breaking of time-translation symmetry is possible. After having observed so-called discrete time-crystals, it would be interesting to consider using this for various quantum information tasks and also to investigate which role MBL in general could play in realizing quantum computers.
Appendices
Appendix A

Consequences of the ETH-ansatz

We will show explicitly that when the ETH ansatz in Eq. (4.32) correctly describes the matrix elements of observables the systems will be thermal. We will also see how fast the discrepancies between the long time average and the thermodynamic ensemble average vanish when we increase the size of our system. Given a system fulfilling ETH we consider the canonical thermal average

$$\langle A \rangle_{\text{th}} = \sum_n e^{-\beta E_n} A(E_n) + \sum_n R_{nn} e^{-\beta E_n} \frac{e^{-S(E_n)}}{Z} g(E_n, 0)$$

We solve these integral through the saddle point approximation. This demands $dS(E) = \beta$, which we expected since this is how we define temperature in the canonical ensemble, this also fixes $E = E_{\text{th}}$. We have $S(E)$ as an extensive quantity which we may write in terms of the entropy per particle $s(e) = S(E, N)/N$ where $e = E/N$ is the energy per particle. This gives us $N(s(e) - \beta e)$ in the exponential. We can now change integration variable to energy per particle use the standard saddle point approximation which may be stated as

$$\int_\gamma dz f(z) e^{\lambda S(z)} = F(S(z_0)) e^{\lambda S(z_0)} [f(z_0) + O(\lambda^{-1})]$$

Where $F(S(z))$ is some function of the eigenvalues of the Hessian matrix for $S(z)$. This gives

$$\langle A \rangle_{\text{th}} \approx A(E_{\text{th}}) + O(N^{-1}) + O(e^{-\frac{S}{2}})$$

Since $N$ is large we can Taylor expand the first term to first order to get that it is $A(E_{\text{th}}) + O(N^{-1})$. We can also show explicitly that the second term in Eq. (A.1) is exponentially small in entropy. However we will simply hold this as obvious, since we know that $R_{nn}$ and
\(\exp(-\beta E_n)/Z\) are both of order \(\mathcal{O}(1)\) and \(g(E_n, 0)\) will certainly not be exponentially large. Therefore the term \(\exp(-S(E)/2)\) dominate the second term and it goes exponentially fast to zero. Where we use that we have assumed highly excited states, which implies that the entropy should be extensive. We thus have

\[
\langle A \rangle_{th} = A(E_{th}) + \mathcal{O}(e^{-\frac{S(E)}{2}}) + \mathcal{O}(N^{-1})
\]  

(A.4)

We now study the time-evolution of the expectation value of \(A\)

\[
\langle A(t) \rangle = \sum_n |C_n|^2 (A(E_n) + g(E_n, 0) e^{-\frac{S(E)}{2}} R_{nn}) + \sum_{n \neq m} C_n^* C_m e^{-i(E_m - E_n)t} g(E, \omega) e^{-\frac{S(E)}{2}} R_{nm}
\]  

(A.5)

We then take the infinite-time average

\[
\Rightarrow \langle A(t) \rangle = \sum_n A(E_n) |C_n|^2 + \mathcal{O}(e^{-\frac{S(E)}{2}})
\]  

(A.6)

We Taylor-expand \(A(E_n)\) about \(E_{th}\) to the second order

\[
\langle A(t) \rangle = A(E_{th}) + \sum_n \left[ (E_n - E_{th}) |C_n|^2 \frac{dA}{dE} \bigg|_{E_{th}} \right] + \left[ (E_n - E_{th})^2 |C_n|^2 \frac{d^2A}{dE^2} \bigg|_{E_{th}} \right] + \mathcal{O}(e^{-\frac{S(E)}{2}}) + \mathcal{O}(N^{-1}) + \mathcal{O}(\Delta^2) + \mathcal{O}(e^{-\frac{S(E)}{2}})
\]  

(A.7)

Where we have used that we are concerned with a closed system for which it must hold that \(E_{th} = \langle E \rangle\). We here quantify in terms of the operator \(A\) what we mean by demanding the uncertainty in energy being small, namely \(\Delta^2 \ll |A/A''|\). All in all, this leaves us with

\[
\langle A(t) \rangle = \langle A \rangle_{th} + \mathcal{O}(N^{-1}) + \mathcal{O}(\Delta^2) + \mathcal{O}(e^{-\frac{S(E)}{2}})
\]  

(A.8)

We have also shown that the infinite time average equals the ensemble average, and this follows entirely from the structure of the matrix elements without any assumptions regarding the initial state, other than that it is away from the edges of the spectrum and that it has relatively small uncertainty in energy.
Appendix B

Calculation of Level Repulsion

We will in the following determine the probability distribution of $r^{(n)}$ as defined in Eq. (6.7). We consider the statistical probability for $r^{(n)}$ to equal $r$

$$P(r) = \left\langle \delta(r - \min \{\Delta_n, \Delta_{n-1}\} / \max \{\Delta_n, \Delta_{n-1}\}) \right\rangle$$  \hspace{1cm} (B.1)

Where the average depends on which statistics $\{E_n\}$ follow. We generally need to use the joint probability distribution for the eigenvalues to calculate $P(r)$. We consider the cases when the spectra obeys Poisson and GOE statistics.

**Poisson statistics**

For the Poisson statistics $P(r)$ can be easily evaluated analytically since the levels are completely randomly distributed and therefore the level spacings must also follow the Poisson distribution

$$P_{\text{Poisson}}(r) = \frac{1}{\delta^2} \int_0^\infty d\Delta_n \int_0^{\Delta_{n-1}} d\Delta_{n-1} \delta(r - \frac{\min \{\Delta_n, \Delta_{n-1}\}}{\max \{\Delta_n, \Delta_{n-1}\}}) e^{-\frac{\Delta_n}{\delta}} e^{-\frac{\Delta_{n-1}}{\delta}}$$  \hspace{1cm} (B.2)

We split the integral into the two parts; one where $\Delta_{n-1} < \Delta_n$ and the other where $\Delta_{n-1} > \Delta_n$.

$$P_{\text{Poisson}}(r) = \frac{1}{\delta^2} \int_0^\infty d\Delta_n \int_0^{\Delta_n} d\Delta_{n-1} \delta(r - \frac{\Delta_n}{\Delta_{n-1}}) e^{-\frac{\Delta_n}{\delta}} e^{-\frac{\Delta_{n-1}}{\delta}}$$

$$+ \frac{1}{\delta^2} \int_0^\infty d\Delta_n \int_0^{\Delta_{n-1}} d\Delta_{n-1} \delta(r - \frac{\Delta_{n-1}}{\Delta_n}) e^{-\frac{\Delta_n}{\delta}} e^{-\frac{\Delta_{n-1}}{\delta}}$$

$$= \frac{2}{\delta^2} \int_0^\infty d\Delta_n \int_0^{\Delta_n} d\Delta_{n-1} \Delta_n \delta(\Delta_n r - \Delta_{n-1}) e^{-\frac{\Delta_n}{\delta}} e^{-\frac{\Delta_{n-1}}{\delta}}$$

$$= \frac{2}{\delta^2} \int_0^\infty d\Delta_n \Delta_n e^{-\frac{\Delta_n(r+1)}{\delta}}$$

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Within GOE it is somewhat more involved to calculate the probability distribution of Gaussian Orthogonal Ensemble Statistics $E$ energies are ordered, i.e. for arbitrary $N$ assume a 3x3-matrix and use the PDF on this matrix. Hopefully this will mimic the behavior probability distribution from Eq. (3.14), but its not tractable to calculate this exactly so we same so it was straightforward to evaluate the distribution. For the GOE we need the joint exactly, due to the level repulsion. In the Poisson-case the statistics of $E$ now we can find the expectation value of $r^{(n)}$ for a spectrum obeying Poisson statistics

\[
\langle r \rangle_{\text{Poisson}} = \int_0^1 dr \, r P_{\text{Poisson}}
\]

\[
= \int_0^1 dr \, \frac{2r}{(1+r)^2}
\]

\[
= \left[ \frac{2}{1+r} + 2 \ln(1+r) \right]_0^1 = 2 \ln(2) - 1 \approx 0.39
\]

\[\text{Gaussian Orthogonal Ensemble Statistics}\]

Within GOE it is somewhat more involved to calculate the probability distribution of $r^{(n)}$ exactly, due to the level repulsion. In the Poisson-case the statistics of $E_n$’s and $\Delta_n$’s are the same so it was straightforward to evaluate the distribution. For the GOE we need the joint probability distribution from Eq. (3.14), but its not tractable to calculate this exactly so we assume a 3x3-matrix and use the PDF on this matrix. Hopefully this will mimic the behavior for arbitrary $N$ well, in the same way that the Wigner surmise was a good approximation for large $N$. We now venture to find the GOE distribution of $r^{(n)}$

\[
P_{\text{GOE}}(r) \propto \int_{-\infty}^{\infty} dE_{n-1} \int_{E_{n-1}}^{\infty} dE_n \int_{-\infty}^{E_{n-1}} dE_{n-2} \, \delta(r - \min\{\Delta_n, \Delta_{n-1}\})
\]

\[
	imes |E_n - E_{n-1}| |E_n - E_{n-2}| |E_{n-1} - E_{n-2}| e^{-\frac{E_{n-1}^2}{4} + \frac{E_n^2}{4} - \frac{E_{n-2}^2}{4}}
\]

Here we once again consider separately the cases where $\Delta_n \leq \Delta_{n-1}$ and exploit that the energies are ordered, i.e. $E_n \geq E_{n-1} \geq E_{n-2}$ which renders the absolute value signs obsolete

\[
P_{\text{GOE}}(r) \propto \int_{-\infty}^{\infty} dE_{n-1} \int_{E_{n-1}}^{\infty} dE_n \int_{-\infty}^{E_{n-1}} dE_{n-2} \, \delta(r - \frac{E_{n-1} - E_{n-2}}{E_{n-1} - E_{n-2}})
\]

\[
	imes (E_n - E_{n-1})(E_n - E_{n-2})(E_{n-1} - E_{n-2}) e^{-\frac{(E_{n-1}^2 + E_n^2 + E_{n-2}^2)}{4}}
\]

\[
+C \int_{-\infty}^{\infty} dE_{n-1} \int_{E_{n-1}}^{\infty} dE_n \int_{-\infty}^{E_{n-1}} dE_{n-2} \, \delta(r - \frac{E_{n-1} - E_{n-2}}{E_n - E_{n-2}})
\]

\[
	imes (E_n - E_{n-1})(E_n - E_{n-2})(E_{n-1} - E_{n-2}) e^{-\frac{(E_{n-1}^2 + E_n^2 + E_{n-2}^2)}{4}}
\]

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We now change variables to
\[ x = E_{n-1} - E_{n-2} \in [0, \infty) \]
\[ y = E_n - E_{n-1} \in [0, \infty) \]
\[ E_{n-1} \in (-\infty, \infty) \]  
(B.9)

For the exponents we then get
\[ E_n^2 + E_{n-1}^2 + E_{n-2}^2 = (E_{n-1} + y)^2 + (x - E_{n-1})^2 + E_{n-1}^2 \]
\[ = 3(E_{n-1}^2 + \frac{1}{3}(x - y))^2 - \frac{1}{3}(x - y)^2 + x^2 + y^2 \]  
(B.10)

We now shift the limits of integration in the integral over \( E_{n-1} \) by \( \frac{1}{3}(x - y) \) which does not cause any problem since our function goes exponentially fast to zero when the arguments go to infinity. This yields
\[ P_{\text{GOE}}(r) \propto \int_{-\infty}^{\infty} dE_{n-1} \int_0^\infty dx \int_0^\infty dy \ \delta(yr - x) y^2 x(x + y) \ e^{-\frac{1}{4}(x^2 + y^2 - \frac{1}{3}(x-y)^2)} e^{-\frac{\pi E_{n-1}^2}{4}} \]
\[ \propto \int_0^\infty dx \int_0^\infty dy \ \delta(yr - x) y^2 x(x + y) \ e^{-\frac{1}{4}(x^2 + y^2 - \frac{1}{3}(x-y)^2)} \]
\[ \propto \int_0^\infty dy \ y^4 (r + r^2) \ e^{-\frac{(1 + r + r^2)}{6} y^2} \]  
(B.11)

We use the known definite integral
\[ \int_0^\infty dx x^{2n} e^{-\alpha x^2} = \frac{(2n)!}{n! 2^{2n+1}} \sqrt{\frac{\pi}{\alpha^{2n+1}}} \]  
(B.12)

to get
\[ P_{\text{GOE}}(r) = K \frac{r + r^2}{(1 + r + r^2)^\frac{3}{2}} \]  
(B.13)

We now find our normalization constant through demanding that our distribution for \( r \) must be normalized to unity
\[ \mathcal{N} \int_0^1 dr \frac{r + r^2}{(1 + r + r^2)^{\frac{3}{2}}} = \mathcal{N} \frac{4}{27} \]  
(B.14)

Where we evaluated the integral in Mathematica. Now we can readily evaluate the expectation value of \( r^{(n)} \) for a GOE spectrum
\[ \int_0^1 r \ P_{\text{GOE}}(r) dr = \int_0^1 dr \frac{27r}{4} \frac{r + r^2}{(1 + r + r^2)^{\frac{3}{2}}} \]
\[ = \left[ -\frac{x^3 + 15x^2 + 12x + 8}{2(x^2 + x + 1)^{\frac{3}{2}}} \right]_0^1 = 4 - 2\sqrt{3} \approx 0.53 \]  
(B.15)

Where we once again used Mathematica. We have thereby showed that \( \langle r^{(n)} \rangle_{\text{Poisson}} \approx 0.39 \) and \( \langle r^{(n)} \rangle_{\text{GOE}} \approx 0.53 \). The GOE prediction is something of an approximation but we find numerically that it is rather accurate. It is easy to find this numerically by considering a large ensemble of Gaussian orthogonal random matrices and estimating the average value of \( r^{(n)} \) over this ensemble.
Appendix C

Kubo Formula for Conductivity Tensor

Using linear response and in particular the Kubo formula we will deduce an expression for the DC part of the electric conductivity tensor. We know that the total current $J(\omega)$ can be expressed as the linear response to an electric field $E(\omega)$ as $J(\omega) = \sigma(\omega)E(\omega)$, and we have the Kubo formula for the conductivity

$$\sigma(\omega) = \frac{1}{\Omega} \int_0^\infty dt e^{-i\omega t} \int_0^\beta d\lambda \langle J(-i\lambda)J(t) \rangle$$ (C.1)

where the current operator $J$ is in the Heisenberg picture, i.e. $J(t) = e^{iHt}Je^{-iHt}$ where $H$ is the unperturbed (without electric field) Hamiltonian. The volume of the system is denoted $\Omega$ and $\langle \cdots \rangle = tr\{Z^{-1}e^{-\beta H} \cdots \}$ denotes the thermal average. We go through the integrals

$$\sigma(\omega) = \frac{1}{Z\Omega} \int_0^\infty dt e^{-i\omega t} \int_0^\beta d\lambda \sum_n \langle n| e^{-\beta H} J e^{\lambda H} e^{-iHt} J e^{iHt} |n \rangle$$

$$= \frac{1}{Z\Omega} \sum_{nm} e^{-\beta E_n} \int_0^\infty dt e^{-i\omega t} e^{i(E_m-E_n)t} \int_0^\beta d\lambda e^{\lambda(E_m-E_n)} \langle n|J|m \rangle \langle m|J|n \rangle$$ (C.2)

Where $\{|n\rangle\}$ are the eigenstates of the unperturbed Hamiltonian and we inserted a complete set of states. Now will we add an infinitesimal convergence factor $\eta t$ in the exponential. This corresponds to an adiabatic switching-on of the electromagnetic field at $t = -\infty$. The limit $\eta \to 0$ should be taken later

$$\sigma(\omega) = \frac{1}{Z\Omega} \sum_{nm} e^{-\beta E_n} \frac{1 - e^{\beta(E_n-E_m)}}{E_n - E_m} \int_0^\infty dt e^{i(E_m-E_n-\omega+i\eta)t} \langle n|J|m \rangle \langle m|J|n \rangle$$

$$= \frac{1}{Z\Omega} \sum_{nm} \frac{i \langle n|J|m \rangle \langle m|J|n \rangle}{E_m - E_n - \omega + i\eta} e^{-\beta E_n} - e^{-\beta E_m}$$ (C.3)

We are only interested in the DC-current and we will therefore be taking the limit of frequency going to zero later. We thus only need to consider the real part of the conductivity. We will
make use of the following identity

$$\frac{1}{\omega \pm i\eta} = \mathcal{P}\frac{1}{\omega} \pm i\pi\delta_\eta(\omega) \quad (C.4)$$

Where $\mathcal{P}1/\omega$ denotes the principal value of $1/\omega$ and the delta function has been regularized, but in the limit $\eta \to 0$ it goes to the standard Dirac delta function.

$$\Re\{\sigma(\omega)\} = \sum_{nm} \frac{\langle n|J|m\rangle\langle m|J|n\rangle}{Z\Omega} e^{-\beta E_n} e^{-\beta E_m} \Re\{i\mathcal{P} \frac{1}{E_m - E_n - \omega} - \pi\delta_\eta(E_m - E_n - \omega)\}$$

$$= \frac{\pi}{Z\Omega} \sum_{nm} \langle n|J|m\rangle\langle m|J|n\rangle \frac{e^{-\beta E_n} - e^{-\beta E_m}}{E_m - E_n} \delta_\eta(E_m - E_n - \omega)$$

$$= -\frac{\pi}{Z\Omega} \sum_{nm} \langle n|J|m\rangle\langle m|J|n\rangle e^{-\beta E_n} \frac{1 - e^{-\beta\omega}}{\omega} \delta_\eta(E_m - E_n - \omega) \quad (C.5)$$

We now take the limit of frequency going to zero. Where we Taylor expand

$$\lim_{\omega \to 0} \frac{1 - e^{-\beta\omega}}{\omega} = \lim_{\omega \to 0} \frac{1 - (1 - \beta\omega + \frac{1}{2}\beta^2\omega^2 + \ldots)}{\omega} = \beta \quad (C.6)$$

We then arrive at

$$\Re\{\sigma(\omega \to 0)\} = \frac{\pi\beta}{Z\Omega} \sum_{nm} \langle n|J|m\rangle\langle m|J|n\rangle e^{-\beta E_n} \delta_\eta(E_m - E_n) \quad (C.7)$$

We finally use that the total current $J$ is given as the sum over all local currents

$$J = \sum_r J_r = \sum_r \dot{e^r} \quad (C.8)$$

This yields our final result

$$\Re\{\sigma(\omega \to 0)\} = \frac{\pi\beta}{Z\Omega} \sum_{rr'} \sum_{nm} \langle n|J_{r+r'}|m\rangle\langle m|J_r|n\rangle e^{-\beta E_n} \delta_\eta(E_m - E_n) \quad (C.9)$$
Appendix D

RG Rules for Random transverse-Field Ising Model

We will here review the renormalization group transformation, as suggested by Fisher [64], to the transverse field Ising Hamiltonian with quenched randomness. The Hamiltonian for the model is

\[ H = -\sum_j J_j S_j^z S_{j+1}^z - \frac{1}{2} \sum_j h_j S_j^x \]  

(D.1)

Where the interaction strengths \( J_j \)'s are drawn independently from a distribution \( \pi_0(J) dJ \) and the transverse fields \( h_j \) from \( \rho_0(h) dh \), where both the \( h_j \)'s and the \( J_j \)'s are positive.

The renormalization transformation consists of the following steps; we find the largest interaction term or field in the system with strength \( \Omega = \Omega_0 \equiv \max\{J_j, h_j\} \) and decimate this away to get rid of the high energy information and focus on the desired low energy physics. There are two possibilities:

- If \( \Omega = J_j \) we make the approximation that the two spins \( S_j \) and \( S_{j+1} \) are rigidly locked together as a spin cluster with an effective field \( \tilde{h} \approx h_j h_{j+1}/\Omega \) and effective magnetic moment \( \tilde{g} = g_j + g_{j+1} = 2 \), obtained from lowest order perturbation theory in \( h_j/J_j \).

- If \( \Omega = h_j \) we simply eliminate this site and get an effective interaction strength \( \tilde{J} = J_j J_{j-1}/\Omega \) between the remaining nearest neighbor sites \( j+1 \) and \( j-1 \).

This procedure is to be iterated. At each step both \( \tilde{J} \) and \( \tilde{h} \) is necessarily smaller than \( \Omega \), so when we iterate the maximal energy scale will gradually decrease. We need to keep track of the distributions \( \pi_\Omega(\tilde{J}) \) and \( \rho_\Omega(\tilde{h}, \tilde{g}) \) at each step. Fisher showed that these distributions will broaden exponentially, which implies that the perturbative decimation process gets better and better.
Appendix E

C++ Code

We include some of the code which was used to produce the results in the thesis in this section. We only list some essential snippets of code and state their utility. We used a binary representation of the non-entangled product states and we used the Eigen library to store the matrices. We show how we find the Hamiltonian matrix and diagonalize it. To find the various expectation values we have more or less straightforwardly utilized the binary representation of the product states.

First we import the necessary libraries and make some initializations:

```cpp
#include<iostream>
#include<bitset>
#include<vector>
#include<math.h>
#include<cstdlib>
#include<ctime>
#include<Eigen>
#include<stdio.h>
#include"lapacke.h"

using namespace std; using namespace Eigen;

const int P = 10; // Number of Particles
const int N = pow(2.0,P); // Dimensionality of Problem
int S = 0; // Our choice of Spin Sector
double J = 1; // Strengths of Interaction
double W = 3; // Strength of Disorder
typedef Triplet<double> T;
vector<T> tripletList;
MatrixXd eig_vecs;
VectorXd eig_vals;
vector<double> r;
vector<int> Sz_sec;
srand(time(NULL));
```
The following code was used to find the product states with total spin in the z-direction equal to $S$ and the calculate the matrix elements of the Hamiltonian:

```cpp
// Finding the Basis - States in Given Spin - Sector
for (int i = 0; i < N; i++) if (spin_x(S, i)) Sz_sec.push_back(i);

// Calculating the matrix elements
double a, b; int i, j = 0;
for(vector<int>::iterator it1 = Sz_sec.begin(); it1 != Sz_sec.end(); ++it1)
{
    i = 0;
    for(vector<int>::iterator it2 = Sz_sec.begin(); it2 != Sz_sec.end(); ++it2)
    {
        x if (i==j) tripletList.push_back(T(j,i,inter_diag_elem(*it1,J)));
        a = *it1^*it2;
        if (check_dyad(a))
        {
            b = a - (abs(*it1 - *it2));
            if(is_integer(log2(b)) && b!=0) tripletList.push_back(T(i,j,J*0.5));
        }
        i++;
    }
    j++;
    if (j >= Sz_sec.size()) break;
}
int N_sector = Sz_sec.size();
SparseMatrix<double> H(N_sector, N_sector);
H.setFromTriplets(tripletList.begin(), tripletList.end());
```

Where we used the following functions:

```cpp
// Function for Checking if a Number is an Integer
bool is_integer(double k) { return floor(k) == k; }

// Calculating the Diagonal Part of the Interaction
double inter_diag_elem(int i, double J)
{
    bitset<P> a(i);
    double elem = 0;
    for (int i = 0; i < P - 1; i++)
    {
        if (a[i] == a[i + 1]) elem += J / 4;
        else elem -= J / 4;
    }
    if (a[0] == a[P - 1]) elem += J / 4;
    else elem -= J / 4;
    return elem;
}
```
// Checking if a binary Number has two adjacent 1’s and the rest 0’s
bool check_dyad(int a)
{
    bitset<P> b(a);
    int dyad = 0;
    int count = 0;
    int last_bit = 1;

    for (int i = 0; i < P; i++)
    {
        if (b[i] == 1)
            count += 1;
    }

    if (count == 2)
    {
        for (int i = 0; i < P; i++)
        {
            if (b[i] == 1 && i != P - 1)
            {
                if (b[i + 1] == 1)
                {
                    dyad = 1;
                    break;
                }
            }
        }
        if (b[0] == 1 && b[P - 1] == 1)
            dyad = 1;
    }

    return dyad;
}

// Checking if Product State has total Spin equal to given S
bool spin_tot(int S, int i)
{
    bitset<P> spins(i);
    bool is_spin_tot = 0;
    int count = 0;
    for (int i = 0; i < P; i++)
    {
        if (spins[i] == 1)
            count ++;
        else
            count --;
    }

    if (count == S)
        is_spin_tot = 1;

    return is_spin_tot;
}

We then calculate the disorder part of the Hamiltonian for each disorder realization as:

// Adding the random Fields
r = gen_ran_vec(W, (double)rand());
for(int i = 0; i < N_sector; i++) A.coeffRef(i,i) += rand_elem(r,Sz_sec[i]);
Where we used the following functions:

```cpp
// Generating the random Part of the i’th diagonal Element
double rand_elem(vector<double> ran_vec, int i) {
    bitset<P> spins(i);
    double elem = 0;
    for (int i = 0; i < P; i++) {
        if (spins[i]) elem += ran_vec[i]*0.5;
        else elem -= ran_vec[i]*0.5;
    }
    return elem;
}

// Generating a Vector of uniformly distributed random Numbers on [-W,W]
vector<double> gen_ran_vec(double W, double seed) {
    vector<double> R;
    srand(seed);
    for (int i = 0; i < N; i++) R.push_back(2*W*(rand()/double(RAND_MAX)-0.5));
    return R;
}
```

We diagonalize the matrix using the dsyev routine from LAPACK:

```cpp
// Diagonalizing
int LDA = N_sector;
int n = N_sector; lda = LDA, info, lw;
double wk;
double* work;
double w[N_sector];
double* L = A.data();
lw = -1;
LAPACK_dsyev((char*)"Vectors", (char*)"Upper", &n, L, &lda, w, &wk, &lw, &info);
lw = (int)wkopt;
work = (double*)malloc(lwork*sizeof(double));
LAPACK_dsyev((char*)"Vectors", (char*)"Upper", &n, L, &lda, w, work, &lw, &info);
if( info > 0 ) {
    printf("The algorithm failed to compute eigenvalues.\n");
    exit( 1);
}
free((void*)work);
eig_vals = Map<VectorXd>(w, N_sector);
eig_vecs = Map<MatrixXd>(L, N_sector, N_sector);
```
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