Entanglement Hamiltonians: From Field Theory to Lattice Models and Experiments

Marcello Dalmonte,* Viktor Eisler, Marco Falconi, and Benoît Vermersch

Results about entanglement (or modular) Hamiltonians of quantum many-body systems in field theory and statistical mechanics models, and recent applications in the context of quantum information and quantum simulation, are reviewed. In the first part of the review, what is known about entanglement Hamiltonians of ground states (vacua) in quantum field theory is summarized, based on the Bisognano–Wichmann theorem and its extension to conformal field theory. This is complemented with a more rigorous mathematical discussion of the Bisognano–Wichmann theorem, within the framework of Tomita–Takesaki theorem of modular groups. The second part of the review is devoted to lattice models. There, exactly soluble cases are first considered and then the discussion is extended to non-integrable models, whose entanglement Hamiltonian is often well captured by the lattice version of the Bisognano–Wichmann theorem. In the last part of the review, recently developed applications in quantum information processing that rely on the specific properties of entanglement Hamiltonians in many-body systems are summarized. These include protocols to measure entanglement spectra, and schemes to perform state tomography.

1. Introduction

Since its inception, quantum mechanics has routinely thrilled theoretical and experimental physicists due to its striking—and, at times, counterintuitive—differences from the classical world one is accustomed to. One of the most remarkable examples of those is certainly entanglement.\cite{1,2} This form of quantum correlations went relatively unnoticed until the last decades of the 20th century, but has experienced renewed and widespread interest over the last thirty years.

Entanglement captures the degree of inseparability between quantum mechanical states: it vanishes in the absence of quantum correlations, and it is instead finite in case separability is not satisfied. It has found numerous and profound applications in quantum information science, ranging from quantum communication to its connection to the capabilities of quantum computing hardware, just to name a few.\cite{1–3}

In parallel to progresses in quantum information and quantum optics, it has emerged that entanglement is not only useful to describe few-body processes at a microscopic level, but that, in addition, it is an invaluable resource in characterizing and classifying genuine quantum mechanical features of many-body systems.\cite{4–9} Nowadays, entanglement is routinely used in a variety of contexts. Since its first applications in many-body lattice models—in particular, focused on critical behavior\cite{4–6} and topological order\cite{7,8}—it has found applications as wide as the characterization of computational methods and their performances,\cite{10} to the classification of many-body behavior and holographic properties in quantum field theory.\cite{9,11–13}

The main actors in determining the relevance of entanglement in many-body systems are measures of bipartite entanglement (and, to a lesser extent, entanglement witnesses and measures of multipartite entanglement).\cite{2,14} Consider as an example the setting described in Figure 1: given a state \( \rho \), one is interested in characterizing the quantum correlations between its partition \( A \) and its complement \( B \). Those are captured by the reduced density matrix (RDM):

\[
\rho_A = \text{Tr}_B \rho = \sum_{\alpha=1}^{r_A} \lambda_\alpha |\alpha\rangle\langle\alpha| \tag{1}
\]

where the last equality represents an eigendecomposition with eigenvalues \( \lambda_\alpha \) and eigenvectors \( |\alpha\rangle \), and \( r_A \) is the rank of \( \rho_A \).\cite{15}
For the case of globally pure states—that is, states for which $\text{Tr} \rho^2 = 1$—a paramount quantity that quantifies the entanglement between $A$ and $B$ is the von Neumann entropy:[15] The von Neumann entropy and other related measures (such as concurrence[14]) have found extensive use in quantum many-body theory—see, for example, the reviews in refs. [4, 9, 11]. One key aspect of the entropy and related quantities is that they are solely sensitive to the eigenspectrum of RDM—the so-called entanglement spectrum.[17,27] The much richer relation between the finer structure of the RDM—that is, the structure of the eigenvectors, and their relation to the corresponding eigenvalues—has instead been initially overlooked, partly because it is considerably more challenging to characterize at the theoretical (and experimental) level. It is this finer structure that is the topic we plan to cover in this review.

Our starting point is the observation that, given the fact that a density matrix is always positive semi-definite, it is always possible to cast it in an exponential form (analogously to a “partition function”) as follows.

$$\rho_A \propto e^{-\mathcal{H}}$$

The dimensionless operator $\mathcal{H}$ is known under two different names: modular Hamiltonian in quantum field theory, and entanglement Hamiltonian (EH) in the context of lattice models and quantum information. Throughout this work, we will follow the second nomenclature. The operator $\mathcal{H}$ is bounded from below, as a consequence of the semi-positive-definiteness of the RDM.

The entanglement Hamiltonian has a somehow interesting history, that proceeds along three parallel routes. Historically, the first appearance of EH was related to developments in the field of axiomatic field theory (1970s): the most relevant achievement presented is the Bisognano–Wichmann theorem.[18,19] That theorem was the von Neumann entropy and other related measures (such as concurrence[14]) have found extensive use in quantum many-body theory—see, for example, the reviews in refs. [4, 9, 11]. One key aspect of the entropy and related quantities is that they are solely sensitive to the eigenspectrum of RDM—the so-called entanglement spectrum.[17,27] The much richer relation between the finer structure of the RDM—that is, the structure of the eigenvectors, and their relation to the corresponding eigenvalues—has instead been initially overlooked, partly because it is considerably more challenging to characterize at the theoretical (and experimental) level. It is this finer structure that is the topic we plan to cover in this review.

Our starting point is the observation that, given the fact that a density matrix is always positive semi-definite, it is always possible to cast it in an exponential form (analogously to a “partition function”) as follows.

$$\rho_A \propto e^{-\mathcal{H}}$$

(2)

Interestingly, the three aforementioned routes proceeded almost independently, with very few intersections, until very recently. In particular, developments in the field of quantum field theory have been almost completely decoupled from lattice statistical mechanics models.

Against this background, the aim of this review is twofold. First, we will present the basic toolkit and most representative results from those three separate research lines, and emphasize the net of connections between all of those that has been developed over the last years. The quantum field theory side of the story will be presented in Sections 2 and 3: we will start from a brief collection of all relevant results in the field, and then proceed with a more mathematical oriented derivation of the main tool available—Tomita–Takesaki theory—on the basis of the Bisognano–Wichmann theorem. In Section 4, we will discuss integrable lattice models: these models allow for considerable analytical insight into the structure of EH, that is of key importance to gauge the relevance of field theory results on the lattice. This discussion will naturally evolve to non-integrable lattice models in Section 5. There, we will emphasize the role played by entanglement Hamiltonians in the context of topological and quantum critical matter. The second aim of this review is to present applications of such knowledge in the broader context of quantum information science. Section 6 will be devoted to this task: in particular, we will review methods to characterize many-body systems in quantum computers and quantum simulations that leverage on the many-body insights based on entanglement Hamiltonians.

While, as mentioned above, different communities have studied entanglement Hamiltonians with limited crosstalks, we have opted to follow a different route. While we have kept the various lines separate in terms of sections, throughout the review, we have often commented on how results in one field have impacted others, both in terms of interpretation of the physical results, and methods. Hopefully, such choice may help the reader interested in a specific section to better appreciate the breadth of the field, and the various intersections.

The entanglement Hamiltonian is of course not disconnected from entropies and the entanglement spectrum: the first are nothing but moments of its expectation value, and the latter is its spectrum. Since these quantities have already been subjects of excellent reviews (see refs. [6, 7, 11]), the stress of our article will be on aspects of the EH that go beyond those—in particular, its structure as a many-body operator. While not extensively, we will refer to properties of the ES whenever useful. In addition to the aforementioned works, we emphasize that generic aspects of modular Hamiltonians have also been reviewed recently in Ref. [9].
2. Entanglement Hamiltonian: Definitions and Insights from Quantum Field Theory

In this section, we will first provide some definitions and basic observations on entanglement Hamiltonians and related quantities. Then, we will review some basic results about EH of quantum field theories. Some of those will then be reframed in a more rigorous mathematical framework in the next section.

2.1. Definitions and Notations

Let us recall here some basic properties of the RDMs we are dealing with. First of all, we will always work with normalized RDMs

\[ \text{Tr} \rho_A = 1, \quad \sum_a \lambda_a = 1 \quad (3) \]

This will require some care in defining the EH: indeed, shifting the operator \( H \) by a constant will change the trace of \( \rho_A \). We will utilize the following normalization

\[ \rho_A = e^{-H/Z} \quad (4) \]

which is related to another one used in the literature as follows:

\[ \rho_A = e^{-(H+C)}, \quad Z = \exp C \quad (5) \]

where \( C \) is a constant. This consideration implies that the absolute values of the entanglement spectrum \( \epsilon \) have little operational meaning, while what is really relevant are the corresponding entanglement energy differences and relative magnitudes. This fact will have direct implications in protocols to measure the ES experimentally.

It is worth noting that, at first sight, it is very unclear whether the EH shall have a meaningful many-body structure. Indeed, one could simply interpret it as the logarithm of a density matrix. In general, the corresponding operator does not have to be local, neither few-body.\cite{18,19}

It is thus a remarkable fact that there are generic circumstances under which the EH of an extended partition \( A \) of a many-body system is extremely well approximated—if not exactly given—in terms of operators \( H(x) \) that are both local and few-body

\[ H = \int_{x \in A} dx \, \beta(x) \, H(x) \quad (6) \]

where \( \beta(x) \) is some weight function. The reason behind this fact traces back to quantum field theory (that naturally encompasses locality, relativistic invariance, and quantum mechanical effects), and is centered around the locality of many-body dynamics.

2.2. Entanglement Hamiltonian and the Bisognano–Wichmann Theorem

The most basic result at the heart of the few-body, local nature of EH is the Bisognano–Wichmann theorem.\cite{18,19,20,26} Consider a relativistic quantum field theory in (\( D+1 \)) dimensions (\( D \) denotes the spatial dimensions only), with Hamiltonian density \( H(x) \), and with coordinates \( x = \{ x_1, x_2, ..., x_3 \} \) labeling spatial coordinates. Given a half-plane partition \( A \) defined by the condition \( x_1 > 0 \), the Bisognano–Wichmann theorem states that the entanglement Hamiltonian of the vacuum state reads

\[ H = \frac{2\pi}{c} \int_{x \in A} dx \, x_1 \, H(x) \quad (7) \]

where \( c \) is the corresponding “speed of light,” which makes the EH dimensionless. This result—applicable irrespective of the particle content of the theory, and compatible with the existence of gauge symmetries—establishes that not only is the EH of such states local and few-body: it is actually nothing but the boost operator, that is, the original Hamiltonian with space-dependent couplings.

A first physical interpretation of the Bisognano–Wichmann theorem is illustrated in Figure 2a: one can either see Equation (7) as describing a thermal state with respect to the boost operator, or, equivalently, describing a thermal state with respect to the original Hamiltonian, but with a position-dependent temperature. In this second interpretation, the entanglement temperature is “high” close to the boundary, and progressively decreases moving away from it. This interpretation immediately suggests that the quantum correlations between \( A \) and its complement are overwhelmingly dominated by the “high” region—the boundary—thus providing a simple picture to understand a conjecture (proved in \( D = 1 \) for the case of gapped theories\cite{24}) known as area law.\cite{5} At the end of next section, we will discuss another physical interpretation in the context of the Unruh effect\cite{25} (somehow related to the first viewpoint above).

2.3. Entanglement Hamiltonians in the Presence of Conformal Symmetry

In the presence of additional global conformal symmetry, the result above can be extended to different systems, and different types of partitions. Here, we review the basic extensions of the BW theorem that have been obtained so far.

2.3.1. D-Dimensional Theories

For a partition \( A \) consisting of a ball of volume \( V \) and radius \( R \) in a \( D \)-dimensional relativistic conformal field theory, it was shown in Ref.\cite{20,26} that the corresponding entanglement Hamiltonian reads

\[ H_A^{CFT} = \frac{2\pi R}{c} \int_{x \in A} dx \, \frac{R^2 - r^2}{2R^2} \, T_{00}(x) \quad (8) \]

where the ball has been placed at the origin, \( r(x) \) is the Euclidean distance from the origin, \( T_{00} \) is the stress-energy tensor of the CFT (in fact analogue of the Hamiltonian density). The physical consequence of this result is that, in the presence of conformal symmetry, the EH retains all its useful locality and few-body properties even in cases of finite partitions, even if it is not correspondent to the boost operator any more. Note that the treatment of more complicated partitions (other than the half-space.
and the sphere in $D > 1$ is very complicated in general, however, some approximate approaches exist for free field theories, predicting the emergence of anisotropic inverse entanglement temperatures.

### 2.3.2. 1D Theories

In $D = 1$ spatial dimension, conformal symmetry becomes even more powerful in allowing the determination of EH for various types of partitions. This was originally explored in ref. [20], and further extended in ref. [28]. We review here the basic cases of interest to equilibrium physics, noting that, remarkably, some extensions to time-dependent problems have also been derived.

The first relevant case deals with a partition of finite size $\ell$, embedded in an infinite chain (Figure 2b). In that case, the EH reads

$$\mathcal{H}^{\text{CFT2}}_A = \frac{2\pi\ell}{c} \int_0^\ell dx \sin \left(\frac{x}{\ell}\right) H(x)$$

(9)

that corresponds to the $(1+1)$-d version of Equation (8). Note that, close to each of the boundaries, the EH is reminiscent of the original BW functional form. By exploiting conformal mappings, it is possible to generalize this formula to the case of a finite interval inside a finite circle of circumference $L$, that then reads

$$\mathcal{H}^{\text{CFT2}}_A = \frac{2L}{c} \int_0^\ell dx \sin \left(\frac{x}{L}\right) H(x)$$

(10)

This last equation will be particularly useful in the numerical studies in Section 5.

Finally, it is possible to consider a finite interval of length $L/2$ at the boundary of an open chain of length $L$ (Figure 2b): in this case, the EH reads

$$\mathcal{H}^{\text{CFT3}}_A = \frac{2L}{c} \int_0^{L/2} dx \sin \left(\frac{\pi x}{L}\right) H(x)$$

(11)

The physical interpretation of these results is immediate, and graphically illustrated in Figure 2: all of them predict $\beta(x)$ to be linear close to the boundaries, that is, the entanglement temperature decreases with the inverse of the distance within the “entangled” region, whereas $\beta(x)$ saturates following a “parabolic-like” shape at large distances, consistently with the generic prediction in ref. [26].

### 2.4. Entanglement Hamiltonian at Finite Temperature

So far, we have only dealt with vacuum states of a field theory—that is, pure states. Remarkably, under some specific conditions, it is also possible to get insights on the structure of the EH for the case of mixed states.

The first examples are thermal states at inverse temperature $\beta$. If one considers a small partition embedded in a large system whose dynamics is described by a non-integrable theory, the corresponding entanglement Hamiltonian is determined by quantum typicality, and it is nothing but the Hamiltonian of the subsystem $\mathcal{H}_A$, that is, $\rho_A \propto e^{-\beta H_A}$, up to corrections due to boundary terms. This formula is expected to work well for temperatures larger than the typical energy scales in the system.

Remarkably, again for the case of $(1+1)$-d conformal field theories, even more precise statements can be made at any temperature. In that case, the EH of a finite region embedded in an infinite system reads

$$\mathcal{H}^{\text{CFT4}}_A = 2\beta \int_0^\ell dx \left[ \left. \frac{\sin \left(\frac{\ell x}{\beta c}\right) \sin \left(\frac{x}{\beta c}\right)}{\sin \left(\frac{\ell x}{\beta c}\right)} \right| H(x) \right]$$

(12)

The result above reproduces Equation (9) in the limit $\beta \gg \ell$, where the EH properties are well captured by the BW theorem. On the other hand, in the limit $\beta \ll \ell$ one approaches a Gibbs state at inverse temperature $\beta$. This prediction would also be obtained, assuming typicality, for pure states of the total system at high energies. We note that some of these results can also be extended to the post-measurement case, as has been show in ref. [32].

Figure 2. Bisognano–Wichmann theorem and entanglement Hamiltonian in quantum field theory. a) For a generic, relativistic quantum field theory, the entanglement Hamiltonian of a half-system is given by the boost operator. The corresponding RDM can be interpreted as either a thermal state with Hamiltonian density $x_1 \mathcal{H}(x)$, where $x_1$ is the distance from the boundary, or with an equilibrium state subject to a position-dependent inverse temperature $\beta(x) = 2\pi x_1/c$, and Hamiltonian density $\mathcal{H}(x)$. In the second interpretation, the entanglement temperature goes from “high” close to the boundary between the partitions (red), to cold away from the boundary (blue). b) Some of the extensions of the Bisognano–Wichmann to $(1+1)$ dimensional conformal field theory, formulated in terms of a space-dependent inverse entanglement temperature: finite interval in an infinite system (top) and half chain with open boundary conditions (bottom).
The examples discussed so far are valid for generic CFTs. Further results exist for some specific field theories, such as the free boson or fermion theories. In particular, the case of multiple disjoint intervals has been derived for the massless Dirac fermion and free scalar theories, where the local part of the EH has a similar structure (6) as in the single interval case. Namely, the energy density is multiplied with an appropriate space-dependent weight function \( \beta(x) \) which can be obtained explicitly. Some of these results can even be extended to finite temperatures. For the Dirac theory the main new feature is a subleading bilocal piece in the EH, which couples only to a single conjugate point in each of the intervals. Similar features have been observed for the massless Dirac fermion in the presence of a boundary or defect. The explicit treatment of the massive case is much more difficult even for a single interval, and the available results suggest the presence of a subleading nonlocal piece in the EH. Remarkably, similar nonlocal terms were found to appear also for massless fermions in the presence of zero modes and in low-lying excited states.

Finally, one should mention that some extensions to inhomogeneous physical systems exist, the simplest example being the massless Dirac fermion with a spatially varying Fermi velocity. This inhomogeneity can be absorbed by a local change of the metric and is equivalent to considering the Dirac theory in a curved background. Considering the EH of a singly connected partition, the effect of the curved metric can be taken into account and leads again to a local expression (6) where \( \beta(x) \) can be obtained explicitly.

We have now provided a full picture about what is known about EH in quantum field theory. The pivotal result throughout our discussion has been the Bisognano–Wichmann theorem, both as a result per se, and as a starting point for extensions to conformal field theory. In the following section, we will address the question: why is the EH taking such a simple functional form? In particular, what is the principle that determines its locality? Those questions are properly answered in the context of axiomatic quantum field theory, that we will try to present in a concise form that is of use to theoretical physics. Readers who are not familiar with QFT can proceed directly to Section 4, where we discuss lattice statistical mechanics models.

### 3. Tomita–Takesaki’s Modular Theory and Bisognano–Wichmann’s Theorem in Algebraic QFT: Mathematics Comes to the Rescue

In this section we will briefly overview two interconnected mathematical aspects of quantum theories that play a crucial, if somewhat unexpected, role in the description of entanglement Hamiltonians: Tomita–Takesaki theory of modular groups, and Bisognano–Wichmann’s theorem in algebraic QFT. The mathematical details will be kept to a minimum, the interested reader may refer to several books and reviews on algebraic quantum field theory, and the references thereof contained; among those, let us single out from which we drew inspiration in writing this review, and the more physically oriented review by Witten.

### 3.1. Tomita–Takesaki’s Theory

This theory has been developed by Tomita in the late sixties [the results were announced in], and then expanded and made known to a broad audience by Takesaki with his 1970 lecture notes.

The theory of Tomita and Takesaki provides a way to construct, given an algebra of quantum observables \( \mathcal{M} \) realized as operators on some Hilbert space \( \mathcal{H} \), and a quantum state realized as a vector \( \Omega \) (with some “reasonable” properties, to be detailed below), two operators associated with the couple \((\mathcal{M}, \Omega)\): the modular operator \( \Delta \), and the modular conjugation \( J \). These operators enjoy some nice properties, again to be detailed below.

The importance of modular operators \( \Delta \) and \( J \) in characterizing algebras of operators, both as pure mathematical objects and as collections of quantum observables, has very soon become apparent. In fact, on the one hand Tomita–Takesaki’s theory is at the heart of Connes’ 1973 classification of “type III” factors in von Neumann algebras, and on the other hand it has been put in relation to the KMS condition describing quantum equilibrium states by Takesaki in his 1970 lecture notes, and by himself and Winnick in 1973, while studying the algebraic properties of KMS states. While the second aspect is the one that is more relevant for this review, let us mention nonetheless that all relativistic local algebras of observables in QFT turn out to be type III factors, in particular of the so-called “type III1” in Connes’ terminology. They are characterized by the fact that the spectrum of the modular operator \( \Delta \) is the whole real line, [see]. Before Connes’ work, inspired as we said by Tomita–Takesaki theory, the type III factors were very poorly understood (in fact, the type III was not even defined), and thus this is another key application of such theory to quantum physics, even if it lies beyond the scope of this review.

To introduce Tomita–Takesaki’s modular operators, and their connection to equilibrium states in quantum statistical mechanics, let us fix some mathematical terminology. Preliminarily, let us remark that Tomita–Takesaki’s theory can be developed in a more general framework than the one presented below, however the setting given here is general enough for the physical applications we have in mind. Let \( \mathcal{H} \) be a separable Hilbert space, and let \( \mathcal{M} \) be a collection of bounded operators (quantum observables). Then \( \mathcal{M} \) is an algebra if and only if it is closed under addition and multiplication (i.e., given any \( A, B \in \mathcal{M} \), then \( A + B \in \mathcal{M} \) and \( AB \in \mathcal{M} \)). The commutant \( \mathcal{M}' \) of an algebra of operators \( \mathcal{M} \) is the collection of all bounded operators on \( \mathcal{H} \) that commute with all elements of \( \mathcal{M} \). Clearly, \( \mathcal{M} \subseteq \mathcal{M}' \) (an algebra is contained in its bicommutant). An algebra of operators \( \mathcal{M} \) is a von Neumann algebra iff

\[
\mathcal{M} = \mathcal{M}' \tag{13}
\]

Given a von Neumann algebra \( \mathcal{M} \), the center is given by \( \mathcal{M} \cap \mathcal{M}' \). Then, \( \mathcal{M} \) is a factor iff \( \mathcal{M} \cap \mathcal{M}' = \mathbb{C}1 \), i.e that is, the only operators in \( \mathcal{M} \) that commute with all the others are multiples of the identity operator. Now, let \( \Omega \in \mathcal{H} \) be a vector (associated to the quantum state \( \Omega \langle \Omega \rangle \)), then \( \Omega \) is cyclic and separating iff both \( \mathcal{M}\Omega \) and \( \mathcal{M}'\Omega \) are dense subsets of \( \mathcal{H} \).
conjugation antilinear operators acting as

\[ S M \Omega = M' \Omega \]

\[ F M' \Omega = (M')^\dagger \Omega \]

They are both densely defined, since \( \Omega \) is cyclic and separating. Tomita proved that their closures exist, they are one adjoint to another (\( F = S' \)), and they can be expressed, as every closed operator, through an antilinear polar decomposition \( S = J \Delta^{1/2} \), that introduces the modular operators. The term “modular” is used exploiting the analogy between the operator polar decomposition and the polar expression of complex numbers (\( \Delta^{1/2} \) plays the role of the modulus, and \( J \) that of the phase, that in addition takes antilinearity into account). Because \( S \) and \( S' \) conjugate the observables in \( M \) and in its commutant, the modular operators inherit very useful properties, described below.

### 3.1. The Modular Operator and the Modular Conjugation

Given a couple \( (M, \Omega) \), where \( M \) is a von Neumann algebra of operators on \( \mathcal{H} \) and \( \Omega \in \mathcal{H} \) is cyclic and separating, a couple of operators \( (\Delta, J) \) on \( \mathcal{H} \) can be defined, called, respectively, the modular operator and the modular conjugation. The couple \( (\Delta, J) \) is identified uniquely by the following properties:

- \( \Delta \) is self-adjoint, positive, and invertible; in addition, the vector \( \Omega \) is invariant under the action of both \( \Delta \) and \( J \)

\[ \Delta \Omega = J \Omega = \Omega \]  

(15)

- The unitary group \( (\Delta^t)_{t \in \mathbb{R}} \) defines a group of automorphisms of the algebra \( M \): for any \( t \in \mathbb{R} \)

\[ \Delta^t M \Delta^{-t} = M \]  

(16)

Let us denote by

\[ \sigma_t(A) = \Delta^t A \Delta^{-t}, \quad A \in M \]  

(17)

such a group of automorphisms, also called the modular group.

- \( \Delta \Omega \subseteq D(\Delta^{1/2}) \), the latter being the domain of self-adjointness of \( \Delta^{1/2} \).

- The operator \( J \) is a conjugation, that is, \( J \) is antilinear, \( J = J^* = J^{-1} \), and it commutes with the modular group: \( [J, \Delta^t] = 0 \). Hence

\[ J \Delta J = \Delta^{-1} \]  

(18)

- The modular conjugation maps the algebra \( M \) into its commutant

\[ JMF = M' \]  

(19)

- The operators \( S = J \Delta^{1/2} \) and \( S^* = \Delta^{1/2} J = J \Delta^{-1/2} \) "conjugate" \( \mathcal{M} \Omega \): \( \forall A \in M, \forall A' \in M' \)

\[ SA \Omega = A' \Omega \]  

(20)

- The function

\[ R \ni t \mapsto \Delta^t A \Omega \in \mathcal{H}, \quad A \in M \]  

(21)

has an analytic continuation in the strip \( \{ z \in \mathbb{C}, -1/2 < \Im z < 0 \} \). The function

\[ R \ni t \mapsto \Delta^t A' \Omega \in \mathcal{H}, \quad A' \in M' \]  

(22)

has an analytic continuation in the strip \( \{ z \in \mathbb{C}, 0 < \Im z < 1/2 \} \). As a consequence, \( \forall A \in M \) and \( \forall A' \in M' \)

\[ \Delta(t t^*) A \Omega = \Delta(t A t^*) \Omega, \quad \Delta(i t^*) A' \Omega = \Delta(i A' t^*) \Omega \]  

(23)

- The function \( t \mapsto \langle \Omega | B \sigma_t(A) \Omega \rangle \), \( A, B \in M \), can be analytically continued in the strip \( \{ z \in \mathbb{C}, -1 < \Im z < 0 \} \). At the lower boundary, the following condition holds

\[ \langle \Omega | B \sigma_{-1}(A) \Omega \rangle = \langle \Omega | \sigma_1(A) B \Omega \rangle \]  

(24)

or equivalently,

\[ \langle \Omega | B \Delta((i t^*) - 1) \Omega \rangle = \langle \Omega | \Delta^{-(i t^*)} B \Omega \rangle \]  

(25)

The last property, 24, is called \( \sigma \)-KMS condition for the state \( \langle \Omega | \Omega \rangle \), and it is related to the notion of equilibrium in quantum statistical mechanics, as we will explain shortly. Let us remark that, even if it may not be apparently so, this KMS condition is by itself sufficient to identify the modular couple \( (\Delta, J) \).

### 3.1.2. Equilibrium States, and their Relation to Modular Groups

What is an equilibrium state in a quantum theory? Given the evolution of a physical system as a unitary operator \( e^{i t H} \) on some Hilbert space, the equilibrium state \( \omega_\beta \) at inverse temperature \( \beta \in \mathbb{R} \) is typically given by the Gibbs prescription

\[ \omega_\beta(A) = \frac{1}{\text{tr} e^{-\beta H + \mu N}} \text{tr}(e^{-\beta H + \mu N} A) \]  

(26)

provided that such a state makes sense mathematically (e.g., \( \text{tr} e^{-\beta H + \mu N} \) could not be finite; this typically happens in taking the thermodynamic limit). A generalization of the notion of Gibbs state, that survives the thermodynamic limit and is applicable to QFTs, is that of a KMS state.[55] Let \( (\tau_t)_{t \in \mathbb{R}} \) be a group of automorphisms of a von Neumann algebra of observables \( M \). Then a state \( \omega_\beta \) on \( M \) is a \( \tau \)-KMS state at inverse temperature \( \beta \in \mathbb{R} \) iff for any \( t \in \mathbb{R} \), and for any \( A, B \in M \)

\[ \omega_\beta(\tau_t (A) B) = \omega_\beta(B \tau_{-t} (A)) \]  

(27)

In other words, the state \( \omega_\beta \) behaves almost like a trace, the correction to being a trace “measured” by \( \tau_t \).

If \( \tau_t (A) = e^{it H} A e^{-it H} \), and the Gibbs state can be defined, then the latter is a \( \tau \)-KMS state at inverse temperature \( \beta \). It can moreover be shown that in a finite volume, the only KMS states are the Gibbs states [see, for example, ([49], Ch. V)].
Comparing (24) with (27), it is clear that \( |\Omega(\Omega) \) is a \( \sigma \)-KMS state at inverse temperature \( \beta = -1 \). It is perhaps convenient to rescale temperature to its physical value \( \beta \) in the units of our choosing. First of all, by functional calculus it is possible to rewrite

\[
\Delta = e^{-K} \tag{28}
\]

where \( K \) is now called the modular Hamiltonian. In 24, let us make the substitution \( t = -\beta^{-1} s \)

\[
\langle \Omega | B_{\sigma^{-1}}(A)\Omega \rangle_{\mathcal{F}} = \langle \Omega | \sigma^{-1}(A)B\Omega \rangle_{\mathcal{F}} \tag{29}
\]

Now, let us define \( H = \beta^{-1} K \), obtaining

\[
\langle \Omega | B_{\tau^{-\beta}}(A)\Omega \rangle_{\mathcal{F}} = \langle \Omega | \tau(A)B\Omega \rangle_{\mathcal{F}} \tag{30}
\]

that is, \( |\Omega(\Omega) \) is a \( \tau \)-KMS state at inverse temperature \( \beta \), with the evolution \( \tau \) generated by \( H = \beta^{-1} K \).

We can therefore draw the following key conclusion:

Any equilibrium state at inverse temperature \( \beta \) can be seen as a faithful state\(^{[56]} \) over the algebra \( \mathcal{M} \) of observables, whose modular group is given by time translations, with a group parameter \( s \) related to physical time by \( t = -\beta s \).

We also have a nice converse piece of information, due to Takesaki.\(^{[52]} \) Let \( \mathcal{M} \) be a von Neumann algebra, and \( \omega \) a faithful state (represented on some \( \mathcal{F} \) by \( |\Omega(\Omega) \), with \( \Omega \) cyclic and separating), whose associated modular group is given by \( \sigma_{t} \in \mathbb{R} \). Are there any other equilibrium states, apart from \( \omega \), with respect to the modular group flow? The answer is as follows. Let \( \rho \) be a density matrix. Then the following two statements are equivalent:

- \( \rho \) is a \( \sigma \)-KMS state
- There exists a (unique) positive operator \( T \) on \( \mathcal{F} \), whose spectral decomposition belongs to the center of \( \mathcal{M} \), such that for all \( A \in \mathcal{M} \)

\[
\text{tr}_{\mathcal{F}} \rho A = \langle T^{1/2} \Omega, AT^{1/2} \Omega \rangle_{\mathcal{F}} \tag{31}
\]

Clearly, if \( \mathcal{M} \) is a factor (as many physically interesting algebras of observables are) then \( T = 1 \), that is, \( \omega \) is the unique equilibrium state for the modular group flow.

### 3.2. The Theorem of Bisognano and Wichmann

Bisognano and Wichmann’s results, published in a couple of papers between 1975 and 1976,\(^{[18,19]} \) concern local algebras of observables in relativistic quantum theories, and in particular the concept of Haag duality for wedge regions, to be explained below. Their results have then been extended to CFTs by Hislop and Longo,\(^{[20]} \) in conformal theories, duality can be proved for more general regions than the wedges considered by Bisognano and Wichmann, in particular for diamonds (see the illustrative examples in Section 2.3).

At a first glance, the result of Bisognano and Wichmann seems unrelated to Tomita–Takesaki’s theory. This is however true only in appearance, in fact Bisognano and Wichmann themselves remarked the connection of their result to the modular theory. We focus on such connection, and on the additional connections with entanglement Hamiltonians, in the last part of this section. For the moment, let us focus on Bisognano–Wichmann’s theorem itself.

#### 3.2.1. Haag Duality in Relativistic Quantum Theories

If one wants to build up a (field) theory of relativistic observables in Minkowski spacetime (or on any other Lorentzian manifold), causality shall be taken into account. However, there are serious mathematical obstructions to the definition of pointwise local relativistic quantum observables: as is well known, quantum fields should be smeared by smooth functions supported on some open region in the Minkowski spacetime, in order for them to make sense as operators. A field that is smeared by a function supported in a region \( \mathcal{O} \), is said to be localized in \( \mathcal{O} \). Causality for localized fields is assumed in the following form: fields that are localized on space-like separated regions of spacetime shall commute between each other. Algebras of observables in QFT are built by the action of fields (and possibly their momenta). Given a spacetime region \( \mathcal{O} \), the local algebra of observables \( \mathcal{M} (\mathcal{O}) \) shall be, intuitively, the one constructed by the action of fields localized in the region \( \mathcal{O} \). In general, Haag’s idea is to start with a collection of abstractly defined local algebras \( \mathcal{M} (\mathcal{O})_{\mathcal{O} \subset \Sigma} \) of observables, each defined on an open region \( \mathcal{O} \) of a given spacetime \( \Sigma \). Such a collection of local algebras shall satisfy certain assumptions that reflect the physical axioms of a relativistic theory, and in particular causality, in the form of commutation of space-like separated observables.

A desirable feature, that is however somewhat difficult to prove, is the following, called Haag duality. Let us denote by \( \mathcal{O}^\prime \) the causal complement of the region \( \mathcal{O} \), that is, the set of space-time points that are at a space-like distance from all the points of \( \mathcal{O} \). Then duality is true for the region \( \mathcal{O} \) iff

\[
\mathcal{M} (\mathcal{O})^\prime = \mathcal{M} (\mathcal{O}^\prime) \tag{32}
\]

In other words,

The commutant of a local algebra of observables in \( \mathcal{O} \) consists precisely of all observables in the causal complement \( \mathcal{O}^\prime \).

We will not discuss all the results about duality available in the literature, let us only mention, in addition to the Bisognano–Wichmann’s papers, the pioneering works by Araki,\(^{[57,58]} \) Dell’Antonio,\(^{[59]} \) and Eckmann–Osterwalder\(^{[60]} \) for special Minkowski regions and free field’s scalar observables.

#### 3.2.2. The Result of Bisognano and Wichmann: Haag Duality on Wedges

Let \( \Sigma \) be the Minkowski spacetime: \( \Sigma = (\mathbb{R}^{3}, \eta_{\mu \nu}) \), where

\[
\eta = \begin{pmatrix}
+ & 0 & 0 & 0 \\
0 & - & 0 & 0 \\
0 & 0 & - & 0 \\
0 & 0 & 0 & -
\end{pmatrix} \tag{33}
\]
is the flat pseudo-Riemannian metric. The right wedge \( W_R \subset \Sigma \) is the open set enclosed by two light rays starting from the origin and propagating in the \( x^1 \) direction:

\[
W_R = \{ x = (x^0, x^1, x^2, x^3) \in \Sigma, \quad x^1 > |x^0| \}
\]  

(34)

Its causal complement is the left wedge \( W_L \):

\[
W'_L = W_L = \{ x = (x^0, x^1, x^2, x^3) \in \Sigma, \quad x^1 < -|x^0| \}
\]  

(35)

Let us now denote by \( \Lambda \) a generic spacetime transformation belonging to the Poincaré group. Of special usefulness will be the following Lorentz transformations: the boosts belonging to the Poincaré group. Of special usefulness will be the boosts \( \Lambda(s) \) in the 1 direction, with parameter \( s \in \mathbb{R} \):

\[
\Lambda(s) = \begin{pmatrix}
\cosh s & \sinh s & 0 & 0 \\
\sinh s & \cosh s & 0 & 0 \\
0 & 0 & 1 & 0 \\
0 & 0 & 0 & 1
\end{pmatrix}
\]  

(36)

and the spatial rotation \( R_\theta(\pi) \), meaning a rotation of spatial coordinates by an angle of \( \pi \) around the 1-axis.

Let us now consider a possibly interacting quantum field theory on Minkowski spacetime. In particular, let us suppose that we can properly define the vacuum representation of the field theory as an algebra of operators acting on a Hilbert space \( \mathcal{H} \) on which the vacuum \( \Omega \in \mathcal{H} \) is represented as a vector, and that there is a unitary representation \( U(\Lambda) \) of every Poincaré transformation, suitably acting on smeared quantum fields, that are defined as self-adjoint operators \( \varphi(f) \), where \( f \) is any smooth test function on \( \Sigma \). The rigorous construction of the “correct” local algebras of observables \( \mathcal{M}(W_R) \) and \( \mathcal{M}(W_L) \), localized on the right and the left wedge, respectively, given by Bisognano and Wichmann is rather technical, and we shall omit it here. It suffices to keep in mind that these algebras (of bounded operators) are suitably related to the (unbounded) operators obtained by forming polynomials of the fields, each one localized in the right or left wedge, respectively.

The (main) theorem of Bisognano and Wichmann then reads as follows:

Haag duality holds for the von Neumann algebras \( \mathcal{M}(W_R) \) and \( \mathcal{M}(W_L) \). More precisely (by taking the commutant of the Haag duality for the right wedge below, one gets Haag duality for the left wedge)

\[
\mathcal{M}(W_R)' = \mathcal{M}(W_L) = \mathcal{M}(W_R')
\]  

(37)

This result positively resonated in the community of mathematical physics: the aforementioned duality theorems of Araki and Eckmann–Osterwalder, despite being far from trivial, were true only for free scalar fields; the results of Bisognano and Wichmann are true for every field theory that can be reasonably defined, be it free or interacting. The only drawback is that one shall restrict to wedges. As already mentioned, their result (and idea of the proof) was later extended to conformal fields by Hislop and Longo, where, thanks to the additional conformal symmetry, more general spacetime regions such as diamonds could be considered. Bisognano–Wichmann’s theorem is still a topic of research in algebraic QFT nowadays [see, e.g., [51–53], and the references therein].

3.2.3. The Link to Modular Operators

How is the Haag duality of Bisognano and Wichmann related to Tomita– Takesaki’s modular theory? The link, although not used directly, was already remarked by Bisognano and Wichmann in their original paper.\(^{[33]}\) In fact, they are able to provide one of the very few concrete and explicit realizations of the modular couple \( (\Delta, J) \). The special properties of the modular couple, explicit in their case, play a crucial role in their proof of Haag duality.

The von Neumann algebras they consider are, clearly, the wedge local algebras \( \mathcal{M}(W_R) \) and \( \mathcal{M}(W_L) \). A very general result of relativistic QFTs, that goes by the name of Reeh–Schlieder’s theorem,\(^{[64,65]}\) ensures that the vacuum \( \Omega \) is cyclic and separating for \( \mathcal{M}(W_R) \) (and \( \mathcal{M}(W_L) \)). Hence, there exists a modular couple \( (\Delta, J) \) associated to the vacuum on the right wedge local algebra (and analogously for the left wedge). To write the couple explicitly, let us consider the unitary realization of the group of boosts \( U(\Lambda(s)) \). As any strongly continuous unitary group, it can be written by Stone’s theorem as

\[
U(\Lambda(s)) = e^{iK}
\]

(38)

with \( K \) some self-adjoint operator on \( \mathcal{H} \), the quantum generator of the Lorentz boost. In addition, let us denote by \( \Theta \) the CPT operator on \( \mathcal{H} \), where CPT stands for charge-parity-time reversal (such an operator can always be defined, both abstractly and when possible concretely, on a relativistic QFT). Then

\[
\Delta_R = e^{-2\pi K}
\]

\[
J_k = \Theta U(R_k(\pi))
\]

(39)

Hence, it also follows that the modular group of the right wedge is given by the adjoint action of the group of boosts \( U(\Lambda(2\pi x_1)) \) on \( \Omega \).

This remarkable aspect of Bisognano–Wichmann’s construction is key for applications: it paved the way to the use of Tomita–Takesaki’s theory as the foundational motivation of important concepts of both theoretical and experimental physics, in particular that of entanglement Hamiltonian.

3.3. The Modular Hamiltonian as an Entanglement Hamiltonian: Relation to Unruh’s Effect

To explain the connection (even more so, the identification) between modular and entanglement Hamiltonians, it is useful to reformulate the Bisognano–Wichmann setting in a slightly different way.

We are given a quantum theory, that is bipartite: one part consists of the right wedge, and the observables \( \mathcal{M}(W_R) \) localized on it, the rest of all other observables. The vacuum state \( \omega_R(\Omega) = |\Omega \rangle \langle \Omega| \) is a “special state,” being the ground state of the whole system. It is also a state when restricted to the subsystem...
of local observables $\mathcal{M}(W_\beta)$, again playing a special role: it is in fact faithful, and thus the corresponding modular couple $(\Delta_\beta, J_\beta)$ can be defined.

However, in making the restriction to the wedge and its local observables, the nature of the vacuum state changes. In fact, as we explained in 3.1.2, $\omega_\beta$ is now a thermal equilibrium state, at temperature $\beta = -1$, for the modular group of the right wedge, that corresponds to the adjoint action of the group of boosts $\left( U(\Lambda(2\pi s)) \right)_{\text{rel}}$, as discussed above. Equivalently, defining the entanglement Hamiltonian as $H = \beta^{-1} K$, and time as $t = -(2\pi \beta)s$, one obtains that $\omega_\beta$ is an equilibrium state at temperature $\beta$ for the entanglement Hamiltonian time flow. We can rephrase the last statement in a more physical fashion:

In a relativistic field theory, tracing out the degrees of freedom other than the ones localized on the right wedge, thermalizes the vacuum. The resulting state is an equilibrium (Gibbs) state, at inverse temperature $\beta < \infty$, with respect to the flow generated by the entanglement Hamiltonian $H$, given by a suitable rescaling of the generator of Lorentz boosts $K$.

The general physical definition of an entanglement Hamiltonian is indeed the Hamiltonian (in a subsystem) with respect to which the partial trace (with respect to the other subsystem) of a ground or equilibrium state on the total bipartite system thermalizes. So, the entanglement Hamiltonian and the modular Hamiltonian of a von Neumann sub-algebra represent the same physical concept (provided that the considered state is faithful on the sub-algebra, but that is mostly a mathematical nuisance).

In a very powerful and almost unique way, Bisognano–Wichmann’s theorem provides an explicit form for the entanglement Hamiltonian on a Minkowski wedge, for the vacuum of any relativistic field theory.

Let us conclude by remarking that the thermalization of the vacuum in Bisognano–Wichmann is intimately related (essentially, a mathematical proof) to Unruh’s effect. In fact, the trajectory of the spacetime point $(0, a^{-1}, 0, 0)$ under the boosts $\Lambda(s)$ is that of a uniformly accelerated motion on the right wedge, with acceleration $a$. For an observer on this trajectory that uses his proper time $\tau = s$ as the time coordinate, the generator of time translations, in the coordinate system at rest with her/him, is none other than $H = aK$. Therefore, by Bisognano–Wichmann’s theorem, the vacuum state $\omega_\beta$ is thermalized for her/him, with temperature

$$T = \frac{a}{2\pi}$$

that, up to restoring the physical constants always omitted by mathematicians, is exactly the Hawking–Unruh temperature. In his book,\(^{[49]}\) Haag goes even further by arguing that the right wedge, populated by observers that are constrained to never leave it, provides the simplest example of an event’s horizon, and this is why Unruh’s temperature coincides with Hawking’s temperature. We merely report this interpretation as a nice little extra feature, to conclude this section, and hopefully to further invite the readers to the exploration of Tomita–Takesaki’s and Bisognano–Wichmann’s beautiful mathematical works.

### 4. Entanglement Hamiltonians of Integrable Models

In the previous section we took a detour in algebraic field theory in order to provide a strict mathematical background to the BW result and its CFT generalizations in Section 2. These are the most important examples where the EH of a system composed by continuous degrees of freedom is exactly tractable. From here on we shall rather turn our attention toward particular lattice models, whose low-energy behavior is known to be described by a relativistic field theory. Then the most important question is how the results derived for the EH in the QFT context generalize to these lattice systems. Indeed, the presence of the lattice breaks the Lorentz invariance of the theory, which is central to the BW theorem. Nevertheless, the continuum results turn out to provide, after a proper discretization, a very accurate description of the lattice EH. In this regard integrable systems play an important role, since they allow for explicit analytical results. These are available for two different subsystem geometries, namely a half-infinite chain corresponding to the BW setting, as well as a finite interval.

#### 4.1. Half Chain

The RDM of a half-chain $\rho \propto \exp(-H_{\text{half}})$ can be related to the corner transfer matrix (CTM) of a corresponding 2D statistical physics model.\(^{[21]}\) The CTM was introduced and studied for integrable models by Baxter,\(^{[22,66,67]}\) showing that they allow for an explicit analytical treatment in the thermodynamic limit. We first consider the ground state of the transverse Ising (TI) chain

$$\hat{H} = -\sum_{j=0}^{\infty} \left( \lambda \sigma_j^x \sigma_{j+1}^x + h \sigma_j^z \right)$$

where the half-chain RDM is related to the CTM of the 2D Ising model.\(^{[21]}\) The EH has different forms in the ordered ($h < \lambda$) and the disordered phase ($h > \lambda$) of the chain and reads\(^{[68,69]}\)

$$H_{\text{half}} = \begin{cases} -2I(k') \sum_{n=1}^{\infty} \left[ n \sigma_n^x \sigma_{n+1}^x + k \left( n - \frac{1}{2} \right) \sigma_n^z \right], & \text{ordered} \\ -2I(k') \sum_{n=1}^{\infty} \left[ k n \sigma_n^x \sigma_{n+1}^x + \left( n - \frac{1}{2} \right) \sigma_n^z \right], & \text{disordered} \end{cases}$$

where $I(k')$ is the complete elliptic integral of the first kind, and the elliptic parameters are defined as $k = \min(\lambda/h, h/\lambda)$ and $k' = \sqrt{1 - k^2}$. Hence $H_{\text{half}}$ has exactly the BW form, the energy density being multiplied with a linear term, albeit with a prefactor that depends explicitly on the ratio $h/\lambda$. The EH can also be diagonalized exactly and has an equidistant single-particle entanglement spectrum.\(^{[21]}\)

$$\epsilon_l = \begin{cases} 2l \epsilon & \text{ordered} \\ (2l + 1) \epsilon & \text{disordered} \end{cases}$$

with $l = 0, 1, \ldots$ and spacing $\epsilon$. Similar results are found for the anisotropic XY chain, where the RDM is related to the CTM of a triangular Ising model.\(^{[70]}\)
The CTM approach can also be applied to the interacting XXZ chain
\[ \hat{H} = \sum_{n=-\infty}^{\infty} \left( \sigma_n^x \sigma_{n+1}^x + \sigma_n^y \sigma_{n+1}^y + \Delta \sigma_n^z \sigma_{n+1}^z \right) \] (44)

in the gapped phase ($\Delta > 1$), where the half-chain RDM is related to the CTM of the six-vertex model, which was studied previously in refs. [22, 71, 72]. This has again a lattice BW form with
\[ \hat{H}_{\text{half}} = c(\Delta) \sum_{n=1}^{\infty} n \left( \sigma_n^x \sigma_{n+1}^x + \sigma_n^y \sigma_{n+1}^y + \Delta \sigma_n^z \sigma_{n+1}^z \right) \] (45)

where the constant $c(\Delta)$ depends on the anisotropy. The EH can again be diagonalized exactly, and the spectrum is similar to the TI case (43) in the ordered phase, with the spacing given by $\varepsilon = \text{arcosh} \Delta$. Note that the above results even generalize to the XYZ chain (related to the eight-vertex model) both at the level of the CTM as well as the corresponding entanglement spectrum.

The CTM method is, however, not restricted to spin chains but allows for the study of continuous variable systems as well, such as the harmonic oscillator chain
\[ \hat{H} = \frac{1}{2} \sum_{n=-\infty}^{\infty} \left[ p_n^2 + \omega^2 q_n^2 + K(q_{n+1} - q_n)^2 \right] \] (46)

Choosing $K = k$ and $\omega = 1 - k$, the RDM is related to the CTM of the 2D Gaussian model with an elliptic parameterization of the couplings. The EH of the half chain then reads
\[ \hat{H}_{\text{half}} = 2i(k') \sum_{n=1}^{\infty} \left[ \left( n - \frac{1}{2} \right) p_n^2 + \left( n - \frac{1}{2} \right)(1 - k)^2 q_n^2 \right. \\
+k n (q_{n+1} - q_n)^2 \left. \right] \] (47)

which has again a BW form, with single-particle spectrum as for the TI chain in (43) in the disordered phase. Note that, via a canonical transformation of the positions and momenta, one can also obtain the result for the parameterization $K = 1$ and arbitrary $\omega$. In this case the elliptic parameter $k$ is the solution of $\omega^2 = (1 - k)^2/k$ and the prefactor of the EH reads $2i(k') \sqrt{k}$.\[ \text{[77]} \]

4.2. Interval

The case of an interval $A = [1, N]$ is more difficult to handle, and a direct calculation of the EH is only possible for free lattice models. In particular, analytical results are available for an infinite hopping chain
\[ \hat{H} = -\sum_n t (c_n^e c_{n+1}^c + c_n^c c_{n+1}^e) + \sum_n d c_n^f c_n^c \] (48)

where $c_n^e$ and $c_n^c$ are fermion creation/annihilation operators. Setting $t = 1/2$ and $d = \cos q_f$, the Hamiltonian (48) is diagonalized by a Fourier transform and the ground state is a Fermi sea with occupied momenta $q \in [-q_f, q_f]$. Due to Wick’s theorem, the RDM is given by $\rho_A = Z^{-1} \exp(-\hat{H})$, where the EH is another free-fermion operator\[ \text{[78-80]} \]
\[ \hat{H} = \sum_{i,j=1}^{N} H_{ij} c_i^c c_j^e \] (49)

The matrix elements are given by
\[ H_{ij} = \sum_{k=1}^{N} \phi_k(i) \varepsilon_k \phi_k(j) \] (50)

where $\varepsilon_k$ and $\phi_k(i)$ are the eigenvalues and eigenvectors of the reduced correlation matrix $C_k$, with matrix elements $C_{ij} = \langle \phi_j^\dagger \phi_i \rangle$ restricted to $i, j \in A$.

The expression (50) allows for a calculation of the EH, which requires very high precision numerics, as the dominant $\varepsilon_k$ contributions originate from eigenvalues $\varepsilon_k$ that lie exponentially close to zero or one. The analytical treatment, however, follows a different route based on the existence of a commuting tridiagonal operator $T$, with matrix elements $T_{ij} = d_i$ and $T_{i,i+1} = T_{i+1,i} = t_i$ given by
\[ t_i = \frac{i}{N} \left( 1 - \frac{i}{N} \right), \quad d_i = -2 \cos q_f \frac{2i - 1}{2N} \left( 1 - \frac{2i - 1}{2N} \right) \] (51)

Thus $T$ describes an inhomogeneous hopping chain, with hopping amplitudes following the exact same parabolic profile as one would obtain from a proper discretization of the CFT result (9). This suggests the relation $H = -N \pi T$, which was also found for the low-lying $\varepsilon_k$ eigenvalues.\[ \text{[82]} \] However, compared to the numerical results, one finds some discrepancies as shown in Figure 3 for half filling ($q_f = \pi/2$). Indeed, one observes a slight deviation from the expected nearest-neighbor hopping profile, as well as nonvanishing hopping to more distant sites (note that only odd distances appear due to the particle-hole symmetry).

The difference can be understood by a proper comparison of the spectra of $h = -H/N$ and $T$, which was studied by Slepian

Figure 3. Hopping matrix elements in $-H/N$ and nearest-neighbor hopping in $\pi T$ (green) as functions of the position for $N = 40$ sites at half filling. The third and fifth neighbor hopping amplitudes are magnified by a factor of 20 for better visibility. Reproduced with permission.\[ \text{[83]} \] Copyright 2017, Institute of Physics.

The third and fifth neighbor hopping amplitudes are magnified by a factor of 20 for better visibility. Reproduced with permission.\[ \text{[83]} \] Copyright 2017, Institute of Physics.
in ref. [81]. It turns out that, in the limit $N \to \infty$, one can relate them via a series expansion

$$h = \sum_{m=0}^{\infty} a_m \beta_m T^{2m+1}$$

(52)

with appropriate constants $a_m$ and $\beta_m$. The leading order has $a_0 \beta_0 = \pi$ and thus gives the expected CFT behavior. However, higher powers of $T$ generate more distant hopping and also modify the nearest-neighbor one. Interestingly, the matrix elements of $h$ can be found in a closed form via generalized hypergeometric functions. In particular, in the limit $i, N \to \infty$ with $x = i/N$ fixed, one has ref. [83]

$$h_{i,i+1} = \pi x(1-x), F(\frac{1}{2}, \frac{3}{4}, \frac{3}{4}; 1, 1; [4x(1-x)]^2)$$

(53)

where the parabolic profile is multiplied by a function which increases smoothly from 1 at $x = 0$ to 1.076 at $x = 1/2$ and thus gives a deviation of roughly 8% in the middle. Similar formulae are found for $h_{i,i+1}$ with $r$ odd, showing a strong suppression of the distant hopping. Nevertheless, the maxima of the profiles at $x = 1/2$ decays only as a power law $r^{-1}$ [83]. Note that the analytical treatment can be extended for general fillings, where the relation (52) between $h$ and $T$ becomes more complicated, [83] and also hopping with even $r$ appear.

A similar approach can also be applied for the study of the EH in a finite chain. In fact, a commuting tridiagonal matrix $T$ exists for both periodic [86] as well as for open chains [85] describing a hopping profile that is again the discretized version of the CFT result in the corresponding geometries [128]. Analogously to the infinite chain case, however, the nearest-neighbor hopping in the EH is slightly larger around its maximum, and longer-range terms with much smaller amplitudes are also present. Numerical investigations suggest a relation similar to (52), where the coefficients depend only on the ratio of subsystem and full system lengths, but their analytical expression is not known explicitly [85].

Note that, based on the theory of bipartritical, commuting tridiagonal matrices have also been identified for particular inhomogeneous free-fermion Hamiltonians [86, 87], the corresponding relation (52) was, however, not yet studied.

Finally, we discuss the case of a non-critical chain, obtained by dimerizing the couplings $t_{2n-1} = (1-\delta)/2$ and $t_{2n} = (1+\delta)/2$ of the hopping chain in (48). Using the spin representation and introducing dual variables [88–91] such a Hamiltonian can be mapped into two interlacing TI chains where $h_i = t_{2n-1}$ and $\lambda_i = t_{2n}$, whereas the couplings in the second chain are interchanged. For a half chain, this would yield an EH of the form (42), where the two terms are now replaced by the even and odd hopping terms, with the elliptic parameter $k = (1-\delta)/(1+\delta)$ multiplying the odd ones for $\delta > 0$. One can now argue that, if the size of the interval is much larger then the correlation length ($N \gg \xi$), then the contributions from the two boundaries should decouple in the RDM. Thus the EH should effectively behave as a half-infinite chain from each end, with the profile given by

$$h_{2i-1,2i} \approx 2i(1/k) \Delta((2i-1)/N), \quad h_{2i,2i+1} \approx 2i(1/k) \Delta(2i/N)$$

(54)

Figure 4. Nearest-neighbor hopping in the EH of the dimerized chain for various $\delta$ and $N = 40$. The hopping across odd bonds (empty symbols) are divided by a factor of $k$. The dashed lines have slopes $2i(k')$ corresponding to the result (54). Reproduced with permission. [77] Copyright 2020, Institute of Physics.

with the triangular function

$$\Delta(x) = \begin{cases} x & 0 \leq x \leq 1/2 \\ 1-x & 1/2 \leq x \leq 1 \end{cases}$$

(55)

The numerical results are shown in Figure 4, and are well described by (54) in the vicinity of the interval edges, with some deviations around the center. The triangular approximation improves with decreasing correlation length $\delta \to 1$, however, toward the critical limit $\delta \to 0$ one observes a crossover to a roughly parabolic form.

4.3. Continuum Limit

The exact form of the EH for an interval in a critical free-fermion chain thus shows characteristic deviations from the continuum CFT results, which need to be properly understood. On the one hand, one could check whether the discretized CFT result (i.e., replacing $H \to N\pi T$ in the RDM) would produce an error in the expectation values of local observables that vanishes in the limit $N \to \infty$. Indeed, using this approximation, one observes that the fermionic correlation matrix $C_i$ can be reproduced to very high accuracy even for smaller values of $N$ (although its translational invariance is lost). Moreover, one can make even stronger statements by considering the trace distance between the two RDMs as a function of $N$, as shown in the next section.

On the other hand, it would be desirable to understand how the proper continuum limit of the lattice EH arises, when we introduce a lattice spacing $s$ and consider the limit $N \to \infty$ and $s \to 0$ with $\ell = N s$ fixed. For the simple homogeneous hopping chain (48), the standard procedure is to linearize the dispersion around the Fermi points $\pm q_{\ell/2}$, and introduce slowly varying fields that describe left- and right-moving fermions. Then one ends up with a massless Dirac Hamiltonian with a single parameter given by the Fermi velocity $v_f$. Let us now add long-range hopping over $2p + 1$ sites with amplitude $t_{2p+1}$, which enters the dispersion with a factor $2\cos(2p+1)q_{\ell/2}$ and thus modifies $v_f$. Finally, assume that the argument holds true for hopping amplitudes that
vary slowly in space $t_{2p+1}(x) = Nh_{-p,i,p+1}$ as in the EH, where $x = i s$ is a continuous coordinate. The space-dependent Fermi velocity then reads\[^{[92]}\]

$$v_F(x) = 2\pi \sum_p (-1)^p (2p + 1) h_{-i,p,i,p+1}$$

(56)

where the alternating factor is due to $q_F s = \pi/2$ at half filling. Inserting (52), the resulting double infinite sum (56) can be carried out exactly and delivers $v_F(x) = 2\pi x (\ell - x) / \ell$, which is identical to the weight function in the CFT result (9).

Hence, the function multiplying the energy density in the continuum treatment has now the interpretation of a local Fermi velocity (instead of a local inverse temperature) that follows from the lattice EH via (56). Remarkably, this relation can also be used in numerical calculations for chains of finite length or at finite temperatures\[^{[92]}\] (see Figure 5), as well as for disjoint segments\[^{[36]}\] perfectly reproducing the corresponding CFT results.\[^{[28,33]}\] Furthermore, the continuum limit of the EH for an interval in the harmonic chain (46) can be dealt with along similar lines of thought. This yields again the renormalized couplings as weighted sums over the diagonals of the lattice EH, perfectly reproducing the CFT result in the massless limit.\[^{[91]}\] In contrast, in the massive case one experiences convergence issues with the sums\[^{[77]}\] which might indicate that the EH becomes nonlocal even in the continuum limit, as suggested by a perturbative treatment in the CFT context.\[^{[36]}\]

5. Entanglement Hamiltonians on the Lattice Beyond Exactly Soluble Models

The insight from integrable lattice systems is remarkable: the functional form of BW is exactly recovered for massive theories on the half chain, albeit with a prefactor that depends on the mass term. For gapless theories on the interval, one shall handle with more care: however, there is typically a large overlap between the exact and BW guessed Hamiltonian.

These works have stimulated the study of generic lattice Bisognano–Wichmann (LBW) entanglement Hamiltonians, to understand whether the latter have predictive power. In this section, we will review the current status of this search. Instead of following a historical perspective (which shall start with Li and Haldane work on topological phases), we prefer to start with critical 1D theories, that bridge more naturally to the previous section, as well as with the CFT predictions in Section 2.3. We will then cover massive phases in 1D (including symmetry-protected topological phases), and dimensions larger than one. Before continuing, it is worth mentioning that, from the perspective of state characterization, entanglement Hamiltonians have also been discussed in the context of representation of tensor network states.\[^{[10]}\] This research line, which has been proved very insightful on its own, is quite distinct from the state characterization of generic Hamiltonian eigenstates that we discuss here, so we refer the reader to specific reviews in the field for a detailed discussion—see, in particular, Ref. [94].

5.1. Bisognano–Wichmann Theorem on the Lattice

As most field theory results, the Bisognano–Wichmann theorem can be adapted to the lattice via space discretization. This procedure is particularly delicate however, as one is dealing with what is a very complicated operator (the logarithm of the RDM, capturing arbitrary correlation functions), whose definition presents challenges close to the boundary. We review here a formulation of such discretization following Ref. [95], that is convenient for both 1D and 2D systems. For the sake of simplicity, we consider a square lattice, with a bipartition as shown in Figure 6. We are interested in characterizing the ground state wave function of a Hamiltonian that reads as

$$H = \Gamma \sum_{x,y,\delta = \pm 1} \left[ h_{(x,y),(x+s,y)} + h_{(x,y),(x,y+s)} \right] + \Theta \sum_{x,y} l_{(x,y)}$$

(57)
suggest the following dependence of the EH couplings:

where the first two terms represent nearest-neighbor contributions along the $x$ and $y$ directions, respectively, and the last term is on-site.

Discretizing at first order in the lattice spacing, the BW theorem leads to the following lattice BW EH:

\[
H_{ABW} = \beta_{EH} \sum_{x,y} \left( \Gamma_x h_{(x+y+1)} + \Gamma_y h_{(x+y)} + \Theta_{x,y} l_{(x,y)} \right)
\]

where the inhomogeneous couplings and on-site terms depend on the distance from the boundary separating subsystem $A$ and $B$ (see Figure 6) according to the geometry of the original system, and the inverse entanglement temperature $\beta_{EH}$ includes both constants and the characteristic velocity on the lattice (that corresponds to the speed of sound for critical theories).

5.1.1. Half-Plane Lattice BW EH

For the case of a system dynamics that is governed by a generic, relativistic field theory (either massive or not), the BW theorem suggest the following dependence of the EH couplings:

\[
\Gamma_x = x\Gamma
\]

\[
\Gamma_y = \left( x - \frac{1}{2} \right)\Gamma
\]

\[
\Theta_{(x,y)} = \left( x - \frac{1}{2} \right) \Theta
\]

(59)

Note that there is an important point to be emphasized, that is, what is the corresponding distance from the boundaries of the various terms (a feature also encountered in the case of the Ising half chain (42) in the previous section). The rule chosen here is as follows: for local terms, we took as distance from the boundary their shortest Euclidean distance. For instance, in the 1D Ising case for $h_n$, this corresponds to $n - 1/2$. For terms defined on bonds (such as kinetic energy and nearest-neighbor spin couplings), one instead utilizes as a distance from the boundary the shortest Euclidean distance from the center of the bond (see Figure 6). While strictly speaking applicable to the infinite half plane only, the formulas above are expected to work equally well for other kinds of partitions with a single interface and no angles—such as half partitions of infinite cylinders. Finally, we note that the same discretization is obtained as a formulation of the lattice Unruh effect. [96]

5.1.2. Finite Partition in a Finite Chain

The formulas above, while very generic, are strictly speaking applicable solely to the case of an infinite partition in an infinite system. For the case of a $(1+1)$-d conformal field theory, it is possible to adapt to the lattice the conformal extensions discussed in Section 2.3. In particular, for a periodic chain of length $L$, the half-partition case reads

\[
\Gamma_x = \frac{L}{2\pi} \sin \left( \frac{2\pi x}{L} \right) \Gamma
\]

\[
\Theta_x = \frac{L}{2\pi} \sin \left( \frac{2\pi x}{L} \left( x - \frac{1}{2} \right) \right) \Theta
\]

while for the open chain

\[
\Gamma_x = \frac{L}{\pi} \sin \left( \frac{\pi x}{L} \right) \Gamma
\]

\[
\Theta_x = \frac{L}{2\pi} \sin \left( \frac{\pi x}{L} \left( x - \frac{1}{2} \right) \right) \Theta
\]

(61)

To connect the above equations to the CFT expressions in Equations (10) and (11), the corresponding inverse entanglement temperature shall satisfy $\beta_{EH} = \frac{\pi}{\nu}$, where $\nu$ is the speed of sound of the lattice model. Note that $\nu$ depends on the model parameters and may differ from the maximal group velocity. For instance, in case of free fermions discussed in Section 4, $\nu$ is simply the Fermi velocity and thus depends on the filling. In principle, the CFT predictions can also be lattice regularized at finite temperature in the same manner. The main questions are then how well such lattice regularizations approximate the exact EH, which can be investigated numerically.

5.2. Numerical Investigations of Lattice Bisognano–Wichmann Entanglement Hamiltonians in 1D Systems

Differently from the direct evaluation of the lattice entanglement Hamiltonian available for certain integrable models, [96,97] for non-integrable systems, as well as for interacting systems under arbitrary boundary conditions, investigations of the lattice BW theorem as formulated in the previous paragraphs are typically carried out in an indirect manner. The reason is that a direct extraction of the terms in the EH is a computationally cumbersome operation, that, to date, can only be performed utilizing a specific ansatz for the couplings. At the exact level, this is typically feasible only on relatively small system sizes. Approximate numerical schemes have been devised, utilizing either parent-Hamiltonian type methods, [98] or quantum Monte

Figure 6. Graphical representation of the 2D lattice with partitions $A$ and $B$, and corresponding notations of the distance from the boundary in units of the lattice spacing: nearest-neighbor terms along the vertical ($\gamma$) direction, and local terms have distance $n - 1/2$, while nearest-neighbor terms along the horizontal ($\nu$) direction have distance $n$. Reproduced with permission. [93] Copyright 2018, American Physical Society.
Carlo sampling.\cite{99} We will comment on those at the end of the next subsection.

The indirect route to study EH of interacting systems relies on validating a posteriori the lattice BW EH. In fact, what one is really checking is whether, for the ground state of a given model Hamiltonian, the true EH and the lattice BW EH have the same properties up to a given threshold error. Various types of tests have been devised for this purpose. The most immediate choice is to directly compare their spectra and their eigenstates; more “coarse-grained” tests include entropies (in this context, construed as moments of the distribution of the entanglement spectrum) and expectation values of order parameters and correlation functions. In some cases, it is also possible to compare the operator distance(s) between the EH and the lattice BW EH: those are by far the strictest tests. In the following, we present the operator distance(s) between the lattice BW EH and the exact EH. Red lines are proportional to $e^{-2}$; the agreement with the fit is extremely good, so that most points are actually beneath the fit line. Reproduced under the terms of the Creative Commons CC-BY license.\cite{101}

$$\rho_{A,BW}$$

5.2.1. Ising Models: Finite-Volumes and Long-Range Interactions

We have seen in the previous section that the EH of the Ising model away from its critical point, for the half-chain partition in an infinite chain, is exactly given via the CTM. At the critical point and for finite partition sizes, an exact solution is not available.

In Ref.\cite{95,100,101}, a systematic comparison between the lattice BW EH and the real EH was performed, at the level of both eigenvalues, eigenvectors, and full RDM. In Figure 7, some of the properties of the RDM are considered, for partition sizes $\ell$ embedded in an infinite chain (left), or finite periodic chain (right) of length $L = 4\ell$. The left panel shows how different moments of the entanglement spectrum distribution (i.e., the traces of the RDM to a given power) become closer and closer as the partition size is increased. This signals that the entanglement spectrum of the lattice BW EH is approaching the same distribution of the real ES.

In the right panel, we show the scaling of several operator distances between the real EH, and the lattice BW one, from Ref.\cite{101}. In particular, $D_n$ are generalized Schatten distances

$$D_n = \frac{1}{2^{1/n}} \|\rho_{A,BW} - \rho_A\|_n$$

$D = D_1$ is the trace distance, and the triangles are related to the Uhlmann fidelity $F = \text{tr} \sqrt{\rho_{A,BW} \rho_A \sqrt{\rho_{A,BW}}}$. All distances scale systematically to zero in the thermodynamic limit. This signals the fact that, while at finite size there are discrepancies between the real and lattice BW EH, the corresponding RDMs converge to the same operator in the limit of large enough partitions. Similar results have been obtained for XY spin chains.

These types of accurate verification can only be performed for quadratic Hamiltonians: it is, for instance, not feasible for the (experimentally relevant) case of long-range Ising models. Still, the latter entanglement spectra have been shown to be accurately reproduced by the lattice BW EH, at least in cases where long-range interactions decay as dipolar or van der Waals cases.\cite{102}

5.2.2. Potts Models

For generic quantum systems, different ways of comparing $\rho_A$ and $\rho_{A,BW}$ have been introduced. One practical method for numerical simulations is to compare their eigendecompositions, both in terms of spectrum and vectors. As far as the former is concerned, the following ratios are typically employed\cite{102}

$$\kappa_{\nu} = \frac{\epsilon_\nu - \epsilon_0}{\epsilon_1 - \epsilon_0}$$

where $\epsilon_\nu$ are the entanglement eigenenergies in increasing order. Such ratios have the practical advantage of being insensitive to the definition of entanglement temperature, as well as to additive constants in $H_A$. In Figure 8, first two columns, we show such ratios corresponding to $\rho_A$ (black line) and $\rho_{A,BW}$ (red dots) for the case of the 1D three-state Potts model. For both OBC and PBC,
as well as at \((g = 1.0)\) or away from \((g = 1.4)\) the critical point, the lattice BW EH is able to accurately predict the full low-lying spectrum down to values smaller than \(10^{-6}\) even for the modest system sizes considered here (the partition \(A\) is of length \(L/2\)). Note that the several degeneracies present outside of the critical point (bottom row) are captured at the percent level.

Eigenvectors can also be compared one-by-one, at least for small system sizes where a full diagonalization of the lattice BW EH can be performed. In the right panels of Figure 8, we show such comparison for a system of size \(L = 14\), where the quantity plotted is

\[
M_{\alpha,\alpha'} = |\langle \psi_{\alpha} | \psi_{\alpha'}^{\text{BW}} \rangle| \tag{64}
\]

Deviations from zero outside of the diagonal are smaller than \(10^{-3}\), while degeneracies in the spectrum appear as finite terms just next to the diagonals (both for \(g = 1.0, 1.4\)). Overall, the agreement is excellent even at such small sizes.

### 5.2.3. Other Critical Chains

The analysis above has been performed for a broad variety of critical points and phases, including Heisenberg chains, spin-1 bilinear-biquadratic, XX and XY models, and Hubbard models. More recent works have also discussed different partitions, as well as ladder systems.\([101]\) Overall, the results support the picture that is also suggested by the exactly soluble models discussed in the previous sections: when not exact, the lattice BW EH often provides a very accurate description of the real EH, both in terms of its generic properties (e.g., locality and types of terms) and of its spectral properties (eigenvalues and eigenvectors) as long as the low-energy field theory is Lorentz invariant.

### 5.3. Entanglement Hamiltonian of Topological Matter and the Li–Haldane Conjecture

As mentioned in the Introduction, the structure of the EH for massive phases in condensed matter systems was motivated by the investigation of the connection between entanglement and topological phases. This connection was first formulated in Ref. [17] in the context of fractional quantum Hall states. In particular, it was conjectured that the low-lying entanglement spectrum of a connected partition features the same properties of the edge modes of the theory corresponding to the given state. This observation was backed up by numerical simulations, and has later been verified for a series of other situations in both 1D and 2D systems. For a review, see Ref. [7].

As far as the entanglement Hamiltonian is concerned, the Li–Haldane conjecture strongly suggests that the corresponding EH is dominated, at low energies, by the edge Hamiltonian. This result is very much consistent with what one would expect from the BW theorem: indeed, this argument was proposed in Ref. [104] as a proof of the connection between edge mode spectra, and entanglement spectra in half-infinite systems. In 1D systems, a further insight can be gathered by directly studying the perturbed CFT: in the case of a relevant perturbation, it has been shown that the corresponding EH is given in terms of the chiral Virasoro operator—that, again, is consistent with BW theorem\([105]\) (this fact can be interpreted similarly to the case of CTM in the context of integrable models). This fact has been confirmed on the lattice on several models, including the Haldane and the Su–Schrieffer–Heeger chain.\([95, 100, 105]\)

### 5.4. Entanglement Hamiltonian of 2D Phases and Critical Points

The Li–Haldane conjecture motivated the study of the structure of EH well beyond the 1D case—in particular, for 2D systems. In those settings, considerably less is known, since there are no available explicit expressions in general field theory for finite volumes.

Direct tests of the accuracy of the lattice BW results in any dimension larger than one are extremely challenging for interacting systems, as they often require the exact diagonalization of the EH. However, it is possible to perform indirect tests based on correlation functions, by performing quantum Monte Carlo
or, in principle, tensor network) simulations of the corresponding model. The first examples in this direction have been the 2D Heisenberg model and Heisenberg bilayer.\cite{95,106} In both cases, it was found that correlation functions obtained utilizing a “thermal” sampling of the lattice BW EH asymptotically converge to the value obtained by sampling the original Hamiltonian, substantiating the validity of the approach (even if a finite size scaling theory on how this works is presently lacking).

Based on these insights, the lattice BW EH was utilized to predict properties that are inaccessible otherwise. One example is the von Neumann entropy, that cannot be computed in conventional MC simulations, and whose computation for 2D models would require—for the case of matrix-product-state wave functions—prohibitive bond dimensions. This demonstrates how leveraging on the BW EH enables computational methods that allows to make predictions in otherwise inaccessible regimes.

### 5.5. Entanglement Hamiltonians Out of Equilibrium

So far, we have solely treated equilibrium systems. Unfortunately, extensions of the BW theorem for real time are not known. The only exception is CFT, where it is possible, for certain quench protocols, to identify the EH correspondent to a finite partition.\cite{28,29,108} Typically, such EH depend on the full stress-energy tensor (not only on its energy density component, $T_{00}$), so their direct application to lattice models requires more caution. Still, some results exist for the harmonic and free-fermion regimes.\cite{109} and we will see in the next section one working example in the context of quantum spin chains.

### 6. Entanglement Hamiltonians in Experiments

We have introduced in the previous sections closed form formulas for the entanglement Hamiltonian of lattice models. The fact that the entanglement Hamiltonian can be expressed in a form that is reminiscent of the original Hamiltonian allows us to devise methods to probe the entanglement Hamiltonian experimentally. Below we review recent theoretical and experimental works that showed how to make use of the concept of entanglement Hamiltonians in order to experimentally extract the entanglement properties of a many-body system.

We organize this section in three parts: In Section 6.1, we show how one can use the physical implementation of the entanglement Hamiltonian in a experiment as a scalable method to measure the entanglement spectrum. We then show in Section 6.2 how one can efficiently measure the entanglement Hamiltonian based on local ansatz that are inspired from the BW theorem. These two approaches are finally combined in Section 6.3. In addition, and in order to help the reader in comparing these methods, we provide in Table 1 a summary for each protocol of the different experimental requirements, and of quantities that can be measured.

---

**Figure 9.** Entanglement entropies of 2D lattice spin models obtained via metadynamics Monte Carlo sampling of the lattice Bisognano–Wichmann entanglement Hamiltonian.\cite{106} The left- and right-most panels show the scaling of bipartite entropy versus size in Heisenberg, XY, and Heisenberg-bilayer models. The central panel singles out logarithmic corrections, whose coefficient is proportional to the number of Goldstone modes in a phase that features spontaneous symmetry breaking of a continuous symmetry.\cite{107} Reproduced with permission.\cite{106} Copyright 2020, Institute of Physics.
Table 1. Summary of methods for probing entanglement Hamiltonians. Ref. [116] combines Ref. [102] and Ref. [117] by considering a variational learning procedure of the EH, where estimations of the EH that are iteratively updated are physically applied on the state \( \rho_A \).

| Method | States implemented | Types of measurement | Measured quantities |
|--------|--------------------|----------------------|--------------------|
| Quantum simulation of EHs [102] | The eigenstates of \( H \) | Spectroscopy | Entanglement spectrum |
| EH tomography [117] | \( |\psi\rangle \) with reduced states \( \rho_A \) | Tomography on \( \rho_A \) | Entanglement Hamiltonian |
| Variational estimation of the EH [116] | \( |\psi\rangle \), with \( \rho_A \) time evolved with estimations of \( H \) | Time-evolved observables | Entanglement Hamiltonian |

6.1. Implementation of Entanglement Hamiltonians

The first motivation behind the approach of quantum simulation of entanglement Hamiltonians presented in Ref. [102] is the measurement of entanglement spectra (ES). The question of the experimental access to the ES is a long-standing challenge for quantum simulation. Measuring the ES allows us, for instance, to understand the structure of symmetry-protected topological phases [110] or more generally to test the Li–Haldane conjecture [17,111]. The measurement of the ES can also be used to extract the entanglement entropies that quantify the presence of entanglement in correlated quantum systems [6].

In the standard scenario of quantum simulation [112], a quantum state of interest \( |\psi\rangle \) is realized based on the physical implementation of the Hamiltonian \( H \) of a lattice model. This implementation can be based on ultracold atoms, ions, or superconducting circuits realizing a quantum computer, etc. For instance, we can be interested in the entanglement properties of the ground state \( |\psi_0\rangle \) of \( H \), and we would like to probe the RDM \( \rho_A = \text{Tr}_{-A}(|\psi_0\rangle \langle \psi_0|) \) of the subsystem \( A \). Having access to the ES, the eigenvalues of \( \rho_A \), is notoriously difficult in quantum simulation or quantum computing. One either needs to measure the full density matrix \( \rho_A \) via quantum state tomography, which requires exponentially many measurements [113]. Alternatively, one can use interferometric methods that have the drawback of requiring a large number of multiple copies of the system [114]. We show for illustration in Figure 10a,b, a measurement of the ES performed in a quantum computer [115] and obtained via quantum state tomography. In Figure 10a, the quantum circuit to create the ground state of a symmetry-protected topological phase...
with \( L = 8 \) is represented. Quantum state tomography is then realized on a RDM of \( \ell = 4 \) qubits, which gives the ES shown in Figure 10b.

Beyond small qubit sizes, full state tomography is no longer an option due to the exponential cost in terms of measurements. In order to access easily ES for large systems, it was proposed in Ref. [102] to physically implement the entanglement Hamiltonian \( H \) in an experiment. Instead of preparing the ground state \( |\psi_0\rangle \) via the implementation of \( H \), as in the “traditional” scenario of quantum simulation, one prepares the eigenstates \( |\psi_\alpha\rangle \) of the entanglement Hamiltonian \( H \). As shown below, such quantum simulation of the entanglement Hamiltonian can be used in particular to extract, in a scalable way, the entanglement spectrum using, well established, spectroscopy techniques.

6.1.1. Implementing the Entanglement Hamiltonian

The idea behind Ref. [102] is that the BW theorem provides an experimentally friendly method to implement entanglement Hamiltonians. In particular, the lattice version of the BW theorem Equation (58) gives an expression of the entanglement Hamiltonian \( H \) in terms of particles operators (spins, bosons/fermions), which we can relate in the context of an experiment to the particles of the system (cold atoms, etc). Also, the locality of \( H \) copes well with the local character of the interactions that can be implemented between atoms, ions, etc. Finally, as these interactions are often mediated using tunable controls, such as external lasers, we can easily engineer inhomogeneous terms following the prescription of the BW theorem. This is illustrated in Figure 10c,d for Rydberg and trapped ions quantum technologies implementing the entanglement Hamiltonian of quantum Ising models. Using cold atoms placed in optical lattices, the EH can be engineered via laser-assisted tunneling,\cite{[102]} or other types of inhomogeneous optical forces.\cite{[118]}

6.1.2. Entanglement Hamiltonian Spectroscopy

Once the EH is implemented in an experiment, one can explore entanglement properties, in a scalable way, and using the standard tools of quantum simulation. In particular, in order to reveal the ES experimentally, one can first prepare the ground state \( |\psi_0\rangle \) with a weak perturbation generated by a time-dependent Hamiltonian \( \hat{h}(t) = \hbar \text{sin}(\nu t) \). When the frequency of the perturbation approaches an entanglement “transition,” \( \nu \approx \epsilon_\alpha - \epsilon_0 \), the corresponding eigenstate \( |\psi_\alpha\rangle \) is resonantly excited, and we observe a response of the system that can be detected by monitoring the dynamics of certain observables. In Figure 10e, this is illustrated for the case of the entanglement spectroscopy of the spin-1 Haldane phase.\cite{[116]} Here, the value of the first entanglement transition vanishes in the thermodynamic limit, corresponding to the presence of topological degeneracies in the entanglement spectrum, see Figure 10f for a subsystem of \( \ell = 40 \) sites. This means that, by monitoring the closure of the entanglement gap in such an experiment, we can reveal the topological nature of the Haldane phase. Utilizing similar ideas, it has been proposed in Ref. [106] that also moments of the ES distribution—such as the von Neumann entropy—can be measured.

6.2. Measurements of Entanglement Hamiltonians

As described above, under the assumptions of the BW theorem, we can reveal entanglement properties by physically implementing the EH \( H \). In certain situations however, for example, away from equilibrium, the BW theorem does not strictly apply. Therefore we may wonder whether an experiment is able to measure the entanglement Hamiltonian, and in particular check the validity of the BW theorem. Here, in contrast to Section 6.1, we have in mind again the “traditional” scenario of quantum simulation where an arbitrary state \( |\psi\rangle \) with RDMs \( \rho_\alpha \) is physically realized, cf., Table. 1. The goal is to measure the corresponding entanglement Hamiltonian.

As the EH \( H \) is the matrix log of the density matrix \( \rho_\alpha \), the measurement of \( H \) is formally equivalent to quantum state tomography, and thus require in principle exponentially many measurements.\cite{[113]} In Ref. [117], it was however shown that one can use the concept of entanglement Hamiltonians to perform quantum state tomography more efficiently.

The entanglement Hamiltonian tomography (EHT) protocol is shown in Figure 11a. The system is subject to various measurements, which provide an estimation of the density matrix \( \rho_\alpha \). This estimation is typically inaccurate because it was made from too few measurements compared to the requirements of tomography. However, this “poor” density matrix estimation is then fitted to an ansatz \( \rho_\alpha \propto e^{-\lambda H} \) that is based on a local EH \( H \), that is, inspired from the result of the BW theorem. As this ansatz is made of a polynomial number of fitting terms in system size, very few measurements are indeed necessary to obtain a faithful reconstruction of the density matrix.

Interestingly, the process of using experimental measurements to reconstruct the entanglement Hamiltonian can be realized in the framework of randomized measurements that provide estimations of density matrices\cite{[120],[121]} (among other quantities, see below). In Ref. [117], this was used to measure entanglement Hamiltonians and spectra, based on existing randomized measurements experimental data from Ref. [119]. In Ref. [122], it was also proposed to use the method Ref. [117] (as well as the one Ref. [116] presented in the next section) to experimentally study the Li–Haldane conjecture. The corresponding experimental reconstructing of the ES is shown in Figure 11b. Here the system under study was a trapped ion spin chain that was evolved via the long-range XY Hamiltonian.

Remarkably, the efficiency of EHT only relies on the fact that the EH is local. In this case, the density matrix can be interpreted as a Gibbs state of a local Hamiltonian (the EH), which, as shown rigorously in Ref [123], can be indeed “learned” from few measurements. Importantly, the method presented in Ref. [117] can be used to prove the locality of the EH, and in particular for ground states to verify the lattice version of the BW theorem. Using a randomized measurement protocol for fidelity measurements ref. [124] as subroutine, one can check that the EH is indeed faithfully estimated. If necessary, additional terms can also be included to “enrich” the EH and to provide a better fit to the experimental data.
Figure 11. Measurement of entanglement Hamiltonians. a) Protocol for EHT ref. [117] based on fitting estimations of the density matrix with an ansatz based on a local EH. b) Experimental reconstruction from EHT of the ES from the randomized measurements data of Ref. [119]. a) Reproduced with permission. Copyright 2021, Springer. b) Reproduced with permission. Copyright 2021, American Physical Society.

Figure 12. Measurement of entanglement Hamiltonians. a) Quantum variational learning of the entanglement Hamiltonian. This is based on a minimization procedure using as input a measurement of certain observables, and as outputs the variables that parameterize the EH. b) Numerical illustration of the protocol, where the local terms of the EH are reconstructed at the end of the minimization procedure. Figure published in Ref. [116].

6.3. Combined Implementations and Measurements of Entanglement Hamiltonians

The protocol[102] physically implements the EH $\mathcal{H}$ according to the prescription of the BW theorem. Instead, the protocol Ref. [117] measures the EH via a tomography of the density matrix $\rho_A$, and can in particular signal deviations from the BW theorem. Recently, the advantages of these two approaches have been combined in a new protocol presented in Ref. [116]. In this approach, one iteratively measures the EH, and implements it on the quantum device in order to perform the entanglement spectroscopy. As shown in Figure 12a, the idea is to run an iterative algorithm, where at each step $n$: i) we first realize a quantum state $|\psi\rangle$, ii) we apply on a subsystem $\rho_A$ time evolution with an estimation $\mathcal{H}^n = \sum_j \mathcal{H}_j[n]$ of the EH, which is parameterized with a polynomial number of local terms $\{\mathcal{H}_j[n]\}$. Note that such parameterization also occurs in the context of Hamiltonian learning.[123–128] iii) we measure the response of the system, and assess how the estimation $\mathcal{H}^n$ of the unknown EH can be improved, see below. The sequence (i–ii–iii) is then repeated, until the algorithm converges. At this point, we obtain a faithful estimation of the EH, which can be then used for entanglement spectroscopy.

The procedure to update the estimation of the EH from a measurement, which is the crucial step (iii) of the protocol, builds on the following observation: If the initial state $\rho_A$ has been time evolved from $\mathcal{H}$, the final state must remain unchanged, that is, $\rho_A(t) = e^{-i\mathcal{H}t}\rho_A e^{i\mathcal{H}t} = \rho_A$, because $\rho_A$ and $\mathcal{H}$ commute. Therefore, we can translate the problem of finding $\mathcal{H}$ into a minimization problem. The cost function to be minimized is defined as $C_{\mathcal{H}}(0) = \langle O(0) \rangle - \langle O(0) \rangle$, where $\langle O(0) \rangle = \text{Tr}(O e^{-i\mathcal{H}t}\rho_A e^{i\mathcal{H}t})$ and $O$ is a well chosen observable. The variables to be adjusted are the local terms $\{\mathcal{H}_j\}$. Such minimization procedure can be executed on a classical computer, using as input the experimental measurements of the cost function $C_{\mathcal{H}}(\mathcal{H}_j(0))$, and as outputs the variables $\{\mathcal{H}_j[n+1]\}$ that parameterize the new EH candidate to
be tested on the experiment. A numerical example of reconstruction of local terms parameterizing the EH is shown in Figure 12b.

6.4. Summary of the Experimental Section and Comparisons with Other Methods

We have presented experimental protocols that take advantage of the concept of entanglement Hamiltonians to probe entanglement properties of many-body systems. These probing methods are scalable in the sense that one requires a number of measurements that scales polynomially in system size. This is much less than the number of measurements required for state tomography, which typically scale as \( r^2D \) for density matrices \( \rho_A \) of rank \( r \) and dimension \( D \).[113] As a direct consequence, while the ES was previously measured based on tomography[115] on moderate system sizes of 4 qubits, the protocols presented above show how to measure the ES in much larger systems.

Having access to the ES and EH also give access to any entanglement property, such as entanglement entropies. Compared to alternative methods based on implementing multiple copies to measure Rényi entropies,[129–132] negativities[133] (and also the ES[114]), the protocols presented above involve single instances of the quantum states and appear thus as more experimentally friendly. Finally, protocols based on measuring statistical correlations between randomized measurements have been also devised to measure entanglement entropies,[119,124–128] and entanglement negativities.[119,129] These protocols have the advantage of not making assumptions on the state, and of not requiring multiple copies. However, they require, in experiments with \( N \) qubits, an exponential number of measurements \( 2^{N^2} \), with a modest exponent \( \alpha \leq 1 \) compared to tomography. As described above, randomized measurement protocols can also be combined with EH tomography.[117]

7. Conclusions and Outlook

The characterization of quantum correlations in quantum many-body systems is of paramount importance to deepen our understanding of physical phenomena, with direct applications to quantum technologies. In this review, we have summarized how the entanglement Hamiltonian constitutes an extremely powerful tool to carry out such characterization. Our presentation has followed the three parallel lines that have characterized its study: quantum field theory, integrable systems, and topological matter. Those three lines—that have had very little overlaps for almost 30 years—have been mutually intersecting over the last decade, providing deep insights on the structure of equilibrium state of matter. We have tried to emphasize as much as possible such intersections, and in particular, the pivotal role played by exact results in both axiomatic quantum field theory and integrable systems along these developments.

The overarching message that this review summarizes is that, for most physical states of interest to many-body theory, the entanglement structure of bipartitions simplifies dramatically, and in fact captured by local (inhomogeneous) operators, featuring only few-body terms, similarly to conventional Hamiltonian dynamics. Such a structure gives direct access to both a physical interpretation of entanglement (via, e.g., Unruh effect), and paves the way to powerful applications in quantum information processing, including methods to measure entanglement spectra and even perform quantum state tomography based on the simplified structure of the entanglement Hamiltonian—methods that have already been demonstrated on experimental data.

There are a number of open questions in the field. A first set of questions is related to the structure of the entanglement Hamiltonian in different setups from the ones described here. One important example is, what is the structure of entanglement Hamiltonians out of equilibrium: here, the present understanding is limited to specific quench protocols in conformal field theory, and a few numerical examples. Deeper insights on the entanglement structure after quench may help shedding light on the origin of the so-called entanglement barrier, and on the evolution of operator entanglement along thermalization, just to name two applications.

Another field where little is known is the case of disconnected partitions: while it is known that entropies of such partitions are more informative than the ones of connected partitions for specific cases,[141–143] very little is known about the corresponding EH, except for free field theories (see Section 2.5). Beyond numerical studies, it would be very intriguing to see whether it is possible to at least partly address such questions in the context of axiomatic quantum field theory: exact results such as the Bisognano–Wichmann theorem have proven pivotal so far, and one expects similar impact on the cases above. This could also help shed light on mixed state entanglement in those settings.[144] Another set of questions concerns applications that leverage on the known (or assumed) structure of entanglement Hamiltonians to empower tools in quantum information processing. While it is challenging to speculate on future applications, the successful example of tomographic reconstruction methods (one, if not the most, challenging characterization of quantum states) is definitely a strong basis to build future applications on.

Until recently, the study of entanglement Hamiltonians had followed separate paths in different branches of mathematics and (theoretical) physics. Now that links between these fields have been established, we expect an even faster and more efficient crossfertilization to take place, aimed at characterizing quantum correlations in many-body systems from an operational viewpoint.

Acknowledgements

The authors thank P. Calabrese, A. Elben, R. Fazio, G. Giudici, C. Kokail, T. Mendes-Santos, S. Murciano, I. Peschel, M. A. Rajabpour, B. Suard, E. Tonni, R. Van Bijnen, V. Vitale, T. Zache, J. Zhang, and P. Zoller for discussions and collaborations over the years on topics related to the present review. V.E acknowledges funding from the Austrian Science Fund (FWF) through Project No. P35434-N. B.V. acknowledges funding from the Austrian Science Fund (FWF, P 32597 N), and from the French National Research Agency (ANR-20-CE47-0005, JCJC project QRand). M.D. was partly supported by the ERC under grant number 758329 (AGEntH), by the MIUR Programme FARE (MEPH), and by European Union’s Horizon 2020 research and innovation programme under grant agreement No. 817482 (Pasquans).

Conflict of Interest

The authors declare no conflict of interest.
