Integrals of motion for one-dimensional Anderson localized systems

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Abstract

Anderson localization is known to be inevitable in one-dimension for generic disordered models. Since localization leads to Poissonian energy level statistics, we ask if localized systems possess ‘additional’ integrals of motion as well, so as to enhance the analogy with quantum integrable systems. We answer this in the affirmative in the present work. We construct a set of nontrivial integrals of motion for Anderson localized models, in terms of the original creation and annihilation operators. These are found as a power series in the hopping parameter. The recently found Type-1 Hamiltonians, which are known to be quantum integrable in a precise sense, motivate our construction. We note that these models can be viewed as disordered electron models with finite-range hopping, where a similar series truncates at the linear order. We show that despite the infinite range hopping, all states but one are localized. We also study the conservation laws for the disorder free Aubry–Andre model, where the states are either localized or extended, depending on the strength of a coupling constant. We formulate a specific procedure for averaging over disorder, in order to examine the convergence of the power series. Using this procedure in the Aubry–Andre model, we show that integrals of motion given by our construction are well-defined in localized phase, but not so in the extended phase. Finally, we also obtain the integrals of motion for a model with interactions to lowest order in the interaction.

1. Introduction

The simplest theoretical model to study localization for non-interacting particles in the presence of disorder was proposed by Anderson [1]. A single particle localized state has a wavefunction that decays exponentially about some point in space over a characteristic localization length. In three-dimensions, localized states exist below a certain energy (the mobility edge) for a given strength of disorder. A disordered electronic system is thus localized if its Fermi energy lies below the mobility edge. In one- and two-dimensions, an infinitesimal amount of disorder is sufficient to localize all single particle states and thus a disordered non-interacting electronic system is always localized [2, 3].

Recent developments in the area of eigenstate thermalization [4–6] relate closely to the above well established notions of Anderson localization. In this context, it is believed that an isolated localized eigenstate does not thermalize, in the sense that no subsystem of it can be brought into thermal equilibrium by exchanging heat with the rest of the system. An analogous statement can be made about information, as defined through an appropriate partial trace of the density matrix. A related feature of such a system is the lack of level repulsion in its energy level spectrum. This can be thought of as arising from the presence of almost degenerate states localized so far apart that they are unable to hybridize to lift the degeneracy.

The effect of interactions on such systems is very interesting. Interactions among the elementary degrees of freedom generically tend to drive the system towards thermalization and delocalization [7, 8]. This tendency competes with the one that causes localization in the presence of disorder. Understanding the resultant
phenomenon of many body localization, that is observed for sufficiently strong disorder, is currently a very active area of research [9–13].

Another class of systems that fail to thermalize are integrable ones. These often contain a variable parameter (such as interaction or external field strength, which we denote here as y) and possess a set of similarly dynamical (i.e. depending on the parameter) integrals of motion. Standard examples of such systems are the one-dimensional Hubbard and XXZ models. In these examples, the integrals of motion \( I_k \) are polynomial in \( y \) with the order of the polynomial [14–21] increasing with \( k \). An arbitrary linear superposition of all integrals

\[
Q = \sum_k a_k I_k
\]

—is an infinite power series in \( y \). Gaudin magnets [22, 23] on the other hand provide examples of integrable models where all conserved charges\(^6\) are linear in \( y \). It should be emphasized that there is no generally accepted precise notion of integrability in quantum mechanics [24, 25] in contrast to classical mechanics where it is unambiguous. However, we do not dwell on this issue in present work\(^7\).

The only aspect important for us here is the existence of parameter-dependent conservation laws.

Conserved charges greatly constrain the dynamics of integrable systems. As a result, when started off from an arbitrary initial state in isolation, these systems do not evolve in a way that causes thermalization in the sense of the above paragraph [6, 31]. Additionally the usual space–time symmetries result in degeneracies in the energy level spectrum, and hence a lack of level repulsion [32]. The addition of perturbations destroys such conservation laws and restores level repulsion, although the strength of the perturbations has a non-trivial finite-size dependence [33–35].

In this context, it is natural to ask in what ways are localized systems similar to integrable ones. In particular we may ask if (parameter dependent) conservation laws, similar to those in integrable systems exist for localized systems. It has been argued in the context of many-body localization that they do, and results related to the growth of entanglement in these systems are predicated on their existence [36–38]. However, obtaining the structure of the conserved charges directly in terms of microscopic parameters remains a challenge and effective renormalization procedures need to be employed instead [10, 39]. The situation is less complicated in the absence of interactions since the Hamiltonian is that of a single particle. Nevertheless, obtaining the charges systematically and analytically in terms of the microscopic parameters of the Hamiltonian is non-trivial. In this paper we outline the procedure to do so. We also elucidate the connection between localization and conserved charges.

In this work we study a general one-dimensional model with on-site disorder that can interpolate between models with long-range hopping and the more standard Anderson-type one. The starting point is a Type-1 Hamiltonian reviewed in [25–30]. This was introduced as the most simple model of quantum integrability in finite dimensional spaces. This model has infinite ranged hopping, and as such has no inbuilt metric or length scale. We first show by calculating its participation ratio (PR) [40, 41] the perhaps surprising result that all states except one are localized. This is done as follows: an eigenstate \( |\psi\rangle \) of the Hamiltonian is expanded in a basis of position eigenstates on the lattice as

\[
|\psi\rangle = \sum_k c_k |k\rangle,
\]

where \( k \) labels the position eigenstates and \( c_k \) are the coefficients in the expansion. The PR for this state is then defined as

\[
\text{PR}_\psi = \frac{\left( \sum_k |c_k|^2 \right)^2}{\sum_k |c_k|^4}.
\]

It is usually understood that \( \text{PR}_\psi \sim O(1) \) indicates localization while \( \text{PR}_\psi \sim O(N) \)–delocalization, where \( N \) is the number of sites. While this definition is valid for a fixed wave function, we may also define the PR at a given energy, as later in the paper, where an averaging over disorder realizations is carried out, at a fixed energy.

The Type-1 model has a known set of conservation laws, which inspire the construction of a generic Anderson-type model having only nearest-neighbor hopping. In 1d it is well known that for this model, all single particle eigenstates are localized for any strength of the disorder. The conserved charges of this model are then systematically and analytically in terms of the microscopic parameters of the Hamiltonian is non-trivial. In this paper we outline the procedure to do so. We also elucidate the connection between localization and conserved charges.

We then turn our focus to a model which contains both localized and delocalized phases (i.e. phases in which all single particles states are either localized or delocalized). This is the Aubry–Andre model [42], in which the random potential is replaced by a quasi-periodic one. This allows us to test our criterion for the convergence of the power series and clearly elucidate the connection of the conserved charges to localization. Thus, the

\[^6\] We use the term conserved charges interchangeably with conservation laws or integrals of motion.

\[^7\] By integrable we will generally mean quantum many-body models colloquially recognized as such, see examples in this paragraph. The only exception are Type-1 Hamiltonians that stem from a recently proposed well-defined notion of quantum integrability [25–30].
convergence (divergence) of the power series representation of conserved charges can indeed be identified with
the presence (or absence) of localization and the localization–delocalization transition can be located using the
charges. Finally, we investigate the effect of interactions and argue that a power series in the interaction becomes
intractable and thus obtain the the conserved charges only to first order in it.

We emphasize that the main feature of our construction is that the conservation laws do not depend
explicitly on the wavefunctions of the single particle energy eigenstates. In fact, the recursion relations we obtain
for the coefficients of the expansions of the conserved charges are the same for all generic one-dimensional
models. Our approach is thus completely model independent requiring no knowledge of exact solutions or
properties of energy eigenfunctions.

Another important aspect of the construction of conservation laws we emphasize here, which has not been
discussed before is ‘gauge freedom’ of a certain kind, defined more precisely later. We show that a judicious
choice of gauge can bring out important features of the conserved charges, such as the truncation of their series
representation at finite order. These features can be obscured in gauges that arise in constructions of these
charges from direct applications of standard methods such as the Rayleigh–Schrödinger series or the locator
expansion.

2. Lattice models

We consider a general Hamiltonian of non-interacting particles hopping on a one-dimensional lattice with an
on-site potential

\[ H = H(y) = \sum_i \epsilon_i n_i - y \sum_{j} t_{ij} c_j^\dagger c_j, \]

where \(c_j^\dagger\) and \(c_j\) are fermionic creation and annihilation operators, \(n_i = c_j^\dagger c_j\) is the number operator, \(\epsilon_i\) is the
on-site disordered potential, and \(t_{ij}\) is the hopping between sites \(i\) and \(j\). The parameter \(y\) is a real number introduced
for convenience, which; it allows us to perform an expansion of the conserved charges in its powers.

Our general strategy to construct conserved charges for this models will be to first consider the
‘unperturbed’ Hamiltonian which only has the on-site potential. The conserved charges for this Hamiltonian are
simply the operators \(n_i\), which are independent and commute with each other and the Hamiltonian. It can also be readily seen that the eigenstates of this Hamiltonian are completely localized on the individual sites. Thus the zeroth order Hamiltonian trivially describes a localized system with conserved charges. We now show that upon introducing the hopping, new conserved charges \(Q_i\) appear, which can still be labeled by the site indices \(i\) while the system remains localized. To do this, we consider different types of hopping parameters \(t_{ij}\).

3. Type-1 Hamiltonians

We now summarize a known set of conserved charges \(Q_i\). We rework the construction in [25, 27, 28], in a fashion
that suggests a natural generalization for short ranged models. These charges are linear in the hopping (or the
parameter \(y\)), and commute exactly with the Hamiltonian of the Type-1 family. The Type-1 Hamiltonian is
obtained from equation (2) by specializing to infinite ranged hopping \(t_{ij} = \gamma_i \gamma_j\), with arbitrary parameters \(\gamma_j\).
Specializing to \(y = 0\) we write down the charge \(Q_0\)

\[ Q_0 = n_0 - y \sum_{k \neq 0} \frac{1}{\epsilon_0 - \epsilon_k} [t_{0k} (c_k^\dagger c_k + c_k c_k^\dagger) - \alpha_0^0 n_0 - \beta_0^0 n_k], \]

where \(\alpha_0^0\) and \(\beta_0^0\) are yet to be determined. The commutator of \(Q_0\) and \(H\) vanishes to linear order in \(y\) by
construction. The surviving term is of \(O(y^2)\) and is given by

\[ [Q_0, H] = y^2 \sum_{j \neq k} \frac{1}{\epsilon_0 - \epsilon_k} [A(0, j, k)(c_j^\dagger c_j - c_j c_j^\dagger) + B(0, j, k)(c_j^\dagger c_j - c_j c_j^\dagger)]= 0, \]

where

\[ A(0, j, k) = t_{0k} t_{kj} - \alpha_0^0 t_{0j}, \]

\[ B(0, j, k) = t_{0j} t_{kj} - \beta_0^0 t_{0k}. \]

A few words on the form of equation (3) are appropriate here. The last two terms \(-\alpha_0^0 n_k - \beta_0^0 n_0\) commute with
\(H\) \((y = 0)\) trivially, since they are expressed in terms of the number operators. These actually represent a
particularly convenient ‘gauge choice’, their presence enables the second order term \(O(y^2)\) to vanish, and thus
the commutator series to truncate exactly for the Type-1 matrices. The requirement that \([Q_0, H] = 0\) is satisfied
by the following form of \(t_{ij}\)
This gives $A(0, j, k) = B(0, j, k) = 0$. It is straightforward to extend this definition to arbitrary $Q_j$, and further to show that $Q_j$ and $Q_j'$ are indeed the conserved charges of the Hamiltonian $H$ [26, 27]. The Hamiltonians described by $t_{ij}$ of the form given in equation (6) are called Type 1 [25, 27], and can also be interpreted geometrically as representing a ‘d-simplex’ [43].

### 3.1. PR for Type-1 Hamiltonians

All single particle states of Type-1 Hamiltonians (6), except possibly the ground state for $y > 0$ or the highest energy state for $y < 0$ are localized, see e.g. figure 1.

This can be understood in more detail from the exact solution for the spectrum of these models [27]. Exact un-normalized single particle eigenstates of the Hamiltonian (6) are

$$|E\rangle = \sum_{i=0}^{N-1} \frac{\gamma_i c_i^+}{E - \epsilon_i} |0\rangle$$

and the corresponding eigenvalues $E$ (energies) are solutions of the equation

$$\sum_{i=0}^{N-1} \frac{\gamma_i^2}{E - \epsilon_i} = -\frac{1}{y}.$$  

Suppose $\epsilon_i$ are ordered in the ascending order. By plotting the left hand side of equation (8) as a function of $E$, one can verify that it has $N - 1$ real roots $E_1, E_2, \ldots E_{N-1}$ located between consecutive $\epsilon_i$, i.e. $\epsilon_{i-1} < E_i < \epsilon_i$. The remaining root $E_0$ is also real and is below $\epsilon_0$ (ground state) for $y > 0$ and above $\epsilon_{N-1}$ for $y < 0$ (highest excited state).

Equations (7) and (8) also provide an exact solution for one fermion (Cooper) pair and one spin flip sectors of the BCS and Gaudin models, respectively

$$H_{BCS} = \sum_{i, \sigma = \uparrow, \downarrow} \epsilon_{i\sigma} c_{i\sigma}^+ c_{i\sigma} - y \sum_{i} c_{i\uparrow}^+ c_{i\uparrow} |s_i \rangle \langle s_i|,$$

$$H_i(y) = s_i^2 - y \sum_{j} \frac{s_i \cdot s_j}{2},$$

where $c_{i\sigma}$ are spin-full fermions and $s_i$ are quantum spins of arbitrary magnitudes $s_i$, see [27] for details. For the BCS (Gaudin) model one needs to replace $c_{i\sigma}^+ \rightarrow c_{i\uparrow}^+ c_{i\downarrow}^+ |s_i \rangle$ in equation (7), set $\gamma_i = 1/\sqrt{2s_i}$ and the corresponding eigenvalue is equal to $2E [2s_i (E - \epsilon_i)^{-1}]$ rather than $E$. Our results for the PR of Type-1 Hamiltonians therefore also apply to these sectors of these models.

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**Figure 1.** PR of eigenstates of a Type-1 Hamiltonian for $y = .004$, $N = 10^3$ in ascending order according to the energy. Each $\epsilon_i$ and $\gamma_i$ is an independent random number uniformly distributed in an interval $(-1, 1)$. Larger circle near the left top corner indicates the ground state, which is extended. Left inset is the same as above, but averaged over $10^3$ realizations of disorder and compared to equation (15) for the same $y$, $N$, $w$. The right inset shows the PR (except the ground state) for $N = 10^3$ equally spaced $\epsilon_i$, $\gamma = 1$ and $y = .004$ similarly compared to equation (15) (the two curves are indistinguishable).
The PR defined through equation (14) reads

$$\text{PR}_E = \frac{\left[ \sum_i \gamma_i^2 \right]^2}{\sum_i \gamma_i^2}, \quad (10)$$

For concreteness we take $y > 0$. Then, the ground state is $E_0 < \epsilon_0$. We assume that most $\gamma_i$ are of the same order of magnitude and consequently the vector with components $\gamma_i$ is delocalized. Further, we take $\epsilon_i$ to lie in a fixed interval that does not scale with $N$, e.g. from $-w$ to $w$.

For excited states $E_k$ is between $\epsilon_{k-1}$ and $\epsilon_k$. The summations in the numerator and denominator of equation (10) both come from $\epsilon_i$ in a small vicinity of $\epsilon_k$ for large $N$ and converge as $\sum_n n^{-2}$ and $\sum_n n^{-4}$, respectively, where $n = |i - k|$. The numerator and the denominator scale as $[\gamma_i^2 / \delta^2]^2$ and $\gamma_i^2 / \delta^4$, where $\delta \propto 1/N$ is the mean level spacing between $\epsilon_i$ in the vicinity of $\epsilon_k$. Therefore, $\text{PR}_E$ is of order 1 (much smaller than $N$) meaning excited states are always localized. Figure 1 shows PR for $N = 10^3$ uncorrelated random $\epsilon_i$ uniformly drawn from an interval $(-1, 1)$ and the same distribution of $\gamma_i$.

Consistent with our numerical results, we estimate the largest PR for excited states to scale as $\ln N$, i.e.

$$\text{PR}_E^\text{max} \approx \alpha \ln N, \quad (11)$$

for large $N$, where $\alpha$ depends on $N$ much weaker than $\ln N$. Such values of PR come from clustering in $\epsilon_i$.

Indeed, suppose spacings $\delta_i = \epsilon_{i+1} - \epsilon_i$ between $m$ of $\epsilon_i$ for $i$ from $k$ to $k + m$ are all much smaller than $\delta_{k-1}$ and, moreover, $\epsilon_{k+m} = \epsilon_k \leq \delta_{k-1}$. It follows from equation (10) that $\text{PR}_E \approx \text{PR}_{E_{k+m}} \approx m$ because the above $\epsilon_i$ contribute most to these PRs. Normalized spacings $s_i = \delta_i / \delta$ are distributed according to the Poisson distribution $P(s) \propto e^{-\delta} s^{\delta}$. The probability of having $m$ spacings between 0 and $s_0 < 1$ is then roughly $s_0^m$. We need $m s_0 \ll 1$ and also $N s_0^m = 1$ so that at least one such clustering occurs. This implies $m \approx \ln N / \ln(\ln N)$ and equation (11) follows. Numerically we find that typical values of $\alpha \approx 1 - 3$ and averaged over disorder $\bar{\alpha} \approx 1.7$, at least for $N = 2^3 - 2^{12}$. Note that according to this argument such large values of PR typically come in pairs spaced by $m + 1$, roughly equal to the value of the PR itself. We also stress that, in contrast to the largest PR, a typical (and average) PR is something between one and three for any $N$ (does not scale) as can be seen from figure 1.

It is interesting to compare this $\ln N$ behavior to the flat band localization studied earlier [44, 45]. The latter leads to a (weakly) divergent PR in the localized regime, a phenomenon that is viewed as corresponding to critical (power law type) localization. The Type-1 Hamiltonian kinetic energy may also be viewed as a ‘flat band’ model, with a flat dispersion for all except one state. Indeed, for $t_{ij} = \gamma_j \delta_{ij}$ all but one eigenvalues of the second term in equation (2) are zero. The non-zero eigenvalue (ground state for $y > 0$) corresponds to the eigenstate $\gamma_i \epsilon_i [0]$.

Let us consider limits $y \to 0$ and $y \to \infty$ separately. When $y \to 0$ all states are localized as expected. Indeed, equation (8) implies $E_k \to \epsilon_k$, summations in equation (10) are dominated by the $i = k$ term and we obtain $\text{PR}_E \approx 1$ for all $k$. When $y \to \infty$ excited states are localized as before because $E_k$ for $k \geq 1$ remains trapped in the interval $(\epsilon_{k-1}, \epsilon_k)$. The ground state energy on the other hand diverges—equation (8) implies $E_0 \to -y \sum_i \gamma_i^2$. Then, $\epsilon_i$ are negligible as compared to $E_0$ in equation (10) and

$$\text{PR}_E = \frac{\left[ \sum_i \gamma_i^2 \right]^2}{\sum_i \gamma_i^2}, \quad (12)$$

which is of order $N$ according to our choice of $\gamma_i$. The ground state is therefore delocalized for $y \to \infty$. It undergoes a localization–delocalization crossover at a certain $\chi$, which we estimate below in this section.

It is possible to evaluate the PR analytically to leading order in $1/N$ for distributions of $\epsilon_i$ and $\gamma_i$ with negligible short range fluctuations (such that the spacing $\delta_i = \epsilon_{i+1} - \epsilon_i$ changes slowly with $i$—$|\delta_{i+1} - \delta_i| / \delta_i$ is of order $1/N$ for all $i$—and similarly for $\gamma_i$). For simplicity, let us take constant $\gamma_i$, which we can set to one with no loss of generality, and equally spaced $\epsilon_i$, i.e. $\delta_i = \delta = 2w/N$.

For excited states, we write $E_k = \epsilon_k - \alpha_k \delta$, where $0 < \alpha_k < 1$, and solve equation (8) for $\alpha_k$ to the leading order in $1/N$ as described in appendix B of [46]. This yields

$$\cot \pi \alpha_k = \frac{\delta}{\pi y} + \frac{1}{\pi} \ln \frac{\epsilon_k + w}{w - \epsilon_k} = f(\epsilon_k). \quad (13)$$

We note that $\lambda = y / \delta$ is the proper dimensionless coupling constant in the sense that it must stay finite in the $N \to \infty$ limit. This is because the second summation in equation (2) scales as $N^2$ for $t_{ij} = \gamma_j \gamma_i$ and our choice of $N$. More precisely, the probability that $m$ of $\epsilon_i$ occur in an interval of length $\delta$ for Poisson distribution is $e^{-1/\delta^2}$, which however still leads to the same estimate (11).
\[ \gamma_i. \text{Therefore, we need } y \propto \delta \propto 1/N \text{ so that both terms in equation (2) are extensive in the thermodynamic limit. For the BCS Hamiltonian in equation (9), so defined } \lambda \text{ is the dimensionless superconducting coupling [47].} \]

Equations (10) becomes to leading order in $1/N$

\[
PR_{E_k} = \sum_{n=0}^{\infty} \left( \frac{1}{(n + \alpha_k)^2} \right)^2 + \left( \frac{1}{(n + 1 - \alpha_k)^2} \right)^2,
\]

which evaluates to

\[
PR_{E_k} = \frac{3}{1 + 2 \cos^2 \pi \alpha_k} = \frac{3}{1 + 3 f^2(\epsilon_k)}.
\]

This answer is in good agreement with numerics already for $N = 20$, see also figure 1. Note that $1 \leq PR_{E_k} \leq 3$.

We saw above that the ground state energy $E_0 \to -\infty$ as $y \to \infty$, while $E_0 \to \epsilon_0$ for $y \to 0$. Let $y$ be large enough that $E_0$ is well separated from $\epsilon_0$. Then, we can replace summation in equation (8) with integration and obtain

\[
\ln \frac{E_0 - w}{E_0 + w} = \frac{\delta}{y} = \frac{2w}{Ny}.
\]

Performing the same replacement in equation (10) and using equation (16), we derive

\[
PR_{E_k} = \frac{3N}{1 + 2 \cosh(\delta/y)}.
\]

Note that in the limit $y \to \infty$, $PR_{E_k} = N$ in agreement with equation (12). This expression also allows us to estimate the value $y_l$, beyond which the ground state becomes extended. We obtain $\lambda_l = \gamma_l / \delta \approx 1/\ln N$. This also corresponds to the coupling for which the gap in the spectrum $\Delta = E_1 - E_0 \approx -w - E_0$ becomes comparable to the spacing $\delta$. For a superconductor described by the BCS model (9) this localized-extended crossover translates into a normal-superconducting one[48, 49]. As $N \to \infty$ this crossover becomes a quantum phase transition at $\lambda = 0$, i.e. any infinitesimal coupling is sufficient to make the ground state extended (superconducting). The localized character of the excited states for the specific case of $\gamma_l = 1$ has been demonstrated in a previous work as well [43].

4. A model with finite-ranged hopping

We now consider the following Anderson-type model in one-dimension with nearest neighbor hopping

\[
H = \sum_i \epsilon_i n_i - y \sum_i (\epsilon_i^+ + h.c.) = H_0 + yH_f.
\]

This corresponds to the case with $t_{ij} = t$ for $|i - j| = 1$ and 0 otherwise for the general Hamiltonian in equation (2). $H_0$ is the zeroth order Hamiltonian with only the on-site potential and $H_f$ contains the hopping. It is known that all single particle eigenstates of this Hamiltonian are localized [1, 3].

4.1. Construction of the conserved charges

Proceeding as for the case of Type-1 Hamiltonians, we focus on the conserved charge $Q_0$, corresponding to the site $i = 0$, which to lowest order is equal to $n_0$. However, in this case $Q_0$ is not simply linear in $y$. In fact, it can be argued that the an expansion of $Q_0$ in the hopping does not truncate at any finite order in the thermodynamic limit. Indeed, as explained in the Introduction, conserved charges are generally infinite power series in $y$. We thus assume $Q_0$ of the form

\[
Q_i = P_{i0} + yP_{i1} + y^2P_{i2} + \cdots,
\]

where $P_{i0} = n_i$ and $P_{i1}, P_{i2}, \ldots$ are operators to be determined in terms of the microscopic parameters subject to the condition $[Q_i, H] = 0$. For concreteness, we first take our one-dimensional system to be a finite-sized ring of $N + 1$ sites going from 0 to $N$.

Since the Hamiltonian $H$ and all the zero order charges $n_i$ are quadratic in the creation and annihilation operators, we take all the operators $P_{i0}, P_{i2} \ldots$ to be similarly quadratic, i.e.
\[ P_{im} = \sum_{jk} \eta_{jk}^{(m)}(i) c_i^\dagger c_j, \]  
(20)

where the symmetric coefficients \( \eta_{jk}^{(m)}(i) = \eta_{kj}^{(m)}(i) \) are to be determined. We have

\[ [Q_i, H] = [P_{00}, H_0] + \sum_m \gamma^{m+1}(i)(P_{0m}, H_0) + [P_{m0}, H_0]. \]

The requirement that the commutator vanishes to all orders in \( \gamma \) requires

\[ [P_{0m}, H_0] + [P_{m0}, H_0] = 0 \]  
(21)

and yields a recursion relation among \( \eta_{ij}^{(m)} \)'s

\[ \eta_{ab}^{(m+1)}(s) = \delta_{ab} R_{aa}^{(m+1)}(s) + \sum_{s} \frac{1}{\epsilon_a - \epsilon_b} \sum_{k,j} \left( \eta_{kj}^{(m)}(s) + \eta_{jk}^{(m)}(s) - \eta_{ab}^{(m)}(s) \eta_{kj}^{(m)}(s) - \eta_{jb}^{(m)}(s) \eta_{ik}^{(m)}(s) \right), \]  
(22)

with initial conditions \( \eta_{00}^{(0)}(s) = \delta_{0k} \delta_{0\ell} \). The diagonal term \( R_{aa}^{(m+1)} \) represents a ‘gauge’ freedom, since the corresponding term in \( P_{0m} \) commutes trivially with \( H_0 \). We further discuss this freedom below. Specializing to the case of nearest neighbor hopping equation (18) and with \( i = 0 \), it can be verified that terms present in \( P_{0m} \) are of the form

* \( \eta_{0m}^{(m)}(s) c_0^\dagger c_m + c_m^\dagger c_0 \),
* \( \eta_{0N,m-1}^{(m)}(s)c_0^\dagger c_{N-1} + c_{N-1}^\dagger c_0 \),
* \( \sum_{\ell,j=m,|\ell-j| - \text{even}} \eta_{00,\ell}^{(m)}(s)(c_{\ell-1}^\dagger c_j + c_j^\dagger c_{\ell+1}) \) (if \( m \) is even),
* \( \sum_{\ell,j=m,|\ell-j| - \text{odd}} \eta_{00,\ell}^{(m)}(s)(c_{\ell-1}^\dagger c_j + c_j^\dagger c_{\ell+1}) \) (if \( m \) is odd).

This is shown schematically in figure 2 for the first few \( P_{0m} \).

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure2.png}
\caption{Schematic diagram showing hopping terms present in the operators \( P_i \). The base site 0 is in the middle and its neighbors are sites 1 and \( N \), since we imposed periodic boundary conditions. Lines connecting pairs of sites indicate the presence of the corresponding hopping term in the operator \( P_{im} \). Note that the range of the hopping in \( P_{im} \) increases with \( m \).}
\end{figure}

The \( Q_i \)'s are related to each other by translating all site indices in the above relations by an appropriate number. By construction, they all commute with \( H \). Since \( H \) is generally non-degenerate, this implies \( Q_i \) also commute among themselves, \( [Q_i, Q_j] = 0 \) \( \forall i, j \). To see this, first recall that for Hermitian matrices \( [A, B] = [A, C] = 0 \) implies \( [B, C] = 0 \) as long as eigenvalues of \( A \) are non-degenerate. All operators involved in the above construction of \( Q_i \) are of the form \( \hat{A} = \sum_{ij} A_{ij} c_i^\dagger c_j \), where \( A_{ij} \) is a Hermitian \( N \times N \) matrix, which represents operator \( \hat{A} \) in the sector with total particle number \( n = 1 \). Moreover, the commutativity of any two such operators is equivalent to that of the underlying matrices. Eigenvalues of the Hamiltonian in the \( n = 1 \) sector at \( y = 0 \) are \( \epsilon_i \), which are assumed to be distinct, i.e. the corresponding matrix is non-degenerate at \( y = 0 \). By continuity of the eigenvalues in \( y \), it remains non-degenerate in some finite interval (until the first level crossing) of the real axis containing \( y = 0 \). Thus, \( [Q_i, Q_j] = 0 \) \( \forall i, j \) in this interval of \( y \). But, as can be seen e.g. from the above construction of \( Q_{00} \), commutativity of \( Q_i \) on any finite interval of values of \( y \) implies that they commute for all \( y \).

We noted above that commutation relations (21) and consequently recursion relations (22) do not constrain the diagonal part of the coefficients \( \eta_{00}^{(m)}(s) \) i.e. \( R_{00}^{(m)}(s) \), for \( m \geq 1 \). The choice of \( R_{aa}^{(m)}(s) \) however does affect the off-diagonal part of \( \eta_{ij}^{(k)} \) for \( k > m \). In our construction of \( Q_i \) we set \( R_{aa}^{(m)}(s) = 0 \) for all \( m \geq 1 \), since this leads to the most compact description of these objects. we will refer to this as the standard gauge. Conserved charges \( \tilde{Q}_i \) resulting from any other choice \( R_{aa}^{(m)}(s) \) uniquely relate to our standard gauge \( Q_i \)'s, a brief calculation shows their relationship is
\[ \tilde{Q}_i = Q_i + \sum_{m} y_m \sum_{r} R_{r}^{(m)}(i) Q_r. \]  
(23)

Another advantage of our choice of a gauge is in a simple relationship between the Hamiltonian (18) and the conserved charges, namely

\[ H = \sum_{i} \epsilon_i Q_i. \]  
(24)

To see this, consider the difference

\[ H - \sum_{i} \epsilon_i Q_i = y W_1 + y^2 W_2 + y^3 W_3 + \ldots, \]  
(25)

where \( W_i \) are \( \gamma \)-independent operators. Note that the zeroth order term cancels in the difference. Since \( H \) commutes with all \( Q_i \), the right-hand side (rhs) of equation (25) must also commute. This implies in particular \[ [W_i, n_i] = 0 \] for all \( i \) (from the coefficient at the lowest power of \( y \) in the commutator of the rhs with \( Q_i \)), which in turn means that \[ W_i = \sum_{j \neq i} r_{ij} n_j. \] Now note that the left-hand side (lhs) has zero diagonal matrix elements, i.e. no terms of the form \( \epsilon_i c_i \). This is because the zeroth order term is absent, while higher order terms have no diagonal matrix elements since \( r_{ij}^{(m)} = 0 \) for all \( m \geq 1 \) in our gauge (and similarly the diagonal is absent in other \( Q_i \)). Then, the diagonal matrix elements must vanish on the rhs as well, to all orders in \( y \). In particular, \( r_{ij} = 0 \) i.e. \( W_i = 0 \) and

\[ H - \sum_{i} \epsilon_i Q_i = y^2 W_2 + y^3 W_3 + \ldots. \]  
(26)

Applying the same argument to the rhs of this equation we similarly obtain \( W_2 = 0 \) etc., until we finally arrive at equation (24).

### 4.2. Type-1 Hamiltonians redux

We have seen above that the conserved charges are power series in the hopping. This is unlike the case of Type-1 Hamiltonians, where the power series truncates after the first term. The gauge where the series truncates corresponds to having distinct terms for \( m = 1 \), one can see in equation (3) (the gauge terms are indicated in the lower braces).

It is an amusing exercise to determine the correct gauge terms that lead to truncation, starting from the recursion relations equation (22). To obtain Type-1 Hamiltonians we set \( t_{ij} = \gamma_i \gamma_j \), so that the recursions simplify to

\[ \eta_{ab}^{(m+1)}(i) = \delta_{ab} R_{a}^{(m+1)}(i) - \frac{1}{\epsilon_a - \epsilon_b} (Y_{ab}^{(m)}(i) - Y_{ba}^{(m)}(i)) \]

\[ Y_{ab}^{(m)}(i) = \sum_{j} \gamma_{ab}(m) \gamma_{jb}. \]  
(27)

With the initial condition \( \eta_{ab}^{(0)}(i) = \delta_{ab} \), we obtain at the first level

\[ \eta_{ab}^{(1)}(i) = \delta_{ab} R_{a}^{(1)}(i) + \frac{\gamma_a \gamma_b (1 - \delta_{ab}) (\delta_{ib} - \delta_{ia})}{\epsilon_a - \epsilon_b}. \]  
(28)

At this point we pause and ask if we can choose the gauge term \( R_{a}^{(1)}(i) \) such that \( \eta_{ab}^{(1)}(i) \) can be made to vanish identically, so that the iterations stop at the first level. From equation (27) we see that the relevant condition is the vanishing of \( (Y_{ab}^{(1)}(i) - Y_{ba}^{(1)}(i)) \). Using equation (27) compute

\[ Y_{ab}^{(1)}(i) = \gamma_a \gamma_b \left\{ R_{a}^{(1)}(i) + \frac{\gamma^2_a}{\epsilon_a - \epsilon_i} (1 - \delta_{ia}) - \delta_{ia} \sum_{j} \frac{\gamma^2_j}{\epsilon_j - \epsilon_i} \right\}. \]  
(29)

We may choose \( R_{a}^{(1)} \) so that the term in braces vanishes, thus leading to the truncation of the iterations. From equation (28) we have the complete first order term, and we can proceed to construct the charge (denoting the currents by the symbol \( \tilde{Q} \))

\[ \tilde{Q}_i = n_i + y \sum_{ab} \eta_{ab}^{(1)}(i) c_a^j c_b, \]  
(30)

which is identical to that in equation (3).

The use of the gauge term here is very special, and guided by our understanding of this model. On the other hand, we could by default set all the gauge terms \( R_{a}^{(m)} \) to zero, giving us the irreducible (i.e. standard gauge) currents. These no longer truncate even for Type-1 Hamiltonians. For completeness we note the second order term for the current in this (standard) gauge
\[ Q_i = n_i + (y + y^2 \sum_j \frac{\gamma_j}{\epsilon_j - \epsilon_i}) \times \sum_{i \neq j} \frac{\gamma_{ij}}{\epsilon_j - \epsilon_i} (c_i^+ c_j + c_j^+ c_i) \]
\[ + y^2 \gamma_i^2 \sum_{a,b} \frac{\gamma_{ab}}{(\epsilon_a - \epsilon_i)(\epsilon_b - \epsilon_i)} c_j^+ c_j + O(y^3). \] (31)

Thus the Type-1 Hamiltonians allow for variety of expressions of the constants of motion. To establish their equivalence in general is a subtle problem, where some surprising results have been found quite recently in [30].

This type of gauge choice, made explicit in our construction could be exploited further to test the possibility that the series can take simpler forms, as compared to a brute force expansions to infinite order. We leave this interesting question for future investigation.

4.3. Currents found from the Rayleigh–Schrödinger (locator) expansion

A natural question that arises is the relationship between the currents found above and those found from a brute force expansion of the projection operators of the Anderson model in powers of the coupling constant \( y \). The model has a formal single particle eigenfunction expansion in the form
\[ |\Psi(y)\rangle = \sum_k u_{0k} (y) c_k^\dagger |0\rangle, \] (32)

with an initial condition localized say at the site 0 as \( u_{0k} (0) = \delta_{k0} \). The projector \( Q = |\Psi(y)\rangle \langle \Psi(y)| \) can be expanded in a series in \( y \)
\[ \hat{Q} = \sum_{j,k} u_{0j} u_{0k}^* c_j^\dagger c_k = \hat{P}(0) + y \hat{P}(1) + y^2 \hat{P}(2) + \cdots \] (33)
so that the basic expansion of the wave functions in a Rayleigh–Schrödinger (RS) series in \( y \) generates the conserved currents. We can use the standard result in text books to write a perturbative expansion for the state at site 0 with standard normalization to \( u_{00} = 1 \) as
\[ |\Psi(y)\rangle = c_0^\dagger |0\rangle + \sum_{k=0} u_{0k} c_k^\dagger |0\rangle, \] (34)
with a power series expansion for \( u_{0k} \)
\[ u_{0k} = -y \frac{f_{0k}}{\epsilon_0 - \epsilon_k} + y^2 \sum_{i=0} \frac{f_{0i} f_{0k}}{(\epsilon_0 - \epsilon_i)(\epsilon_0 - \epsilon_i)} - y^2 \frac{f_{00} f_{0k}}{(\epsilon_0 - \epsilon_k)^2} + O(y^3). \] (35)

Using this expansion, we may generate the series equation (33), the result is explicitly stated below in equation (38). From this series we can verify to second order, that this series differs from that in the standard gauge equation (19) by specific gauge terms. The advantage of equation (19) is that this gauge invariance is manifest in the construction by the nested commutators. On the other hand, equations (33)–(35), corresponds to a particular gauge picked out by the R–S method, and the currents found here are some linear combinations of the ones in equation (19) as in equation (23).

It seems to us that the series in equation (19) possesses an essential simplicity relative to the Rayleigh–Schrödinger series equations (33)–(35). The R–S perturbation expansion simultaneously determines the energy eigenvalue, and for this purpose very specific gauge terms are needed. On the other hand all terms in equation (19) are generated by completely off diagonal terms, those terms that avoid multiple visits to any site. This leads to simpler recursion relations, as in equation (22), relative to the RS series. For this reason our numerical work in this paper uses the series in equation (19).

4.4. Locator expansion for Type-1 Hamiltonians

The Rayleigh–Schrödinger series can also be constructed for Type-1 Hamiltonians using the exact eigenstates \( |E\rangle \) with eigenvalues \( E \) as given in equations (7) and (8). The projector \( |E\rangle \langle E| = \sum_{\gamma} \frac{\gamma \gamma^\dagger}{(E - \epsilon_k)(E - \epsilon_k)} \) can be expanded in \( y \) as shown in equation (33). In the limit \( y \to 0 \), the roots of equation (8) tend to \( \epsilon_i \). We take the root \( E \to \epsilon_0 \) to obtain \( \tilde{Q}_0 \) the conserved charge corresponding to site 0 calculated using the Rayleigh–Schrödinger gauge. Other roots yield other \( \tilde{Q}_i \). Expanding equation (8) for \( E \) in \( y \) near \( E = \epsilon_0 \), we get
\[ E = \epsilon_0 - y\gamma_0^2 + y^2 \gamma_0^2 \sum_{i=0} \frac{\gamma_i^2}{\epsilon_0 - \epsilon_i} + O(y^3). \] (36)

Since, the projector diverges in \( y \to 0 \) limit, we define our conserved charge as \( \tilde{Q}_0 = \frac{(E - \gamma_0^2)}{\gamma_0^2} |E\rangle \langle E| \) to make it well behaved. \( \tilde{Q}_0 \) is given by

9 For example, see equation (5.1.44) Sakurai 1994 Modern Quantum Mechanics (London: Pearson Education).
The conserved charges constructed above depend on the microscopic parameters of the Hamiltonian, i.e. the hopping and on-site energies. As we shall show later, the same Hamiltonian can have a localized and delocalized phase depending on the values of these parameters. It is thus important to understand if and how the conserved charges themselves differ in the two phases. More precisely, how do the conservation laws 'know' whether a particular choice of microscopic parameters produces a localized or delocalized phase?

The answer has to do with their convergence since they are expressed as power series in the microscopic parameters and particle operators. We thus need to state in what sense the power series are convergent. A reasonable condition for convergence is a sufficiently rapid decay of the coefficients $\beta_m^m$ with increasing $m$. However, this is complicated by the fact that there are energy difference denominators in the coefficients $\beta_m^m$ that can cause them to blow up when the on-site energies at two different sites are equal. To avoid this, we restrict ourselves to a particular type of disorder that may be termed 'non-resonant'. By this we mean any ensemble of $\epsilon_i$ which shows 'level repulsion', i.e. the probability of finding $\epsilon_i$ very close to each other is very small.

From the random matrix theory, we know that the eigenvalues of a generic matrix display level repulsion in their eigenvalues of various degree, the Gaussian orthogonal ensemble (GOE) [50] of real symmetric matrices has the least level repulsion. This condition ensures that perturbative resonances from small denominators, that would otherwise cause individual terms in the expansions of the conserved charges to diverge, are prohibited. This choice is similar to the one involving limited level attraction recently adopted in the context of many-body localization [51].

We have verified that this distribution of onsite energies gives us localization (as indicated from a calculation of the PR) immediately upon switching on the hopping term. Thus, this particular choice of onsite energies, which is of great convenience from the point of view of calculations, is also not unphysical. The on-site energies $\epsilon_i$ are drawn from the eigenvalues of a real symmetric matrices whose elements are taken from a Gaussian random distribution with fixed variance. The eigenvalues of these matrices are assigned randomly to different sites. Different random assignments then constitute different realizations of disorder, which can then be averaged over to check for convergence. The result of this procedure is shown in figure 3, where $\epsilon_i$ are drawn from the eigenvalues of real symmetric matrices whose elements are taken from a Gaussian distribution of variance $\sigma = 0.1, 0.25$ and 0.4. It can be seen that the $\beta_m^m$ decrease rapidly with increasing order of power series $m$.

Other $\tilde{Q}_k$ can be obtained with the replacement $0 \to k$. Unlike $Q_k$, there is no indication of the series truncating at any finite order for $\tilde{Q}_k$.

4.5. Convergence of the power series

The conserved charges constructed above depend on the microscopic parameters of the Hamiltonian, i.e. the hopping and on-site energies. As we shall show later, the same Hamiltonian can have a localized and delocalized phase depending on the values of these parameters. It is thus important to understand if and how the conserved charges themselves differ in the two phases. More precisely, how do the conservation laws 'know' whether a particular choice of microscopic parameters produces a localized or delocalized phase?

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Figure 3. Plot indicating convergence of conserved charges (see equations (19) and (20)) of the Anderson model (18) for $N = 500$. $[\eta]$ represents a typical $m$th coefficient $\eta_m^m$ averaged over a distribution of the on-site disorder $\epsilon_i$, see the end of section 4.5. The plot shows the logarithm of the average as a function of $m$, $\epsilon_i$ are drawn from the eigenvalues of real symmetric matrices whose elements are Gaussian random variables of variance $\sigma = 0.1, 0.25$ and 0.4, and we set $y = 1$. 

\[
\tilde{Q}_0 = n_0 + \frac{E - \epsilon_0}{\gamma_0} \sum_{j=0}^{\infty} \gamma_j (c_j^+ c_j + c_j^+ c_j) + \frac{(E - \epsilon_0)^2}{\gamma_0^2} \sum_{i,j=0}^{\infty} \frac{\gamma_i \gamma_j c_i^+ c_j}{(E - \epsilon_i)(E - \epsilon_j)}. \tag{37}
\]

Then, replacing $\frac{(E - \epsilon_0)^2}{\gamma_0^2} \to y^2 \gamma_0^2$ and then $E \to \epsilon_0$, we have obtained $\tilde{Q}_0$ as a combination of $Q_i$ (see equation (3)) as follows

\[
\tilde{Q}_0 = Q_0 - y \sum_{i=0}^{\infty} \frac{\gamma_i^2 Q_i + \gamma_i^2 Q_0}{\epsilon_0 - \epsilon_i} + O(y^3). \tag{38}
\]

Other $\tilde{Q}_k$ can be obtained with the replacement $0 \to k$. Unlike $Q_k$, there is no indication of the series truncating at any finite order for $\tilde{Q}_k$. 

\[
\tilde{Q}_0 = n_0 + \frac{E - \epsilon_0}{\gamma_0} \sum_{j=0}^{\infty} \gamma_j (c_j^+ c_j + c_j^+ c_j) + \frac{(E - \epsilon_0)^2}{\gamma_0^2} \sum_{i,j=0}^{\infty} \frac{\gamma_i \gamma_j c_i^+ c_j}{(E - \epsilon_i)(E - \epsilon_j)}. \tag{37}
\]
indicating convergence. We have also checked the convergence of the power series for \( \epsilon_i \) drawn from the eigenvalues of non-integrable \( t - t' - V \) model, which also follow a GOE distribution [33, 52]. Since \( \eta_j^m \) contain more than one term for each \( m \), we checked the convergence of a typical term, which is of the form \( \sum_{(\epsilon_{a_1} - \epsilon_{b_1})... (\epsilon_{a_m} - \epsilon_{b_m})} e^{\sigma} \). Recall that the \( m^{th} \) order term in the calculation of \( \tilde{Q}_0 \) involves sites with labels between \( N - (m - 1) \) and \( m \) as can be seen from figure 2. Thus, the only values of \( \epsilon_i \) involved are those chosen from \( [\epsilon_{N - (m - 1)}, \epsilon_{m}] \) (\( \epsilon_0 \) is at the center) such that \( \epsilon_{a_i} = \epsilon_{b_i} \) \( \forall i \) and \( \max|a_i - b_i| = m \).

As the aim of this work is to construct conserved charges in localized systems, it is legitimate to ask whether this slightly non-standard choice of disorder distribution produces localization. We have verified this through numerical exact diagonalization by calculating the PR. We find that the PR for different eigenstates is indeed close to zero for systems of size \( N = 500 \) as shown in figure 4, consistent with localization. We thus conclude that our model with on-site energies taken from a GOE distribution does indeed produce a localized phase. A similar exercise to construct the conservation laws for the above model has been carried out in [53]. In that work too, the conserved charges have been constructed as infinite operator series but whose coefficients correspond to the amplitudes of a particle to be on the sites of a square lattice whose sides are the physical one-dimensional lattice. The recursion relation obtained is between conserved charges on different sites and the convergence of the series is assumed to follow from the exponential decay of the eigenfunctions of the Hamiltonian. In our calculations, we construct the conserved charges directly in terms of the microscopic parameters of the Hamiltonian and our convergence criterion is not based on any assumption about the nature of the eigenstates of the Hamiltonian. In fact, as we show in the next section, the convergence of the series for the conserved charges can be used to identify the delocalized and localized phases instead of the eigenfunctions.

5. Aubry–Andre model

Having constructed the conserved charges for a model with finite-range hopping and defined a condition for convergence of the power series for them, we can further investigate the meaning of our convergence criterion. In particular, since our goal is to identify the validity of our construction of the conservation laws with the presence of localization, the power series should fail to converge according to our criterion in a delocalized phase.

We thus require a non-interacting model with disorder in one-dimension which has a delocalized phase. While any model with finite-range hopping and an on-site random potential in one-dimension always produces localization [1, 3], a quasi-periodic potential can produce localized and delocalized phases. Such a model is the Aubry–Andre model [42] given by the Hamiltonian

\[
H = h \sum_j \cos(2\beta \pi j) c_j^\dagger c_j - \frac{1}{2} \sum_j (c_j^\dagger c_{j+1} + \text{h.c.}),
\]

(39)

where \( \beta \) is an irrational number. The parameter \( h \) can be tuned to effect a transition from a localized phase (for \( h > 1 \)) to a delocalized phase (for \( h < 1 \))[42]. We note that this model is usually studied with an additional term that introduces a \( p \)-wave pairing gap [54], but we set it equal to zero for our analysis.

The localized phase here is one in which all single particle states are localized and similarly all single particle states are delocalized in the delocalized phase. The transition between these phases happens at \( h = 1 \). Since the Hamiltonian in equation (39) is also of the form (18), we can use the expressions obtained for the \( \eta_j^m \) in the
previous section to construct the conserved charges. These will now depend on the parameter $h$ (i.e. $y \rightarrow (2 \ h)^{-1}$ in the previous section) and if the criterion for convergence postulated by us is a valid one to detect localization, we should observe the power series to converge in the localized phase ($h > 1$) and diverge in the delocalized phase ($h < 1$). This is indeed the case as we see e.g. from figure 5, which shows that a typical matrix element of $\eta^m$ goes to zero quite rapidly with increasing $m$ for $h > 1$ but diverges for $h < 1$. Thus, we have established that our convergence criterion is valid for identifying the localization–delocalization transition.

6. Interactions

We now turn to systems with interactions. The simplest way to introduce interactions to models we studied here is through a nearest neighbor density–density term. Let us, for example, add such a term to equation (18)

$$H = \sum_i \epsilon_i n_i - ty \sum_i (c_i^\dagger c_{i+1} + \text{h.c.}) + V \sum_i n_i n_{i+1}$$

where we redefined $H_0$ as compared to equation (18).

We assume that the particles here are spineless fermions. It is tempting to try a construction of the conserved charges starting from a zeroth order Hamiltonian that combines the on-site and interaction terms since they commute with each other and their eigenstates are localized at every site. However, the interaction term is quartic in creation and annihilation operators and so the conserved charges can no longer be assumed to be power series in the hopping with each term quadratic in the creation and annihilation operators. Such an assumption leads to no solution for the coefficients since the commutators keep producing terms with increasingly longer trails of creation and annihilation operators as one goes to higher orders in the hopping. A more profitable exercise is to try to obtain the conserved charges as power series in the hopping but only to the first order in the interaction. While these are not exact, they offer a reasonable approximation in the limit of small interaction strength. Weak interactions typically should not destroy the localization present in the non-interacting limit and thus conserved charges should continue to exist.

We know from our previous calculation that the operator of the form $Q_0 = n_0 + \sum_{ijm} \eta^m_{ij} y^m c_i^\dagger c_j^\dagger$ commutes with $H_0$. Let us now define a new operator $Q = Q_0 + V \delta Q$ to linear order in $V$ and calculate the commutator

$$[Q, H] = [Q_0 + V \delta Q, H_0 + V \delta H]$$

$$= V [\delta Q, H_0] + [Q_0, \delta H] + O(V^2).$$

We choose $\delta Q$ such that $[\delta Q, H_0] + [Q_0, \delta H] = 0$, so that $Q$ and $H$ commute to $O(V)$. We assume the form $\delta Q = \sum_{rst} \psi_{rst} c_r^\dagger c_s^\dagger c_t$. Note this is quartic in the creation and annihilation operators since the interaction term is as well. Thus

$$[\delta Q, H_0] + [Q_0, \delta H] = 0$$
\[
\sum_{k,m} y^m (\eta_{k+1,1}^m c^i_k c_{k+1} + \eta_{k-1,1}^m c^i_k c_{k+1}) = y \sum_{r,s} \psi_{r,s} (1 - \psi_{r,s} - 1^v + \psi_{r,s} \psi_{r+s} - \psi_{r+s} - \psi_{r+s}^2)
\]

We now assume that \(\psi_{r,s}\) can be written as a power series in \(y\), i.e. \(\psi_{r,s} = \sum_{m} A^r_{m,s} y^m\). Equating the coefficients at different orders of \(y\), one can in principle obtain \(A^r_{m,s}\) in terms of the \(\eta^r_{t,s}\) for the case with \(V = 0\). In fact, it can be seen that at a given order \(m\), the \(A^r_{m,s}\) are linear combinations of the \(\eta^r_{t,s}\) and the \(A^m_{m,s}\). One can also impose constraints arising from the anti-commutation of the fermionic operators, the Hermitian nature of the conservation laws and the number of non-zero components of the \(A^m_{m,s}\) to severely constrain the number of non-zero components of \(A^m_{m,s}\).

Let us, for example, derive \(\delta Q\) to the first order in \(y\), i.e. we set \(m = 1\). We have

\[
\sum_{k,i} (\eta_{k+1,i}^1 c^i_k c_{k+1} + \eta_{k-1,i}^1 c^i_k c_{k+1}) = \sum_{r,s} A^{1}_{r,s} (-c^i_e + c^i_f) c^i_r c^i_s c^i_v.
\]

Since, only \(\eta_{k,i+1}^1\) and \(\eta_{k,i-1}^1\) are non-zero, the non-zero \(A^{1}_{r,s}\) are given by the following equations:

\[
A^{1}_{k+1,k+1,k+1} = A^{1}_{k,k+1,k+1} = \frac{\eta_{k+1,k}^1}{\epsilon_{k+1} - \epsilon_k}
\]

\[
A^{1}_{k-1,k+1,k+1} = A^{1}_{k-1,k,k+1} = \frac{\eta_{k-1,k}^1}{\epsilon_{k-1} - \epsilon_k}
\]

\[
A^{1}_{k,k+1,k} = A^{1}_{k,k,k+1} = \frac{\eta_{k,k+1}^1}{\epsilon_k - \epsilon_{k+1}}
\]

\[
A^{1}_{k,k+2,k+1} = A^{1}_{k,k+1,k+2} = \frac{\eta_{k+2,k+1}^1}{\epsilon_{k+2} - \epsilon_{k+1}}
\]

The corresponding expression for \(\delta Q\) to order \(y\) is

\[
\delta Q = y V \sum_{r,s} A^{1}_{r,s} c^i_r c^i_s c^i_v
\]

\[
= y V \sum_{k} \left[ \frac{\eta_{k+1,k}^1}{\epsilon_{k+1} - \epsilon_k} (c^i_{k+1} c^i_k + c^i_{k+1} c^i_{k+1}) n_{k+1} + \frac{\eta_{k-1,k}^1}{\epsilon_{k-1} - \epsilon_k} (c^i_{k-1} c^i_k + c^i_{k-1} c^i_{k-1}) n_{k+1}
\]

\[
+ \frac{\eta_{k,k+1}^1}{\epsilon_k - \epsilon_{k+1}} n_k (c^i_{k+1} c^i_k + c^i_{k+1} c^i_{k+1})
+ \frac{\eta_{k+2,k+1}^1}{\epsilon_{k+2} - \epsilon_{k+1}} n_k (c^i_{k+2} c^i_{k+1} + c^i_{k+1} c^i_{k+2}) \right].
\]

Other approaches to construct conservation laws for interacting systems have been proposed including a recent one where the interacting problem is mapped onto a non-Hermitian problem on a lattice in operator space [53]. A convergence criterion for the resultant series based on the operator norm is then used to identify localized and delocalized phases.

7. Conclusions and discussion

Inspired by the Type-1 Hamiltonian system, we have demonstrated a scheme to obtain the conserved charges for non-interacting disordered models displaying localization in one-dimension. One of our motivation was an observation of similarities between localized and integrable systems, such as the absence of level repulsion and the absence of thermalization. Our conserved charges are exhibited as a power series in the hopping, and using a suitable convergence criterion, we show that the convergence (or divergence) of conserved charges tracks the presence (or absence) of localization. An interesting issue of ‘gauge dependence’ of the conserved charges is unearthed and explored. It is shown that a full understanding of the gauge dependence leads to considerable simplifications of the charges in some cases. On the other hand, straightforward Rayleigh–Schrödinger perturbation theory or equivalent schemes, commit one to a particular gauge that is often inconvenient.

This work provides a novel link between the concepts of localization and integrability. Our results hold within the context of the 1d Anderson model, where all states are localized, and the Andre–Aubry model, where (all) states undergo a transition tuned by a coupling constant. It is not immediately obvious how to extend these
results to a higher dimensional Anderson model with a mobility edge separating the two classes of states. The Aubry–André model exhibits an interesting kind of duality which allows the localized and delocalized phases to be mapped onto each other with the roles of the hopping and onsite potential exchanged. The duality transformation is expressed in terms of new fermionic operators given by $c_{\xi} = \frac{1}{\sqrt{L}} \sum_{n} e^{i(2\pi \tilde{k} \xi)} c_{n}$, which are eigenstates of the momentum operator with eigenvalue: $k = \tilde{k} F_{n-1} \mod F_{n}$, where $F_{n}$ is the $n$th Fibonacci number and $L = F_{n}$ [55, 56]. In terms of these fermionic operators the Hamiltonian (39) becomes

$$\frac{H}{h} = \frac{1}{h} \sum_{\tilde{k}} \cos(2\beta \pi \tilde{k}) n_{\tilde{k}} - \frac{1}{2} \sum_{\tilde{k}} (c_{\tilde{k}}^\dagger c_{\tilde{k}+1} + \text{h.c.}).$$ (42)

The Hamiltonian satisfies the duality relation: $H(h)/h = H(1/h)$. We have shown that for the Aubrey-Andre Hamiltonian written in real space, one can construct a set of conserved charges that converge for $0 < h < 1$. Because of the duality of the model one can construct similar conserved charges in terms of $c_{\xi}$ and $c_{\xi}^\dagger$. The power series of these charges converge when $0 < 1/h < 1$ and both sets of charges diverge at $h = 1$. Thus, the duality of the model allows us to explicitly construct conservation charges in one phase given that they exist in the other.

This can be better understood by noting that localization is a basis dependent concept. We have been using localization (as is the standard practice) to mean localization in real space. To obtain the conserved charges for such a localized phase, we start from a Hamiltonian whose eigenstates are perfectly localized in real space and then add terms perturbatively in the hopping. Similarly, the delocalized phase of the Aubry–André model is localized in momentum space and one can then obtain its conserved charges by starting with a Hamiltonian perfectly localized in momentum space (tight binding model) and then add terms perturbatively in the on-site potential. This is the essence of the duality outlined above. Thus, the conserved charges also carry labels indicating the space (real or momentum) where the system is localized. What is important though is that once the basis in which the system is localized is identified and the conserved charges are constructed accordingly, they are sensitive to the onset of delocalization in that basis and can be used to locate localization–delocalization transitions.

The importance of the basis can be further understood when one compares the behavior hard-core bosons with that of spinless fermions in the Aubry–André model [57, 58]. The duality between the localized and delocalized phases is destroyed for hard-core bosons. As a result, the relaxation of real space local observables in the localized phase is different from their conjugates in momentum space in the delocalized phase. This feature is absent for spinless fermions where the duality holds and as a consequence, conserved charges of the type derived in this work exist in both phases.

While it is only possible to construct these charges to lowest order in the interaction using our procedure, their fate upon the introduction of interactions can in principle be investigated numerically, which we defer to a future work.\[10\]

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Note that the introduction of interactions destroys the duality of the model since it no longer has the same form in real and momentum space under the duality transform.
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