Unveiling and exemplifying the unitary equivalence of
discrete time quantum walk models

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Abstract

The two major discrete time formulations for quantum walks, coined and scattering, are unitarily equivalent for arbitrary position-dependent transition amplitudes and any topology (Andrade \textit{et al} 2009 \textit{Phys. Rev. A} \textbf{80} 052301). Although the proof explicitly describes the mapping obtention, its high technicality may hinder relevant physical aspects involved in the equivalence. Discussing concrete examples—the most general constructions for the line, square and honeycomb lattices—here we unveil the similarities and differences of these two versions of quantum walks. We moreover show how to derive the dynamics of one from the other by means of proper projections. We perform calculations for different probability amplitudes such as Hadamard, Grover, discrete Fourier transform and the uncommon in the area (but interesting) discrete Hartley transform, comparing the evolutions. Our study illustrates the models’ interplay, an important issue for implementations and applications of such systems.

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1. Introduction

Since their introduction [1–3], quantum walks (QWs), a paradigmatic and relatively simple class of quantum systems [4], have found many applications in different areas of science (see, e.g., the recent review in [5]). At the first glance, such usefulness may be attributed to QWs distinct possible formulations. For instance, although in all cases the dynamics take place in discrete spatial structures, graphs (or lattices), time can be either a continuous [6] or a discrete variable. Moreover, in discrete time versions, the evolution can be dictated by inner states

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called ‘coins’ [7] (coined QWs, CQWs) or resulting from scattering-like processes [8–10] (scattering QWs, SQWs).

Regarding uses for QWs, some aspects of the general searching problem [11] seem to be very appropriate for SQWs [12], including identification and comparison of parts of a graph [13, 14]. Also, SQWs are frequently considered in the study of scattering in semi-infinite graphs [9, 10, 15], a relevant configuration to implement universal quantum computation through QWs [16]. On the other hand, the literature proposes a much larger number of applications for CQWs [5, 17–19]. Examples are transport in biological systems [20] (in a continuous time context, see [21]), Bose–Einstein condensates redistribution [22], quantum phase transition in optical lattices [23], decoherence processes [24] and even quantum games [25]. In particular, CQWs are widely discussed in quantum computation as a tool for the development of quantum algorithms [26].

As concerns implementations, distinct experimental setups based on distinct physical phenomena can be used to actually build QWs. For CQWs, protocols based on trapped atoms in optical lattices [27], quantum dots [28], photons orbital angular momentum [29] and QED cavities [30], to cite just a few, have been devised. Concrete lab realizations were constructed with waveguide lattices [31] (refer to [32] for waveguides in the continuous time case), passive optical elements [33] and liquid-state nuclear magnetic resonance [34], again only mentioning a few examples (an overview is given in [35]). In turn, SQWs could be associated with optical networks [36, 37] and eventually may ensemble just with linear optical elements [12, 13], similarly as in [38] to fabricate a quantum version of Galton’s quincunx (a classical mechanical machine which at certain locations ‘chooses’—with 50%:50% probability—between two directions to go for a traveling ball).

The above-mentioned models’ distinctions for applications (and in to a lesser extent for implementations) are, however, due to practical and operational instead of fundamental reasons. Indeed, continuous time and coined QWs are closely related, since they lead to the same dynamics in proper limits [39, 40] (although the limits may not be so direct as to achieve [40]). Furthermore, it has been rigorously proven that CQWs and SQWs are unconditionally (i.e. for arbitrary topologies and spatially-dependent transition probabilities) unitary equivalent [41].

These kindred relations open important perspectives in the employment of QWs, especially for the discrete time versions. They should be equally appropriate in any application and in principle (see comments in section 4), the same physical implementation would be capable of simulating both CQWs and SQWs. But to benefit from such connections, it is necessary to have a clear understanding of the similarities and differences of the two versions of the system as well as the correct way to obtain one from another. The proof in [41], having as the main goal to formally demonstrate their equivalence, relies on general and very technical arguments, making it difficult to fully appreciate the conceptual features associated with the QWs reciprocities. Thus, knowing that CQWs and SQWs are always equivalent, we can focus on the representative cases, discussing in detail the interplay between the discrete time formulations. Hence, here we illustrate that different QWs are in fact closely tied and a specific model choice might be much more a matter of practicality than of unfeasibility of the corresponding sibling construction.

The work is organized as follows. From a parallel with the usual classical random walks, in section 2 we discuss for the one-dimensional (1D) case the main characteristics of CQWs and SQWs, also addressing their mapping. We do so assuming the most general situation of complete arbitrary position-dependent transition probabilities at the 1D lattice sites. In section 3, the same type of analysis is carried out for two 2D lattices, the square and the honeycomb. The models are formulated in detail and simulations, exemplifying their
Figure 1. (a) A classical random walk faced as alternate stochastic and deterministic processes along a 1D lattice. (b) Example of a possible path for $t = 6\tau$, whose probability is $p^2(1 - p)^4$. Here, $p[1 - p]$ is the probability to go to the right [left].

distinctions for the time evolution along the graphs structures, are presented. Importantly, we show how the spatial probabilities distributions for one model can be obtained from the other through proper projections. Finally, results for different coin (and the equivalent scattering) matrices operators are compared. It includes the more common Hadamard, Grove and discrete Fourier transform (DFT), as well as the not so usual (but interesting) discrete Hartley transform (DHT) and few others. A short conclusion is drawn in section 4.

2. Discrete quantum walks on the line: two different views

To better understand the main ideas underlying the construction of the existing discrete QWs models, we first recall an interesting way to view the classical case on the line (1D).

Consider a classical walker, starting in $x = 0$ at $t = 0$, that takes steps of fixed length $L$ and moves with constant speed $v$. Its simple dynamics is described as the following. Each time the walker reaches a position $x(t = n\tau) = x_n = \pm jL (j = 0, 1, \ldots$) for $n = 0, 1, \ldots$ and $\tau = L/v$, it randomly (and instantaneously) chooses a new direction to go, either to the right ($\sigma = +1$), with probability $p$, or to the left ($\sigma = -1$), with probability $1 - p$. So, this 1D continuous system can be characterized by two processes taking place in an ‘effective lattice’: (i) one purely stochastic (choosing directions), occurring at the ‘lattice sites’ $\pm jL$; and (ii) other purely deterministic (ballistic motion, with $x(t) = x_{n-1} + \sigma v(t - t_{n-1})$ for $(n - 1)\tau \leq t \leq n\tau$), occurring along the ‘lattice bonds’. Such a view is schematically represented in figure 1.

The construction of a discrete time QW closely follows the previous picture. Essentially, there are two possible implementations, based on what we assume as the primary process, either (i) or (ii) above, to describe the quantum states. Indeed, in the first case (associated with CQWs) the quantum states describe the system at the lattice site positions: classically, the locations where a probabilistic choice about the next step direction is made. In the second case (associated with SQWs), the quantum states are defined on the bonds: classically, corresponding to the deterministic locomotion along the lattice. This is an important distinction since each model has a different state representation with a different possible interpretation.

One does not need to relate any ‘lattice’ structure to QWs. Nevertheless, it is instructive to go on with such an analogy and to view quantum walks in 1D as a dynamics defined on a ‘Hilbert lattice’ [42], depicted in figure 2. In this way, we can associate the classical step
Figure 2. For QWs in 1D, the associated ‘Hilbert lattice’, but which is not necessarily a spatial structure once the states do not need to be position eigenvectors. The states are defined (a) on the sites, coined and (b) along the bonds, scattering, formulations. For SQWs, details in (b) show the scattering quantum amplitudes defined on each site.

length $L$ with the characteristic unit distance $\Delta j = 1$ between two consecutive ‘sites’ of the Hilbert lattice. Moreover, in both cases the discrete time evolution is given by a single step unitary operator $U$, acting on states belonging to this Hilbert (lattice) space, so that $|\Psi(n+1)\rangle = U|\Psi(n)\rangle$.

2.1. The coined quantum walk model

Quantum states are defined on the sites $j$, figure 2(a). Then, we consider the basis states $\{|j\rangle\}$, which span the Hilbert subspace $H_p$. Moreover, to any $j$ we associate an inner two-level Hilbert subspace $H_c$, whose ‘coin’ states are $|\pm\rangle$. They give the correct unitary ‘stochastic’ character for the problem [2, 43]. Also, their role is somehow similar to that of choosing directions in the classical case: once $|j\rangle$ represents only spatial ‘location’ (sites), then the ‘direction’ must be played by the auxiliary coin states.

The appropriate orthonormal basis to describe the entire space $H = H_p \otimes H_c$ is $\{ |j\rangle \otimes |\sigma\rangle \}$, with $\sigma = \pm$ and $j = 0, \pm 1, \pm 2, \ldots$. Defining a shift operator as $S|j\rangle = |j+1\rangle$ (and $S^\dagger|j\rangle = |j-1\rangle$), we write

$$U_c = (S \otimes |+\rangle \langle +| + S^\dagger \otimes |-\rangle \langle -|) \left( \sum_j |j\rangle \langle j| \otimes C^{(j)} \right).$$

(1)

$U_c$ is unitary if the coin-flip operators $C^{(j)}$ are also unitary. There are many possible choices for the $C$ [44], a common one being that of Hadamard, for which $\forall j, C^{(j)} = H_2$ with $H_2|\sigma\rangle = (1 - \sigma) \otimes |\sigma\rangle)/\sqrt{2}$. However, the most general case is to consider the matrix elements $\langle \sigma''|C^{(j)}|\sigma'\rangle = c^{(j)}_{\sigma''\sigma}$ and to ensure unitarity just by imposing

$$|c^{(j)}_{++}|^2 + |c^{(j)}_{-+}|^2 = |c^{(j)}_{+-}|^2 + |c^{(j)}_{--}|^2 = 1, \quad |c^{(j)}_{+\pm}|^2 = |c^{(j)}_{-\pm}|^2,$$

$$c^{(j)}_{++} [c^{(j)}_{-\pm}]^* + c^{(j)}_{-+} [c^{(j)}_{-\pm}]^* = 0.$$  

(2)

Thus, we have

$$U_c |j\rangle \otimes |\sigma\rangle = c^{(j)}_{\sigma\sigma} |j+\sigma\rangle \otimes |\sigma\rangle + c^{(j)}_{\sigma\sigma} |j-\sigma\rangle \otimes |-\sigma\rangle.$$  

(3)

One of the most important applications of CQWs is in quantum computation. So, it is usual to define the above matrix elements in a way easy to associate with quantum gate operators. It explains, e.g., the frequent choice in the literature of Hadamard’s coin (fundamental to manipulate $q$-bits). This practice, however, leads to constructions of coin matrices in a format...
not common among physicists, but popular in mathematics and computer science (as nicely explained and exemplified in [45, e.g., p 23]). Indeed, the states $|\pm\rangle$ are written as the column matrices
\begin{align}
|\pm\rangle = \begin{pmatrix} 1 \\ 0 \end{pmatrix}, \quad |+\rangle = \begin{pmatrix} 0 \\ 1 \end{pmatrix}.
\end{align}

opposite to the usual spin-up and spin-down conventions. Following this representation, the most general form for the coin operator obeying equation (2) and having the Hadamard as a particular case is (see also [46–48])
\begin{align}
C^{(j)} = \exp[i\gamma_j] \begin{pmatrix} \exp[i\xi_j] \cos[\theta_j] & \exp[i\xi_j] \sin[\theta_j] \\ \exp[-i\xi_j] \sin[\theta_j] & -\exp[-i\xi_j] \cos[\theta_j] \end{pmatrix},
\end{align}

with $0 < \gamma_j, \xi_j, \theta_j < 2\pi$. Then, equation (3) reads
\begin{align}
U_{\tau} |j\rangle \otimes |\sigma\rangle &= -\sigma \exp[i(\gamma_j - \sigma \xi_j)] |j + \sigma\rangle \otimes |+\sigma\rangle \\
&+ \exp[+i(\gamma_j + \sigma \xi_j)] |j - \sigma\rangle \otimes |-\sigma\rangle.
\end{align}

2.2. The scattering quantum walk model

For our purposes, it is easier to formulate the scattering model in a slightly different (but completely equivalent) way than usually done in the literature [10]. Along each bond, joining two consecutive sites, say $j - 1$ and $j$ (see figure 2(b)), we assume two possible states $|+\rangle = |j + 1\rangle$ and $|--\rangle = |j - 1\rangle$. Hence, in contrast to the coined model, the quantum number $\sigma = \pm 1$ is now associated with a ‘direction’ along a bond in the Hilbert lattice. The full Hilbert space $\mathcal{H}$ is no longer a direct product of two subspaces. Each basis element $|\sigma, j\rangle$ belongs to $L^2(\mathbb{Z} \times \mathbb{Z}_2)$ and satisfies $(j', \sigma'|\sigma, j\rangle = \delta_{j,j'} \delta_{\sigma,\sigma'}$.

Defining the operators $T$ and $R$ by
\begin{align}
T|\sigma, j\rangle = t^{(j)}_{\sigma} |\sigma, j + \sigma\rangle, \quad R|\sigma, j\rangle = r^{(j)}_{\sigma} |- \sigma, j - \sigma\rangle,
\end{align}
and
\begin{align}
T^\dagger|\sigma, j\rangle = t^{(-j-\sigma)^\ast}_{\sigma} |\sigma, j - \sigma\rangle, \quad R^\dagger|\sigma, j\rangle = r^{(-j-\sigma)^\ast}_{\sigma} |- \sigma, j + \sigma\rangle,
\end{align}
the one-step time evolution is simply
\begin{align}
U_{\tau} = T + R,
\end{align}
so that
\begin{align}
U_{\tau} |\sigma, j\rangle = t^{(j)}_{\sigma} |\sigma, j + \sigma\rangle + r^{(j)}_{\sigma} |- \sigma, j - \sigma\rangle.
\end{align}

The unitarity of $U_{\tau}$ implies [49]
\begin{align}
|t^{(j)}_{\sigma}|^2 + |r^{(j)}_{\sigma}|^2 = 1, \quad |r^{(j)}_{\sigma}|^2 = |r^{(j)}_{\sigma}^\ast|^2, \quad r^{(j)}_{\sigma} t^{(j)}_{\sigma} + r^{(j)}_{\sigma}^\ast t^{(j)}_{\sigma}^\ast = 0,
\end{align}
which are exactly the relations satisfied by the reflection and transmission amplitudes in a quantum scattering problem in 1D [49, 50], resulting from the unitarity of the $S$ scattering matrix.

Equation (11) automatically holds if for any $j$ (with $0 < \rho_j < 1$ and $0 < \lambda_j, \varphi_j < 2\pi$)
\begin{align}
t^{(j)}_{\sigma} &= \exp[i\lambda_j](\sqrt{1 - \rho_j} \exp[i\sigma \rho_j]), \\
r^{(j)}_{\sigma} &= \exp[i\lambda_j](\sigma \sqrt{\rho_j} \exp[i\sigma \varphi_j]).
\end{align}

If $\lambda_j = \lambda$ for all $j$, without loss of generality we can set $\lambda = 0$. We should mention that the expressions in equation (12) are not the only possibility [42]. There is a certain arbitrariness in
signals convention. The present choice, however, has a direct physical motivation. For time-
reversal invariant systems, $t^{(j)}_\sigma = t^{(j)}_{-\sigma}$ [50]. Then, if we also assume real amplitudes, the signals for the $r$ must be opposite, exactly the case in equation (12). Furthermore, for $j$-independent scattering coefficients, the above (with $\phi_j = \varphi_j = 0$) are the relations adopted in the original work introducing SQW models [8].

2.3. Obtaining the probabilities

At this point we should emphasize that the above constructions are more general than simply quantizing the dynamics of a common classical random walk. Here, by common we mean that each time a new direction needs to be chosen, we use the same probabilities ($p$ and $1 - p$) to decide between right and left. Quantum mechanically to allow the parameters to depend on $j$ (cf, equations (5) and (12)) implies that we explicitly assume position-dependent probability amplitudes. Obviously, by making such parameters $j$-independent, we recover the relation with the usual classical case.

Now, suppose that we shall determine which is the probability $P^{(j)}(n)$ to be in the ‘position’ state $j$ (which means a site (bond) state in the coined (scattering) model) at time $n$, regardless of the value of the coin (direction) quantum number $\sigma$. We then define

$$
\mathcal{P}_c^{(j)} = |j\rangle \langle j| \otimes \sum_\sigma |\sigma\rangle \langle \sigma|,
$$

$$
\mathcal{P}_s^{(j)} = \sum_\sigma \left| j + \frac{\sigma - 1}{2}, \sigma \right\rangle \left\langle j + \frac{\sigma - 1}{2}, \sigma \right|,
$$

respectively, the coin and scattering position projector operators. The sought probability is the expected value

$$
P^{(j)}(n) = \langle \Psi(n) | \mathcal{P}^{(j)}(n) | \Psi(n) \rangle
$$

for $\mathcal{P}^{(j)}$, one of the expressions in equation (13).

Projection is thus an essential ingredient in defining a QW model, which differs from classical walk systems by the typical interference effects due to equation (14) (see, e.g., [51]).

2.4. The unitary equivalence of the two 1D quantum walk models

Finally, in the present 1D topology, the unitary equivalence between CQWs and SQWs is straightforwardly established.

First, note the one-to-one correspondence between their full Hilbert spaces, $L^2(\mathbb{Z}) \otimes L^2(\mathbb{Z}_2) \equiv L^2(\mathbb{Z} \times \mathbb{Z}_2)$. So, it follows directly the existence of an isomorphic unitary operator [8] $E : \mathcal{H} \rightarrow \mathcal{H}$, given by

$$
E |\sigma, j\rangle = |j\rangle \otimes |\sigma\rangle,
$$

$$
E^\dagger |j\rangle \otimes |\sigma\rangle = |\sigma, j\rangle.
$$

Second, (i) inspecting equations (3) and (10), (ii) considering the most general conditions for the models quantum coefficients, represented by equations (2) and (11) (observe that equations (5) and (12), more usual in the literature, already obey particular conventions) and (iii) taking into account equation (15), one finds that the following coefficients play complete similar roles

$$
c^{(j)}_{\sigma \sigma} \leftrightarrow t^{(j)}_{\sigma}\quad \text{and} \quad c^{(j)}_{-\sigma \sigma} \leftrightarrow r^{(j)}_{\sigma}.
$$

Hence, setting $c^{(j)}_{\sigma \sigma} = t^{(j)}_{\sigma}$ and $c^{(j)}_{-\sigma \sigma} = r^{(j)}_{\sigma}$ for all $j$, we have in both models exactly the same probability amplitudes for their time evolutions.

Third, from equations (3), (10), (15) and assuming the above equalities, the resulting dynamics are unitary equivalent once

$$
U_j = E^\dagger U_c E.
$$
As is known [8], even when different QW models are unitary equivalent, the associated probabilities—obtained through direct projections—can be distinct. This is so because we choose contrasting physical representations, sites (coined) and bonds (scattering), to describe the problem. The specific states which characterize one of these ‘spatial’ configurations are not akin to the states for the other. For instance, the two states corresponding to the site \( j \) in CQWs, \( |j\rangle \otimes |\sigma = \pm\rangle \), are mapped to states at different bonds in SQWs. Therefore, the probability to be in a unique site is not equal to the probability to be in a unique bond (cf, equations (13) and (14)). Mathematically, it is related to the fact that considering equations (13) and (15), we obtain \( P_s^{(j)} \neq E\dagger P_c^{(j)} E \) and \( P_c^{(j)} \neq E P_s^{(j)} E\dagger \) (inequalities holding true in any topology given the results in [41]). Thus, for \( |\Psi_1\rangle_s \) and \( |\Psi_1\rangle_c = E |\Psi_1\rangle_s \), we have from equation (14) that generally \( P_s^{(j)} \neq P_c^{(j)} \).

Nevertheless, due to the QWs unitary equivalence we can recover the probabilities from each other version by constructing proper cross projector operators. Indeed, defining

\[
P_s^{(j)}|c\rangle = E\dagger P_c^{(j)} E, \quad P_c^{(j)}|s\rangle = E P_s^{(j)} E\dagger, \tag{18}
\]

one readily obtains the probabilities for the coined (scattering) model by applying \( P_s^{(j)}|c\rangle \) to the state \( |\Psi(n)\rangle_s |\Psi(n)\rangle_c \), evolved according to the scattering (coined) formulation.

The numerical examples for 2D lattices in section 3 will clearly illustrate all the above observations.

3. More general topologies: two illustrative examples

As already mentioned, it has been proven in [41] that CQWs and SQWs are always related by an unitary transformation. In the previous section, profiting from its relative simplicity (although considering the most general situation, hence extending the results of [8]), we have explicitly illustrated so for the 1D case.

However, the benefits in being able to map CQWs and SQWs become really evident when one considers more complex topologies. Thus, we next present a detailed analysis of the correspondence between CQWs and SQWs in two particular, but very instructive, 2D examples, square and honeycomb lattices.

3.1. Quantum walks on a square lattice

The square lattice, represented in figure 3, is the natural 2D extension of the 1D topology. In the following, we describe in such a case the coined and scattering QW formulations, as well as their unitary equivalence.

3.1.1. The coined formulation. In figure 3(a), we schematically represent the coined Hilbert lattice, whose basis states are \( \{|j\rangle \otimes |k\rangle \otimes |\sigma\rangle\} \), with \( \sigma = 1, 2, 3, 4 \) and \( j, k = 0, \pm 1, \pm 2, \ldots \). For notational simplicity, we do not write the ‘spatial’ states in the form \( |j\rangle_x \otimes |k\rangle_y \) (and operators as \( A_x \otimes B_y \)) to distinguish between the two distinct 1D dimensions. Also, we assume the natural ordering convention \( \langle A \otimes B \otimes C |\alpha\rangle \otimes |\beta\rangle \otimes |\gamma\rangle = A|\alpha\rangle \otimes B|\beta\rangle \otimes C|\gamma\rangle \).

For the inner coin states \( |\sigma\rangle \), written as

\[
|1\rangle = \begin{pmatrix} 0 \\ 1 \\ 0 \\ 0 \end{pmatrix}, \quad |3\rangle = \begin{pmatrix} 1 \\ 0 \\ 0 \\ 0 \end{pmatrix}, \quad |2\rangle = \begin{pmatrix} 0 \\ 0 \\ 0 \\ 1 \end{pmatrix}, \quad |4\rangle = \begin{pmatrix} 0 \\ 0 \\ 1 \\ 0 \end{pmatrix}, \tag{19}
\]
Figure 3. A QW on a square lattice. (a) At each site \((j, k)\), the arrows schematically represent the four inner states, \(\sigma = 1, 2, 3, 4\), of the coined formulation. (b) For the scattering formulation, it is shown the four ‘incoming’ states propagating to the lattice sites marked in (a). The two possible states defined on the bond common to the regions A and B (A and C) are \(|++\rangle, (j, k)\rangle\) and \(|−−\rangle, (j, k−1)\rangle\) and \(|−+\rangle, (j, k)\rangle\) and \(|+−\rangle, (j, k−1)\rangle\).

the coin operator matrix (at each \((j, k)\)) reads

\[
C^{(j,k)} = \begin{pmatrix}
  c_{13}^{(j,k)} & c_{31}^{(j,k)} & c_{41}^{(j,k)} & c_{21}^{(j,k)} \\
  c_{14}^{(j,k)} & c_{34}^{(j,k)} & c_{44}^{(j,k)} & c_{24}^{(j,k)} \\
  c_{12}^{(j,k)} & c_{32}^{(j,k)} & c_{42}^{(j,k)} & c_{22}^{(j,k)} \\
  c_{11}^{(j,k)} & c_{31}^{(j,k)} & c_{41}^{(j,k)} & c_{21}^{(j,k)}
\end{pmatrix}.
\]

(20)

The unitarity of \(C\), directly extending the relations in equation (2), leads to

\[
\sum_l c_{il}^{(j,k)} c_{jm}^{(j,k)*} = \sum_l c_{il}^{(j,k)*} c_{jm}^{(j,k)} = \delta_{im}.
\]

(21)

As the one-step time evolution, there are two common possibilities. The first is the so-called natural choice, since single steps displacements follow the natural topology of the lattice [52], i.e. from a given site representing a state, it can go either to right, left, up or down [7, 53] (figure 4(a)). The second is the diagonal version [54]. In this case, single steps are those in which ‘moves’ take place along the diagonals of the Hilbert lattice (figure 4(b)).
Figure 4. Schematics of the one-step time evolution for the coined model in a square lattice in the natural (a) and the diagonal (b) cases. The system, initially at the state represented by the site \( \times \), in the first step can reach the states represented by the sites ■. In the second step, the states which can be visited are those represented by the sites ■ and \( \times \). The diagonal version (dotted lattices) can be thought as the natural one rotated by \( \pi/4 \) and rescaled by a factor \( \sqrt{2} \).

Let us define (for \( S \) as in section 2.1)

\[
N = S \otimes I \otimes |1\rangle \langle 1| + S^\dagger \otimes I \otimes |3\rangle \langle 3| + I \otimes S \otimes |2\rangle \langle 2| + I \otimes S^\dagger \otimes |4\rangle \langle 4|, \quad (22)
\]

with \( I \) being the identity operator. The time evolution for the natural version, an extension of equation (1), yields

\[
U_c = N \times \left( \sum_{j,k} |j\rangle \langle j| \otimes |k\rangle \langle k| \otimes C^{(j,k)} \right). \quad (23)
\]

For the diagonal version, we first define the operator

\[
D = I \otimes S \otimes |1\rangle \langle 1| + I \otimes S^\dagger \otimes |3\rangle \langle 3| + S^\dagger \otimes I \otimes |2\rangle \langle 2| + S \otimes I \otimes |4\rangle \langle 4|. \quad (24)
\]
So, the time evolution is given by
\[ U_{c \text{- diagonal}} = DU_c. \] (25)
Equation (25), with \( D \) as in equation (24), has a very simple interpretation (if we maintain that equation (25) represents a single step). Indeed, the diagonal case is essentially the natural one rotated by \( \pi/4 \) (anti-clockwise) in the Hilbert lattice. Moreover, quantities associated with any norm (length scale) defined on the lattice, e.g., diffusiveness, should be re-scaled by a factor \( \sqrt{2} \), which is just the ratio between the diagonal and side of the unitary cell, figure 4. In all our further analysis, we will consider only the natural version, observing that the diagonal would easily follow from equation (25).

Finally, for the natural coined model the projector operator is
\[ P^{(j,k)} = \left| j \right\rangle \left\langle j \right| \otimes \left| k \right\rangle \otimes \sum_{\sigma=1}^{4} \left| \sigma \right\rangle \left\langle \sigma \right| \] (26)

3.1.2. The scattering model. For \( \sigma_x, \sigma_y = \pm 1 \) (± for short) and \( j, k = 0, \pm 1, \pm 2, \ldots \), the scattering model basis states are \( \{|\sigma_x, \sigma_y, (j,k)\rangle\rangle \), figure 3(b). The time evolution is again \( U_j = T + R \), with
\[ T|\sigma_x, \sigma_y, (j,k)\rangle = \sum_{[\alpha, \beta] \neq [-\sigma_y, -\sigma_x]} t^{(j,k)}_{[\alpha, \beta], \sigma_x, \sigma_y} |\alpha, \beta, \left( j + \alpha \sigma_x + \beta \sigma_y \right) / 2, k + \beta \sigma_y - \alpha \sigma_x / 2 \rangle \], \[ R|\sigma_x, \sigma_y, (j,k)\rangle = r^{(j,k)}_{[-\sigma_x, -\sigma_y, \sigma_x, \sigma_y]} |\sigma_x - \sigma_y, (j + \sigma_x | \sigma_x + \sigma_y | / 2, k - \sigma_y | \sigma_x + \sigma_y | / 2 \rangle \]. (27)
The corresponding actions of \( T^\dagger \) and \( R^\dagger \) are a straightforward generalization of equation (8), in a format similar to equation (27). Defining
\[ \Gamma^{(j,k)} = \begin{pmatrix} t^{(j,k)}_{++}, & -r^{(j,k)}_{++}, & t^{(j,k)}_{+-}, & -r^{(j,k)}_{+-} \\ r^{(j,k)}_{++}, & +t^{(j,k)}_{++}, & r^{(j,k)}_{+-}, & +t^{(j,k)}_{+-} \\ t^{(j,k)}_{-+}, & -r^{(j,k)}_{-+}, & t^{(j,k)}_{-,+}, & -r^{(j,k)}_{-,+} \\ r^{(j,k)}_{-+}, & +t^{(j,k)}_{-+}, & r^{(j,k)}_{-,+}, & +t^{(j,k)}_{-,+} \end{pmatrix}, \] (28)
the unitarity of \( U_j \) implies in the unitarity of the scattering matrices \( \Gamma^{(j,k)} \) (once more a direct extension of previous results, in this case of equation (11)). So, the elements of \( \Gamma^{(j,k)} \) must satisfy relations completely analogous to those in equation (21).

For the projector operators, we need to distinguish between the horizontal \( (\sigma_x \times \sigma_y = +) \) and the vertical \( (\sigma_x \times \sigma_y = -) \) bonds. Hence, we define
\[ P^{(j,k; +)} = \sum_{\sigma_x} |\sigma_x, \sigma_x, (j + \sigma_x / 2, k)\rangle \left\langle (j + \sigma_x / 2, k) \right|, \sigma_x, \sigma_x \], \[ P^{(j,k; -)} = \sum_{\sigma_x} |\sigma_x, \sigma_x, (j, k + \sigma_x / 2)\rangle \left\langle (j, k + \sigma_x / 2) \right|, -\sigma_x, -\sigma_x \]. (29)

3.1.3. The mapping. To map the two models, we first make the following identification between the scattering directions and the inner coin quantum numbers \( (\sigma_x, \sigma_y \leftrightarrow \sigma) \):
\[ + + \leftrightarrow 1, -+ \leftrightarrow 2, -- \leftrightarrow 3, ++ \leftrightarrow 4, \] (30)
which can be cast as \( \sigma = (5 - (2 + \sigma_x) \sigma_y) / 2 \).
Using the procedure in the last section (or likewise, the rigorous construction in [41]), i.e. to associate the scattering states ‘incoming’ to a certain site (see figure 3(b)) with the coin states at that site, we can set \( E \) as (taking into account equation (30))

\[
E |\sigma, \sigma_j (j, k)\rangle = |j\rangle \otimes |k\rangle \otimes \left\{ \frac{5}{2} - \frac{(2 + \sigma_j)\sigma_j}{2} \right\}.
\]

(31)

Furthermore, assuming the coefficients in equation (20) equal to those in equation (28) (which is consistent with the relation in equation (30)), again we find that the two models are unitary equivalent, since for equations (23), (27) and (31), the relation in equation (17) holds.

To illustrate the above general constructions, we analyze some particular cases for the probability amplitudes (see, for instance, [7, 52, 53]). For simplicity, we suppose all the coin (and therefore the scattering) matrices to be independent on the quantum numbers \( j \) and \( k \). We choose the following four coin operators (whose corresponding scattering matrices are written in exactly the same form):

(a) \( x-y \) decoupled Hadamard

\[
C_{\text{HdH}} = \left( \begin{array}{cc} H_x & 0 \\ 0 & H_x \end{array} \right) = \frac{1}{\sqrt{2}} \left( \begin{array}{cccc} +1 & +1 & 0 & 0 \\ +1 & -1 & 0 & 0 \\ 0 & 0 & +1 & +1 \\ 0 & 0 & +1 & -1 \end{array} \right);
\]

(32)

(b) full \( 4 \times 4 \) Hadamard

\[
C_{\text{H}} = \frac{1}{2} \left( \begin{array}{cccc} +1 & +1 & +1 & +1 \\ +1 & -1 & +1 & -1 \\ +1 & +1 & -1 & -1 \\ +1 & -1 & -1 & +1 \end{array} \right);
\]

(33)

(c) \( 4 \times 4 \) Grover

\[
C_{\text{G}} = \frac{1}{2} \left( \begin{array}{cccc} -1 & +1 & +1 & +1 \\ +1 & -1 & +1 & +1 \\ +1 & -1 & -1 & -1 \\ +1 & -1 & +1 & -1 \end{array} \right);
\]

(34)

(d) \( 4 \times 4 \) DFT

\[
C_{\text{DFT}} = \frac{1}{2} \left( \begin{array}{cccc} +1 & +1 & +1 & +1 \\ +1 & +i & -i & -i \\ +1 & -i & +i & -i \\ +1 & -i & -i & +i \end{array} \right).
\]

(35)

We compute \( U^n |\Psi(0)\rangle \), with \( n = 20 \), for both QW models and the above matrices. For each \((j, k)\), we use proper projectors to calculate the probability \( P^{(j, k)}(n = 20) = \langle \Psi(20)|P^{(j, k)}|\Psi(20)\rangle \). As \( |\Psi(0)\rangle \), we take

\[
|\Psi(0)\rangle = \frac{1}{2} [1| + + (0, 0)\rangle + i| - (0, 0)\rangle + | - + (0, 0)\rangle + i| + - (0, 0)\rangle],
\]

\[
|\Psi(0)\rangle = E |\Psi(0)\rangle = \frac{1}{2} |0\rangle \otimes |0\rangle \otimes [|1\rangle + i|3\rangle + |2\rangle + i|4\rangle].
\]

(36)

The \( P^{(j, k)} \) are displayed as 3D density plots. So, in each graph the heights on the z-axis correspond to the probabilities values. Moreover, here and for the honeycomb lattice in
Figure 5. Quantum walks in the square lattice considering the decoupled Hadamard transition coefficients, equation (32). The density plots represent the probabilities to be in the different states (as defined by the projector operators) after \( n = 20 \) time steps for \( |\Psi(0)\rangle \) given in the main text. The results are for the: (a) coined, (b) scattering obtained from the coined, (c) scattering and (d) coined obtained from the scattering models. The graphs (a) and (d), and (b) and (c) are completely identical.

section 3.2, each pair of labels \( j \) and \( k \) in the \( x-y \) plane indicates the \((j,k)\) lattice site spatial location. Thus, for the coined formulation, the \( P(j,k) \) are marked just over the sites. For the scattering formulation, since the states are defined along the bonds, we mark the \( P \) exactly over the middle points of the corresponding bonds. Hence, we show the probability patterns of each QW version in its own state representation, but in a way which makes it easy to qualitatively compare the two models.

The results for the decoupled Hadamard, full Hadamard, Grove and DFT operators are presented, respectively, in figures 5–8. The plots are organized as follows. For the coined formulation, figures 5–8(a) [(b)] show the probabilities obtained from the projectors in equation (26) [the projectors \( P(j,k) \)]. On the other hand, for the scattering formulation, figures 5–8(c) [(d)] show the \( P(j,k) \) from the projectors in equation (29) [the projectors \( P_{s}(j,k) \)].

By inspecting figures 5–8, some points become evident. (i) Due to the decoupled form of the transition amplitude matrix, equation (32), in figure 5 we actually have two independent 1D evolutions. Hence, each direction is a good example of the 1D results in section 2. (ii) Despite the fact that the two QW versions are completely equivalent by unitary transformations, they lead to distinct probability patterns (compare (a) or (d) with (b) or (c)). Indeed, as already emphasized (for instance, the discussion just after equation (17) in section 2.4), the one-to-one
Figure 6. The same as in figure 5, but for the $4 \times 4$ Hadamard, equation (33).

correspondence between the two formulations pinpoints a dynamical similarity. However, the states in each case represent different (although correlated) aspects of the Hilbert lattice; thus, the distributions of the $P$ do not need to coincide. (iii) Moreover, such patterns display richer structures in the scattering case. To understand so, assume the simpler 1D lattice, for which the initial state $|+\rangle_0 = |0\rangle \otimes |+\rangle$ evolves one time step. For the SQW (CQW), we obtain

$$|\Psi_1(1)\rangle = r(0)|-\rangle + t(0)|+\rangle$$

and

$$|\Psi(1)\rangle = c(0)_{-}|-\rangle \otimes |+\rangle + c(0)_{+}|+\rangle \otimes |+\rangle.$$ 

Considering the Hilbert lattice picture, note that for the former we have two ‘neighbor’ bond states in the expansion of $|\Psi_1(1)\rangle$, those ‘attached’ to the site 0. On the other hand, for the latter the site states composing $|\Psi(1)\rangle$ are $j = -1$ and $j = +1$, but not $j = 0$. It illustrates a very typical situation in any topology, namely the SQW dynamics tends to excite ‘contiguous’ spatial (bond) basis states, whereas CQWs may skip some ‘successive’ (site) basis states. Thus, interference [51] is usually more recurrent in SQWs than in CQWs (due to this difference in the spreading of $|\Psi(1)\rangle$), explaining the behavior observed in the plots. (iv) Nevertheless, we can recover the $P^{(j,k)}$ from each other model by correctly defining the cross projectors (e.g., SQWs in (b) from CQWs in (a) and CQWs in (d) from SQWs in (c)), since the information about one model is always encoded in the other.

We finally mention that we have analyzed the diagonal CQW and its corresponding scattering version. We have obtained one formulation $P$ from the other by correctly defining the cross projectors. Furthermore, as expected the probabilities plots (not shown) are exactly the ones here, just rotated by $\pi/4$ and rescaled by a factor $\sqrt{2}$.
3.2. Quantum walks on a honeycomb lattice

Our last example is a QW defined on a honeycomb lattice, whose structure is depicted in figure 9(a). It is far more involving than the previous square case mainly because now the coordination number, equal to 3, is odd [47] (for coordination 3 in 1D see, e.g., [55]). We note that although this topology was recently investigated in the context of continuous time models [56], and few discrete time implementations do exist for torus-like boundary conditions [57] and for a similar three-state QW [58] (but not in a truly honeycomb geometry), the present is the first general treatment for the problem.

The labeling of the sites in both formulations is indicated in figure 9(b). Without loss of generality, we adopt the convention: in the $y$-direction the quantum number $k$ is chosen such that if $k$ is even, then the corresponding infinite row of sites have the bonds along north (up), southeast and southwest. On the other hand, for $k$ odd, the bonds configuration at each site is south (down), northeast and northwest.

An interesting aspect of the honeycomb lattice, consequence of how it imposes the relation between the quantum number $\sigma$ and directions, is that any sequence

$$(j_0, k_0)_\sigma \xrightarrow{U} (j_1, k_1)_\sigma \xrightarrow{U} (j_2, k_2)_\sigma \ldots \xrightarrow{U} (j_{N-1}, k_{N-1})_\sigma \xrightarrow{U} (j_N, k_N)_\sigma; \quad (37)$$

i.e. a particular $N$ steps evolution for which the value of $\sigma$ remains the same, is not naturally (i.e. necessarily) a ballistic-like trajectory. By natural we mean those cases (like regular lattices with an even coordination number, e.g., our square lattice) where we always can associate
quantum states and directions such that $\sigma \rightarrow \sigma$ leads to evolution along straight lines. In such cases, the above dynamics, equation (37), would be ballistic in the Hilbert lattice space regardless the specific $\sigma$. As we are going to see for the construction adopted here, successive transitions $\sigma \rightarrow \sigma$ yield a return to a same site (in a round trip) after visiting six sites. Hence, diffusion throughout the lattice implies transitions of the type $\sigma \rightarrow \sigma' \neq \sigma$.

3.2.1. The coined model. For the inner coin states $|\sigma\rangle$, we have three possibilities, namely $\sigma = 0, 1, 2$ (we also could call them 1, 2, 3, but the use of 0 instead of 3 simplifies the notation). Due to the honeycomb particular topology, some care is necessary in defining the dynamics in terms of the coin variable. Indeed, states with the same $\sigma$ but at different sites do not always evolve to the same direction. So, to properly associate the $\sigma$ with the system evolution under $U$, we consider the following prescription (which, however, is not the only one possible\(^3\)). Let $[x]_N \equiv x \mod N$, i.e. $[x]_N$ is the remainder of $x/N$. Then, for a given site $(j, k)$, the quantum numbers 0, 1, 2 are associated with directions in the lattice as indicated in figure 9(c). An example of the resulting configuration is displayed in figure 10 (in particular, note from figure 10 that leaving from a certain site, say $(j, k)$, and always evolving to a same $\sigma$, mandatorily will get back to $(j, k)$ in exactly six steps). The rules in figure 9(c) establish in a unique and self-consistent way—for the entire lattice—how single steps $(j, k) \rightarrow (j', k')$ are determined by the values of $\sigma$.

\(^3\) A full classification of equivalent constructions for the honeycomb lattice will appear elsewhere.
Figure 9. (a) The honeycomb lattice and (b) the convention for the sites labeling. (c) For the coined formulation, the association between the quantum number $\sigma$ and directions. Here, $[x]_3$ denotes $x \mod 3$, i.e. the remainder of $x/3$.

Figure 10. An illustration of the resulting association between coin quantum numbers and directions according to the rules in figure 9(c). Here $k$ is even and $[j]_3 = 0$.

The basis states are written as $|(j, k)\rangle \otimes |\sigma\rangle$, with a column vector representation for $|\sigma\rangle$ given by

$$
|0\rangle = \begin{pmatrix} 1 \\ 0 \\ 0 \end{pmatrix}, \quad |1\rangle = \begin{pmatrix} 0 \\ 1 \\ 0 \end{pmatrix}, \quad |2\rangle = \begin{pmatrix} 0 \\ 0 \\ 1 \end{pmatrix}.
$$

(38)
Figure 11. For the honeycomb lattice, the SQW basis states labeling rules. The \( \sigma \) values here (\( \sigma_a \), \( \sigma_b \) and \( \sigma_c \)) are defined in equation (44). The upper details summarize the convention in figure 9(c) to label the CQW inner states.

So, the coin operator at any site \((j, k)\) is

\[
C^{(j,k)} = \begin{pmatrix}
  c_{00}^{(j,k)} & c_{01}^{(j,k)} & c_{02}^{(j,k)} \\
  c_{10}^{(j,k)} & c_{11}^{(j,k)} & c_{12}^{(j,k)} \\
  c_{20}^{(j,k)} & c_{21}^{(j,k)} & c_{22}^{(j,k)}
\end{pmatrix},
\]

(39)

which we suppose to be an unitary matrix.

Assuming the above construction, after a little lengthy but straightforward analyses one finds that the one-step time evolution operator for the coined formulation reads

\[
U_c = \left( \sum_{\sigma=0}^{2} S_\sigma \otimes |\sigma\rangle\langle\sigma| \right) \times \left( \sum_{j,k} |(j,k)\rangle\langle(j,k)| \otimes C^{(j,k)} \right),
\]

(40)

where

\[
S_\sigma |(j,k)\rangle = |(f(j,k;\sigma), g(j,k;\sigma))\rangle,
\]

\[
S_\sigma^\dagger |(j,k)\rangle = |(f(j,k;\phi_k(\sigma)), g(j,k;\phi_k(\sigma)))\rangle,
\]

(41)

for (with \( \text{sgn}[x] = x/|x|, \) if \( x \neq 0, \) and \( \text{sgn}[0] = 0 \))

\[
f(j,k;\sigma) = j + (-1)^{\sigma-\lceil j+k \rceil}\text{sgn}[\sigma - \lceil j+k \rceil],
\]

\[
g(j,k;\sigma) = k + (-1)^{\sigma} (1 - 2\text{sgn}[\sigma - \lceil j+k \rceil]),
\]

\[
\phi_k(\sigma) = [\sigma - (-1)^{\lceil j+k \rceil}].
\]

(42)

By construction \( S_\sigma S_\sigma^\dagger = S_\sigma^\dagger S_\sigma = I \), leading to a unitary \( U_c \).

Finally, the corresponding projector operator is

\[
\mathcal{P}_c^{(j,k)} = |(j,k)\rangle\langle (j,k)| \otimes \sum_{\sigma=0}^{2} |\sigma\rangle\langle\sigma|.
\]

(43)

3.2.2. The scattering model. For the scattering formulation, we consider the schematics in figure 11. We denote all the basis state incoming to the site \((j,k)\) (continuous arrows in figure 11) by \(|\sigma, (j,k)\rangle\). Since now the lattice coordination is 3, we need more than the usual \( \pm \) to label the direction quantum number \( \sigma \). Therefore, we assume that \( \sigma \) has the values 0, 1, 2, in analogy with the coined model. But in each bond we can ascribe only two of these possible values to the two states representing opposite directions. In this way, for the general situation
Figure 12. Quantum walks in the honeycomb lattice considering the $3 \times 3$ typical unbiased transition coefficients, equation (51). The density plots represent the probabilities to be in the different states (as defined by the projector operators) after $n = 20$ time steps for $|\Psi(0)\rangle$ given in the main text. The results are for the: (a) coined, (b) scattering obtained from the coined, (c) scattering, and (d) coined obtained form the scattering, models. The graphs (a) and (d), and (b) and (c) are completely identical.

At each bond, for a state $|\sigma,j,k\rangle$ incoming to $(j,k)$ there is a corresponding outgoing state, which is itself an incoming state to another site $(j',k')$ (dashed arrows in figure 11). From the protocol used to label the $\sigma$—figure 11 and equation (44)—it is not difficult to show that $\sigma' = \phi_k(\sigma)$, $j' = f(j,k;\phi_k(\sigma))$ and $k' = g(j,k;\phi_k(\sigma))$, for $f$, $g$ and $\phi$ given in equation (42). For instance, in figure 11 for $k$ even and $[j]_3 = 0$, $\sigma_a = 0$ and the other state in the same bond as $|0, (j, k)\rangle$ is $|2, (j + 1, k - 1)\rangle$, as verified by direct inspection. Likewise, this follows directly from $\sigma' = \phi_{k_{\text{even}}}(0) = 2$, $j' = f(j,k;2)|k_{\text{even}},[j]_3 = 0 = j + 1$ and $k' = g(j,k;2)|k_{\text{even}},[j]_3 = 0 = k - 1$.

Thus, the evolution operator is $U_s = T + R$, with

$$T|\sigma,j,k\rangle = \sum_{\alpha \neq \sigma} \langle j,k|\phi_{\alpha}(\sigma), f(j,k;\phi_{\alpha}(\sigma)), g(j,k;\phi_{\alpha}(\sigma))\rangle,$$

$$R|\sigma,j,k\rangle = r_{\phi_{\alpha}(\sigma),\sigma} |\phi_{\alpha}(\sigma), f(j,k;\phi_{\alpha}(\sigma)), g(j,k;\phi_{\alpha}(\sigma))\rangle,$$

(45)
Figure 13. The same as in figure 12, but for the $3 \times 3$ biased real matrix, equation (52).

where furthermore

$$T^\dagger |\sigma, (j, k)\rangle = \sum_{\alpha \neq \phi_k(\sigma)} [t^{(f(j,k;\phi_k(\sigma)),g(j,k;\phi_k(\sigma)))}]^* |\alpha, (f(j,k;\phi_k(\sigma)), g(j,k;\phi_k(\sigma)))\rangle,$$

$$R^\dagger |\sigma, (j, k)\rangle = [r^{(f(j,k;\phi_k(\sigma)),g(j,k;\phi_k(\sigma)))}]^* |\phi_k(\sigma), (f(j,k;\phi_k(\sigma)), g(j,k;\phi_k(\sigma)))\rangle.$$

Now, if we define

$$\Gamma^{(j,k)} = \begin{pmatrix} \Gamma_{00}^{(j,k)} & \Gamma_{01}^{(j,k)} & \Gamma_{02}^{(j,k)} \\ \Gamma_{10}^{(j,k)} & \Gamma_{11}^{(j,k)} & \Gamma_{12}^{(j,k)} \\ \Gamma_{20}^{(j,k)} & \Gamma_{21}^{(j,k)} & \Gamma_{22}^{(j,k)} \end{pmatrix},$$

and identify $(\alpha, \beta = 0, 1, 2)$

$$r_{\phi_k(\sigma), \sigma} = \Gamma_{\phi_k(\sigma), \sigma}^{(j,k)}, \quad t_{\beta, \alpha} = \Gamma_{\beta, \alpha}^{(j,k)} \quad \text{(for } \beta \neq \phi_k(\alpha)\text{)},$$

then one has that the unitarity of $\Gamma$ in equation (47) guarantees that $U_s$ is also unitary.
Finally, the projector operator reads

$$P_{\sigma, (j, k)} = |\phi_k(\sigma), f(j, k; \phi_k(\sigma)), g(j, k; \phi_k(\sigma))\rangle \langle (f(j, k; \phi_k(\sigma)), g(j, k; \phi_k(\sigma))), \phi_k(\sigma)|,$$

$$+ |\sigma, (j, k)\rangle \langle (j, k), \sigma|.$$ (49)
analyze the following five different examples for the transition probabilities (assumed to be the same at all the lattice sites):

(a) $3 \times 3$ typical unbiased operator

$$C_{\text{unb}} = 1 \sqrt[3]{-1 \exp[-\pi i/3]} \exp[-\pi i/3] \exp[-\pi i/3] \exp[-\pi i/3] \exp[-\pi i/3] \exp[-\pi i/3] ;$$

(b) $3 \times 3$ biased (and real) operator

$$C_{\text{bias}} = \frac{1}{3} \begin{pmatrix} 1 & 1 - \sqrt{3} & 1 + \sqrt{3} \\ 1 + \sqrt{3} & 1 & 1 - \sqrt{3} \\ 1 - \sqrt{3} & 1 + \sqrt{3} & 1 \end{pmatrix} ;$$

(c) $3 \times 3$ DHT [59]

$$C_{\text{DHT}} = \frac{1}{2\sqrt{3}} \begin{pmatrix} 2 & 2 & 2 \\ 2 & -1 + \sqrt{3} & -1 - \sqrt{3} \\ 2 & -1 - \sqrt{3} & -1 + \sqrt{3} \end{pmatrix} ;$$

(d) $3 \times 3$ Grover

$$C_{G} = \frac{1}{3} \begin{pmatrix} -1 & 2 & 2 \\ 2 & -1 & 2 \\ 2 & 2 & -1 \end{pmatrix} ;$$
Figure 16. The same as in figure 12, but for the 3 × 3 DFT, equation (55).

(e) 3 × 3 DFT

\[ C_{\text{DFT}}^{3} = \frac{1}{\sqrt{3}} \begin{pmatrix} \exp[2\pi i/3] & 1 & \exp[-2\pi i/3] \\ 1 & \exp[-2\pi i/3] & 1 \\ \exp[2\pi i/3] & 1 & \exp[2\pi i/3] \end{pmatrix} . \]  

(55)

Few comments about the matrices above are in order. The first case, equation (51), is a typical (complex) unbiased operator since the coefficients have the same value for their modulus square, \(|c_{\sigma'}\sigma|^2 = 1/3\). Therefore, the resulting transition probabilities are equal to 1/3. On the other hand, the second, equation (52)—here with all entries real numbers—is strongly biased because the transition probabilities are very different (but obviously summing up to 1). For instance, for the scattering formulation, \(k\)-even and \([j]_3 = 0\) in figure 11 (cf equation (48)), we have \(|t_{0,0}|^2 = 1/9 \approx 0.11\), \(|t_{1,0}|^2 = (2 + \sqrt{3})/9 \approx 0.83\), \(|r_{2,0}|^2 = (2 - \sqrt{3})/9 \approx 0.06\).

The discrete Hartley transform matrix [59], equation (53), although not usually considered in QWs, is an interesting example to study due to its usefulness in signal processing [60], furthermore always being real. The other two are just the 3 × 3 versions of the Grover and DFT. Finally, note there is not a 3 × 3 Hadamard matrix.

Assuming as the initial states

\[ |\Psi(0)\rangle_s = |1, (0, 0)\rangle, \quad E |\Psi(0)\rangle_c = |0, (0, 0)\rangle \otimes |1\rangle, \]  

(56)

we evolve the models 20 time steps. The resulting probability patterns are displayed in figures 12–16. The graphs are organized exactly as in section 3.1.4 (figures 5–8).
Again, certain facts are clearly observed from figures 12–16. First, as it should be, we can obtain one evolution from the other by correct projections. Second, like in the square lattice, in the honeycomb the CQW probabilities are in general more sparse and somewhat smoother than those for the SQW. And third, it is interesting to notice the particular pattern in figure 13, with a tendency of three preferable directions of propagation along the lattice. This is a consequence of the probabilities bias resulting from the matrix \( C_{\text{bias}} \), equation (52), for which three transition elements \( c_{02} = c_{10} = c_{21} = (1 + \sqrt{3})/3 \) are considerable higher than the other six.

Finally, comparing the Grover (DFT) probability amplitudes for the square and honeycomb topologies, respectively, figures 7 and 15 (figures 8 and 16) we can have an idea of the influence of a regular lattice coordination number to the QW dynamics. In fact, note that the \( |\Psi(0)\rangle \) used for the square lattice examples is equally ‘distributed’ among the four \( \sigma \) values basis states, equation (36), which is not the case for the honeycomb lattice whose \( |\Psi(0)\rangle \) is written in terms of just one value of \( \sigma \) (from the three possible), equation (56). Even then, we see that in the latter the resulting probabilities are more uniformly distributed, specially in the Grover case. This illustrates the intricate process of quantum transitions at each lattice site as a function of the number of bonds attached to it.

4. Conclusion

By means of detailed analysis of specific examples, in this work we have compared the different aspects associated with the construction of the discrete time coined and scattering quantum walks. Specifically, we have presented a complete formulation for the line, square and the involving honeycomb lattice topologies. In all the examples studied, we have clarified how to map one model version to the other. Also, we have illustrated the particularities in the probability pattern distributions along the graph structures, resulting from the different time evolution of each formulation. Finally, distinct coin (and corresponding scattering) matrices operators were considered. Besides the usual Hadamard, Grover and discrete Fourier transform, we have addressed the discrete Hartley transform as well as a few other cases.

Our aim here was to show through concrete situations that in fact the CQWs and SQWs are closely related. Moreover, the use of a specific QW version in different applications may be more a matter of practicality (or even of taste, as recently stressed [14]) than due to prohibitive conceptual differences between the models.

Regarding implementations, the issue may be a little subtle considering that usually distinct experimental architectures are proposed for CQWs and SQWs (see the introduction section). However, a given realization of a specific QW model conceivably could also be used to obtain results from the other model. But then, the experimental setup somehow should allow the effective construction of proper cross projectors (as defined in sections 2.3 and 2.4). The important point is that efforts in this direction certainly would pay off since an actual implementation of one version of QWs having proper built-in projections to the other version—thus ‘carrying’ the features of both type of QWs—would be much more flexible and useful in applications.

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