Quantum Simulation of Dissipative Processes without Reservoir Engineering

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We present a quantum algorithm to simulate general finite dimensional Lindblad master equations without the requirement of engineering the system-environment interactions. The proposed method is able to simulate both Markovian and non-Markovian quantum dynamics. It consists in the quantum computation of the dissipative corrections to the unitary evolution of the system of interest, via the reconstruction of the response functions associated with the Lindblad operators. Our approach is equally applicable to dynamics generated by effectively non-Hermitian Hamiltonians. We confirm the quality of our method providing specific error bounds that quantify its accuracy.

While every physical system is indeed coupled to an environment¹,², modern quantum technologies have succeeded in isolating systems to an exquisite degree in a variety of platforms³–⁶. In this sense, the last decade has witnessed great advances in testing and controlling the quantum features of these systems, spurring the quest for the development of quantum simulators⁷–¹⁰. These efforts are guided by the early proposal of using a highly tunable quantum device to mimic the behavior of another quantum system of interest, being the latter complex enough to render its description by classical means intractable. By now, a series of proof-of-principle experiments have successfully demonstrated the basic tenets of quantum simulations revealing quantum technologies as trapped ions¹¹, ultracold quantum gases¹², and superconducting circuits¹³ as promising candidates to harbor quantum simulations beyond the computational capabilities of classical devices.

It was soon recognised that this endeavour should not be limited to simulating the dynamics of isolated complex quantum systems, but should more generally aim at the emulation of arbitrary physical processes, including the open quantum dynamics of a system coupled to an environment. Tailoring the complex nonequilibrium dynamics of an open system has the potential to uncover a plethora of technological and scientific applications. A remarkable instance results from the understanding of the role played by quantum effects in the open dynamics of photosynthetic processes in biological systems¹⁴,¹⁵, recently used in the design of artificial light-harvesting nanodevices¹⁶–¹⁸. At a more fundamental level, an open-dynamics quantum simulator would be invaluable to shed new light on core issues of foundations of physics, ranging from the quantum-to-classical transition and quantum measurement theory¹⁹ to the characterization of Markovian and non-Markovian systems²⁰–²². Further motivation arises at the forefront of quantum technologies. As the available resources increase, the verification with classical computers of quantum annealing devices²³,²⁴, possibly operating with a hybrid quantum-classical performance, becomes a daunting task. The comparison between different experimental implementations of quantum simulators is required to establish a confidence level, as customary with other quantum technologies.
Facing the high dimensionality of the Hilbert space of the composite system made of a quantum device embedded in an environment, recent developments have been focused on the reduced dynamics of the system that emerges after tracing out the environmental degrees of freedom. The resulting nonunitary dynamics is governed by a dynamical map, or equivalently, by a master equation\(^2\). In this respect, theoretical\(^{26-28}\) and experimental\(^{29}\) efforts in the simulation of open quantum systems have exploited the combination of coherent quantum operations with controlled dissipation. Notwithstanding, the experimental complexity required to simulate an arbitrary open quantum dynamics is recognised to substantially surpass that needed in the case of closed systems, where a smaller number of generators suffices to design a general time-evolution. Thus, the quantum simulation of open systems remains a challenging task.

In this Letter, we propose a quantum algorithm to simulate finite dimensional Lindblad master equations, corresponding to Markovian or non-Markovian processes. Our protocol shows how to reconstruct, up to an arbitrary finite error, physical observables that evolve according to a dissipative dynamics, by evaluating multi-time correlation functions of its Lindblad operators. We show that the latter requires the implementation of the unitary part of the dynamics in a quantum simulator, without the necessity of physically engineering the system-environment interactions. Moreover, we demonstrate how these multi-time correlation functions can be computed with a reduced number of measurements. We further show that our method can be applied as well to the simulation of processes associated with non-Hermitian Hamiltonians. Finally, we provide specific error bounds to estimate the accuracy of our approach.

Consider a quantum system coupled to an environment whose dynamics is described by the von Neumann equation \(i\hbar\frac{d\rho}{dt} = [H, \rho]\). Here, \(\rho\) is the system-environment density matrix, \(H = H_S + H_I\), where \(H_S\) and \(H_I\) are the system and environment Hamiltonians, while \(H_I\) corresponds to their interaction. Assuming weak coupling and short time-correlations between the system and the environment, after tracing out the environmental degrees of freedom we obtain the Markovian master equation

\[
\frac{d\rho}{dt} = \mathcal{L}^t\rho, \tag{1}
\]

being \(\rho = \text{Tr}_E(\rho)\) and \(\mathcal{L}^t\) the time-dependent superoperator governing the dissipative dynamics\(^1\). Notice that there are different ways to recover Eq. (1)\(^2\). Nevertheless, Eq. (1) is our starting point, and in the following we show how to simulate this equation regardless of its derivation. Indeed, our algorithm does not need to control any of the approximations done to achieve this equation. We can decompose \(\mathcal{L}^t\) into \(\mathcal{L}^t = \mathcal{L}_{\text{H}}^t + \mathcal{L}_{\text{D}}^t\). Here, \(\mathcal{L}_{\text{H}}^t\) corresponds to a unitary part, i.e. \(\mathcal{L}_{\text{H}}^t\rho \equiv -i[H(t), \rho]\), where \(H(t)\) is defined by \(H_S\) plus a term due to the lamb-shift effect and it may depend on time. Instead, \(\mathcal{L}_{\text{D}}^t\) is the dissipative contribution and it follows the Lindblad form\(^3\)

\[
\mathcal{L}_{\text{D}}^t\rho \equiv \sum_{i=1}^N \gamma_i(t) \left\{ L_i \rho L_i^\dagger - \frac{1}{2} \{L_i^\dagger L_i, \rho\} \right\},
\]

where \(L_i\) are the Lindblad operators modelling the effective interaction of the system with the bath that may depend on time, while \(\gamma_i(t)\) are nonnegative parameters. Notice that, although the standard derivation of Eq. (1) requires the Markov approximation, a non-Markovian equation can have the same form. Indeed, it is known that if \(\gamma_i(t) < 0\) for some \(t\) and \(\int_0^t dt' \gamma_i(t') > 0\) for all \(t\), then Eq. (1) corresponds to a completely positive non-Markovian channel\(^2\). Our approach can deal also with non-Markovian processes of this kind, keeping the same efficiency as the Markovian case. While we will consider the general case \(\gamma_i = \gamma_i(t)\), whose sign distinguishes the Markovian processes from the non-Markovian ones, for the sake of simplicity we will consider the case \(H = H(t)\) and \(L_i = L_i(t)\) (in the following, we will denote \(\mathcal{L}_{\text{H}}^t\) simply as \(\mathcal{L}_H\)). However, the inclusion in our formalism of time-dependent Hamiltonians and Lindblad operators is straightforward.

One can integrate Eq. (1) obtaining a Volterra equation\(^3\)

\[
\rho(t) = e^{i\mathcal{L}_H\rho(0)} + \int_0^t ds \ e^{i(t-s)\mathcal{L}_H} \mathcal{L}_{\text{D}}^s \rho(s), \tag{2}
\]

where \(e^{i\mathcal{L}_H\rho} \equiv \sum_{k=0}^{\infty} t^k \mathcal{L}_H^k / k!\). The first term at the right-hand-side of Eq. (2) corresponds to the unitary evolution of \(\rho(0)\) while the second term gives rise to the dissipative correction. Our goal is to find a perturbative expansion of Eq. (2) in the \(\mathcal{L}_D^t\) term, and to provide with a protocol to measure the resulting expression in a unitary way. In order to do so, we consider the iterated solution of Eq. (2) obtaining

\[
\rho(t) = \sum_{i=0}^{\infty} \rho_i(t). \tag{3}
\]

Here, \(\rho_i(t) = e^{i\mathcal{L}_H\rho(0)}\), while, for \(i \geq 1\), \(\rho_i(t)\) has the following general structure: \(\rho_i(t) = \int_i^{i+1} \Phi_j \ e^{i\mathcal{L}_H\rho} (0) \Phi_j^\dagger\), \(\Phi_j\) being a superoperator acting on an arbitrary matrix \(\xi\) as \(\Phi_j \xi = \int_0^{s_j} ds_j \ e^{i(s_j-j)\mathcal{L}_H} \mathcal{L}_{\text{D}}^{s_j} \xi\), where \(s_0 \equiv t\). For instance, \(\rho_2(t)\) can be written as
\[
\rho_2(t) = \Pi_{i=1}^{2} \Phi_i e^{i\xi^2 t \epsilon^2 \mathbf{u}} \rho(0) = \Phi_1 \Phi_2 e^{i\xi^2 t \epsilon^2 \mathbf{u}} \rho(0)
\]

\[
= \int_0^t ds_i e^{i(t-s_i) \mathbf{L}_i^2} \sum_{j=1}^n ds_j e^{i(s_j-t_z) \mathbf{L}_j^2} \rho_0(t).
\]

In this way, Eq. (3) provides us with a general and useful expression of the solution of Eq. (1). Let us consider the truncated series in Eq. (3), that is \(\tilde{\rho}_n(t) = e^{i\xi^2 t \mathbf{u}} \rho(0) + \sum_{i=1}^n \rho_i(t)\), where \(n\) corresponds to the order of the approximation. We will prove that an expectation value \(\langle O \rangle_{\rho(t)} \equiv \text{Tr} [O \rho(t)]\) corresponding to a dissipative dynamics can be well approximated as

\[
\langle O \rangle_{\rho(t)} \approx \text{Tr} [O e^{i\xi^2 t \epsilon^2 \mathbf{u}} \rho(0)] + \sum_{i=1}^n \text{Tr} [O \rho_i(t)].
\] (4)

In the following, we will supply with a quantum algorithm based on single-shot random measurements to compute each of the terms appearing in Eq. (4), and we will derive specific upper-bounds quantifying the accuracy of our method. Notice that the first term at the right-hand-side of Eq. (4), i.e. \(\text{Tr} [O e^{i\xi^2 t \epsilon^2 \mathbf{u}} \rho(0)]\), corresponds to the expectation value of the operator \(O\) evolving under a unitary dynamics, thus it can be measured directly in a unitary quantum simulator where the dynamics associated with the Hamiltonian \(H\) is implementable. However, the successive terms of the considered series, i.e. \(\text{Tr} [O \rho_i(t)]\) with \(i \geq 1\), require a specific development because they involve multi-time correlation functions of the Lindblad operators and the operator \(O\).

Let us consider the first order term of the series in Eq. (4)

\[
\langle O \rangle_{\rho_1(t)} = \int_0^t ds_1 \text{Tr} [O \xi^2 t \mathbf{L}_1 e^{i\xi^2 t \epsilon^2 \mathbf{u}} \rho_0(s_1)]
\]

\[
= \sum_{i=1}^N \int_0^t ds_1 \gamma_i(s_1) \left\{ L_i^1(s_1) O(t) L_i^1(s_1) \right\} - \frac{1}{2} \left\{ \{O(t), L_i^1 L_i^1(s_1)\} \right\},
\] (5)

where \(\xi(s) \equiv e^{iHt_0^2 \epsilon^2 \mathbf{u}}\) for a general operator \(\xi\) and time \(s\), and all the expectation values are computed in the state \(\rho(0)\). Note that the average values appearing in the second and third lines of Eq. (5) correspond to time correlation functions of the operators \(O\), \(L_i^1\), and \(L_i^1 L_i^1\). In the following, we consider a basis \(\left\{ Q_j \right\}_{j=1}^d\), where \(d\) is the system dimension and \(Q\) are Pauli-kind operators, i.e. both unitary and Hermitian (see supplemental material for more details). The operators \(L_i\) and \(O\) can be decomposed as \(L_i = \sum_{k=1}^{M_i} q_i^k Q_i^k\) and \(O = \sum_{i=1}^{M_i} q_i^{iO} Q_i^{iO}\), with \(q_i^{iO} \in \mathbb{C}\). \(Q_i^{iO} \in \left\{ Q_j \right\}_{j=1}^d\), and \(M_i M_j \leq d^2\). We obtain then

\[
\{L_i^1(s_1) O(t) L_i^1(s_1)\} = \sum_{i=1}^{M_i} \sum_{k=1}^{M_i} q_i^{iO} q_i^{s_1} q_i^{s_1} \{Q_i^k(s_1) Q_i^{s_1}(s_1)\},
\] (6)

that is a sum of correlations of unitary operators. The same argument can be easily extended to higher-order corrections. Indeed, for the \(n\)-th order, we have to evaluate the quantity

\[
\langle O \rangle_{\rho_n(t)} = \int dV_n \text{Tr} [O e^{i\xi^2 t \epsilon^2 \mathbf{u}} \mathbf{L}_D^{t_1} \ldots \mathbf{L}_D^{t_n} e^{-i\epsilon^2 \mathbf{u} H} \rho(0)] \equiv \sum_{i_1, \ldots, i_n} \int dV_n \{A_{i_1, \ldots, i_n}(\vec{s})\}.
\] (7)

Here,

\[
A_{i_{11}, \ldots, i_{nn}}(\vec{s}) \equiv e^{i\xi^2 t \epsilon^2 \mathbf{u} \mathbf{L}_D^{t_n} \ldots \mathbf{L}_D^{t_1} e^{i\xi^2 n-t_{n-1}} \mathbf{L}_n^2 \ldots \mathbf{L}_n^2} \mathbf{L}_D^{t_n} e^{i\epsilon^2 \mathbf{u} H} O_{i_{11}, \ldots, i_{nn}}(\vec{s}),
\]

where

\[
\mathbf{L}_D^s \xi = \gamma_1(s) \left\{ L_s \xi^1 - \frac{1}{2} \left\{ L_s \xi^1 \right\} \right\}, \quad \vec{s} = (s_1, \ldots, s_n) \int dV_n = \int_0^t \ldots \int_0^{t_{n-1}} \int_{s_1} \ldots ds_{n-1} ds_n
\]

and \(\mathbf{L}_D^s \xi = (\xi^{s})\) for a general superoperator \(\mathbf{L}\). As in Eq. (5), the above expression contains multi-time correlation functions of the Lindblad operators \(L_{i_1}, \ldots, L_{i_n}\) and the observable \(O\), that have to be evaluated in order to compute each contribution in Eq. (4).

Our next step is to provide a method to evaluate general terms as the one appearing in Eq. (7). The standard approach to estimate this kind of quantities corresponds to measuring the expected value \(\langle A_{i_{11}, \ldots, i_{nn}}(\vec{s}) \rangle\) at different random times \(\vec{s}\) in the integration domain, and then calculating the average. Nevertheless, this strategy involves a huge number of measurements, as we need to estimate an expectation value at each chosen time. Our technique, instead, is based on single-shot random measurements, and as we will see below, it leads to an accurate estimate of Eq. (7). More specifically, we will prove that

\[
\sum_{i_{11}, \ldots, i_{nn}} \int dV_n \{A_{i_{11}, \ldots, i_{nn}}(\vec{s})\} \approx \frac{N \int dV_n}{\sum_{i_{11}, \ldots, i_{nn}} \{A_{i_{11}, \ldots, i_{nn}}(\vec{s})\}}
\]

(8)
where $\tilde{A}_n(\vec{t})$ corresponds to a single-shot measurement of $A_\omega(\vec{t})$, being $[\vec{\omega}, \vec{t}] \in \Omega_n \subset \{|\vec{\omega}, \vec{t}\}|, [\vec{\omega} = [i_1, \ldots, i_N], i_k \in [1, N], \vec{t} \in \mathbb{R}^N\}, |\Omega_n|$ is the size of $\Omega_n$, and $[\vec{\omega}, \vec{t}]$ are sampled uniformly and independently. As already pointed out, the integrand in Eq. (7) involves multi-time correlation functions. In this respect, we note that a quantum algorithm for their efficient reconstruction has recently been proposed\cite{SupplementalMaterial}. Indeed, the authors in Ref. [34] show how, by adding only one ancillary qubit to the simulated system, general time-correlation functions are accessible by implementing only unitary evolutions of the kind $e^{i\mathcal{L}_n}$, together with entanglement operations between the ancillary qubit and the system. It is noteworthy to mention that these operations have already experimentally demonstrated in quantum systems as trapped ions\cite{TrappedIons} or quantum optics\cite{QuantumOptics}, and have been recently proposed for cQED architectures\cite{CQED}. Moreover, the same quantum algorithm allows us to measure single-shots of the real and imaginary part of the corresponding correlation function. Finally, in the supplemental material\cite{SupplementalMaterial} we prove that

$$\sum_{i_1, \ldots, i_N} \int dV_n \left\{A_{\{i_1, \ldots, i_N\}}(\vec{t})\right\} - \frac{1}{n!} \left[\sum_{i \in \Omega_n} \tilde{A}_n(\vec{t})\right] \leq \delta_n$$

with probability higher than $1 - e^{-\delta}$, provided that $|\Omega_n| > \frac{36 M_0^2 (2 + \beta) (2\pi MN)^{2n}}{n!}$, where $n \equiv \max_{i \in \Omega_n} |\gamma_i(s)|$ and $M \equiv \max_i M_i$. Equation (9) means that the quantity in Eq. (7) can be estimated with arbitrary precision by random single-shot measurements of $A_{\{i_1, \ldots, i_N\}}(\vec{t})$, allowing, hence, to dramatically reduce the resources required by our quantum simulation algorithm. Notice that the required number of measurements to evaluate the order $n$ is bounded by $3^n |\Omega_n|$ and the total number of measurements needed to compute the correction to the expected value of an observable up to the order $K$ is bounded by $\sum_{n=0}^K 3^n |\Omega_n|$ In the following, we discuss at which order we need to truncate in order to have a certain error in the final result.

So far, we have proved that we can compute, up to an arbitrary order in $\mathcal{L}_D$, expectation values corresponding to dissipative dynamics with a unitary quantum simulation. It is noteworthy that our method does not require to physically engineer the system-environment interaction. Instead, one only needs to implement the system Hamiltonian $H$. In this way we are opening a new avenue for the quantum simulation of open quantum dynamics in situations where the complexity on the design of the dissipative terms exceeds the capabilities of quantum platforms. This covers a wide range of physically relevant situations. One example corresponds to the case of fermionic theories where the encoding of the fermionic behavior in the degrees of freedom of the quantum simulator gives rise to highly delocalized operators\cite{FermionicEncoding,FermionicEncoding2}. In this case a reliable dissipative term should act on these non-local operators instead of on the individual qubits of the system. It is noteworthy that our method does not require to physically engineer the system-environment interaction. Instead, one only needs to implement the system-environment interaction. Instead, one only needs to implement the system Hamiltonian $H$. In this way we are opening a new avenue for the quantum simulation of open quantum dynamics in situations where the complexity on the design of the dissipative terms exceeds the capabilities of quantum platforms. This covers a wide range of physically relevant situations. One example corresponds to the case of fermionic theories where the encoding of the fermionic behavior in the degrees of freedom of the quantum simulator gives rise to highly delocalized operators\cite{FermionicEncoding,FermionicEncoding2}. In this case a reliable dissipative term should act on these non-local operators instead of on the individual qubits of the system. Our protocol solves this problem because it avoids the necessity of implementing the Lindblad superoperator. Moreover, the scheme allows one to simulate at one time a class of master equations corresponding to the same Lindblad operators, but with different choices of $\gamma_i$, including the relevant case when only a part of the system is subjected to dissipation, i.e. $\gamma_i = 0$ for some values of $i$.

We shall next quantify the quality of our method. In order to do so, we will find an error bound certifying how the truncated series in Eq. (3) is close to the solution of Eq. (1). This error bound will depend on the system parameters, i.e. the time $t$ and the dissipative parameters $\gamma_i$. As figure of merit we choose the trace distance, defined by

$$D_1(\rho_1, \rho_2) \equiv \frac{1}{2} \left\| \rho_1 - \rho_2 \right\|_1,$$

where $\|A\| \equiv \sum \sigma_i(A)$, being $\sigma_i(A)$ the singular values of $A^{\otimes\omega}$. Our goal is to find a bound for $D_1(\rho(t), \bar{\rho}_n(t))$, where $\bar{\rho}_n(t) \equiv \sum_{i=0}^n \rho_i(t)$ is the series of Eq. (3) truncated at the $n$-th order. We note that the the following recursive relation holds

$$\bar{\rho}_n(t) = \mathcal{L}_D \rho(0) + \int_0^t ds e^{(t-s)\mathcal{L}_D} \mathcal{L}_D \bar{\rho}_{n-1}(s).$$

From Eq. (11), it follows that

$$D_1(\rho(t), \bar{\rho}_n(t)) = \frac{1}{2} \left\| \int_0^t ds e^{(t-s)\mathcal{L}_D} \mathcal{L}_D \left(\rho(s) - \bar{\rho}_{n-1}(s)\right) \right\|_1$$

$$\leq \int_0^t ds \left\|\mathcal{L}_D\right\|_{-1} D_1(\rho(s), \bar{\rho}_{n-1}(s)),$$
where we have introduced the induced superoperator norm $\|A\|_{-1} \equiv \sup_{\rho} \frac{\|e^{tA}\rho\|}{\|\rho\|}$. For $n = 0$, i.e. for

$$\bar{\rho}_n(t) = e^{tA}\rho(0),$$

we obtain the following bound

$$D_1(\rho(t), \bar{\rho}_0(t)) \leq \frac{1}{2} \int_0^\infty ds \left\| e^{tA}\rho(0) \right\| \leq \sum_{n=1}^N |\gamma_i(\varepsilon_i)| \left\| L_i \right\|^2_{\infty} t,$$

(13)

where $0 \leq \varepsilon_i \leq t^{37}$, and $\|A\|_{\infty} \equiv \sup_{\rho} \sigma_i(A)$. Notice that, in finite dimension, one can always renormalize $\gamma_i$ in order to have $\|L_i\|_{\infty} = 1$, i.e. if we transform $L_i \rightarrow L_i/\|L_i\|_{\infty}$, $\gamma_i \rightarrow \|L_i\|_{\infty} \gamma_i$, the master equation remains invariant. Using Eq. (12)–(13), one can show by induction that for the general $n$-th order the following bound holds

$$D_1(\rho(t), \bar{\rho}_n(t)) \leq \Pi_{k=0}^{n-1} \left[ 2\sum_{\varepsilon_k(\varepsilon_k)} |\gamma_k(\varepsilon_k)| \right]^{\frac{t^{n+1}}{2(n+1)}},$$

(14)

where $0 \leq \varepsilon_k \leq t$ and we have set $\|L_i\|_{\infty} = 1$. From Eq. (14), it is clear that the series converges uniformly to the solution of Eq. (1) for every finite value of $t$ and choices of $\gamma_i$. As a result, the number of measurements needed to simulate a certain dynamics at time $t$ up to an error $\varepsilon < 1$ is $O\left( \left( t + \log \frac{1}{\varepsilon} \right)^2 \frac{1}{\varepsilon^2} \right)$, where $t = \pi N t^{37}$. Here, a discussion on the efficiency of the method is needed. From the previous formula, we can say that our method performs well when $M$ is low, i.e. in case where each Lindblad operator can be decomposed in few Pauli-kind operators. Moreover, as our approach is perturbative in the dissipative parameters $\gamma_i$, the method is more efficient when $|\gamma_i|$ are small. Notice that analytical perturbative techniques are not available in this case, because the solution of the unperturbed part is assumed to be not known. Lastly, it is evident that the algorithm is efficient for certain choices of time, and the relevance of the simulation depends on the particular cases. For instance, a typical interesting situation is a strongly coupled Markovian system. Let us assume a site-independent couple parameter $g$ and a dissipative parameter $\gamma$. We have that $e^{Mt} \leq 1 + 12eM\bar{t}$ if $t \leq \frac{1}{12eMg} \equiv t_0$. In this period, the system oscillates typically $C \equiv gt_0 = \frac{g}{12eM} \bar{t}$ times, so the simulation can be considered efficient for $N \sim g/C$, which, in the strong coupling regime, can be of the order of $10^3$. Notice that, in most relevant physical cases, the number of Lindblad operators $N$ is of the order of the number of system particles.

All in all, our method is aimed to simulate a different class of master equations with respect to the previous approaches, including non-Markovian quantum dynamics, and it is efficient in the range of times where the exponential $e^{Mt}$ may be truncated at some low order. A similar result is achieved by the authors of Ref. [27], where they simulate a Lindblad equation via Trotter decomposition. They show that the Trotter error is exponentially large in time, but this exponential can be truncated at some low order by choosing the Trotter time step $\Delta t$ sufficiently small. Our method is qualitatively different, and it can be applied also to analogue quantum simulators where suitable entangled gates are available.

Lastly, we note that this method is also applicable to simulate dynamics under a non-Hermitian Hamiltonian $J = H - i\Gamma$, with $H = H^\dagger$, $\Gamma = \Gamma^\dagger$. This type of generator emerges as an effective Hamiltonian in the Feshbach partitioning formalism [41], when one looks for the evolution of the density matrix projected onto a subspace. The new Schrödinger equation reads

$$\frac{d\rho}{dt} = -i[H, \rho] + [\Gamma, \rho],$$

(15)

This kind of equation is useful in understanding several phenomena, e.g. scattering processes [42] and dissipative dynamics [43], or in the study of PT-symmetric Hamiltonians [44]. Our method consists in considering the non-Hermitian part as a perturbative term. As in the case previously discussed, similar bounds can be easily found (see the supplemental material [37]), and this proves that the method is reliable also in this situation.

In conclusion, we have proposed a method to compute expectation values of observables that evolve according to a generalized Lindblad master equation, requiring only the implementation of its unitary part. Through the quantum computation of $n$-time correlation functions of the Lindblad operators, we are able to reconstruct the corrections of the dissipative terms to the unitary quantum evolution without reservoir engineering techniques. We have provided a complete recipe that combines quantum resources and specific theoretical developments to compute these corrections, and error-bounds quantifying the accuracy of the proposal and defining the cases when the proposed method is efficient. Our technique can be also applied, with small changes, to the quantum simulation of non-Hermitian Hamiltonians. The presented method provides a general strategy to perform quantum simulations of open systems, Markovian or not, in a variety of quantum platforms.

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R.D.C. did the calculations. R.D.C., J.S.P., A.D.C., E.S. and J.C. contributed to the developing of the ideas, obtention of the results and writing the manuscript.

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