Enhanced Convergence and Robust Performance of Randomized Dynamical Decoupling

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(Dated: April 1, 2022)

We demonstrate the advantages of randomization in coherent quantum dynamical control. For systems which are either time-varying or require decoupling cycles involving a large number of operations, we find that simple randomized protocols offer superior convergence and stability as compared to deterministic counterparts. In addition, we show how randomization may allow to outperform purely deterministic schemes at long times, including combinatorial and concatenated methods. General criteria for optimally interpolating between deterministic and stochastic design are proposed and illustrated in explicit decoupling scenarios relevant to quantum information storage.

PACS numbers: 03.67.Pp, 03.65.Yz, 05.40.Ca, 89.70.+c

Counterintuitive roles of randomness in physical processes have long been acknowledged. A paradigmatic example is stochastic resonance [1], whereby the transmission of a weak signal is maximized at an optimum noise level. Within quantum information science, a number of applications benefitting from randomness in quantum states and operations have been identified recently. Suggestive evidence is offered by dissipation-assisted quantum computation schemes [2] and by the idea that chaos may stabilize quantum algorithms [3]. In the quantum communication setting, the use of random operations has been shown to decrease the communication cost of achieving remote state preparation and of constructing efficient quantum data-hiding schemes [4]. Finally, random unitary operators may pave the way to efficient estimation methods for open quantum systems [5].

In the context of quantum control theory, the fragility of quantum coherence against uncontrolled noise and randomness hindered the exploration of stochastic control methods till recently [2, 3, 8, 9]. General control-theoretic arguments supporting the usefulness of controlled randomness in dynamical decoupling (DD) schemes for switching off unwanted evolution in interacting quantum systems were provided in [8], and have been validated in specific examples [8] (see also [2]). These studies leave, however, several key questions largely unanswered. In particular: For given control resources, under what circumstances (if any) can randomized DD match or outperform the best deterministic schemes known to date? Can randomized design be exploited in general to further push the efficiency and viability of DD schemes?

In this Letter, we present conclusive evidence of the benefits of randomization in light of the above questions. To isolate the factors responsible for different DD performance, we focus on the simplest control benchmark: complete refocusing of Hamiltonian evolution by means of a restricted but otherwise perfect set of control operations. In the process, we verify conjectures made in [8], stretch the analysis into unexplored domains, and offer criteria for constructing new highly efficient DD protocols. For time-independent systems, we demonstrate the superior averaging of randomized protocols in control scenarios involving sufficiently long control cycles and/or evolution times. For time-varying systems, randomized DD provides a robust strategy in the presence of system uncertainties. The combination of three key ingredients – concatenated control, symmetry, and randomization – emerges as a general principle for design optimization.

**DD setting.** DD methods modify a target dynamics $H_0(t)$ through the addition of a control field $H_c(t)$. We assume that $||H_0(t)||_2 = \text{max} |\text{eig}(H_0(t))| < \kappa$, for $\kappa > 0$ and all $t$. Let $U_c(t) = T \exp[-i \int_0^T H_c(u) du]$ denote the control propagator, with $T$ indicating time ordering. In a logical frame which explicitly removes $H_c(t)$, the controlled evolution is described by the propagator $\bar{U}(t) = U(t)\bar{U}(t)^\dagger = T \exp[-i \int_0^T \bar{H}(u) du]$, where $U(t)$ is the total propagator in the physical frame and $\bar{H}(t) = U_c(t)^\dagger H_0(t) U_c(t)$ is the logical Hamiltonian [10]. If $H_0(t)$ is time-independent and $U_c(t)$ periodic with a cycle time $T_c$, then $\bar{U}(T_n) = U(T_n)$, $T_n = nT_c$, $n \in \mathbb{N}$, and both frames coincide stroboscopically. The DD objective here is to make $\bar{U}(T)$, for final time $T > 0$, as close as possible to $I$ according to a metric of choice.

Let the control resources be specified by a discrete set of unitary operators corresponding to a (projective) representation of a group $G = \{g_j\}$, $j = 0, \ldots, |G|-1$, in the Hilbert space $H$ of the system, $\dim(H) = d < \infty$. Both the basic deterministic and random DD protocols may be understood as effecting an appropriate symmetrization of $H_0(t)$ according to $G$. Assume a time-independent system first. In standard deterministic DD, average Hamiltonian theory (AHT) allows the logical evolution to be represented as $\bar{U}(T_n) = [\exp(-i\bar{H}T_n)]^{n}$, where $\bar{H} = \sum_{k=0}^{\infty} \bar{H}^{(k)}$ and each term $\bar{H}^{(k)}$ is computed from the Magnus expansion [10]. Under the convergence condition $\kappa T_c < 1$, the leading zero-th order contribution $\bar{H}^{(0)}$ may be mapped into a group contribution $\bar{H}_G = \sum_{j} g_j^\dagger H_0 g_j$ by sequentially steering $U_c(t)$ through all the group elements $\{g_j\}$ – which corresponds to a train of “bang-bang”
instantaneous pulses $P_k = g_k g_{k-1}^\dagger$, $k = 1, \ldots, |\mathcal{G}|$, separated by $\Delta t > 0$ and with $T_c = |\mathcal{G}| \Delta t$. A cyclic protocol based on a fixed, predetermined control path within $\mathcal{G}$ will be referred to as periodic DD, henceforth (PDD), provided first-order averaging is achieved, $\hat{H}^{(0)} = 0$. The simplest randomized version of such a DD procedure is obtained by sampling $U_c(t)$ over $\mathcal{G}$ uniformly at random according to the Haar measure $\mathbb{P}$ leading to what we call naive random DD (NRD). Let $\mathbb{E}\{\cdot\}$ denote expectation over all possible control realizations; $\mathcal{G}$-symmetrization still emerges on average at each $t$ via a quantum operation $\mathbb{E}\{U_c(t) H_0(t) U_c^\dagger(t)\} = H_G$, and convergence may be rigorously established in the limit $T \Delta t k^2 \ll 1$.

Lower bounds for the expected logical-frame fidelity of an arbitrary pure state $|\psi\rangle$, $\mathbb{E}\{F_G(T)\} = \mathbb{E}\{|\langle \psi| \hat{U}(T)|\psi\rangle|^2\}$, are given by $\mathbb{E}\{F_G(T)\} \geq 1 - \mathcal{O}(T^2 T_c^2 k^4)$ for PDD, and $\mathbb{E}\{F_G(T)\} \geq 1 - \mathcal{O}(T \Delta t k^2)$ for NRD, respectively. Within their regime of validity, these bounds indicate the potential for NRD to outperform PDD when $|\mathcal{G}|^2 (T \Delta t k^2) \gg 1$. In order to quantitatively compare DD schemes, a control metric which is both efficiently computable and state-independent is desirable. Here, we consider average (unlike worst-case as in $\mathbb{P}$) performance, and remove the dependence upon $|\psi\rangle$ by invoking the (expected) entanglement fidelity $\mathbb{F}_{\psi}$, evaluated as $\mathbb{E}\{F_G(T)\} = \mathbb{E}\{|\langle \psi| \hat{U}(T)|\psi\rangle|^2\}$. $F_G(T)$ is linearly related to the average of $\mathbb{F}_{\psi} (T)$ over all $|\psi\rangle$. Perfect DD corresponds in this metric to $\mathbb{E}\{F_G(T)\} \rightarrow 1$. In practice, Monte Carlo simulations of the controlled dynamics are used to estimate $\mathbb{E}\{F_G(T)\}$ through a statistical average $\langle \langle F_G(T) \rangle \rangle$ over control realizations.

**Time-independent case: Convergence improvement.**

By way of illustration, consider $N$ spin-1/2 particles (qubits), described by the following model Hamiltonian:

$$
H = \sum_{i=1}^{N} \frac{\omega_i}{2} \sigma_i^{(z)} + \sum_{i < j} \sum_{a=x,y,z} J_{ij}^{(a)} (r_{ij}) \sigma_i^{(a)} \otimes \sigma_j^{(a)},
$$

where $\sigma^{(x,y,z)} = \sigma_X, \sigma_Y, \sigma_Z$ are Pauli operators, and $\omega_i, J_{ij}^{(a)}$ denote the frequency of the $i$th qubit, and the coupling strength of the $ij$th pair in the $a$ direction, respectively. Heisenberg interactions exponentially decaying with the distance $r_{ij}$ are typical of quantum dot arrays [13], whereas cubic decays describe dipolarly coupled spins in systems ranging from nuclear magnetic resonance (NMR) crystals and liquid-crystals [10, 11] to electrons on Helium [15]. We assume here that $\omega_i \approx \omega$, and work in a logical-rotating frame whereby the effective Hamiltonian $\hat{H}_R(t) = U_c(t) H U_c^\dagger(t) - \omega \sum_i Z_i^z / 2 |U_R(t) U_c(t), U_R(t) = \exp[-\text{i} \omega t] \sum_i Z_i / 2]$ [13, 17].

PDD protocols capable of refocusing $H$ for arbitrary parameter values (and in fact, arbitrary interactions of the form $\sigma_i^{(a)} \otimes \sigma_j^{(b)}$) may be built by recursively nesting DD sequences based on the group $\mathcal{G}_i = \{\mathbb{I}, X_i, Z_i, Y_i\}$ for each added qubit, $i = 2, \ldots, N$. Although this scheme is not efficient, as the number of pulses per cycle grows as $4^N - 1$, it allows the effect of large $|\mathcal{G}|$ to be studied in moderately small systems. Following [18], the PDD sequence we implement corresponds to a path over the Pauli group $\mathcal{G}_P = \otimes_i \mathcal{G}_i$ which avoids simultaneous pulses. In NRD, $U_c(t)$ is picked uniformly at random over $\mathcal{G}_P$, which may involve collective rotations on up to $R = N - 1$ qubits $\mathcal{G}_R$. Numerical results are shown in Fig. 1. As seen in the main panel, the fidelity for NRD decays substantially slower than for PDD. Irrespective of the validity of the short-time condition underlying the lower bounds of $\mathbb{P}$, these results confirm the faster convergence of stochastic DD when $|\mathcal{G}|$ is large. While $\Delta t$ is kept fixed for NRD, PDD protocols with decreasing $\Delta t$ are considered. NRD eventually surpasses all PDD curves, showing that, for sufficiently long time, the constraints on $\Delta t$ for random DD may be relaxed – which can prove advantageous in practice. Also note that in situations where control constraints make it unfeasible to complete a cycle (inset), NRD outperforms the selected PDD sequence at all intra-cycle times $t_n = n \Delta t < T_c$ – a finding which indicates the existence of better control paths in this time domain.

**Time-independent case: Long-time improvement.**

While different DD protocols are equivalent in the short-time limit under ideal averaging, performance for finite $\Delta t$ and long time depends critically on the way in which residual errors accumulate. We address this point by extending our comparison to higher-level DD schemes.

In the deterministic domain, two strategies exist for improving over PDD. One, motivated by the Carr-Purcell sequence of NMR, consists of a time-symmetrization of the control path, such that all odd order terms in $\hat{H}$ are also canceled [17, 11]. This leads to symmetric DD (SSD). The other, inspired to NMR iterative techniques [10], implements concatenated DD (CDD) [21]. CDD relies on a temporal recursive structure, the $(\ell + 1)$-level pulse sequence being determined by $C_{\ell+1} = C_\ell P_{\ell} C_\ell P_{\ell} \ldots C_\ell P_{N}$,
Dashed and dot-dashed lines correspond to different inner sequences characterizing EMD, GEMD irreducible typically ensures superior performance, and SRPD is outperformed by EMD. While full derivations will be presented in [17], we find $\langle F_n(T) \rangle \geq 1 - O(T^2(|G|\Delta t)^{2+\alpha}κ^{4+\alpha}),$ where $\alpha = 0.2$ for PDD and SDD, respectively. Error bounds for CDD depend on both the system and the concatenation level, but already at $\ell = 2$ CDD is more efficient than SDD in reducing $H^{(k)}$, $k \geq 2$. Notice, however, that unlike the single-qubit DD analyzed in [21], no guarantee of superpolynomial convergence exists when CDD is based on a reducible group. For randomized DD, the basic error bound is contained in [3] for NRD, further adapted in [8] to EMD. While full derivations will be presented in [17], we find $\langle F_n(T) \rangle \geq 1 - O(T^2(|G|\Delta t)^{2+\alpha}κ^{4+\alpha}),$ where $\alpha = 0$ for EMD and RPD, while $\alpha = 2$ for SRPD. As applicability of the above bounds is strictly confined to the regime where $T(|G|\Delta t)κ^2 \ll 1,$ we proceed to a numerical comparison.

For bilinear couplings as in Eq. (1), efficient combinatorial PDD schemes with quadratic complexity are well known [18]. First-order DD sequences for up to $N = 4^m$ qubits, $m \geq 1,$ involve $N$ simultaneous pulses. For $N = 8,$ the basic PDD scheme is derived from the group $G_8 = \{1, Z_3Z_4Y_6Z_7X_8, Z_2Y_3X_4Z_6Y_7X_8, Z_2X_1Y_4X_5Z_6X_7Z_2, Y_2Y_3X_5Z_6X_7, Y_2Z_3X_4Z_5X_6Y_8, Z_2X_3Y_4Z_5X_6Z_7Y_8, X_2Y_3Z_5Y_6Z_7\}.$

As $G_8$ is reducible, saturation in the performance of CDD is observed when $\ell > 2.$ Numerical results are summarized in Fig. 1. As expected, randomized DD surpasses deterministic DD at long times, the crossing being evident between protocols that have comparable efficiency at short times. The dependence of EMD upon the underlying inner path is very pronounced for this system. Remarkably, CDD is outperformed by SRPD. Although CDD is known to minimize sensitivity to control faults not included here, preliminary results on rotation angle imperfections indicate that the above conclusions remain unchanged.

Exploitation of randomized DD in actual devices will vary depending on whether probabilistic pulse generation capabilities are available or not. In the former case, randomization appears especially promising wherever a large number of experimental runs is required for observation, such as in quantum dot ensemble experiments [24]. Even in the absence of dedicated pulsing capabilities, randomized DD may still be useful (e.g., in solid-state NMR systems) once the procedure is de-randomized, that is, randomization is used off-line to post-select optimal pulse realizations for given physical parameters. Such an option is no longer viable for time-varying control systems.

**Time-varying systems and robust performance.**– For simplicity, let us specialize Eq. (1) to a situation where only nearest-neighbor couplings are relevant, but a time-dependent anisotropy is effectively present, $J_{i+1}^{x,y} = J_i, J_{i+1}^{z} = J\Delta(t).$ For arbitrary $N,$ an efficient first-order PDD protocol may be constructed by alternating two collective rotations around perpendicular axes – one acting on all odd qubits, the other on the even ones, e.g., if $N$ is even, $G = \{1, Z_1Z_3\ldots Z_{N-1}, Z_1Y_2Z_3Y_4\ldots Z_{N-1}Y_N, Y_2Y_4\ldots Y_N\}.$

While under mild assumptions on the time dependence of $\Delta(t),$ the NRD fidelity bound is still valid [3].
no general result on deterministic performance is avail-
able. Suppose that the system fluctuation entails sign changes over time, with a potentially unknown period \( \tau \). Then if \( \tau/\Delta t \approx p, p \in \mathbb{N} \), adversarial situations may arise where a pre-established control action is inhibited or even reversed, resulting in surprisingly poor performance of cyclic DD – often worse than NRD or free evolution, see main panel of Fig. 3. Stochastic methods are intrinsically more protected, on average, against such interferences, resulting in more stable DD performance throughout.

**Conclusion.**– We have shown that randomized decoupling can offer distinctive advantages over deterministic methods in terms of faster convergence, long-time improvement, and robust performance. Together with concatenation and symmetry, randomization provides a versatile toolbox for matching different control needs. Both the analysis of fault-tolerance properties and of the interplay between probabilistic design and robustness deserve a closer scrutiny, also in view of known randomized algorithms for classical control systems [25]. It is our hope that the present results will prompt experimental verification in available control devices.

Partial support from Constance and Walter Burke’s Special Projects Fund in QIS and from the NSF through grant No. PHY-0555417 is gratefully acknowledged. Thanks to O. Kern for valuable feedback.

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