Variational quantum dynamics of two-dimensional rotor models

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Abstract
We present a numerical method to simulate the dynamics of continuous variable quantum many-body systems. Our approach is based on custom neural-network many-body quantum states. We focus on dynamics of two-dimensional quantum rotors and simulate large experimentally-relevant systems with using state-of-the-art sampling approaches based on Hamiltonian Monte Carlo. We demonstrate the method can access quantities like the return probability and vorticity oscillations after a quantum quench in two-dimensional systems of up to 64 (8 × 8) coupled rotors. Our approach can be used to perform previously unexplored non-equilibrium simulations bridging the gap between simulation and experiment.

1 Introduction
Non-equilibrium quantum many-body physics has been at the center of physics and chemistry research for over a decade [1, 2]. The field is driven by remarkable progress in our ability to control matter at the atomic scale [3–5]. Such quantum control [6–9] of modern experiments and hardware is becoming increasingly limited by numerical simulation of the real-time evolution of quantum systems. Fast entanglement growth out of equilibrium forces one to keep track of many-body correlations. Challenges remain despite recent progress [10–19], especially in cases of realistic systems with continuous-variables. Practically, calculations are roadblocked by numerical instabilities resulting from a combination of Monte Carlo noise and flatness of the quantum geometry of modern neural network wave functions [20, 17, 21–23].

In this work, we present an approach for capturing long-time dynamics of continuous-variable 2D lattice models, using a combination of methods that were previously unexplored in the field – the Hamiltonian Monte Carlo sampler, a tailored variational ansatz and novel regularization of projected dynamics. We focus on the quantum rotor model with direct applications to arrays of coupled Josephson junctions and explore previously unreachable system sizes and evolution times.

2 Model and Methods
Consider a system of planar rotors, whose angles \( \theta_k \) with respect to an arbitrary axis on a lattice \( \Lambda \) with \( N \) sites. We use the basis \( |\theta\rangle \equiv |\theta_1, \ldots, \theta_N\rangle \) for the Hilbert space \( \mathcal{H} \). We start with an effective Hamiltonian that captures the relevant physics of superconducting Josephson junctions [24–26] such
Convolution

Direct solution of the relevant time-dependent Schrödinger equation (TDSE)

\[ \text{i} \frac{\partial}{\partial t} \psi(t) = -\beta H \psi(t) \]

of practical applications in the study of dynamics of arrays of coupled Josephson junctions [30]. However, QRM real-time dynamics have not been explored despite prospects (QRM). Its equilibrium properties [27] have been studied using Quantum Monte Carlo (QMC) methods. However, QRM real-time dynamics have not been explored despite prospects of practical applications in the study of dynamics of arrays of coupled Josephson junctions [30].

Direct solution of the relevant time-dependent Schrödinger equation (TDSE) \( i \frac{\partial}{\partial t} \psi = H \psi \) is prohibitively expensive even for a handful of interacting rotors. The continuous nature of the |θ| basis exacerbates the problem.

We represent a quantum state using a convolutional neural network (CNN) wavefunction \( \psi_\alpha(\theta) \) where \( \alpha \in \mathbb{C}^P \) is a set of \( P \) complex variational parameters. The full neural-network quantum state (NQS) then reads \( |\psi_\alpha\rangle = \int \text{d}\theta \, \psi_\alpha(\theta) |\theta\rangle \) where \( \text{d}\theta = \text{d}\theta_1 \ldots \text{d}\theta_N \). The integral is performed over the cube \( [-\pi, \pi]^N \).

Our simulation of the real-time dynamics of NQS is based on the time-dependent Variational Monte Carlo (t-VMC) method [31, 16, 15]. The core assumption that allows us to approximately solve the TDSE is that of time dependence of neural-network parameters \( \alpha = \alpha(t) \). Trajectories \( \alpha(t) \) can be derived by substituting \( \partial_t \psi_\alpha = \dot{\alpha} \cdot \nabla_\alpha \psi_\alpha \) into the TDSE. Evolution equations [22] then read

\[
S_{\mu\nu} = \langle O^\dagger_\mu O_\nu \rangle - \langle O^\dagger_\mu \rangle \langle O_\nu \rangle \quad \text{and} \quad g_\mu = \langle O^\dagger_\mu H \rangle - \langle O^\dagger_\mu \rangle \langle H \rangle .
\]

Averages \( \langle \cdot \rangle \equiv \langle \psi_\alpha | \psi_\alpha \rangle / \langle \psi_\alpha | \psi_\alpha \rangle \) are performed at time \( t \) (i.e. for \( \alpha = \alpha(t) \)). Parameters \( \alpha \) are indexed by \( \{\mu, \nu, \ldots\} \) in our notation. Operator \( O_\mu \) is defined by \( \partial_{\alpha_\mu} |\psi_\alpha\rangle = O_\mu |\psi_\alpha\rangle \). The matrix \( S \) is commonly called the quantum geometric tensor (QGT) [32, 21, 23] and corresponds to the metric tensor of the neural-network parameter manifold.

Since quantum averages over an exponentially large Hilbert space \( \mathcal{H} \) in Eq. 2 cannot be computed exactly, Markov chain Monte Carlo (MCMC) sampling methods are often employed [33, 34]. In VMC calculations, it is common to rewrite quantum averages such as those in Eq. 2 using the local operator trick. We refer the interested reader to an excellent summary in Refs. [22, 31].

\[
\langle H \rangle = \frac{\langle H \rangle}{\langle \psi_\alpha | \psi_\alpha \rangle} = \frac{\int \text{d}\theta \, \psi_\alpha^*(\theta) H \psi_\alpha(\theta) \text{d}\theta}{\int \text{d}\theta \, |\psi_\alpha(\theta)|^2} = \mathbb{E}_{\theta \sim |\psi_\alpha|^2} \left[ H \psi_\alpha(\theta) / |\psi_\alpha(\theta)|^2 \right].
\]

After computing the matrix \( S \) and the vector \( g \) at time \( t \), one can formally define \( \dot{\alpha} = -i S^{-1} g \) and use any ordinary differential equation (ODE) integrator to obtain the next set of parameters at time...
We introduce a heuristic regularization (pseudoinverse) that allows us to propagate the TDSE longer in time. The inverse $S^{-1}$ is often ill-defined. One reason is noisy Monte Carlo estimates of matrix elements make small eigenvalues vanish. Therefore, quickly and efficiently obtaining many uncorrelated samples from $\langle \psi_{\alpha(t)} | \psi_{\alpha(t)} \rangle$ is crucial. In addition, overparametrizing $\psi_{\alpha}$ introduces redundancy, producing linearly dependent or vanishing rows/columns in $S$. Therefore, choosing a smaller CNN trial wavefunction is equally important. In practice, adding more parameters to the wavefunction can sometimes unexpectedly reduce accuracy by making $S$ ill-conditioned.

We introduce a heuristic regularization (pseudoinverse) that allows us to propagate the TDSE longer in time. The $S$ matrix is diagonalized $S = U \Sigma U^{-1}$ at each time step. Having obtained eigenvalues $\sigma_\mu^2$ such that $\Sigma = \text{diag}(\sigma_1^2, \ldots, \sigma_P^2)$, we define the pseudoinverse as $S^{-1} \approx U \Sigma^{-1} U^\dagger$ with $\Sigma^{-1}_{\mu\nu} = \frac{1}{\sigma_\mu^2} \delta_{\mu\nu}$. We heuristically find that such smooth cutoff is superior to traditional pseudoinverses when using adaptive integrators for updating parameters $\alpha$.

We employ Hamiltonian Monte Carlo (HMC) [38, 39] to estimate Hilbert space averages in Eq. 2 at each time step $t$. We make this choice because HMC offers a systematic way of making large steps in MCMC proposals while still keeping acceptance probabilities high. This results in a Markov chain with considerably lower autocorrelation times.

For a generic probability distribution $p(\theta)$, HMC augments the configuration space with artificial momentum variables $\pi = (\pi_1, \ldots, \pi_P)$. Hamiltonian dynamics in the resulting $2N$-dimensional space conserves energy (probability). Monte Carlo updates can be defined through numerical integration of Hamilton’s equations. Given initial conditions $\theta(0)$, $\pi(0)$ and a small step size, a common choice is the symplectic leapfrog integrator [38, 37] because it conserves energy (probability) exactly, allowing for large jumps in the $\theta$-space while keeping high acceptance probabilities. Randomness is injected by sampling the normal distribution $\pi(0) \sim \mathcal{N}(0, M)$. Due to the space limitation, we refer interested readers to the excellent review of HMC in Ref. [39].

We use the CNN architecture [40, 41] to model $\psi_{\alpha}(\theta)$, similar to Refs. [17, 42] and employ automatic differentiation (AD) techniques to obtain all derivatives $\partial_\theta$ in Eqs. 2. We concatenate $h_0 = \{ (\cos n \theta_k, \sin n \theta_k) \mid n = 1, \ldots, K \}$ along the input channel axis (Fig. 1) to enforce periodicity in $\theta$. We choose a simple 2-layer CNN model to control the number of parameters $P$. In addition nontrivially affecting the QGT inverse, the diagonalization cost grows as $O(P^4)$. Heuristically, introducing more parameters $\alpha$ requires more Monte Carlo samples to correctly resolve the averages in Eq. 2 and does not significantly contribute to simulation accuracy in our case.

![Figure 2: Results for different quenches from initial value $g_0 = 3$ on a two-dimensional 8 × 8 square lattice. Left: Potential energy, magnetization and angular variance as functions of real time. Right: A parametric plot of the mean rotor direction, parametrized by the real time $t$.](image)

$t + \delta t$. This update rule strongly resembles the quantum natural gradient optimization scheme [21, 23]. We use the Bogacki-Shampine Runge Kutta 3(2) solver. [35–37].
We simulate the effects instantaneous quenches – we approximate a ground state of the Hamiltonian in Eq. 1 using standard VMC methods and then preform time evolution under a different value of $g$. In Fig. 2, we choose a square $8 \times 8$ lattice, tracking the dynamics of the potential energy density $\epsilon_p(t) = -\frac{1}{N} \sum_{(k,l)} \langle \hat{n}_k \cdot \hat{n}_l \rangle_t$ and the average magnetization magnitude $M(t) = \frac{1}{N} \langle |\sum_k \hat{n}_k| \rangle_t$. Averages $\langle \cdot \rangle_t$ are performed with respect to the trial state at time $t$.

These observables were chosen as a proxy for thermalization. Across a wide range of quenches we observe convergence to their respective equilibrium values at $g = g_f$, see Fig. 2. We observe two dynamical regimes in relation to the quantum critical point $g_c \approx 4.25$, when $g < g_c$. For small quenches (left column of Fig. 2) we see slower equilibration with only fluctuations around the magnetization direction. For moderate to large quenches in Fig. 2, we observe a (transient) demagnetization of the sample and convergence to a new equilibrium state.

One also has access to global observables such as the Loschmidt echo with applications in the context of dynamical phase transitions [43] and quantum chaos [44]. It is defined as the (normalized) overlap $F(t) = |\langle 0 | \Psi(t) \rangle|^2$. Following Refs. [45, 46], we evaluate Monte Carlo estimators of $F(t)$. Expectedly, we find that $F$ decays with time, (see Fig. 3, left panel). For smaller quenches, the fidelity shoots back up to a nonzero value suggesting a finite overlap between the initial state and the long time “equilibrium” state after the quench. The latter may be interpreted as a signature of quenching between two Hamiltonians in the ordered phase.

We introduce another time scale $\tau_{1/2}$ defined as the time needed for the fidelity to decrease by 50%. We observe that $\tau_{1/2}$ increases linearly with the quench $g_f$. This result matches basic estimates given by the uncertainty relation $\Delta E \Delta t > 1/2$. Therefore, fidelity decay time can be lower bounded by $\Delta E^{-1}$, estimated using samples from the initial state $\psi_{\alpha(0)}$ [47]. The comparison in Fig. 3 (left, inset) demonstrates that the t-VMC method can be used to estimate quantities of experimental interest for system sizes unreachable by other wavefunction methods.

We compare our results to tensor-network time-evolving block decimation (TEBD) [48, 49] simulations for a one- and two-dimensional versions of the model. For all benchmarks, states were initialized to the coherent superposition of all basis states $|\psi(0)\rangle \propto \int d\theta |\theta\rangle$. Following Refs. [17, 50], we use $r(t) = D(\psi(t+\delta t),e^{-i\delta t\hat{H}}\psi(t))/D(\psi(t),e^{-i\delta t\hat{H}}\psi(t))$ as a figure of merit where $D(\cdot,\cdot)$ is the Fubini-Study distance on the Hilbert space $\mathcal{H}$. Intuitively, $r^2(t)$ measures an appropriately normalized measure of deviation between the full state $e^{-i\hat{H}\delta t}|\psi(t)\rangle$ after one time step $\delta t$ and its projection $|\psi_{\alpha(t+\delta t)}\rangle$. We plot the integrated error $R^2(t) = \int_{t_0}^{t_f} r^2(s) \, ds$ as an upper bound on the square of the integrated error $R(t) = \int_{0}^{t_f} r(s) \, ds$ due to the triangle inequality.

In Fig. 3 (right), we show that this algorithm performs well on a one-dimensional system of $N = \text{64}$ rotors where the growth of the so-called bond dimension $\chi$ is limited. The integrated residual $R^2(t)$ grows more rapidly for lower values of $g$ because the initial state $\psi(0)$ representing a more typical state in the disordered phase. In contrast to the 1D case, we observe that the TEBD method...
exponentially grows the MPS bond dimension $\chi$ past the cutoff $\chi_{\text{max}} = 1000$ at relatively short times. We see qualitative agreement between the two methods for early times, before $\chi$ grows to the point where further simulation is numerically prohibitively expensive.

Overall, both t-VMC and TEBD algorithms predict similar dynamical behavior where the comparison is possible. However, the number of parameters in the MPS grows exponentially due to entropy build-up, blocking tensor-network algorithms [51–53, 12] from scaling to higher dimensions and longer times. Continuous degrees of freedom amplify the problem due to infinite local basis.

4 Conclusion

We present a method to approximate unitary dynamics of continuous-variable quantum many-body systems, based on custom neural-network quantum states. The approach employs Hamiltonian Monte Carlo sampling and custom regularization of the quantum geometric tensor. Our calculations are able to access nontrivial local and global observables. Good agreement was found with tensor-network-based TEBD simulations for the case of one-dimensional systems of comparable size. Our approach paves the way for accurate non-equilibrium simulations of continuous systems at previously unexplored system sizes and evolution times, bridging the gap between simulation and experiment.

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Software libraries

The code used in this work has been packaged into an installable library and is (anonymously) available to reproduce any results in this work or explore new ones: github.com/Matematija/continuous-vmc.

It was built on JAX [54] for array manipulations, automatic differentiation for sampling and optimization and GPU support, Flax [55] for neural-network construction and manipulation and NumPy [56] and SciPy [57] for CPU array manipulations. Matplotlib [58] was used to produce figures.

References


