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Root-*N* Krylov-space correction-vectors for spectral functions with the density matrix renormalization group

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We propose a method to compute spectral functions of generic Hamiltonians using the density matrix renormalization group (DMRG) algorithm directly in the frequency domain, based on a modified Krylov space decomposition to compute the correction-vectors. Our approach entails the calculation of the root-N (N = 2 is the standard square root) of the Hamiltonian propagator using Krylov space decomposition, and repeating this procedure N times to obtain the actual correction-vector. We show that our method greatly alleviates the burden of keeping a large bond dimension at large target frequencies, a problem found with conventional correction-vector DMRG, while achieving better computational performance at large N. We apply our method to spin and charge spectral functions of t-J and Hubbard models in the challenging two-leg ladder geometry, and provide evidence that the root-N approach reaches a much improved spectral resolution compared to conventional correction-vector.

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I. INTRODUCTION

In condensed matter physics, several unusual properties of strongly correlated quantum materials are unveiled using *spectroscopic* techniques, such as angle resolved photoemission spectroscopy $(ARPES)^1$, inelastic neutron scattering (INS), and resonant inelastic x-ray scattering $(RIXS)^2$. These experimental probes do not provide a direct access to the ground state, but rather explore the low energy excitations of the system. Excitations spectra are esperimentally measured looking at the energy and momentum exchanged by the probe of each technique with the material: photo-emitted electron for ARPES, neutron for INS, photon for RIXS, and are theoretically encoded in *spectral functions*. The progressive improvement in momentum and energy resolution in experimental spectroscopic apparatus calls on the theory side for an equally significant improvement of the spectral functions calculations accuracy.

For a one-dimensional (1D) lattice Hamiltonian of size L, a generic spectral function can be defined as

$$O(q,\omega) = \frac{1}{L} \sum_{i,j} e^{iq(i-j)} \int_0^\infty dt e^{i(\omega+E_g)t} \langle \psi | \hat{O}_i e^{-i\hat{H}t} \hat{O}_j | \psi \rangle,$$
(1)

where $|\psi\rangle$ is the ground state of the system Hamiltonian \hat{H} , E_g is the ground state energy, q and ω are the momentum and frequency (or energy) of the electron in the material, and \hat{O}_j is the relevant operator involved in the scattering process of the specific technique acting locally on site "j" $(\hat{O}_j = \hat{c}_{j\sigma} \text{ for ARPES}, \hat{O}_j = \hat{S}_j^z \text{ for INS}, \text{ while special}$ care is needed for RIXS, as written in Ref.³).

In 1D, the most powerful method to compute spectral functions of arbitrary strongly correlated Hamiltonians is the density matrix renormalization group (DMRG)^{4,5}; the

DMRG is a variational but systematically exact algorithm to find a matrix product state (MPS) representation for the ground state of the system⁶. Spectral functions can be computed in the time-space domain using time dependent matrix product state methods^{7–10}. (For a recent review of the different variants, see Ref.¹¹) When using time evolution, the problem is to find an efficient MPS representation of the time-evolved vector

$$|x_j(t)\rangle = e^{-iHt}\hat{O}_j|\psi\rangle,\tag{2}$$

where the ground state of the Hamiltonian \hat{H} is locally modified by the \hat{O}_j , and the resulting state is evolved up to a very large (in principle "infinite") time. This evolution always grows the entanglement of the state, and thus spoils the compression of the MPS representation. Simulations are therefore typically stopped at some large or maximum time, and linear prediction⁹ or recursion methods¹² are needed to obtain a well behaved Fourier transform in frequency.

In this paper, we are concerned with the complementary approach of computing the spectral functions *directly in the frequency domain*. To discuss this case, it helps to rewrite the spectral function as

$$O(q,\omega) = \lim_{\eta \to 0} \frac{1}{L} \sum_{i,j} e^{iq(i-j)} \times \\ \times -\frac{1}{\pi} \operatorname{Im} \left[\langle \psi | \hat{O}_i \frac{1}{\omega - \hat{H} + E_g + i\eta} \hat{O}_j | \psi \rangle \right], \quad (3)$$

where one writes down the Hamiltonian propagator explicitly, and $\eta > 0$ is an arbitrary small extrinsic spectral broadening. Three are the approaches that are typically used by DMRG practitioners. Historically, the hybrid DMRG-Lanczos-vector methods were first introduced¹³

(refined using MPS more recently^{14,15}), then afterwards the correction-vector (CV) method^{16–21} and Chebyshev polynomial methods^{22–25} were proposed. In the CV method, one computes the real and imaginary part of the correction-vector

$$|x_j(\omega + i\eta)\rangle = \frac{1}{\omega - \hat{H} + E_g + i\eta} \hat{O}_j |\psi\rangle \tag{4}$$

at fixed frequency ω , finite broadening η , and then computes the spectral function in real-frequency space as a stardard overlap $\langle \psi | \hat{O}_i | x_j (\omega + i\eta) \rangle$. The real and imaginary part of the correction-vector are typically obtained by solving for coupled matrix equations using conjugategradient methods¹⁶, or by minimizing a properly defined functional^{18,19}. Ref.²⁰ formulated the algorithm in MPS language.

In 2016, we proposed²¹ an alternative method to compute directly the correction-vectors using a Krylov space expansion of the Hamiltonian operator constructed starting from the locally modified MPS $|\phi\rangle = \hat{O}_j |\psi\rangle$. In all these cases, the entanglement content of the correctionvectors is large, and it can be very large for large frequencies. This makes standard CV DMRG simulations very expensive for Hamiltonians beyond spin systems or for large lattices.

In 2011, Holzner et al.²² proposed a MPS method to compute a Chebyshev polynomial expansion (truncated at some order N) of the spectral function (CheMPS). In this approach, the Chebyshev momenta can be obtained from overlaps of a properly defined series of Chebyshev vectors. The main advantage of CheMPS lies in the small entanglement that each Chebyshev vector has, because the method *redistributes* the large entanglement of the correction-vectors $|x_j(\omega + i\eta)\rangle$ for different frequencies (or alternatively the time-evolved state $|x_j(t)\rangle$) over the entire series of Chebyshev vectors.

Inspired by this idea, we here propose a method to compute a generalized correction-vector with smaller entanglement content, the root-N correction vector, defined as

$$|x_j^{1/N}(\omega+i\eta)\rangle = \left(\frac{1}{\omega-\hat{H}+E_g+i\eta}\right)^{1/N} \hat{O}_j|\psi\rangle.$$
 (5)

The idea is to construct the actual correction-vector as the final vector of the series $\{|x_j^{p/N}(\omega + i\eta)\rangle\}_{p\in[1,N]}$ after N applications of the root-N propagator. At first sight, it seems that, if N is sufficiently large, constructing the entire series of vectors just adds a computational overhead compared to the standard DMRG CV algorithm, because only the final vector of the series is actually needed for the spectral function calculation. Yet we will show that the entanglement content of the series slowly builds up with p, and therefore, going through many intermediate steps is more efficient than the conventional DMRG CV algorithm, which tries to compute the last element of the series in one step only. The paper is organized as follows. Section II.A introduces the main steps of the algorithm; section II.B analyzes the algorithm's computational performance, and the entanglement content of the root-N correction-vectors in the test case of a Heisenberg model in the two-leg ladder geometry. Section II.C applies our root-N method to compute spin and charge spectral functions of doped t-Jand Hubbard models in the challenging two-leg ladder geometry, showing how our method improves the spectral resolution and increases the signal-to-noise ratio at large frequencies. Finally, we present our conclusions and outlook.

II. METHOD AND RESULTS

A. root-N CV method algorithm

The algorithm follows five steps. We assume a standard DMRG approach but provide the main step of the algorithm in MPS language in Appendix A.

1. Compute the ground state wave function with the DMRG.

For each frequency ω , repeat the steps 2–4 to cover the desired interval $[\omega_{\min}, \omega_{\max}]$ with some step $\Delta \omega > 0$:

2. Apply the operator O_j at the center of the chain and build the p = 1 root-N correction vector $|x_j^{p/N}(\omega + i\eta)\rangle$ as in Eq. (5). This can be done using conventional DMRG as described in Ref.²¹. Appendix A describes in detail the algorithm in MPS language. In this stage, as in the conventional CV method, the sources of error are two: the Lanczos error in the tridiagonal decomposition of the Hamiltonian (or effective Hamiltonian in MPS language) indicated below by $\epsilon_{\text{Tridiag}}$; the SVD error of the multi-targeting DMRG procedure (state-averaging in MPS language).

Repeat step 3 until the Nth root-N correction vector is constructed and optimized, then go to step 4.

- 3. Build the p+1 root-N correction vector $|x_j^{(p+1)/N}(\omega + i\eta)\rangle$ from the previous one assuming it as a starting point for the Krylov space decomposition of the Hamiltonian. A few DMRG sweeps are performed until a desired convergence is reached.
- 4. Measure the spectral function in real-frequency space as the overlap $\langle \psi | \hat{O}_i | x_j (\omega + i\eta) \rangle$; this part is the same as in conventional DMRG CV.
- 5. Fourier Transform the overlap $\langle \psi | \hat{O}_i | x_j(\omega + i\eta) \rangle$ to get the dynamical structure factor in momentum space, $O(\mathbf{q}, \omega) = \frac{1}{L} \sum_{i,j} e^{iq(i-j)} \langle \psi | \hat{O}_i | x_j(\omega + i\eta) \rangle$.



FIG. 1: Convergence analysis of the root-N Krylov space Correction-Vector (CV) DMRG method: the Heisenberg ladder. Panels (a)-(b) report the $q_y = 0, \pi$ components of the $S(q_x, q_y, \omega)$ using the standard Krylov space CV approach. A ladder with $J_y = 2J_x$ is simulated. Length is $L = 50 \times 2$, broadening $\eta = 0.1$, and resolution step $\delta \omega = 0.1$ (units are set by $J_x = 1$). Panels (c)-(d) report the $q_y = 0, \pi$ components of the $S(q_x, q_y, \omega)$ using the root-N Krylov space CV method using N = 20. Panels (e)-(f) show specific momentum-energy line cuts of the dynamical spin structure computed in panels (a)-(d) for different values of the *root-exponent* N. Numerical fluctuations and instabilities are removed, and quality of the spectra is clearly improved as N is increased. Panels (g)-(h) show the same line cuts as in Panels (e)-(f) but with at fixed N, and increasing the number of DMRG states. When using the root-N CV method with N = 4, a substantially smaller number of states m = 200 - 650 < 1000 suffices to get better quality results than with the standard Krylov space DMRG CV approach.

To clarify the main steps of the algorithm, we draw an analogy with the adaptive time-step targeting approach introduced for time-dependent DMRG in the seminal paper by Feiguin and White²⁶. In this case, one constructs the time-evolved vector $|x_j(\delta t)\rangle = U(\delta t)\hat{O}_j|\psi\rangle$ only for small time step intervals of length $\delta t = t/N$, and where $U(\delta t) = e^{-i\hat{H}\delta t}$ is the time evolution propagator. To get the final time-evolved vector at time t, one repeatedly applies $U(\delta t)$ to the MPS. In practice one does not build the evolution operator $U(\delta t)$ in the local basis but rather directly construct the vector $|x_j(\delta t)\rangle = U(\delta t)\hat{O}_j|\psi\rangle$ using a Krylov space decomposition of the Hamiltonian (or effective Hamiltonian in MPS language).

In our proposed root-N Krylov space approach, we introduce a propagator in a fictitious time space s as $|x_j(\delta s)\rangle = W(\delta s)\hat{O}_j|\psi\rangle$, where $\delta s = 1/N$, and where $W(\delta s) = e^{-\hat{K}\delta s}$, with $\hat{K} = \ln [\omega + E_g + i\eta - \hat{H}]$. Clearly, if we apply $W(\delta s) N$ times to the initial vector $\hat{O}_j|\psi\rangle$ we obtain the desired standard correction vector. In other words, we formally define and solve an auxiliary differential equation

$$\frac{d}{ds}|x_j^s(\omega+i\eta)\rangle = -\ln\left[\omega + E_g + i\eta - \hat{H}\right]|x_j^s(\omega+i\eta)\rangle, \ (6)$$

such that at time s we have the solution

$$|x_j^s(\omega+i\eta)\rangle = \left(\frac{1}{\omega-\hat{H}+E_g+i\eta}\right)^s |x_j^0(\omega+i\eta)\rangle, \quad (7)$$

with the initial condition being $|x_j^0(\omega + i\eta)\rangle = \hat{O}_j|\psi\rangle$.

In this construction, 1/N plays the role of a *small* parameter to compute the resolvent in the standard CV approach. We will show that this method is especially useful at large target frequencies, where large bond dimensions (or DMRG states) are typically needed.

B. Convergence analysis and Computational performance: the Heisenberg model on a two-leg ladder as a case study

We begin by testing our root-N CV method by applying it to an isotropic Heisenberg model on a two-leg ladder geometry. (The supplemental material²⁷ provides computational details for all the models considered in this work.) The Heisenberg Hamiltonian on a two leg ladder with open boundary conditions and size $L = L_x \times 2$ is defined as

$$H_{\text{Heis}} = J_x \sum_{i=1;\gamma=1,2}^{L_x - 1} \mathbf{S}_{i,\gamma} \cdot \mathbf{S}_{i+1,\gamma} + J_y \sum_{i=1}^{L_x} \mathbf{S}_{i,1} \cdot \mathbf{S}_{i,2}, \quad (8)$$

where $\mathbf{S}_{i,\gamma} \equiv \{S_{i,\gamma}^x, S_{i,\gamma}^y, S_{i,\gamma}^z\}$ describe the spin 1/2 operators on site *i* and ladder leg γ . In this work, antiferromagnetic exchange interactions along both the leg and rung directions are assumed, with $J_y = 2J_x$. The spin structure factor $S(\mathbf{q}, \omega)$ with $\mathbf{q} \equiv (q_x, q_y)$ can be defined



FIG. 2: Entanglement Entropy of the root-N Krylov correction vectors and computational performance of the method. Panel (a): Entanglement Entropy computed with conventional CV (dark green) compared to the same quantity computed with the root-N CV; N = 20 and different p values (indicated by orange lines, and thickness increasing with p), as a function of ω . We have used a Heisenberg two-leg ladder with $J_y = 2J_x$, length $L = 50 \times 2$, as in Fig. 1. (b) Accumulated CPU time in hours (obtained summing all the CPU times of the CV simulations in the frequency interval $\omega \in [2.5, 6]$) for different values of the DMRG truncation error ϵ_{Trunc} and (c) for different values of the Krylov space threshold accuracy parameter $\epsilon_{\text{Tridiag}}$. as a function of the root-exponent N. The simulations were run on a single Intel Xeon CPU E5-1620 CPU node. When compared with the standard Krylov space Correction-Vector method, the performance of the root-N approach is superior for sufficiently large N.

as

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$$S(\mathbf{q},\omega) = \frac{1}{2L_x} \sum_{j=1;\gamma}^{L_x} e^{i[q_x(j-i)+q_y(\gamma-\gamma')]} \times \langle \psi | \mathbf{S}_{j,\gamma} \frac{1}{\omega - H + i\eta} \mathbf{S}_{i,\gamma'} | \psi \rangle, \qquad (9)$$

where the *center* point is chosen in the middle of the leg 1, $(i, \gamma') = (L_x/2, 1)$.

Figure 1a-b reports spectral maps of the two components $q_y = 0, \pi$ of the dynamical spin structure factor $S(\mathbf{q},\omega)$ as a function of the momentum transfer q_x along the leg direction, and of the frequency. These are obtained with conventional CV as in $\operatorname{Ref}^{21,28}$ on a system size of length $L = 50 \times 2$ and with an extrinsic broadening parameter $\eta = 0.1 J_x$. By comparison, Fig. 1c-d reports results obtained using the root-N CV method with N = 8. In both cases, we have used a maximum $m = m_{\text{max}} = 1000$ DMRG states and a minimum $m_{\min} = 200$, keeping the truncation error below 10^{-7} . Our DMRG calculations were carried out with the DMRG++ code²⁹. (Please see the description around Eq. (A4) in Appendix A for the definition of the extended MPS which is optimized by SVD in the root-N CV algorithm.) We clearly notice an overall improved spectrum in this case with respect to the conventional CV method. We analyze below the spectral features in more detail.

Figure 1e-f shows momentum $q_x = 0.52\pi$ line cuts of the spin spectra for the $q_y = 0, \pi$ components in the root-N CV method. The data shows that by increasing the *root-exponent* N numerical fluctuations and instabilities are removed with respect to the conventional CV results. The red curve in Fig. 1e-f shows that the conventional CV approach can yield negative values for certain frequencies. As finite size effects are small for a $L = 50 \times 2$ ladder,

these are clearly artifacts of the CV method which might spoil important properties of the spectral functions such as sum rules. On the contrary, the root-N CV approach shows always positive values which progressively improve upon increasing the root-exponent N. Figure 1g-h shows how well the root-N method converges with respect to the number of DMRG states. Contrary to panels (a)-(d), in these panels the data for m < 1000 was obtained by imposing zero truncation error in the DMRG SVDs, therefore setting $m = m_{\text{max}} = m_{\text{min}}$. Our data shows that at fixed root-exponent N = 4, a substantially smaller number of DMRG states m = 200-650 < 1000 is sufficient to get better quality results than with the conventional CV approach. As we will show next, this improvement can be understood by the much lower entanglement content of the root-N correction-vectors.

Figure 2a shows indeed that the entanglement content of the root-N correction vectors is smaller than the actual (conventional) correction-vector. In this calculation, to compute the entanglement entropy of the expanded MPS for root-N correction vectors (Appendix Eq. (A4) has the definition), we have used a maximum m = 2000DMRG states (and a minimum $m_{\min} = 200$), keeping the truncation error below 10^{-8} in both methods. It is nice to see that the entanglement entropy of the extended MPS in the root-N CV method is very close to that of the conventional CV in the lower frequency range investigated $\omega \in [0, \omega^*]$, with $\omega^* \simeq 4.5$. For larger frequencies, the root- $N \, \mathrm{CV}$ approach truncates the entanglement contained in the conventional CV vector, showing that a larger rootexponent N or a larger number of DMRG states should be considered. Yet we highlight that this truncation does not show instablibilities or fluctuations as in the conventional CV approach.

In the same range of frequencies ($\omega \in [2.5, 6]$), we have

monitored the accumulated CPU times taken for the simulations to complete in the two methods (Fig. 2b). We observe that, for moderately small root-exponent N, the root-N CV method can be actually slower than the conventional CV method, assuming the same truncation error ϵ_{Trunc} and truncation threshold $\epsilon_{\text{Tridiag}}$ for the Hamiltonian matrix decomposition in the Krylov space (see Appendix B for a more detailed definition of $\epsilon_{\text{Tridiag}}$). If the root-exponent N is sufficiently large, the root-N CV method is seen to be computationally more efficient than the conventional CV method. Indeed, when the entanglement is decomposed in smaller chunks by considering a larger N, the root-N method becomes faster even though many more optimizations and Lanczos decompositions are actually performed. Eventually, however, if the rootexponent N is very big, the increased number of DMRG sweeps and iterations required to compute the larger number of root-N vectors becomes naturally detrimental for computational performance, as the accumulated CPU times are seen to increase linearly with N.

Figure 2b further shows that, at fixed the root-exponent N, as one decreases the requested DMRG truncation error ϵ_{Trunc} the accumulated CPU times are bigger. This is because of the larger computational load of the SVD decomposition of the MPS tensors in the multitargeting or state averaging approach.

Figure 2c ends this subsection by showing further how the accumulated CPU times vary as a function of the truncation thresold $\epsilon_{\text{Tridiag}}$ for the Hamiltonian matrix decomposition in the Krylov space, at fixed DMRG truncation error ϵ_{Trunc} . A smaller threshold requires a larger number of steps and thus a bigger Krylov space, requiring a larger CPU time to converge and complete the simulations.

C. Correlation functions of t-J and Hubbard models

In this section, we apply the root-N CV method to the more challenging t-J and Hubbard models on a two leg ladder geometry.

The t-J Hamiltonian is defined as

$$H_{t-J} = -t_x \sum_{i=1;\gamma=1,2;\sigma}^{L_x-1} \left(c_{i,\gamma,\sigma}^{\dagger} c_{i+1,\gamma,\sigma} + \text{h.c.} \right)$$
$$- t_y \sum_{i=1;\sigma}^{L_x} \left(c_{i,1,\sigma}^{\dagger} c_{i,2,\sigma} + \text{h.c.} \right)$$
$$+ J_x \sum_{i=1;\gamma=1,2}^{L_x-1} \left(\mathbf{S}_{i,\gamma} \cdot \mathbf{S}_{i+1,\gamma} - \frac{n_{i,\gamma} n_{i+1,\gamma}}{4} \right)$$
$$+ J_y \sum_{i=1}^{L_x} \left(\mathbf{S}_{i,1} \cdot \mathbf{S}_{i,2} - \frac{n_{i,1} n_{i,2}}{4} \right), \quad (10)$$

where $c_{i,\gamma,\sigma}^{\dagger}$ $(c_{i,\gamma,\sigma})$ is the electron creation (annihilation) operator on site *i*, ladder leg γ with spin polarization σ , while $n_{i,\gamma} = \sum_{\sigma} c_{i,\gamma,\sigma}^{\dagger} c_{i,\gamma,\sigma}$ is the electron number operator. The Hubbard Hamiltonian is

$$H_{\text{Hub}} = -t_x \sum_{i=1;\gamma=1,2;\sigma}^{L_x-1} \left(c_{i,\gamma,\sigma}^{\dagger} c_{i+1,\gamma,\sigma} + \text{h.c.} \right) - t_y \sum_{i=1;\sigma}^{L_x} \left(c_{i,1,\sigma}^{\dagger} c_{i,2,\sigma} + \text{h.c.} \right) + U \sum_{i=1;\gamma=1,2}^{L_x} n_{i,\gamma,\uparrow} n_{i,\gamma,\downarrow}.$$
(11)

For both models defined above, the spin structure factor $S(\mathbf{q}, \omega)$ is defined as in the Heisenberg model case (Eq. (9)). Analogously, the charge structure factor is

$$N(\mathbf{q},\omega) = \frac{1}{2L_x} \sum_{j=1;\gamma}^{L_x} e^{i[q_x(j-i)+q_y(\gamma-\gamma')]} \times \langle \psi | \delta n_{j,\gamma} \frac{1}{\omega - H + i\eta} \delta n_{i,\gamma'} | \psi \rangle, \qquad (12)$$

where $\delta n_{j,\gamma} \equiv n_{j,\gamma} - \langle \psi | n_{j,\gamma} | \psi \rangle$, where $|\psi \rangle$ is the ground state of the system.

We start discussing the *t-J* model results, comparing the root-*N* method against the results obtained to the conventional CV approach. We calculate both spin and charge dynamical structure factors for a doped ladder with $N_{\rm el} = 0.88L$, corresponding to 12% hole doping, and with lattice size $L = 50 \times 2$. In this case, we use a maximum m = 1200 DMRG states for both methods (and a minimum $m_{\rm min} = 200$), in order to keep the truncation error below 10^{-8} .

Figure 3a-d shows the comparison for the dynamical spin structure factor $S(\mathbf{q}, \omega)$. We note that for N = 8 the root-N CV method yields results that are practically identical to those obtained with the CV method. Yet



FIG. 3: Dynamical structure factors for a t-J two-leg ladder. Panels (a)-(b) ((e)-(f)) report the $q_y = 0, \pi$ components of the $S(q_x, q_y, \omega)$ $(N(q_x, q_y, \omega))$ using the standard Krylov space correction-vector approach. A ladder with $t_y = t_x = t = 1$, $J_x = J_y = 0.5t$ is simulated. Length is $L = 50 \times 2$, number of electrons $N_{\rm el} = 0.88L$, broadening $\eta = 0.1$, and resolution step $\delta\omega = 0.1$ (units are set by t = 1). Panels (c)-(d) ((g)-(h)) report the $q_y = 0, \pi$ components of the $S(q_x, q_y, \omega)$ ($N(q_x, q_y, \omega)$) using the root-N Krylov space correction-vector using N = 8.

 $\check{q_x}$



FIG. 4: Dynamical structure factors for a Hubbard two-leg ladder. The $q_y = 0$ and $q_y = \pi$ components of the dynamical spin structure factor $S(q_x, q_y, \omega)$ using, in (a)-(b) the conventional Krylov space DMRG CV approach, against in (c)-(d) the root-N Krylov space CV method with N = 4 (a maximum of m = 3000 DMRG states were used.) The $q_u = 0$ and $q_y = \pi$ components of the dynamical charge correlation $N(q_x, q_y, \omega)$, using, in (e)-(f) the conventional Krylov space DMRG CV, against in (g)-(h) the root-N Krylov space CV with N = 4 (a maximum of m = 2000 DMRG states were used.). A ladder $L = 50 \times 2$ ladder with $t_y = t_x = t = 1$, U = 8t is simulated with $N_{\rm el} = 0.88L$ electrons, broadening $\eta = 0.1$, and resolution step $\delta\omega = 0.025$ for $S(q_x, q_y, \omega)$, and $\delta\omega = 0.1$ for $N(q_x, q_y, \omega)$. Units are set by t = 1.

the root-N method provides a much better frequency resolution or larger signal-to-noise ratio for the more challenging dynamical charge structure factor $N(\mathbf{q}, \omega)$, where we also obtain quantitative agreement.

In summary, the dynamical spin structure factor $S(\mathbf{q}, \omega)$ is practically identical in the two methods, while when considering the dynamical charge structure factor $N(\mathbf{q}, \omega)$ besides obtaining qualitative agreement, the root-N provides a much better frequency resolution (or a larger signal-to-noise ratio, as we recall here that in both methods the same broadening η was used).

We now focus on the Hubbard model where minor differences in the results between the two methods can be observed when a moderately small root-exponent N is used in the root-N CV method.

As in the *t-J* case, we consider spin as well as charge dynamical structure factors for a doped ladder ($N_{\rm el} = 0.88L$, 12% hole doping) with system size $L = 50 \times 2$. We consider an isotropic ladder with parameters $t_x = t_y = t = 1$ and U = 8t. Spin and charge structure factors for Hubbard ladders were already studied and discussed by us in Refs.^{30–34}, where the conventional Krylov space CV method was used. Figure 4 uses a maximum m = 3000 DMRG states in both methods for $S(\mathbf{q}, \omega)$ while a maximum of m = 2000 was used for $N(\mathbf{q}, \omega)$. In both cases, the minimum number of DMRG states was $m_{\rm min} = 200$, and the truncation error was kept smaller than 10^{-7} .

Figure 4a-d shows the comparison for the dynamical spin structure factor $S(\mathbf{q}, \omega)$. For N = 4 the root-N CV method gives results practically identical to the CV method, and only minor quantitative differences can be observed. For example, in the root-N method, the broad two-triplon excitation band in the spin structure factor $S(q_x, q_y = 0, \omega)$ appears to be sharper than in the conventional CV method. In the $q_y = \pi$ component, instead, the main spectral features at the incommensurate wave-vector $q_x \simeq 0.88\pi$ appear slightly broader in the conventional CV method as a function of frequency, at low frequencies. From this analysis, we conclude that even a moderately small root-exponent N is sufficient to get a better converged spin spectral function using the root-N CV method.

These observations are relevant when comparing DMRG spectral data with RIXS³⁵ and INS³⁶ experiments in the challenging "telephone number" cuprates, experimental data that recently has became available for the doped regime³⁴.

Finally, we discuss the dynamical charge structure factor, which is of interest in RIXS measurements of the charge-transfer band excitations in ladder cuprates. When a Hubbard ladder is doped with holes with respect to half-filling, we observe two branches in the $N(\mathbf{q}, \omega)$: the first one at low-energy corresponds to in-band particlehole excitations across the Fermi level. The high-energy band describes charge-transfer electronic exitations above the Mott gap. Figure 4e-h shows that the root-N CV method provides high quality spectral data with no appreciable shifts (downwards or upwards) of the main features. (Please remember that we are using the same η for both methods.) Yet some spectral weight redistribution can be noted: spectral intensity on the high-energy charge-transfer band appears more intense in the root-N CV method compared to the conventional CV method. We conclude that in this case, even though very good results can be obtained with a modest root-exponent N, one should prefer simulations with the largest possible N in order to get the best results from our root-N method.

III. DISCUSSIONS AND CONCLUSIONS

In this work, we have proposed a method to compute generic spectral functions of strongly correlated Halmiltonians using generalized correction-vectors with smaller entanglement content: the root-N CV method. The idea behind the root-N CV draws inspiration in part from time dependent MPS methods, and in part from the Chebyshev MPS approach. The CheMPS method helps in computing spectral functions but, as was highlighted recently²⁵, while resolving accurately the low-energy part of the spectral functions, CheMPS cannot resolve the high-energy spectrum accurately because an energy-truncation of the Chebyshev vectors is in general required. To avoid this issue, Xie et al.²⁵ have proposed a reorthogonalization scheme for the Chebyshev vectors (ReCheMPS). Nevertheless, if the target frequency window for the spectral function is chosen to be much smaller than the many body width of the system (this should be in general done to increase the frequency resolution), an energy truncation might still be required. There is evidence that the energytruncation procedure severely limits the applicability of the CheMPS or ReCheMPS methods in challenging cases as in Hubbard or t-J models, as in these cases it likely becomes a necessary step of the algorithm, mainly because the many-body bandwidth is in general much larger than the spectral support of typical spectral functions. When the energy truncation is performed, several Krylov space projections as Chebyshev recurrence steps are required, rendering the method as computationally demanding as the conventional CV method.

Going back to the root-N CV, this publication has showed that when the root-exponent N is sufficiently large, the root-N CV performance becomes better than that of the conventional CV, because the former method handles much less entangled correction-vectors. In particular, we have shown evidence that in the Heisenberg and t-J models the root-N CV method improves even the quality of the spectral functions, and provides a better frequency spectral resolution (or a larger signal-to-noise ratio). Larger N values in the root-N CV method require more sweeping of the lattice, but do not affect much CPU times, because each sweep is faster than using smaller Nvalues.

Finally, the challenging Hubbard model requires a careful use of our root-N CV method: while moderately small root-exponents N give very good results for the main spectral features, our data shows only minor differences with respect to the conventional CV method, which however should be taken into account when high-precision experimental results are available.

We believe that root-*N* correction-vector DMRG will become a much used method, not only when high precision spectral data is sought, but also when high performance is required, performance better than the computationally expensive conventional CV method.

The root-N method should also facilitate high precision spectral function calculations in finite width cylinders, where better computational methods are currently needed. These cylinders try to approach the two-dimensional models that are at the frontier of what DMRG can do, and they need a very large computational effort to simulate.

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Appendix A: MPS algorithm to build the root-N correction-vector

Let us introduce a Matrix Product State representing the ground state of the system for L sites and open boundary conditions (we use a notation similar to Ref.¹¹)

$$|\psi\rangle = \sum_{\substack{\sigma_1...\sigma_L \\ m_0...m_L}} M_{1;m_0,m_1}^{\sigma_1} ... M_{L;m_{L-1},m_L}^{\sigma_L} |\sigma_1...\sigma_L\rangle, \quad (A1)$$

where m_i are the bond dimensions or virtual indices (with m_0 and m_L 1-dimensional dummy indices), and σ_i represent the physical indices of the many-body state of the system. Formally, let us define the tensors $\bar{\psi}_{L,j-1} \equiv (M_1^{\dagger}, ..., M_{j-1}^{\dagger})$ and $\bar{\psi}_{R,j+1} \equiv (M_{j+1}^{\dagger}, ..., M_L^{\dagger})$ which constitute a left and right map, respectively, from the joint Hilbert space on sites 1 through j-1 onto the bond space m_{i-1} , and from the joint Hilbert space on sites j + 1 through L onto the bond space m_j . If we apply these maps to the MPS $|\psi\rangle$, we can obtain the effective state at site j, $|\psi_i^{\text{eff}}\rangle$; see Fig. 5a. When $|\psi\rangle$ is in a MPS mixed-canonical form, $|\psi_j^{\text{eff}}\rangle$ equals the 3-rank tensor $M_{j,m_{j-1}m_j}$ in the MPS at site j, which is often interpreted as a vector of dimensions $(d_j \times m_{j-1} \times m_j)$, where d_i is the local physical Hilbert space dimension. Similarly, the Hamiltonian \hat{H} , in matrix product operator (MPO) form, acts between the maps defined above (and their conjugates, $\psi_{L,j-1} \psi_{R,j+1}$; see Fig. 5b) to yield an effective single site Hamiltonian \hat{H}_{j}^{eff} . This procedure can also be defined in the space of two-sites. A computer program never needs to explicitly construct \hat{H}_{i}^{eff} , but only evaluates its action on $|\psi_i^{\text{eff}}\rangle$.

Using $|\psi_j^{\text{eff}}\rangle$ and \hat{H}_j^{eff} , we construct three local MPS tensors. The first one is obtained by applying the operator \hat{O}_j on $|\psi_j^{\text{eff}}\rangle$, yielding $|\phi\rangle = \hat{O}_j |\psi\rangle$. The MPS $|\phi\rangle$ has all the tensors equal to those of $|\psi\rangle$ except for the one at site $j, M'_{j;m_{j-1},m_j}$

$$M_{j;m_{j-1},m_{j}}^{\prime \sigma_{j}} = \sum_{\sigma_{j}^{\prime}} O_{j}^{\sigma_{j}\sigma_{j}^{\prime}} M_{j;m_{j-1},m_{j}}^{\sigma_{j}}$$
(A2)

We then construct the (real and imaginary part of the) root-N correction-vector by Krylov space decomposition of the Hamiltonian \hat{H}_{j}^{eff}

$$[X(\omega+i\eta)]_{j;m_{j-1},m_{j}}^{\sigma_{j}} = \sum_{\substack{l,l'nn'\\\sigma'_{j},m'_{j-1},m'_{j}}} T^{\dagger}_{l;\sigma_{j},m_{j-1},m_{j}} P^{\dagger}_{ln} \frac{1}{[\omega-\epsilon_{j}^{\text{eff}}\delta_{nn'}+E_{g}+i\eta]^{1/N}} P_{n'l'} T_{l';\sigma'_{j},m'_{j-1},m'_{j}} M'_{j;m'_{j-1},m'_{j}}$$
(A3)



FIG. 5: Effective local state vector and Hamiltonian. Panel (a): Effective state $|\psi_j^{\text{eff}}\rangle$ obtained by projecting the MPS by the maps $\bar{\psi}_{L,j-1} \equiv (M_1^{\dagger},...,M_{j-1}^{\dagger})$ and $\bar{\psi}_{R,j+1} \equiv$ $(M_{i+1}^{\dagger},...,M_L^{\dagger})$. If $|\psi\rangle$ is a mixed-canonical MPS representation, then simply $|\psi_j^{\text{eff}}\rangle = M_j$. Panel (b): Effective (onesite) Hamiltonian obtained by projecting \hat{H} using the maps $\{\bar{\psi}_{L,j-1}, \psi_{L,j-1}, \bar{\psi}_{R,j+1}, \psi_{R,j+1}\}$ defined above. Analogous definitions can be given in the two-site case. Panel (c): (left) Graphical representation of the diagonal effective Hamiltonian projected onto the Kryrol space. Representation of the Krylov projection operator (triangular yellow tensor) $T_{l\alpha}$ where $\alpha = \{\sigma_j, m_{j-1}, m_j\}$ joins three indices so that it can be represented as a matrix (right). \hat{T} tridiagonalizes \hat{H}_{j}^{eff} of panel (b), $\hat{H}_i^{\text{Tridiag,eff}} = T^{\dagger} \hat{H}_i^{\text{eff}} T$, to the smaller Krylov space spanned by the index l, dim $[l] \ll \dim[\alpha] = d_j \times \dim[m_{j-1}] \times \dim[m_j]$. P_{ln} (green tensor) diagonalizes $\hat{H}_j^{\text{Tridiag,eff}}$.

where $T_{l';\sigma_j,m_{j-1},m_j}$ tridiagonalizes \hat{H}_j^{eff} , $\hat{H}_j^{\text{Tridiag,eff}} = T^{\dagger}\hat{H}_j^{\text{eff}}T$, to the smaller Krylov space spanned by the index l, $\dim[l] << d_j \times \dim[m_{j-1}] \times \dim[m_j]$. P_{ln} diagonalizes $\hat{H}_j^{\text{Tridiag,eff}}$, $\hat{H}_j^{\text{diag,eff}} = P^{\dagger}\hat{H}_j^{\text{Tridiag,eff}}P$, where ϵ_j^{eff} are the eigenvalues of $\hat{H}_j^{\text{diag,eff}}$. How is the Krylov space tridiagonalization of \hat{H}_j^{eff} stopped? In practice, we compare the lowest eigenvalue of $\hat{H}_j^{\text{diag,eff}}$, $\epsilon_{\min} = \{\epsilon_j^{\text{eff}}[k]\}_{\min}$ at iteration k and k + 1, and exit the loop when the error breaks below a certain threshold. In this work, we have set $\epsilon_{\text{Tridiag}}$ to a value not too small, in order to avoid the

proliferation of Krylov vectors (and thus Lanczos iterations), and their reorthogonalizations. In general, the three states $|\phi\rangle$, $|X^{\text{Re}}\rangle$, $|X^{\text{Im}}\rangle$ will be represented in a bad basis of the environments ψ_L and ψ_R which are optimized to represent the original state $|\psi\rangle$. To expand these bases, we use state-averaging of the four states, which is equivalent to targeting more than one state in conventional DMRG language. In MPS language, as explained in Ref.³⁷, the state- averaging is done by creating an extra index which labels the states involved. One formally considers an expanded MPS representing a mixed state

$$\begin{pmatrix} |\psi\rangle\\|\phi\rangle\\|X^{\text{Re}}\rangle\\|X^{\text{Im}}\rangle \end{pmatrix} = \sum_{\sigma_1...\sigma_L} A_1^{\prime,\bar{\sigma}_1}...C_j^{\prime,\bar{\sigma}_j}...B_L^{\prime,\bar{\sigma}_L} |\sigma_1...\sigma_L\rangle$$

$$= \sum_{\sigma_1...\sigma_L} \begin{pmatrix} A_1^{\sigma_1}[\psi] & 0 & 0 & 0\\ 0 & A_1^{\sigma_1}[\phi] & 0 & 0\\ 0 & 0 & A_1^{\sigma_1}[X^{\text{Re}}] & 0\\ 0 & 0 & 0 & A_1^{\sigma_1}[X^{\text{Im}}] \end{pmatrix} ... \begin{pmatrix} C_j^{\sigma_j}[\psi]\\C_j^{\sigma_j}[\phi]\\C_j^{\sigma_j}[X^{\text{Re}}]\\C_j^{\sigma_j}[X^{\text{Im}}] \end{pmatrix} ... \begin{pmatrix} B_L^{\sigma_L}[\psi] & 0 & 0 & 0\\ 0 & B_L^{\sigma_L}[\phi] & 0 & 0\\ 0 & 0 & B_L^{\sigma_L}[X^{\text{Re}}] & 0\\ 0 & 0 & 0 & B_L^{\sigma_L}[X^{\text{Im}}] \end{pmatrix} |\sigma_1...\sigma_L\rangle$$

where $C_{j;m'_{j-1},m'_j}^{\prime,\bar{\sigma}_j}$ has four components (representing the four *targeted* vectors) and it has extended bond dimensions $m'_{j-1} = m_{j-1}^{[\psi]} + m_{j-1}^{[\phi]} + m_{j-1}^{[X^{\text{Im}}]} + m_{j-1}^{[X^{\text{Im}}]}$, $m'_j = m_j^{[\psi]} + m_j^{[\phi]} + m_j^{[X^{\text{Re}}]} + m_j^{[X^{\text{Im}}]}$. Here, the notation in terms of A and B tensors underlines a mixed-canonical representation of all the MPSs. By SVD compression, one has

$$C_j^{\prime,\sigma_j} = U_j^{\prime,\sigma_j} S_j^{\prime} V_j^{\prime,\dagger}.$$
 (A4)

As in conventional DMRG, one can also introduce different weights in the direct sum and perform a SVD of the weighted sum of the reduced density matrix $\rho' = \sum_{k=0}^{3} w_k \rho_k$ Once this procedure is performed at site j, one can proceed updating all the tensors at site j + 1. In formulas,

$$C_{j+1}' = \begin{pmatrix} C_{j+1}^{\sigma_{j+1}}[\psi] \\ C_{j+1}^{\sigma_{j+1}}[\phi] \\ C_{j+1}^{\sigma_{j+1}}[X^{\mathrm{Re}}] \\ C_{j+1}^{\sigma_{j+1}}[X^{\mathrm{Im}}] \end{pmatrix} = \begin{pmatrix} U_{j}^{[\prime,\dagger,\sigma_{j}]}C_{j}^{\sigma_{j}}[\psi]B_{j+1}^{\sigma_{j+1}}[\psi] \\ U_{j}^{[\prime,\dagger,\sigma_{j}]}C_{j}^{\sigma_{j}}[\phi]B_{j+1}^{\sigma_{j+1}}[\phi] \\ U_{j}^{[\prime,\dagger,\sigma_{j}]}C_{j}^{\sigma_{j}}[X^{\mathrm{Re}}]B_{j+1}^{\sigma_{j+1}}[X^{\mathrm{Re}}] \\ U_{j}^{[\prime,\dagger,\sigma_{j}]}C_{j}^{\sigma_{j}}[X^{\mathrm{Im}}]B_{j+1}^{\sigma_{j+1}}[X^{\mathrm{Im}}] \end{pmatrix}$$

$$(A5)$$

where the $U_j^{[\prime,\dagger,\sigma_j]}$ from Eq. A4 is common to all the four vectors. After sweeping back and forth through the lattice, a *good* representation of the correction-vectors is obtained.