Local master equations may fail to describe dissipative critical behavior

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Local quantum master equations provide a simple description of interacting subsystems coupled to different reservoirs. They have been widely used to study nonequilibrium critical phenomena in open quantum systems. We here investigate the validity of such a local approach by analyzing a paradigmatic system made of two harmonic oscillators each in contact with a heat bath. We evaluate the steady-state mean occupation number for varying temperature differences and find that local master equations generally fail to reproduce the results of an exact quantum-Langevin-equation description. We relate this property to the inability of the local scheme to properly characterize intersystem correlations, which we quantify with the help of the quantum mutual information.

I. INTRODUCTION

Quantum master equations have been instrumental in the study of open quantum systems since their introduction by Wolfgang Pauli in 1928 [1]. They offer powerful, yet approximate, means to describe the time evolution of the reduced density operator of quantum systems coupled to external environments [2,3]. They allow the analysis of the dynamics of both diagonal density matrix elements (populations), involved in thermalization processes, and of nondiagonal density matrix elements (coherences), associated with dephasing phenomena. As a consequence, they have found widespread application in many different areas, ranging from quantum optics [6] and condensed matter physics [7] to nonequilibrium statistical mechanics [8] and quantum information theory [9].

In the past decade, quantum master equations have become a popular tool to investigate nonequilibrium phase transitions that occur between (detailed-balance breaking) steady states [10–28]. Special attention has been given to two broad classes of out-of-equilibrium phase transitions: (i) those induced by external driving fields in systems interacting with a single bath (driven-dissipative processes) [10–16] and (ii) those generated by the coupling of a system to several baths (boundary-driven processes) [17,28]. Remarkably, nontrivial exact analytical steady-state solutions of local quantum Lindblad master equations have been obtained for various many-body spin-chain models [17,18,21,24,29,25], thus offering new insight into boundary-driven critical systems.

However, the form of the quantum master equations employed in these studies is often postulated. Their validity is thus not completely clear a priori. This is especially true for boundary-driven processes where the system of interest is coupled to several reservoirs. In this case, it has recently been shown that local master equations, that are commonly used to examine nonequilibrium phase transitions [17,28], may violate the second law of thermodynamics [29] and give rise to nonphysical results, such as incorrect steady-state distributions or nonzero currents for vanishing bath interactions [30,31], even in the limit of small bath couplings. These inconsistencies are related to the fact that local quantum master equations, whose total dissipator is simply the sum of the single-bath dissipators, incorrectly neglect bath-bath correlations, which are induced by intersystem interactions, in contrast to global quantum master equations [29,40]. Interestingly, the local approach has been shown to provide a better description of quantum heat engines than the global approach in some parameter regimes [33]. Meanwhile, the validity of Lindblad quantum master equations has, for example, been discussed in the context of quantum transport [41,42], quantum relaxation [43,44], and entanglement generation [45]. But these results cannot be straightforwardly extended to nonequilibrium phase transitions as the considered models do not exhibit critical behavior.

In this paper, we examine the accuracy of a quantum-master-equation description of dissipative critical phenomena by analyzing an exemplary system consisting of two interacting harmonic oscillators, each weakly coupled to a thermal reservoir. This system naturally appears in many areas, most notably in cavity optomechanics [46]. Many-body superradiant phase-transition models, such as the Dicke model [47] and the Tavis-Cummings model [48], can also be mapped onto such a system after a Holstein-Primakoff transformation [49,50]. We concretely compare local and global quantum master equations, with and without rotating-wave approximation for the oscillator-oscillator interaction, to exact results provided by a quantum-Langevin-equation description [42].
We explicitly evaluate the stationary mean occupation number of one oscillator for various nonequilibrium temperature differences. We find that the local master equation generally fails to reproduce the results of the quantum Langevin equation especially for large temperature differences, while the global approach exhibits better agreement. We show that this feature is directly related to the inability of the local description to correctly capture intersystem correlations, which we quantify with the help of the quantum mutual information [9].

II. COUPLED-OSCILLATOR MODEL

We consider a system of two interacting harmonic oscillators with Hamilton operator,

\[ H = \sum_{j=1,2} \omega_j a_j^\dagger a_j + \lambda (a_1 a_2^\dagger + a_2 a_1^\dagger) + \kappa (a_1 a_2 + a_1^\dagger a_2^\dagger), \]

(1)

where \(a_j^\dagger\) and \(a_j\) are the usual ladder operators and \(\omega_j\) the respective frequencies. We will examine two different types of intersystem interactions: (i) a position-position interaction, \(x_1 x_2\), corresponding to \(\kappa = \lambda\), and (ii) its rotating-wave version, obtained for \(\kappa = 0\). Two important points should be stressed: First, the position-position coupling \(x_1 x_2\) leads to critical behavior above a critical interaction strength \([51, 53]\), in contrast to the commonly treated Hookian interaction \((x_1 - x_2)^2\) \([11, 45]\). In addition, while the rotating-wave approximation is usually associated with a weak-coupling condition, \(\lambda/\omega_i \ll 1\), it has recently been shown that counter-rotating terms may be effectively suppressed in modulated systems, even in the ultrastrong regime \([54, 55]\). This opens the possibility to experimentally study critical behavior in strongly interacting rotating-wave models.

The isolated Hamilton operator (1) may be diagonalized exactly for both intersystem interactions, yielding two uncoupled modes with respective energies \([29, 51, 52]\),

\[ \omega_k^{pp} = \left[ (\omega_1^2 + \omega_2^2) \pm \sqrt{(\omega_1^2 - \omega_2^2)^2 + 16\lambda^2 \omega_1 \omega_2} \right] / 2, \]

(2)

\[ \omega_k^{rw} = (\omega_1 + \omega_2) \pm \sqrt{(\omega_1 - \omega_2)^2 + 4\lambda^2} / 2. \]

(3)

These energies display critical behavior at the respective critical couplings \(\lambda_{pp}^c = \sqrt{\omega_1 \omega_2} / 2\) and \(\lambda_{rw}^c = \sqrt{\omega_1 \omega_2}\). Above these points, the eigenfrequencies of the Hamilton operator (1) become imaginary or negative. The energy spectrum is thus no longer bounded from below. These critical values are in agreement with those of the Dicke and Tavis-Cummings models \([56, 58]\).

We next attach each quantum harmonic oscillator to a heat bath with respective temperature \(T_j\) (Fig. 1). As commonly done, we model these reservoirs by an ensemble of harmonic oscillators \([2, 9]\). We further assume that the system-bath coupling is weak, so that the rotating-wave approximation is applicable to that coupling (see details in the Supplemental Material \([59]\)). We emphasize that the interaction between the two harmonic oscillators of the system (1) might be strong.

Most studies of dissipative phase transitions consider complex interacting many-body systems \([17, 28]\). The direct comparison between global and local master equation descriptions is thus extremely difficult in these systems. By contrast, the coupled-oscillator model is complicated enough to exhibit steady-state critical behavior and, at the same time, simple enough to allow for (i) a detailed comparison between global and local approaches, and (ii) the evaluation of intersystem correlations.

III. QUANTUM-MASTER-EQUATION DESCRIPTION

In the usual Born-Markov limit, the density operator \(\rho\) of the joint quantum system obeys a Lindblad master equation of the form \([2, 9]\) (we set \(\hbar = 1\) throughout),

\[ \dot{\rho} = -i[H, \rho] + \sum_{k=1,2} \sum_{i=1,2,3,4} \sum_{j=1,2,3,4} D_k(A_i, A_j), \]

(4)

where the dissipators are given by \(D_k(A_i, A_j) = \Gamma_k(A_i, A_j)\langle A_i \rho A_j - \{A_j A_i, \rho\} / 2\rangle\). The coefficients

![Equilibrium (ΔT= 0)](image_url)

FIG. 2. Steady-state mean occupation number \(\langle a_1^\dagger a_1 \rangle_{ss}\) of the first oscillator as a function of the dimensionless interoscillator interaction strength, \(\lambda/\lambda_c\), for the equilibrium (high-temperature) case \(\Delta T = 0\). For the position-position interaction [see Eq. (1)], the results of the global quantum master equation (blue dots) perfectly agree with those of the quantum Langevin equation (green line) as well as those of the Gibbs state \(\rho_{eq} = \exp(-\beta H) / Z\) (yellow line), while those of the local quantum master equation deviate more and more as the critical point is approached. For the rotating-wave interaction (inset), the global approach still perfectly matches the predictions of the quantum Langevin equation, while the local scheme does not display any critical behavior. Parameters are \(\gamma_1 = \gamma_2 = 1.5 \cdot 10^{-4}\), \(\omega_1 = 5\), \(\omega_2 = 2\), and \(T_1 = T_2 = 98\).
\( \Gamma_k(A_i, A_j) \), as well as the operators \( A_i \), depend on the local or global type of the quantum master equation [29].

In the local approach, each oscillator interacts with its heat bath (labelled by \( k = 1 \) or \( 2 \)) as if it were not coupled to the other oscillator. As a result, the quantum master equation may be derived as usual in the local eigenbasis of one oscillator [29, 43]. The operators \( A_i \) are here the standard ladder operators, \( (a_1, a_1^\dagger, a_2, a_2^\dagger) \), and the dissipators are given by \( \Gamma_k(a_i, a_j^\dagger) = \delta_{kj}\delta_j\gamma_k[N(\omega_j, \beta_k) + 1] \) and \( \Gamma_k(a_i^\dagger, a_j) = \delta_{kj}\delta_j\gamma_k N(\omega_j, \beta_k) \), where \( \gamma_k \) is the damping coefficient of bath \( k \) and \( N(\omega_j, \beta_k) = 1/\exp(\beta_k\omega_j) - 1 \) denotes the thermal occupation number [2–5]. These formulas evidently hold for the two kinds of oscillator-oscillator interaction in Eq. (1).

On the other hand, the global master equation is derived in the global eigenbasis of the combined two-oscillator system [29, 43]. The diagonalization of the joint Hamilton operator (1) accounts for the indirect subsystem-reservoir and reservoir-reservoir correlations which are generated by their coupling to the system. Such correlations are ignored in the local approach. This is the reason why the local master equation may violate the second law of thermodynamics [29]. The explicit (and lengthy) expressions for the dissipators are summarized for both the position-position and rotating-wave interactions in the Supplemental Material [59]. In this situation, they depend on operators \( A_i \) that are given by properly rotated ladder operators [59].

In the following, we will solve the four different quantum master equations (local or global forms with/without rotating-wave interaction) by applying a characteristic function method in symplectic Fourier space [59] and evaluate the steady-state mean occupation number \( \langle a_j a_j^\dagger \rangle_{ss} = \text{tr}(\rho_{ss} a_j^\dagger a_j) \), where \( \rho_{ss} \) is the stationary density operator.

IV. QUANTUM-LANGEVIN-EQUATION DESCRIPTION

In order to assess its validity, both for equilibrium and nonequilibrium conditions, we shall compare the steady-state properties of the approximate quantum-master-equation treatment to those of the exact quantum-Langevin-equation approach [3]. To this end, we will extend the results obtained for Hookian coupling [42, 44] to the position-position and rotating-wave interactions of Eq. (1). The quantum Langevin equations read [3],

\[
\dot{a}_j = -i[a_j, H] - \gamma_j a_j + \sqrt{2\gamma_j} a_{j,\text{in}},
\]

where the noisy input operators \( a_{j,\text{in}} \), stemming from the interaction with the respective baths, are characterized by the correlation function in Fourier space,

\[
\langle a_{j,\text{in}}(\nu_j^\dagger) a_{j,\text{in}}(\nu_j^\prime) \rangle = 2\gamma_j |N(\nu_j, \beta_j) + 1| \delta(\nu_j - \nu_j^\prime),
\]

\[
\langle a_{j,\text{in}}(\nu_j^\dagger) a_{j,\text{in}}(\nu_j) \rangle = 2\gamma_j N(\nu_j, \beta_j) \delta(\nu_j - \nu_j^\prime),
\]

\[
\langle a_{j,\text{in}}(\nu_j^\dagger) a_{j,\text{in}}(\nu_j^\prime) \rangle = 2\gamma_j |N(\nu_j, \beta_j) + 1| \delta(\nu_j - \nu_j^\prime).
\]

The coupled quantum Langevin equations [3] can be solved by matrix inversion in Fourier space [59]. In particular, the mean occupation number is here equal to,

\[
\langle a_1^\dagger a_1 \rangle_{\text{Langevin}} = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} \langle a_1^\dagger(\nu_j) a_1(\nu_j^\prime) \rangle e^{i(\nu_j - \nu_j^\prime)t} d\nu_j d\nu_j^\prime.
\]

Equation (8) is independent of time in the steady-state regime and we will set \( t = 0 \) in the following. Steady-state mean occupation numbers may be evaluated exactly (without any approximations) in the quantum-Langevin-equation formalism in contrast to the quantum-master-equation approach [32].

V. RESULTS

Figure 2 presents the steady-state mean occupation number \( \langle a_1^\dagger a_1 \rangle_{ss} \) of the first oscillator as a function of the reduced interaction strength \( \lambda/\lambda_c \) in the equilibrium (high-temperature) case \( \Delta T = T_2 - T_1 = 0 \). We observe perfect agreement between the global quantum master equation (blue dots), the quantum Langevin equation (green line) and the equilibrium (Gibbs) state \( \rho_{eq} = \exp(-\beta H)/Z \) (yellow line) [59] for all values of \( \lambda/\lambda_c \), for both the position-position and the rotating-wave (interaction) interactions. By contrast, the local quantum master
The success/failure of the quantum-master-equation description of dissipative critical phenomena may be understood both physically and mathematically. To first address the physical aspect, we consider the quantum mutual information between the two harmonic oscillators, \( I(\rho_{ss}) = S(\rho_1) + S(\rho_2) - S(\rho) \), where \( S(\rho_i) = -\text{tr}\{\rho_i \ln \rho_i\} \) is the von Neumann entropy and \( \rho_i = \text{tr}_i \rho \) are the reduced density operators of the respective harmonic oscillators. The quantum mutual information is a measure of the total (classical and quantum) correlations between two subsystems and has been used broadly to characterize critical transitions. Figure 5 shows that the stationary quantum mutual information \( I(\rho_{ss}) \) displays a very similar dependence on the interaction strength \( \lambda \) as the average occupation number \( \langle a_1^\dagger a_1 \rangle_{ss} \) represented in Fig. 1, both for the position-position and rotating-wave interoscillator interactions. The shortcomings of the quantum-master-equation approach, especially in its local version, may thus be traced to its inability to correctly capture intersystem correlations close to the critical point. This feature can be confirmed mathematically by looking at the way the respective Lindblad quantum master equations are obtained. The dissipation in the local master equation is indeed derived in the local eigenbasis of each separate harmonic oscillator, while those of the global master equation follow from a diagonalization of the interacting two-oscillator system (the unitary evolution given by the von Neumann term in Eq. (4) describes coupled dynamics in both cases). The global scheme thus better accounts for intersystem correlations than the local one, and should therefore be preferred. Such intersystem
correlations are indeed crucial for an accurate description of many-body critical systems, and should not be incorrectly omitted. Yet, despite these deficiencies, local quantum master equations have been a tool of choice in numerous studies on dissipative critical behavior.\[17\][28].

VI. CONCLUSIONS

We have examined the ability of global and local quantum master equations to accurately describe dissipative critical phenomena using an illustrative system of two interacting, damped harmonic oscillators, with and without rotating-wave interaction. This model provides a transparent, yet generic, example to perform such a study. We have found that while the global master equation reproduces the results of the quantum Langevin equation reasonably well, the local version usually fails to do so, especially in the far-from-equilibrium regime; it generally fails in the case of the rotating-wave interaction.

We have related these properties to the inability of the local approach to correctly apprehend oscillator-oscillator correlations that we have quantified with the help of the quantum mutual information. The latter quantity could be easily determined in the present two-oscillator model, in contrast to more complex interacting many-body systems. Our findings show that approximate local quantum master equations in general, and their exact analytical solutions in particular, should be used with caution when studying dissipative critical behavior, and that the more complicated global approach should be favored instead.

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APPENDIX A: QUANTUM MASTER EQUATIONS

In the following, we provide details about the dissipators of the four quantum master equations that we consider in our study (global/local forms with/without rotating-wave interaction) as well as their solutions. The standard dissipators of the local quantum master equation are given below Eq. 4 in the main text. They are derived in the local eigenbasis of each oscillator [2][5] and thus hold for both the position-position and rotating-wave intersystem interactions. By contrast, the global master equations are derived in the global eigenbasis of the combined two-oscillator system obtained by diagonalizing the quadratic Hamilton operator \(H\). The respective dissipators are then computed by expanding the system-bath interaction in this basis. We concretely consider the total system-bath Hamilton operator,

\[H_{\text{tot}} = H + H_{SB1} + H_{SB2} + H_{B1} + H_{B2},\]  

with harmonic thermal baths \(H_{Bi} = \sum_j \omega_i b_i^\dagger b_i\) and local system-bath couplings \(H_{SBi} = \sum_{k,l} \kappa_{ij}(a_i b_i^\dagger + \text{h.c.})\) with coupling constants \(\kappa_{ij}(i = 1, 2)\) [2][5].

In the case of the rotating-wave interaction, the diagonalization of \(H\) leads to \(H_{\text{rw}} = \omega_{\pi}^\text{ww} d_i^\dagger d_i + \omega_{\pi}^\text{ww} d_i d_i\ldots\) with the eigenfrequencies \(\omega_{\pi}^\text{ww}\) given in Eq. (3) of the main text and the rotated operators \(d_i = a_i \cos \theta - a_i^\dagger \sin \theta\) and \(d_i^\dagger = a_i \cos \theta + a_i^\dagger \sin \theta\), where the angle \(\theta\) satisfies \(\cos^2 \theta = (\omega_{\pi}^\text{ww} - \omega_{\pi}^\text{ww})/\omega_{\pi}^\text{ww}\) [29]. The global Lindblad dissipators then follow as [29],

\[
\Gamma(a_1, a_1^\dagger) = \gamma_1^+ c^4 + \gamma_2^- s^4 + (\gamma_2^+ + \gamma_2^-) c^2 s^2 \quad (10)
\]

\[
\Gamma(a_1^\dagger, a_1) = \gamma_1^- c^4 + \gamma_1^+ s^4 + (\gamma_1^+ + \gamma_2^-) c^2 s^2 \quad (11)
\]

\[
\Gamma(a_2, a_2^\dagger) = \gamma_2^+ c^4 + \gamma_2^- s^4 + (\gamma_1^- + \gamma_2^-) c^2 s^2 \quad (12)
\]

\[
\Gamma(a_2^\dagger, a_2) = \gamma_2^- c^4 + \gamma_2^+ s^4 + (\gamma_1^- + \gamma_2^+) c^2 s^2 \quad (13)
\]

\[
\Gamma(a_1, a_2^\dagger) = \gamma_1^+ c^3 s - \gamma_1^- s^3 c + \gamma_2^+ c s^3 - \gamma_2^- c^3 s \quad (14)
\]

\[
\Gamma(a_1^\dagger, a_2) = \gamma_1^- c^3 s - \gamma_1^+ s^3 c + \gamma_2^- c s^3 - \gamma_2^+ c^3 s \quad (15)
\]

with \(c = \cos \theta\), \(s = \sin \theta\), \(\gamma_1^\pm = \gamma_1^\pm\), \(\gamma_1^\pm = \gamma_1^\mp e^{-\beta_1 \omega_{\pi}^\text{ww}}\) and \(\Gamma(a_1, a_2) = \Gamma(a_1^\dagger, a_1), \Gamma(a_1, a_2^\dagger) = \Gamma(a_2, a_1^\dagger)\).

In the case of the position-position interaction, the diagonalization of \(H\) is more involved as it couples all four ladder operators with each other, \((a_1, a_2, a_1^\dagger, a_2^\dagger) = S(c_1, c_2, c_1^\dagger, c_2^\dagger)\) [31]. The 4 x 4 diagonalization matrix \(S\) is partitioned into four blocks with the 2 x 2 matrix \(A\) on the diagonal blocks and 2 x 2 matrix \(B\) on the off-diagonal blocks,

\[
A = \frac{(\omega_{\pi}^\text{pp} + \omega_1) \cos \theta - (\omega_{\pi}^\text{pp} + \omega_1) \sin \theta}{2\sqrt{\omega_{\pi}^\text{pp} \omega_1}} \frac{-(\omega_{\pi}^\text{pp} + \omega_1) \sin \theta}{2\sqrt{\omega_{\pi}^\text{pp} \omega_1}}
\]

\[
B = \frac{(-\omega_{\pi}^\text{pp} + \omega_2) \cos \theta}{2\sqrt{\omega_{\pi}^\text{pp} \omega_2}} \frac{-(\omega_{\pi}^\text{pp} - \omega_2) \sin \theta}{2\sqrt{\omega_{\pi}^\text{pp} \omega_2}} \frac{-(\omega_{\pi}^\text{pp} - \omega_2) \sin \theta}{2\sqrt{\omega_{\pi}^\text{pp} \omega_2}} \frac{-(\omega_{\pi}^\text{pp} - \omega_2) \sin \theta}{2\sqrt{\omega_{\pi}^\text{pp} \omega_2}} \end{array},\]  

with the eigenfrequencies \(\omega_{\pi}^\text{pp}\) given in Eq. (2) of the main text. The global Lindblad dissipators are then \(\Gamma(A_i, A_j) = \Gamma_{11}^{ij} + \Gamma_{12}^{ij} + \Gamma_{21}^{ij} + \Gamma_{22}^{ij}\), with

\[
\Gamma_{11}^{ij} = \gamma_1 N(\omega_{+}, \beta_1) S_1^i S_1^j W_1^3 W_1^3
\]

\[
+ \gamma_1 N(\omega_{-}, \beta_1) S_1^i S_1^j W_1^3 W_1^3
\]

\[
+ \gamma_1 N(\omega_{+}, \beta_1) + 1) S_1^i S_1^j W_1^3 W_1^3
\]

\[
+ \gamma_1 N(\omega_{-}, \beta_1) + 1) S_1^i S_1^j W_1^3 W_1^3
\]  

for the quantum oscillator 1 coupled to bath 1 at inverse temperature \(\beta_1\) with \(W = S^{-1}\). Here the indexes \(i, j\) run over 1-4, corresponding to the elements of \((a_1, a_2, a_1^\dagger, a_2^\dagger)\) and \(k, l\) run over all combinations of 1,3. The indexes \(k, l\) correspond to the initially chosen local coupling terms in the derivation of the master equation before the diagonalization is applied, which can be ordered either as \(a_i a_i^\dagger\) or \(a_i^\dagger a_i\). Thus, there are 32 different terms corresponding
to the 16 unique operator orderings $A_i A_j$ in the dissipators. Expressions for second bath at inverse temperature $\beta_2$ are analogous with $k, l$ now combinations of 2, 4.

We explicitly solve the linear local and global quantum master equations by computing the first and second moments of $\rho$ in symplectic space [2]. The symmetric characteristic function is defined by $\chi(\alpha_1, \alpha_2) = \langle D_1(\alpha_1) \otimes D_2(\alpha_2) \rangle$, where $D_i(\alpha_i) = \exp(\alpha_i a_i^\dagger - \alpha_i^* a_i)$ is the displacement operator. The (symmetric) moments are then obtained by differentiation [65],

$$\langle a_i^k a_j^l \rangle_s = \frac{d^k}{d\alpha_i^k} \frac{d^l}{d\alpha_j^l} \chi(\alpha_1, \alpha_2)|_{\alpha_1 = \alpha_2 = 0}, \quad (18)$$

where $\langle \cdot \rangle_s$ is the expectation value of the symmetrized version of the operators $a_i^k a_j^l$. The evolution of the characteristic function is derived from the master equation

$$\frac{d}{dt} \chi(\alpha_1, \alpha_2) = \text{Tr}\{D_1(\alpha_1) \otimes D_2(\alpha_2) \dot{\rho}\}, \quad (19)$$

together with the identities,

$$D_i a_i^\dagger = \left( -\frac{\alpha_i^*}{2} + \frac{d}{dx_i} \right) D_i, \quad D_i a_i = \left( \frac{\alpha_i}{2} - \frac{d}{dx_i} \right) D_i, \quad (20)$$

$$a_i^\dagger D_i = \left( \frac{\alpha_i^*}{2} + \frac{d}{dx_i} \right) D_i, \quad a_i D_i = \left( \frac{\alpha_i}{2} - \frac{d}{dx_i} \right) D_i. \quad (20)$$

with $\alpha_i = x_i + i p_i$ and $\frac{d}{dx_i} = (d/dx_i - i d/dp_i)/2$ using the Gaussian ansatz $\chi(x_1, p_1, x_2, p_2) = \exp(i \vec{p} \vec{y} - \bar{B}' \vec{\sigma} \vec{B}'/2)$ with $\vec{y} = (y_1, z_1, y_2, z_2)^T$. Since the Hamiltonian is purely of quadratic order, the steady state values for the first moments always vanish $\bar{y}_i = 0 = \bar{z}_i$ and the system is completely described by the second moments. Writing these second moments in vector form $\vec{\sigma} = (\sigma_{x1x1}, \sigma_{x1p1}, \sigma_{x1x2}, \sigma_{x1p2}, \sigma_{p1p1}, \sigma_{p1x2}, \sigma_{p1p2}, \sigma_{x2x2}, \sigma_{x2p2}, \sigma_{p2p2})$, one may write the steady-state set of equations as $\vec{\sigma} = \Delta \vec{\sigma}$.

The $10 \times 10$ matrix $\Delta$ can be written down row-wise using $2\Gamma(i, j, k, l, m, n, o, p) = \lambda^\beta(\alpha_1, \alpha_2) \equiv (-1)^k \Gamma(1, 2, 0) + (-1)^k \Gamma(2, 1, 0) + (-1)^k \Gamma(2, 2, 0) + (-1)^k \Gamma(1, 0, 2) + (-1)^k \Gamma(0, 2, 0) + (-1)^k \Gamma(0, 1, 0) + (-1)^k \Gamma(0, 0, 1) + (-1)^k \Gamma(0, 0, 0) + (-1)^k \Gamma(0, 0, 0), 0)$.

$$\Delta_1 = (\Gamma(1, 0, 0, 0, 0, 0, 0, 0, 0, 0) - (\kappa + \lambda), 0, 0, 0, 0, 0, 0, 0, 0, 0)$$

$$\Delta_2 = (\omega_1, 0, 0, 0, 0, 0, 0, 0, 0, 0)$$

$$\Delta_3 = (\Gamma(1, 0, 0, 0, 0, 0, 0, 0, 0, 0) - (\kappa + \lambda), 0, 0, 0, 0, 0, 0, 0, 0, 0)$$

$$\Delta_4 = (\omega_2, 0, 0, 0, 0, 0, 0, 0, 0, 0)$$

$$\Delta_5 = (0, 0, 0, 0, 0, 0, 0, 0, 0, 0)$$

$$\Delta_6 = (0, 0, 0, 0, 0, 0, 0, 0, 0, 0)$$

$$\Delta_7 = (0, 0, 0, 0, 0, 0, 0, 0, 0, 0)$$

$$\Delta_8 = (0, 0, 0, 0, 0, 0, 0, 0, 0, 0)$$

$$\Delta_9 = (0, 0, 0, 0, 0, 0, 0, 0, 0, 0)$$

$$\Delta_{10} = (0, 0, 0, 0, 0, 0, 0, 0, 0, 0)$$

(21)
for both the system matrix $\Lambda$ and steady-state vector $\vec{G}$. Solving
this system of equations (numerically) leads to the
symplectic covariance matrix. The actual covariance ma-
trix is obtained after symplectic transformation: $\sigma_{zz} =$
$\sigma_{pp}/2$, $\sigma_{pp} = \sigma_{zz}/2$, $\sigma_{zj} = -\sigma_{jp}/2$. The
steady-state occupation numbers are finally calculated
via $\langle a^\dagger a \rangle_{ss} = (\sigma_{z1} + \sigma_{p1p} - 1)/2$.

**APPENDIX B: QUANTUM MUTUAL INFORMATION**

The quantum mutual information for a Gaussian
system can be calculated from the covariance matrix as
$I(\sigma) = f(a) + f(b) - f(n_{-}(\sigma)) - f(n_{+}(\sigma))$ \[66\],
with $a = \sqrt{\text{det}(\sigma)}$, $b = \sqrt{\text{det}(\beta)}$, $f(x) = (x +$
$1/2)\ln(x + 1/2) - (x - 1/2)\ln(x - 1/2)$, $n_{+}(\sigma) =$
$\sqrt{\left(\Delta(\sigma) - \sqrt{\Delta(\sigma)^2 - 4\text{det} \sigma}\right)/2}$,
for the covariance matrix defined as $\sigma_{ij} = \langle x_{i}x_{j} \rangle/2$, $x_{i} = (x_{1}, p_{1}, x_{2}, p_{2})$.
In this form, the submatrices of interest are
$\sigma_{\alpha\beta} = \langle (\alpha, \gamma), (\gamma, \beta) \rangle$.

**APPENDIX C: QUANTUM LANGEVIN EQUATIONS**

The quantum Langevin equation is derived in the
Heisenberg picture \[3\]. This approach has the advantage
that it does not involve strong approximations as is
the case for quantum master equations \[12\]-\[15\]. On
the other hand, the drawback is that it cannot easily
be solved in general as the corresponding differential
equations are operator differential equations in Hilbert
space. For Gaussian systems, it can however be solved
using matrix methods \[3\]. The steady-state solution
can thus be obtained by matrix inversion in Fourier
space \[12\], $M(\nu)\tilde{a}(\nu) + \tilde{a}_{in}(\nu) = 0$, with the two vec-
tors $\tilde{a}(\nu) = (\tilde{a}_{1}(\nu), \tilde{a}^\dagger_{1}(\nu), \tilde{a}_{2}(\nu), \tilde{a}^\dagger_{2}(\nu))$ and $\tilde{a}_{in}(\nu) =$
$(\sqrt{2\gamma_{1}}a_{1in}(\nu), \sqrt{2\gamma_{1}}a^\dagger_{1in}(\nu), \sqrt{2\gamma_{2}}a_{2in}(\nu), \sqrt{2\gamma_{2}}a^\dagger_{2in}(\nu))$.
The matrix $M$ is explicitly given by

$$M(\nu) = \begin{pmatrix} -i\nu + i\omega_{1} & 0 & i\kappa & i\lambda \\ 0 & -i\nu - i\omega_{1} & -i\lambda & -i\kappa \\ i\kappa & i\lambda & -i\nu + i\omega_{2} & 0 \\ -i\lambda & -i\kappa & 0 & -i\nu - i\omega_{2} \end{pmatrix} + \tilde{\gamma}$$

with $\tilde{\gamma} = \text{diag}(\gamma_{1}, \gamma_{1}, \gamma_{2}, \gamma_{2})$. Inverting $M^{-1} = m$, the
second moment $\langle a^\dagger a \rangle_{1}$ in the algebraic space is

$$\langle a^\dagger a \rangle_{1} = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} |a_{1}(\nu)a_{1}(\nu')|^2 e^{i(\nu - \nu')t}/2\pi d\nu d\nu' = \frac{1}{\pi} \int_{-\infty}^{\infty} \gamma_{1}(m_{11})^2 N(\nu, \beta_{1}) + |m_{12}|^2(N(\nu, \beta_{1}) + 1) + \gamma_{2}(m_{13})^2 N(\nu, \beta_{2}) + |m_{14}|^2(N(\nu, \beta_{2}) + 1) d\nu$$

and similar expressions for all the other second moments.

The Gibbs state expectation values may in addition
be evaluated by the diagonalization of the Hamilton-
ian given above. In general, any quadratic expectation
value in the $a_{1}$ algebraic space may be calculated via
$\langle A_{1}A_{j} \rangle = \sum_{k} S_{ik}S_{j}(C_{k}C_{j})$ with $A_{i} = (a_{1}, a_{2}, a_{1}^\dagger, a_{2}^\dagger)$ and $C_{i} = (c_{1}, c_{2}, c_{1}^\dagger, c_{2}^\dagger)$. The $\langle C_{i}C_{j} \rangle$ are then given by
the uncoupled oscillators with eigenfrequencies, Eq. \[3\]
of the main text, and corresponding temperature $T$.

**APPENDIX D: NONEQUILIBRIUM STEADY STATES**

The deviation of the local (and, to a lesser extent,
global) quantum master equations from the quantum
Langevin equation does not only depend on the magni-
tude of the temperature difference $\Delta T$ but also on its
sign (Figs. \[4ab\]). We first note that the local master
equation leads to larger (smaller) mean occupation num-
bers for weak (strong) coupling, both for the position-
position and the rotating-wave interactions. This in-
crease of the mean occupation number at small coupling
is caused by the frequency difference between the oscilla-
tors ($\omega_{1} > \omega_{2}$), which leads to a relatively larger occupation
number in the second oscillator (modulated by the tem-
perature difference), whereas its decrease is induced
by strong-coupling effects. In addition, the global mas-
ter equation completely matches the Langevin equation,
for all $\Delta T$ for the rotating-wave interaction, while this
is not the case for the position-position interaction: the
mean occupation number is larger (smaller) than that
for the quantum Langevin for $\Delta T < 0$ ($\Delta T > 0$).

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FIG. 6. Ratio of the steady-state mean occupation numbers $\langle a^\dagger a \rangle_{\text{ss}} / \langle a^\dagger a \rangle_{\text{Langevin}}$ of the quantum master equation and the quantum Langevin equation as a function of $\lambda \lambda_C$, for various nonequilibrium temperature differences $\Delta T$, for a) position-position and b) rotating-wave interactions. Temperatures, for positive $\Delta T > 0$, are $T_1 = 98$, $T_3 = T_1 + \Delta T$, and for negative $\Delta T < 0$, $T_1 = T_2 - \Delta T$, $T_2 = 98$.

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