Entanglement concentration with quantum nondemolition Hamiltonians

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We devise and examine two procrustean entanglement concentration schemes using quantum nondemolition (QND) interaction Hamiltonians in the continuous variable regime, applicable for light, for atomic ensembles or in a hybrid setting. We thus expand the standard entanglement distillation toolbox for use of a much more general, versatile, and experimentally feasible interaction class. The first protocol uses Gaussian ancillary modes and a non-Gaussian postmeasurement, the second a non-Gaussian ancillary mode and a Gaussian postmeasurement. We explicitly calculate the density matrix elements of the non-Gaussian mixed states resulting from these protocols using an elegant Wigner-function-based method in a numerically efficient manner. We then quantify the entanglement increase calculating the logarithmic negativity of the output state and discuss and compare the performance of the protocols.

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I. INTRODUCTION

One of the biggest hurdles quantum communication has yet to overcome is the difficulty of generating highly entangled states between distant nodes. Due to the inherent noise in any optical setup, in reality it is only ever possible to generate partially entangled states at distant locations. The remedy to this problem is entanglement distillation [1–3]. Local operations and classical communication (LOCC) are used to distill at least one maximally entangled bipartite state from multiple weakly entangled states.

One of the latest developments in quantum information science is the use of infinite-dimensional systems or continuous variables (CV), such as the amplitude and phase quadratures of optical modes or the collective spin of macroscopic atomic ensembles. CV systems offer the advantage of cheap resources, easy generation, and control, and off-the-shelf telecommunication components can be used for information processing. The downside of it is that it is in principle impossible to have maximally entangled states, and consequently, there is an additional source of imperfections. Furthermore, finding concentration schemes in the continuous variable regime is made particularly difficult by a no-go theorem that states that distillation of Gaussian states cannot be carried out using Gaussian operations alone [4–6]. One has to resort to challenging non-Gaussian operations, like strong high-order nonlinear interactions or photon counting.

It is worth noting that in the continuous variable regime, most or less all of the proposed schemes for entanglement distillation contain variations of a common theme: beam-splitter interactions. On the theory side, Opatrný et al. [7] showed how beam splitters and photon subtraction can be used to increase the entanglement in a two-mode squeezed vacuum (TMSV) as part of a teleportation scheme. The distillation aspect of this was seized upon and improved [8–11], and a rigorous theoretical description was given by Kitagawa et al. [12]. Experimentally, there have been some notable successes [13,14], including with non-Gaussian noise [15]. In another proposal [16] a complete CV entanglement “distillery” has been proposed, harnessing limited physical space for storing quantum states and distilling entanglement. There, clever manipulation of an imperfect quantum memory complements the beam-splitter-based entanglement concentration scheme.

However, the net could be cast wider. Theoretical work has been carried out utilizing other interactions between an entangled state and ancillary modes, notably the cross-Kerr effect [17,18], in which the nonlinear interaction plays the role of the necessary non-Gaussian element, and weak values [19]. Still, very little is known about entanglement distillation protocols using so-called quantum nondemolition (QND) interaction Hamiltonians, although they lend themselves readily to many experimental systems where distillation could be useful. For example, the off-resonant dipole interaction between heavily polarized light and the macroscopic spin properties of an atomic ensemble can be modeled as QND, and two atomic ensembles were successfully entangled, albeit weakly, using this interaction [20]. Furthermore, for light it was also shown that it is possible to create a QND Hamiltonian using optical instruments such as biased beam splitters and squeezers in combination [21].

The quantum nondemolition interactions [22,23] are well known in quantum optics for quantum metrology, entanglement generation, and other purposes. They are described by Hamiltonians of the form, e.g.,

$$\hat{H}_{\text{int}} = \kappa \hat{P}_S \hat{P}_A,$$

where $\kappa$ is the interaction strength and $\hat{P}_S/A$ are the momentum quadratures of the system and ancillary mode. The system and ancillary mode interact in such a way as to leave one quadrature component of each subsystem intact while phase shifting the conjugate components.

As with the beam-splitter interaction, the QND interaction has been examined in conjunction with continuous variable quantum teleportation [24,25], and a feasible setup for continuous variable quantum nondemolishing interaction of two optical modes at a distance has also been proposed [26]. Motivated by successes in developing quantum memory devices, a protocol for using quantum nondemolition interactions to manipulate atomic spin states in order to produce arbitrary...
superpositions of atomic Dicke states has also been proposed [27], and a further protocol for the Gaussionization of traveling light beams has been explored [28]. In this paper, we suggest and explore two procrustean entanglement distillation schemes that use QND interactions and quantify their performance showing the (probabilistic) increase of the logarithmic negativity of a two-mode squeezed vacuum, defined in the Fock basis as

\[ |\Psi\rangle = \sqrt{1 - \lambda^2} \sum_{n=0}^{\infty} \lambda^n |nn\rangle, \]

where \(|n\rangle\) is the photon number and \(\lambda = \tanh r\) is the squeezing. For the quantitative analysis of these schemes, we amalgamate ideas on using the well-known formalism of Gaussian states to represent non-Gaussian states [29] (used more recently in Ref. [30]) and calculate density matrix elements using multivariate Hermite polynomials [31–33].

Our approach is versatile, making additional interactions and double passes of the interacting modes as simple as multiplying 8 \(\times\) 8 matrices and allowing for detector inefficiencies to be considered in a simple manner. Hence numerical calculations are made fast and accurate. Thus the novelty of the proposed approach to distillation lies both in the scheme design (QND-based) and in the methodology of the calculations. Notably, the suggested scheme is not bound to a certain physical system and can be applied to the distillation of entanglement in light modes, atomic ensembles, or in hybrid systems, the essential requirement being solely the form of the Hamiltonian underlying the interaction.

The paper is organized as follows. In Sec. II we remind the reader of the important properties of Gaussian states required. In Sec. III we study the first scheme, a procrustean entanglement concentration protocol replacing the conventional beam-splitter interaction with the QND interaction between a Gaussian ancilla and a Gaussian entangled state. More precisely, we calculate the logarithmic negativity of the non-Gaussian mixed state resulting from the QND interaction between the two-mode squeezed vacuum and a Gaussian ancillary mode subsequently subjected to a non-Gaussian measurement. In Sec. IV, we explore an alternative scheme where the ancillary mode is non-Gaussian and the postmeasurement is Gaussian. In both cases the entanglement of the system is shown to increase, demonstrating a successful continuous variable entanglement distillation based on the QND Hamiltonian. In Sec. V we discuss the entanglement of the Gaussian states having the same covariance matrices as the non-Gaussian mixed states output from the two protocols. We conclude in Sec. VI.

II. PRELIMINARIES

An \(N\)-mode Gaussian state can be completely defined by the first- and second-order moments of the quadrature operators \(\hat{R}_j\), with \(\mathbf{R} = (x_1, p_1, \ldots, x_N, p_N)^T\). The Wigner function for such a state is given by

\[ W(x_1, p_1, \ldots, x_N, p_N) = \frac{\exp[-(\mathbf{R} - \mathbf{d})^T \gamma^{-1}(\mathbf{R} - \mathbf{d})]}{\pi^N \sqrt{\det \gamma}}, \]

where \(\mathbf{d}\) is the displacement vector \(\mathbf{d} = (\langle x_1 \rangle, \langle p_1 \rangle, \ldots, \langle x_N \rangle, \langle p_N \rangle)^T\) and \(\gamma\) is the covariance matrix of the state in question, defined as

\[ \gamma_{\alpha \beta} = \langle \hat{R}_\alpha \hat{R}_\beta \rangle - 2d_\alpha d_\beta. \]

The two-mode squeezed state that we wish to distill has the following covariance matrix:

\[ \gamma_{\alpha \beta}^{TMSS} = \begin{pmatrix} \cosh(2r) & 0 & \sinh(2r) & 0 \\ 0 & \cosh(2r) & 0 & -\sinh(2r) \\ \sinh(2r) & 0 & \cosh(2r) & 0 \\ 0 & -\sinh(2r) & 0 & \cosh(2r) \end{pmatrix}. \]

This Gaussian distribution is centered at the origin in phase space, and so \(\mathbf{d}\) of Eq. (3) is the zero vector. The squeezing parameter is denoted \(r\).

As a well-known consequence of the Stone–von Neumann theorem, any quadratic unitary operation at the Hilbert space level corresponds to a symplectic operation at the phase space level that preserves the symplectic form \(\Omega = \oplus_{j=1}^N \sigma_j\). In what follows, the following symplectic operations are relevant. First, the QND interaction between two modes (system and ancilla) can be described by its action on phase space, e.g.,

\[ S_{\alpha \beta}^{QND} = \begin{pmatrix} 1 & 0 & 0 & \kappa \\ 0 & 1 & 0 & 0 \\ 0 & \kappa & 1 & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix}, \]

which yields \(\gamma_{QND} = \text{diag}(e^{2s}, e^{-2s})\) when applied to the vacuum covariance matrix \(\gamma_{\text{vac}} = 1\). If \(s > 0\) then the mode is squeezed in \(p\), and if \(s < 0\) then the mode is squeezed in \(x\).

Noisy quantum channels and phase-insensitive amplifiers cannot be modeled by Gaussian unitary transformations. One of the biggest losses by detectors is when they erroneously fail to register a detection. For this reason we model inefficient detectors as being perfect detectors behind a beam splitter and noiseless. For the quantitative analysis of these schemes, we amalgamate ideas on using the well-known formalism of Gaussian states to represent non-Gaussian states [29] (used more recently in Ref. [30]) and calculate density matrix elements using multivariate Hermite polynomials [31–33].
B is given by the measurement, the total Wigner function of modes \( AB \) = \( \gamma_{AB} \) of the two-mode squeezed state increases.

If successful, the entanglement of the ancillary modes \( \gamma_{ab} \) can be calculated by the straightforward integration of \( W_{AB} \), which can be thought of as the overlap of modes \( A \) and \( B \) with \( \mathcal{W}_2 \). Consequently,

\[
P^{(ij)} = \frac{2^{2-i-j} \sqrt{\det \Gamma}}{\sqrt{\det \Gamma_{AB}} \sqrt{\det \Gamma_{ab}}}.
\]

At this point, we have specified all the quantities in the expression (11) for the Wigner function of the two-mode squeezed state after QND interactions with vacuum modes that are later detected for the presence or absence of photons on inefficient detectors. Hence we are able to describe the entangled state modified by the QND interaction by use of the Wigner function (11). However, to show that the entanglement in the two modes has increased, we require the density matrix elements of the state. These can be calculated with the help of the Husimi Q function and multidimensional Hermite polynomials [31–33].

The Q function is defined as the convolution of the Wigner function with the vacuum [36] and so

\[
Q^{(ij)}_{AB}(\alpha, \beta) = \frac{\exp[-R \mathcal{R}]}{\pi^2 \sqrt{\det (R^{(ij)} + \mathbb{1})}}.
\]

where \( R = (\alpha, \alpha^\dagger, \beta, \beta^\dagger) \) and \( \mathcal{R} \) is the unitary transformation between \( (x, p_A, x_B, p_B)^T \to (\hat{a}, \hat{a}^\dagger, \hat{b}, \hat{b}^\dagger)^T \), which in matrix form is given by

\[
U = \frac{1}{\sqrt{2}} \begin{pmatrix} 1 & i & 0 & 0 \\ 1 & -i & 0 & 0 \\ 0 & 0 & 1 & i \\ 0 & 0 & 1 & -i \end{pmatrix}.
\]

The Q function acts as a generating function for the density matrix elements, \( \rho_{\text{imp}}^{(ij)} \)

\[
\rho_{\text{imp}}^{(ij)} = \langle |i \rangle \langle m | \rho^{(ij)} | n \rangle |p\rangle
\]

\[
= \frac{(2\pi)^2}{\sqrt{\prod_{m,n} p!}} \left( \frac{q^{l+n}}{\partial q^l \partial q^n} \right) \left( \frac{\beta_0^{m+p}}{\partial \beta_0^m \partial \beta_0^p} \right) \times [Q^{(ij)}_{AB}(\alpha, \beta) e^{\alpha^\dagger \beta + \beta^\dagger \alpha}]_{\alpha = \beta = 0}.
\]

The matrix element is most easily expressed in terms of the four-dimensional Hermite polynomials \( H_{\text{imp}} \) (see Appendix). After successful detection the total density matrix is given by

\[
\rho_{\text{imp}} = \frac{1}{2\pi} \left( \sum_{l,m,n} \rho_{\text{imp}}^{(ij)} q^l \beta_0^m \beta_0^p \right) \left( \frac{1}{\sqrt{\prod_{m,n} p!}} \right) \left( \frac{1}{\sqrt{\prod_{m,n} p!}} \right)
\]

\[
\times [Q^{(ij)}_{AB}(\alpha, \beta) e^{\alpha^\dagger \beta + \beta^\dagger \alpha}]_{\alpha = \beta = 0}.
\]
\( \rho \) with matrix elements

\[
\rho_{\text{map}} = \frac{4M}{\sqrt{\text{Vol}(H)}} \sum_{i,j=0}^{1} \frac{(-1)^{j+i}}{\sqrt{\text{det}(\gamma^{ij}+1)}} P^{ij} H_{\text{map}}^{(i,j,0)},
\]

(18)

where for brevity we have defined \( H_{\text{map}}^{(i,j,0)} = H_{\text{map}}^{(i,j,0)}(0,0,0,0) \). The matrices \( C^{(ij)} \) are derived from the matrix equation

\[
C^{(ij)} = B (\gamma^{ij} U + 1)^{-1} \frac{1}{2} D
\]

(19)

with

\[
B = \begin{pmatrix}
1 & 0 & 0 & 0 \\
0 & 0 & 1 & 0 \\
0 & 1 & 0 & 0 \\
0 & 0 & 0 & 1
\end{pmatrix},
D = \begin{pmatrix}
0 & 0 & 1 & 0 \\
1 & 0 & 0 & 0 \\
0 & 0 & 0 & 1 \\
0 & 1 & 0 & 0
\end{pmatrix}.
\]

(20)

The purpose of \( B \) and \( D \) is to rearrange the elements of the matrix in such a way as to make the matrix compatible with the form given in Eq. (A1) and to make \( \rho_{\text{map}} \) proportional to \( H_{\text{map}}^{(i,j,0)} \). For single QND passes the matrix \( C^{(ij)} \) is always real. With the density matrix elements of modes \( A \) and \( B \) we can now show that the entanglement of the system has increased. To quantify this we use the additive logarithmic negativity [37], known to be an upper bound on the distillable entanglement, defined as

\[
\mathcal{E}(\hat{\rho}_{AB}) = \ln \| \hat{\rho}_{AB}^{PT} \| = \ln [2N(\hat{\rho}_{AB}) + 1],
\]

(21)

where \( \| \hat{\rho}_{AB}^{PT} \| \) is the trace norm of the partial transpose of \( \hat{\rho}_{AB} \). The negativity \( N \) is defined as the sum of the negative eigenvalues of \( \| \hat{\rho}_{AB}^{PT} \| \).

The logarithmic negativity of the mixed state resulting from this protocol is shown in Fig. 2. The graph demonstrates the performance of the protocol in different regimes depending on the initial entanglement content in the two-mode squeezed vacuum and the QND interaction strength (see caption). Successful entanglement concentration is achieved for weak to medium values of \( \kappa \), i.e., for a moderate QND coupling between the TMSV and ancilla modes, for a wide range of the initial squeezing in the TMSV. If the interaction is too strong (e.g., \( \kappa = 1 \)), then the effect of the QND interaction on the entanglement content is negative, but as the interaction strength decreases, the logarithmic negativity of the resultant state increases and the performed QND operation is beneficial at least when the initial squeezing of the TMSV is weak. That is, as the interaction strength decreases, it becomes more likely that just a single photon is subtracted from each mode of the two-mode squeezed vacuum and these photons are then detected. As expected, this compares well to the case where beam splitters are used in place of QND interactions, investigated numerically in Ref. [12] for light TMSV and in Ref. [11] for TMSV-entangled atomic ensembles. After all, if the pure state resulting from a single photon subtraction in both modes of the entangled state is more entangled than the initial state, then it should not matter which interaction exactly is used to subtract the photons.

The QND interaction \( S_{QND}^{QND}(x,\hat{X}) \) between one of the system modes and an ancillary vacuum mode can be decomposed into two unbalanced beam splitters and two squeezing operations as shown in Fig. 3. The beam-splitter interaction between two modes can be modeled by the symplectic operation \( S_{BS}(T) \), where

\[
S_{BS}(T) = \begin{pmatrix}
T & 0 & \sqrt{1-T^2} & 0 \\
0 & T & 0 & \sqrt{1-T^2} \\
\sqrt{1-T^2} & 0 & -T & 0 \\
0 & \sqrt{1-T^2} & 0 & -T
\end{pmatrix}.
\]

(22)

For \( \kappa > 0 \) the first beam splitter has a transmission coefficient of \( T = \cos \kappa \ln(\sqrt{\kappa^2+1} + 1) \) and the second beam splitter has a transmission coefficient of \( T = \sin \kappa \ln(\sqrt{\kappa^2+1} + 1) \). Between the two beam splitters are situated two squeezers, \( S_{SQ}(\ln(\sqrt{\kappa^2+1} + 1)) \) and \( S_{SQ}(\ln(\sqrt{\kappa^2+1} + 1)) \). As \( \kappa \) is increased the effect of the squeezers dominates and the uncertainty of the photon number in each mode increases. Consequently, as \( \kappa \) increases, the measurement performed at the end measuring device becomes increasingly noisy. For this reason, the results of Fig. 2 are to be expected.

In contrast to the beam-splitter interaction as modeled by Kitagawa et al. [12], in which highly transmissive beam splitters are favored, in the decomposition of the QND
interaction, more balanced beam splitters are preferred due to the interference of the squeezed modes before measurement.

IV. PROTOCOL II: NON-GAUSSIAN ANCILLARY MODE AND GAUSSIAN MEASUREMENT

Whereas protocol I relies on a Gaussian ancillary mode interacting with the Gaussian TMSV via QND interaction, and a subsequent non-Gaussian measurement, we here consider a protocol which uses a non-Gaussian ancilla and a Gaussian measurement after the QND interaction (Fig. 4). The advantage of this new protocol is that probabilistic non-Gaussian measurements are performed in the preparatory stage and, if successful, the QND interaction can be carried out in the next step. Also, the detection part of the scheme outperforms the first protocol as the Gaussian measurement, homodyne detection, can be highly efficient. The same methods as above can be used to calculate the logarithmic negativity of this scheme. The feasibility of this approach is supported by the results presented in Refs. [38,39], where efficient low-noise methods were proposed for transferring quantum states of light onto matter systems. The proposed methods of [38,39] are, for example, capable of uploading squeezed single-photon states into an atomic memory using QND interactions.

A. Preparatory step

In the protocol of Fig. 4 the ancillary modes a and b are in a quantum state with a non-Gaussian Wigner function. Specifically, they are the result of squeezed vacuum states conditioned on an indeterminate number of photons having been detected. We consider first the preparation of the ancillary modes. In contrast to the previous scheme the modes a and b are initially in a squeezed vacuum state, \( \gamma_{ab} = \gamma_{SQ,a}(s) \otimes \gamma_{SQ,b}(s) \). The removal of photons from the mode is modeled in the following way. The modes a and b pass through a beam splitter of transmittance \( T^2 \) and any subtracted photons are subsequently detected with efficiency \( \eta \). As the transmittance \( T^2 \) increases, the model approaches the limit of just a single photon being subtracted from the mode. In what follows, in all numerical calculations we take \( T = 0.95 \).

If photon subtraction is successful, the resultant non-Gaussian Wigner function becomes a superposition of Gaussian Wigner functions in the form
\[
\mathcal{W}_{ab} = M_{ab} \sum_i (-1)^{i+j} P(i) \gamma_{ab}^{(i)}
\]
with probability \( M_{ab} = [\sum (-1)^{i+j} P(i)]^{-1} \). The corresponding covariance matrices that make up the Wigner functions \( \mathcal{W}_{ab}^{(i)} \) are given by \( \mathcal{W}_{ab}^{(i)} = \gamma_{ab}^{(i)} \otimes \gamma_{ab}^{(j)} \), where
\[
\gamma_{ab}^{(i)} = \left( \begin{array}{cc} \tilde{\vartheta}_+(k) & 0 \\ 0 & \tilde{\vartheta}_-(k) \end{array} \right)
\]
and
\[
\tilde{\vartheta}_\pm(k) = 1 + \frac{(2 - k)(e^{\pm 2s} - 1)T^2}{1 + (1 - k)[1 + \eta(e^{\pm 2s} - 1)(1 - T^2)]}
\]
The coefficients \( P(i) \) are calculated as
\[
P(i) = \frac{2^{2 - i - j} \sqrt{\tau_+(i) \tau_-(i) \tau_+(j) \tau_-(j)}}{\sqrt{\tau_+(i) \tau_-(i) \tau_+(j) \tau_-(j)}},
\]
with
\[
\tau_\pm(k) = 1 - k - \frac{1 + (e^{\pm 2s} - 1)T^2}{(1 - \eta)(e^{\pm 2s} - 1)(1 - T^2) - e^{\pm 2s}T^2}.
\]
At this point we have a non-Gaussian state \( \mathcal{W}_{ab} \), consisting of a superposition of Gaussian states. If the preparatory step could be performed independently and only passed to the interaction and homodyne measurement if successful, then we could consider \( \eta = 1 \).

B. Interaction and detection

A non-Gaussian state produced in the preparatory step further interacts with the TMSV via QND coupling and the ancilla is subsequently passed on for homodyning on a detector of efficiency \( \xi \). An angle \( \theta \) describes the phase of the homodyne measurement, that is the angle at which homodyning is performed in phase space. An angle of \( \theta = 0 \) corresponds to a measurement of position quadrature \( x_{\theta=0} = x \) (the \( x \) marginal distribution), whereas an angle of \( \pi/2 \) describes the momentum quadrature measurement \( x_{\theta=\pi/2} = p \) (the \( p \) marginal distribution). Thus the homodyne measurement is characterized by the generalized quadratures.
Hamiltonians, e.g., QND interactions and so the effects induced by interaction momentum correlations in the TMSV are not mixed by the sensitive to which interactions are used. The position and be calculated as before.

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where $B$ described by a $6 \times 4$ matrix (modes $A$ and $B$), $B^{(ij)}$ is a $2 \times 2$ matrix describing the $z$ correlations, and $C^{(ij)}$ contains the cross correlations. The probability of projection of modes $a$ and $b$ is given by

$$q^{(ij)}_z = \frac{\sqrt{\det(B^{(ij)} - C^{T(j)}(A^{(j)} - 1)C^{(j)})}}{\pi} \times \exp(-\pi/2(z,b)(B^{(ij)} - C^{T(j)}(A^{(j)} - 1)C^{(j)}) (z,b)),$$

and the $Q$ function describing modes $A$ and $B$ reads

$$Q^{(ij)}_{AB}(\alpha,\beta) = \frac{\sqrt{\det(\Phi^{(ij)})}}{\pi^2} e^{-\frac{\Lambda^{(j)}}{\gamma} - \frac{\Lambda^{(j)}}{\gamma}} \times \exp(-\mathcal{R}^T \Phi^{(ij)} \mathcal{R} - \Lambda^{(j)} \mathcal{R}),$$

where

$$\Phi^{(ij)} = U(A^{(j)} - 1 + 1)^{-1} U^\dagger,$$

$$\Lambda^{(j)} = 2(z,b)C^{(j)T}(A^{(j)} - 1 + 1)^{-1} U^\dagger.$$

By defining $\Phi^{(ij)} = B^\dagger \Phi^{(ij)} - (1/2) I D$ and $\Lambda' = \Lambda D$ we can write the density matrix elements as

$$\rho_{\text{hom}} = \frac{4 M_{\text{hom}}}{\sqrt{\det(\Phi^{(ij)} - \Lambda^{(j)/2})}} \sum_{i,j=0}^1 (-1)^{i+j} P^{(ij)} q^{(ij)}_z \times \sqrt{\det(\Phi^{(ij)} - \Lambda^{(j)/2})} H^{(\Phi^{(ij)},\Lambda^{(j)/2})}_{\text{hom}},$$

with

$$M_{\text{hom}} = \left( \sum_{i,j=0}^1 (-1)^{i+j} P^{(ij)} q^{(ij)}_z \right)^{-1}.$$

From this, the logarithmic negativity of modes $A$ and $B$ can be calculated as before.

In contrast to the first protocol, the second protocol is sensitive to which interactions are used. The position and momentum correlations in the TMSV are not mixed by the QND interactions and so the effects induced by interaction Hamiltonians, e.g., $H^{(k \times P)}_{\text{int}}$ and $H^{(k \times P)}_{\text{int}}$, are equivalent. Correspondingly, the choice of measurement on the ancillary modes is important. If, for example, the interaction $H^{(k \times P)}_{\text{int}}$ is used, then the momentum quadratures of the ancillary modes are, by definition, unaffected, but information about the TMSV is imprinted on the position quadrature distribution. If homodyne measurements on the ancillas are performed on $\rho (\theta = \pi/2)$, then the resulting outcome can reveal nothing about the state of the TMSV and the probabilities $q^{(ij)}_z$ (30) are independent of $\kappa$ and $r$. As nothing can be learned probabilistically about modes $A$ and $B$, procrustean entanglement concentration cannot occur. The effects on the TMSV can be transformed away by local Gaussian operations and ancillary modes and so the entanglement of the TMSV is unchanged. The above is true irrespective of whether the ancillary modes are initially squeezed in position or momentum.

If homodyne measurements were taken of the position quadratures ($\theta = 0$), then information about the TMSV will have been probabilistically imprinted on this measured distribution. If $H^{(k \times P)}_{\text{int}}$ is used, then the momentum quadratures of the TMSV contain information about the momentum quadratures of the ancillary modes. That is, some noise has been added to modes $A$ and $B$ which can assist or disrupt entanglement concentration. If the ancillary modes are squeezed in momentum, then only a little noise is added to the $p$ quadratures of the TMSV. The result is an increase in entanglement dependent on $s$ and $\kappa$ (Fig. 5). If the ancillary modes are squeezed in position, then the momentum quadratures are antisqueezed and so a lot of noise is added to the TMSV, having a detrimental effect on the entanglement.

As can be seen in Figs. 5 and 6, successful entanglement concentration is achieved if the ancillary modes are squeezed in $p$ quadrature, the QND interaction is of the form $H^{(k \times P)}_{\text{int}}$, and homodyne measurement is performed in $x$ quadratures. The protocol is noticeably insensitive to ancillary mode

![FIG. 5. Performance of protocol II (Fig. 4). Logarithmic negativity of TMSV before (solid line) and after a QND interaction for different initial squeezing of the ancilla modes. The QND interaction is of the form $k \tilde{X} \tilde{P}$ and has strength $\kappa = 0.5$; homodyne outcome $\gamma \approx 0$. The ancillary modes are squeezed by $s = 0.1(0.87)$ dB (black dotted), $s = 0.5(4.34)$ dB (black dashed), and $s = 1(8.69)$ dB (thick black dashed). The improvements of entanglement in the TMSV gained by increasing the squeezing of the ancillary modes is small when the initial squeezing of the TMSV $r$ is small. The vertical lines from left to right indicate where the initial squeezing of the TMSV $r$ is 1 dB, 3 dB, 5 dB, and 7 dB. $\eta = \xi = 1$.](image-url)
squeezing—the logarithmic negativity of the resulting state is largely unaffected for low levels of TMSV initial squeezing.

Unlike protocol I, a weak interaction strength ruins the entanglement in the system. Similarly, if the strength is too strong then the entanglement decreases. If \( \kappa \approx 0.5 \) then concentration successfully occurs.

V. ENTANGLEMENT IN THE CLOSEST GAUSSIAN STATES

Wolf et al. [40] proved that for any non-Gaussian state \( \hat{\rho} \) and a Gaussian state \( \hat{\rho}_G \) possessing the same first and second moments as \( \hat{\rho} \), the entanglement \( E(\hat{\rho}) \) of \( \hat{\rho} \) is bounded from below by the entanglement of the Gaussian state \( E(\hat{\rho}_G) \).

However, this only holds if the measure \( E \) used to quantify the entanglement of the state is continuous and strongly superadditive.

Notoriously, the logarithmic negativity \( E \) is not strongly superadditive (although it is additive) and so it does not always hold that \( E(\hat{\rho}) \geq E(\hat{\rho}_G) \). However, for the non-Gaussian mixed states’ output from protocols I and II, this does indeed hold.

Figure 7 shows the entanglement of the Gaussian mixed state with the same covariance matrix as that of the state output from protocol I and can be compared directly with Fig. 2. The covariance matrix of this Gaussian state is given simply by

\[
M = \sum_{ij} \left( -1 \right)^{i+j} \rho^{(i)j} \gamma_A^{(i)} \gamma_B^{(j)}
\]

and the logarithmic negativity can be calculated easily.

As can be seen from the figure, no matter the interaction strength, the Gaussian state is less entangled than the input two-mode squeezed vacuum. Indeed, for low values of squeezing \( r \), the Gaussian state is separable!

A similar result is seen in Fig. 8 in which the logarithmic negativity of the closest Gaussian state to the output state of protocol II is plotted. Figure 8 can be compared directly with Fig. 6.

These results suggest that the entanglement is primarily provided by the higher-order moments of the non-Gaussian output states of these two protocols. Although the closest Gaussian states may contain entanglement for some values of squeezing \( r \), the Gaussian state can never contain more entanglement than the two-mode squeezed vacuum at the same level of squeezing.

VI. DISCUSSION AND CONCLUSIONS

We have presented here two procrustean entanglement concentration schemes utilizing quantum nondemolition (QND)
interactions and photon detectors. The first scheme relied upon QND interactions between Gaussian ancillary modes and a TMSV to successfully subtract photons from the TMSV as heralded by on-off detectors. We have shown how to efficiently calculate the density matrix elements of the resulting quantum state, which can then be used to calculate the logarithmic negativity of the state. This is a nontrivial task. In the asymptotic limit of the QND interaction strength $k \to 0$, one would find that a single photon is subtracted from each arm of the TMSV, although the probability of heralding entanglement concentration approaches zero. This is intuitively correct, as the behavior mimics the protocol of [9] in which highly transmissive beam splitters are used in place of weak QND interactions. There, as the transmittance approaches 1, one finds that a single photon is subtracted from each arm, which is the optimal outcome for entanglement concentration. The results of Browne et al. [41] can be revisited by replacing $S_{\text{QND}}$ with $S_{\eta}(T)$ in Eq. (10), where $S_{\eta}(T)$ is the symplectic operation corresponding to a beam-splitter transformation.

We find that the method we have used for calculating density matrix elements for this protocol offers an improvement in numerical speed and efficiency over calculations in the Fock basis directly.

Double-pass schemes, i.e., letting the ancillae interact with the TMSV twice, do not give any advantage in protocol I. For example, $S_{\text{QND}}^{(2)} S_{\text{QND}}^{(1)} = S_{\text{QND}}^{(3)}$, but the logarithmic negativity is highest for weak interaction strength. By altering the interaction between passes, e.g., $S_{\text{QND}}^{(2)} S_{\text{QND}}^{(1)}$, there is some advantage over a double-pass scheme using the same interaction twice, but this is still eclipsed by the single-pass schemes.

As to detector efficiencies, at low levels of initial TMSV squeezing $r$, there can still be an increase in entanglement for weak interactions such as $k = 0.1$ when the detector efficiency exceeds approximately 50%, $\eta > 0.5$.

The second protocol relied on QND interactions to mix a TMSV with photon-subtracted, squeezed vacuum modes, that is, with non-Gaussian ancillae. The non-Gaussian ancillary modes are then detected after the interaction using homodyne detectors. As was stated in Sec. IV, the success of this scheme is dependent on the initial squeezing of the ancillary modes, the choice of interaction, and the angle $\Theta$ of homodyning. It is most likely that a homodyne measurement yields a result $z = 0$, and the logarithmic negativity shows an improvement as long as the measurement outcome does not stray too far from this.

The benefit of the second protocol is that the probabilistic non-Gaussian state preparation could be performed off-line and therefore the efficiency $\eta$ of the on-off detectors could be assumed to be high. If this were the case, then only the homodyne detector efficiency needs to be taken into account.

Regarding the implementations of the protocols presented here, in a purely optical setup in which the TMSV is created by parametric down-conversion and the QND operations are performed with beam splitters and squeezers, protocol I is probably the easiest to implement, although other effects such as a nonzero dark count rate of the detectors would need consideration. However, protocol II seems to be the best choice in the atomic case. In the atomic systems of [20], the entangled macroscopic spin states of two cesium gas samples represent the TMSV and off-resonant dipole interactions with strongly polarized light from the QND interactions. The vacuum ancillary modes are not true vacuum modes but modes of strongly polarized light. This will represent one of the main experimental challenges for protocol I, as the heralding of the entanglement concentration comes from detecting photons for which the interaction has altered the polarization, which would then require heavy filtering. Otherwise, the probabilities of success in both cases are comparable with those for the beam-splitter-based light schemes, already demonstrated [13] in the laboratory. The possibility of distilling entanglement in atomic ensembles represents the main motivation for using a general QND Hamiltonian for entanglement concentration.

To circumvent the problem of the non-Gaussian outputs in the presented protocols, iterative probabilistic Gausification procedures similar to [4,41] can be used to drive many copies of the output non-Gaussian entangled states towards fewer copies of Gaussian entangled states.

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APPENDIX: MULTIVARIATE HERMITE POLYNOMIALS

In order to calculate the density matrix elements of a Gaussian state, the multivariable Hermite polynomials [42] are an invaluable tool. The four-dimensional Hermite polynomials are defined by

\[
H_{r,s,t,v}^{(\Theta,\Delta)}(y_1,y_2,y_3,y_4) = (-1)^{t+s+r+v} \exp(y^T \Theta y + \Delta y) \frac{\partial^r}{\partial y_1^r} \frac{\partial^s}{\partial y_2^s} \frac{\partial^t}{\partial y_3^t} \frac{\partial^v}{\partial y_4^v} \exp(-y^T \Theta y - \Delta y),
\]

where $y = (y_1,y_2,y_3,y_4)^T$, $\Theta$ is a $4 \times 4$ matrix, and $\Delta = (\Delta_1,\Delta_2,\Delta_3,\Delta_4)$. The matrix $\Theta$ is related to the inverse of the covariance matrix of the Gaussian state, and the matrix $\Delta$ is related to any displacements. The density matrix elements are found by evaluating the Hermite polynomials at $y = 0$. A recursion formula can be derived to help speed up the numerical calculation of the multivariable Hermite polynomials by directly using (A1) and substituting there, e.g., $r + 1$. This gives

\[
H_{r+1,s,t,v}^{(\Theta,\Delta)} = \Delta_1 H_{r,s,t,v}^{(\Theta,\Delta)} - 2r \Theta_{11} H_{r-1,s,t,v}^{(\Theta,\Delta)} + 2s \Theta_{22} H_{r,s-1,t,v}^{(\Theta,\Delta)} + 2t \Theta_{33} H_{r,s,t-1,v}^{(\Theta,\Delta)} + 2v \Theta_{44} H_{r,s,t,v-1}^{(\Theta,\Delta)}.
\]

In the above, $H_{r,s,t,v}^{(\Theta,\Delta)} \equiv H_{r,s,t,v}^{(\Theta,\Delta)}(0,0,0,0)$. Similar formulas can be derived for $H_{r,s,t+1,v}^{(\Theta,\Delta)}$, etc. by replacing the coefficients.
in Eq. (A2). There are some symmetries which can be exploited. As the resulting density matrix is necessarily Hermitian, \( H_{r,s,t,v}^{(x,\Delta)} = H_{r,s,t,v}^{(x,\Delta)} \). Furthermore, if \( \Delta = 0 \) as in protocol I or in protocol II if the homodyne projection yields \( z = 0 \), then only Hermite polynomials for which the sum of the indices is an even number need to be calculated, as whenever the sum is odd, the corresponding polynomial is zero.