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Geometry of perturbed Gaussian states and quantum estimation

Marco G Genoni1, Paolo Giorda2 and Matteo G A Paris3

1 QOLS, Blackett Laboratory, Imperial College London, London SW7 2BW, UK
2 Institute for Scientific Interchange Foundation, I-10133 Torino, Italy
3 Dipartimento di Fisica, Università degli Studi di Milano, I-20133 Milano, Italy

E-mail: m.genoni@imperial.ac.uk, giorda@isi.it and matteo.paris@fisica.unimi.it

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Abstract
We address the non-Gaussianity (nG) of states obtained by weakly perturbing a Gaussian state and investigate the relationships with quantum estimation. For classical perturbations, i.e. perturbations to eigenvalues, we found that the nG of the perturbed state may be written as the quantum Fisher information (QFI) distance minus a term depending on the infinitesimal energy change, i.e. it provides a lower bound to statistical distinguishability. Upon moving on isoenergetic surfaces in a neighbourhood of a Gaussian state, nG thus coincides with a proper distance in the Hilbert space and exactly quantifies the statistical distinguishability of the perturbations. On the other hand, for perturbations leaving the covariance matrix unperturbed, we show that nG provides an upper bound to the QFI. Our results show that the geometry of non-Gaussian states in the neighbourhood of a Gaussian state is definitely not trivial and cannot be subsumed by a differential structure. Nevertheless, the analysis of perturbations to a Gaussian state reveals that nG may be a resource for quantum estimation. The nG of specific families of perturbed Gaussian states is analysed in some detail with the aim of finding the maximally non-Gaussian state obtainable from a given Gaussian one.

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(Some figures in this article are in colour only in the electronic version)

1. Introduction

Non-Gaussianity (nG) is a resource for the implementation of continuous variable quantum information in bosonic systems [1]. Several schemes to generate non-Gaussian states from Gaussian ones have been proposed, either based on nonlinear interactions or on conditional
measurements [2–25]. In many cases the effective nonlinearity is small, and so it is the resulting nG. It is thus of interest to investigate the nG of states in the neighbourhood of a Gaussian state, i.e. the nG of slightly perturbed Gaussian states. Besides the fundamental interest [26], this also provides a way to assess different de-Gaussification mechanisms, as well as nG itself as a resource for quantum estimation. Indeed, in an estimation problem where the variation of a parameter affects the Gaussian character of the involved states, one may expect the amount of nG to play a role in determining the estimation precision.

Quantum estimation deals with situations where one tries to infer the value of a parameter λ by measuring a different quantity X, which is somehow related to λ. This often happens in quantum mechanics and quantum information where many quantities of interest, e.g. entanglement [27, 28], do not correspond to a proper observable and should be estimated from the measurement of one or more observable quantities [29]. Given a set {ϱi} of quantum states parametrized by the value of the quantity of interest, an estimator ḡ for λ is a real function of the outcomes of the measurements performed on ϱi. The quantum Cramer–Rao theorem [30–33] establishes a lower bound for the variance Var(λ) of any unbiased estimator, i.e. for the estimation precision. Var(λ) ≥ (MH(λ))−1 in terms of the number of measurements M and the so-called quantum Fisher information (QFI), which captures the statistical distinguishability of the states within the set. Indeed, the QFI distance itself is proportional to the Bures distance ⟨ϱi|log ⟩⟨ϱi| of infinitesimally close states corresponding to infinitesimally close values of the parameter, i.e. in terms of metrics, H(λ) = 4gλ = 2∑nm(⟨en + ϱ∥⟩|⟨ψm|δλϱλ|ψn⟩|^2 where dρ(ϱλ, ϱλ) = gλ dλ^2, and we have used the eigenbasis ϱλ = ∑n⟩⟨en| of a quantum state ϱλ.

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2. Gaussian states and a measure of non-Gaussianity

Let us consider a single-mode bosonic system described by the mode operator a with commutation relations [a, a†] = 1. A quantum state ϱ is fully described by its characteristic function χ[ϱ] = Tr[ϱ D(λ)] where D(λ) = exp[λ a† − λ∗ a] is the displacement operator. The canonical operators are given by q = (a + a†)/√2 and p = (a − a†)/√2i with commutation relations given by [q, p] = i. Upon introducing the vector R = (q, p)T, the covariance matrix σ and the vector of mean values X of a quantum state ϱ are defined as σij = 1/2⟨Ri Rj⟩ϱ − ⟨Ri⟩ϱ ⟨Rj⟩ϱ and Xj = ⟨Rj⟩ϱ, where ⟨O⟩ϱ = Tr[O ϱ] is the expectation value of the operator O on the state ϱ. A quantum state is said to be Gaussian if its characteristic function has a Gaussian form. Once the CM and the vectors of mean values are given, a Gaussian state is fully determined.

The amount of nG δϱ[ϱ] of a quantum state ϱ may be quantified by the quantum relative entropy [34] S(ϱ∥ϱ0) = Tr[ϱ log ϱ − log ϱ0] between ϱ and its reference Gaussian state ϱ0, which is a Gaussian state with the same covariance matrix σ as ϱ. As for its classical counterpart, the Kullback–Leibler divergence, it can be demonstrated that 0 ≤ S(ϱ∥ϱ) < ∞ when it is definite, i.e. when supp ϱ ⊆ supp ϱ0. In particular, S(ϱ∥ϱ0) = 0 iff ϱ = ϱ0 [35, 36]. Since τϱ is Gaussian Tr[(τϱ − ϱ) log τϱ] = 0, i.e. S(ϱ∥ϱ0) = S(τϱ) − S(ϱ), we may write δϱ[ϱ] = S(τϱ) − S(ϱ) where S(ϱ) = −Tr[ϱ log ϱ] is the von Neumann entropy of ϱ. Finally, since the von Neumann entropy of a single-mode Gaussian state may be written as h(√det σϱ), we have

\[ \delta \rho[\rho] = h(\sqrt{\det \sigma}) = S(\rho). \]

A generic single-mode Gaussian state may be written as τ = USρ US† where US is a symplectic operation, i.e. a unitary US = exp(−iH) resulting from a Hamiltonian H at most
quadratic in the field operators, and \( n \) is a chaotic (maximum entropy) state with \( n_v = \text{Tr}[\rho_a+a_a\rho] \) average thermal quanta, i.e. \( \psi = \sum_k p_k |k\rangle \langle k|, \quad p_k = n_k^2/(1+n_k) \) in the Fock number basis.

3. Classical perturbations to a Gaussian state

An infinitesimal perturbation of the eigenvalues \( p_k \) of a Gaussian state \( \tau \), i.e. \( p_k \to p_k + d p_k \) results in a perturbed state \( \varrho = \sum_k (p_k + d p_k) U_k |k\rangle \langle k| U_k^\dagger \) which, in general, is no longer Gaussian. Since the nG of a state is invariant under symplectic operations we have \( \delta[\varrho] = \delta[\eta] \), where \( \eta = U_\varrho \rho U_\varrho = \sum_k (p_k + d p_k) |k\rangle \langle k| \) is diagonal in the Fock basis. The Gaussian reference \( \tau_\eta \) of \( \eta \) is a thermal state with \( n_\eta = n_v + d n_\varrho = \sum_k k (p_k + d p_k) \) average quanta and the nG may be evaluated upon expanding both terms in \( \delta[\eta](= \delta[\varrho]) \) up to the second order,

\[
\delta[\varrho] = \sum_k \frac{d p_k^2}{2 p_k} - \frac{d n^2}{2 n_v (1+n_v)}.
\]

The nG of perturbed states is thus given by the sum of two contributions. The first term is the Fisher information of the probability distribution \( \{ p_k \} \), which coincides with the classical part of the Bures distance in the Hilbert space. The second term is a negative contribution expressed in terms of the infinitesimal change of the average number of quanta. When travelling on surfaces at constant energy the amount of nG coincides with a proper distance in the Hilbert space and, in this case, it has a geometrical interpretation as the infinitesimal Bures distance. At the same time, since Bures distance is proportional to the QFI one, it expresses the statistical distinguishability of states, and we conclude that moving out from a Gaussian state towards its non-Gaussian neighbours is a resource for estimation purposes. Similar conclusions can be made when comparing families of perturbations \( \{ d p \} \) corresponding to the same infinitesimal change of energy \( d n^2 \): in this case the different amounts of nG induced by the perturbations are quantified by the Bures distance minus a constant term depending on \( d n^2 \) and the initial thermal energy \( n_v \), i.e. the initial purity \( \mu = \text{Tr}[\rho^2] = \text{Tr}[\tau^2] = (2n_v + 1)^{-1} \). We summarize the above statements in the following.

Theorem 1. If \( \tau_\eta \) is a Gaussian state and an infinitesimal variation of the value of \( \lambda \) drives it into a state \( \varrho_{\lambda+\lambda_0} \) with the same eigenvectors, then the QFI distance is equal to the nG \( \delta[\varrho_{\lambda+\lambda_0}] \) plus a term depending both on the infinitesimal variation of energy \( d n^2 \) and on the initial purity

\[
H(\lambda) \, d \lambda^2 = \delta[\varrho_{\lambda+\lambda_0}] + \frac{2 \mu^2 \, d n^2}{1 - \mu^2}.
\]

In particular, for perturbations that leave the energy unperturbed, the nG of the perturbed state coincides with the QFI distance, whereas, in general, it provides a lower bound.

3.1. Examples of finite perturbations

In order to explore specific directions in the neighbourhood of a Gaussian state \( \tau \), let us write the perturbation to the eigenvalues as \( d p_k = \epsilon \mu_k \) where \( \{ \mu_k \} \) is a given distribution. In this case the nG of the perturbed state is given by

\[
\delta[\varrho] = \epsilon^2 \left( \sum_k \frac{(p_k - \mu_k)^2}{2 p_k} - \frac{\Delta n_k^2}{2 n_v (1+n_v)} \right) + O(\epsilon^3),
\]

where \( \Delta n_k = \sum_k (p_k - \mu_k) k |a a| k \). Let us now consider the families of states generated by the convex combination \( \varrho = (1 - \epsilon) \tau + \epsilon \varrho_\mu \) of the Gaussian states \( \tau \) with the target state...
\( \rho_\mu = \sum_k \mu_k \ket{k} \bra{k} U_S^\dagger \), which itself is obtained by changing the eigenvalues \( p_k \) of the initial Gaussian state to \( \mu_k \). Again we exploit invariance of \( \delta[\rho] \) under symplectic operations and focus attention to the diagonal state \( \eta = U_S^\dagger \rho U_S = (1-\epsilon)\eta + \epsilon \eta_\mu \) which has the same nG of \( \rho \). This is the generalization to a finite perturbation of the analysis reported in the previous section, and it is intended as a mean to find the maximally non-Gaussian state obtainable starting from a given Gaussian. The nG of this kind of states can be written as 

\[
\delta[\eta] = S(\tau_\eta) - S(\eta) = h[n_\eta + 1/2] - H[q^{(\epsilon)}],
\]

where \( H[q^{(\epsilon)}] = -\sum q^{(\epsilon)}_k \log q^{(\epsilon)}_k \) denotes the Shannon entropy of the distribution \( q^{(\epsilon)}_k = (1-\epsilon)p_k + \epsilon \mu_k \) and \( n_\eta = \text{Tr}[\eta a^\dagger a] = (1-\epsilon)n_t + \epsilon n_\mu \) is the average number of quanta of \( \eta \). Note that for a thermal state with \( n_t \) quanta we have \( H[p] = h[n_t+1/2] \).

Using the concavity of the Shannon entropy we obtain an upper bound for the nG

\[
\delta[\eta] \leq h[n_\eta + 1/2] - h[n_t + 1/2] + \epsilon (h[n_t + 1/2] - H[\mu]). \tag{4}
\]

In particular, if the two distributions have the same number of quanta, \( n_\mu = n_t \), and thus \( n_\eta = n_t \), the bound on \( \delta[\eta] \) only depends on the difference between the entropy of the initial and target distributions.

Let us now consider perturbations towards some relevant distributions, i.e. Poissonian \( \mu_k^{(p)} = n_k^\epsilon e^{-n_t} \), thermal \( \mu_k^{(t)} \), and Fock \( \mu_k^{(d)} = \delta_{k,n_\mu} \) and evaluate the nG of states obtained as a convex combination of a thermal state with \( n_t \) quanta and a diagonal quantum state with a Poissonian, thermal or Fock distributions and \( n_\mu \) quanta.

In figure 1, we plot the nG of the convex combination \( \rho \) as a function of \( n_\mu \) for different values of \( \epsilon \): if we consider the convex combination with a Fock state, the nG simply increases monotonically with the energy of the added state. For combinations with Poissonian and thermal distributions, we have a maximum for \( n_\mu \to 0 \); then a local minimum (which for the thermal distribution corresponds trivially to \( \delta[\rho] = 0 \)) for \( n_\mu = n_t \), and then the nG increases.
again for higher values of $n_\lambda$. This implies that in order to increase $n_G$ the best thing to do is to perturb the initial Gaussian either with a highly excited state $n_\mu \gg 1$ or with the vacuum state $|0\rangle\langle 0|$. If we choose $n_\rho = n_\mu = n_\tau$, we know from equation (2) that $n_G$ for small perturbations is equivalent to the Bures infinitesimal distance between the two probability distributions $\mu_k$ and $p_k$. In figure 1 (lower-right panel), we show the $n_G$ as a function of $\epsilon$: as is apparent from the plot, the $n_G$ obtained by adding a Fock state is always much larger than the one for a Poissonian profile. We observe that in the latter case, the expansion at the second order obtained in equation (3) is still accurate for values of $\epsilon$ approaching 1, while it fails to be accurate for $\epsilon \gtrsim 0.2$ for a Fock state, becoming an upper bound on the exact amount of $n_G$. We have also investigated what happens by considering a target distribution randomly chosen on a finite subspace of the infinite Hilbert space: again we have obtained that, at fixed energy of the target state, perturbing with a Fock state yields the biggest increase of $n_G$. This may be easily generalized to a system of $d$ bosonic modes, where the most general Gaussian state is described by $d(2d + 3)$ independent parameters.

4. Perturbations at a fixed covariance matrix

Gaussian states are known to be extremal states at a fixed covariance matrix for several relevant quantities, e.g. channel capacities and entanglement measures [37]. Therefore, one may wonder whether perturbing a Gaussian state at a fixed covariance matrix may be quantified in a convenient way for the purposes of quantum estimation. This indeed is the case: the $n_G$ provides an upper bound to the QFI distance at a fixed covariance matrix and thus have an operational interpretation in terms of statistical distinguishability. This is more precisely expressed by the following theorem [1].

**Theorem 2.** If $\tau_\lambda$ is a Gaussian state and an infinitesimal variation of the value of $\lambda$ drives it into a state $\varrho_{\lambda+d\lambda}$ with the same covariance matrix, then the $n_G$ of $\delta[\varrho_{\lambda+d\lambda}]$ provides an upper bound to the QFI distance

$$H(\lambda) d\lambda^2 \leq \delta[\varrho_{\lambda+d\lambda}].$$

**Proof.** If $\varrho_{\lambda+d\lambda}$ and $\tau_\lambda$ have the same CM, then the $n_G$ of $\varrho_{\lambda+d\lambda}$, $\delta[\varrho_{\lambda+d\lambda}] = S(\varrho_{\lambda+d\lambda}||\tau_\lambda) = \tilde{H}(\lambda) d\lambda^2$, where the so-called Kubo–Mori–Bogolubov information $H(\lambda)$ [32, 38] provides an upper bound for the QFI $H(\lambda) \leq \tilde{H}(\lambda)$ [39], thus proving the theorem.

The above theorem says that a larger $n_G$ of the perturbed state may correspond to a greater distinguishability from the original one, thus allowing a more precise estimation. Of course, this is not ensured by the theorem, which only provides an upper bound to the QFI. One may wonder that when $\varrho_{\lambda+d\lambda}$ is itself a Gaussian state the theorem requires $H(\lambda) = 0$, i.e. no reliable estimation is possible. Indeed, this should be the case since Gaussian states are uniquely determined by the first two moments, and thus the requirement that the perturbed $\tau_{\lambda+d\lambda}$ and the original state $\tau_\lambda$ are both Gaussian and have the same covariance matrix implies that they are actually the same quantum state.

5. Conclusions

In conclusion, we have addressed the $n_G$ of states obtained by weakly perturbing Gaussian states and have investigated the relationships with quantum estimation. We found that $n_G$ provides a lower bound to the QFI distance for classical perturbations, i.e. perturbations to eigenvalues leaving the eigenvectors unperturbed, and an upper bound for perturbations...
leaving the covariance matrix unperturbed. For situations where the CM is changed by the perturbation, we have no general results. On the other hand, it has already been shown that non-Gaussian states improve quantum estimation of both unitary perturbations as the displacement and the squeezing parameters \[40\] and nonunitary ones as the loss parameter of a dissipative channel \[41\]. Overall, our results show that the geometry of non-Gaussian states in the neighbourhood of a Gaussian state is definitely not trivial and cannot be subsumed by a differential structure. Despite this fact, the analysis of perturbations to a Gaussian state may help in revealing when, and to which extent, nG is a resource for quantum estimation. We have also analysed the nG of specific families of perturbed Gaussian states with the aim of finding the maximally non-Gaussian state obtainable from a given Gaussian one.

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