

University of the Basque Country

PhD Thesis - 2016



LOWER BOUNDS ON QUANTUM METROLOGICAL PRECISION



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Prologue

This work is part of doctoral project which I started on the sumer of 2013. This work collects almost all the research I have done on those previous years. I try to be as clear as possible, writing from the very basics concepts of modern Quantum Physics until the most recent results on Quantum Metrology and Quantum Information in a clear and concise way, such that it is understandable for a more general public.

The research work I collect in this thesis was done in collaboration with members of the group I belong to. Even if not all of them are at the present day investigating on our group, the collaborators namely were Dr. Philipp Hyllus, Dr. Giuseppe Vitagliano, Dr. Iñigo Urizar-Lanz, Dr. Zóltan Zínboras, Dr. Matthias Kleinmann and Prof. Géza Tóth. Apart from the group GEDENPQOT, based on Bilbao, this thesis also collects some works done in collaboration with the group Theoretical Quantum Optics lead by Prof. Otfried Gühne of the University of Siegen, in Germany, and the group of Prof. Carsten Klempt at the Universitu of Hannover, in Germany too.

Publications

lagoba Apellaniz *et al, New J. Phys.* **17** 083027 (2015) Detecting metrologically useful entanglement in the vicinity of Dicke states

Géza Tóth and Iagoba Apellaniz, *J. Phys. A: Math. Theor.* **47** 424006 (2014) Quantum metrology from a quantum information science prespective

Preprints

Out of the scope of this thesis

Giuseppe Vitagliano *et al* 2014 *Phys. Rev. A* **89** 032307 Spin squeezing and entanglement for an arbitrary spin

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SLD - Symmetric logarithmic derivative

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QFI - Quantum Fisher information

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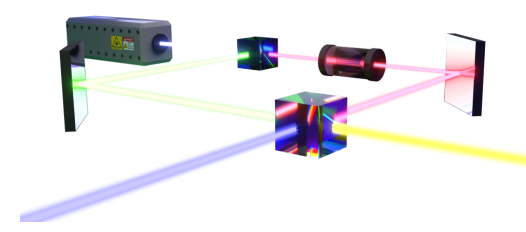
Lower bounds on quantum metrological precision

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September 7, 2016

To my parents, my family and to all the people
I have had around those years.

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I have been visiting several research groups during those years and I don't forget them, they have accept me as one more between them. So I want to thank a very especial research group for me, the Theoretical Quantum Optics group at the University of Siegen. I would really like to mention the names of all of them but I think it would be quite heavy for the average reader of this thesis. Thank you guys! Also I want to thank people from the group QSTAR at Florence, Italy.

On the other hand I also felt very comfortable at my university, the University of the Basque Country, but I want to thank especially the people that make me grow in all ways as person.

1 Introduction

N the recent years...

The figure of merit for the precision is the inverse of the variance normalized with the number of particles, $(\Delta\Theta)^{-2}/N$. It has the following properties:

- (i) The bigger it is the bigger is the precision
- (ii) It is normalized so for the best separable state it is 1. For greater values than 1 it would be a non-classical sign.

SQL

$$(\Delta\Theta)^{-2} \le N \tag{1.1}$$

HL

$$(\Delta\Theta)^{-2} \le N^2 \tag{1.2}$$

This thesis consists of 4 well differentiated parts, apart from the current introduction, on which different topics are developed. In the first part, we will introduce the reader onto the research field

of quantum metrology.

Brief comments on the notation: c_{Θ} and s_{Θ} stand for $cos(\Theta)$ and $sin(\Theta)$ trigonometric functions respectively. We will skip on writing the tensor-product notation \otimes if it is not strictly necessary for the correct comprehension of the text.

Background on Quantum Metrology

M ETROLOGY, as the science of measuring, has played an essential role for the development of the technology as we know it today. It studies several aspects of the estimation process, such as which strategy to follow in order to improve the precision of an estimation. One of the most important figure of merit in this context is the achievable precision for a given system, independently of the strategy. We will show how to characterize it in the subsequents sections. And we will show as well different strategies to achieve the desired results. The metrology science also covers from the design aspects of a precise measuring device, until the most basic concepts of nature which lead in ultimate instance to the better understanding of the whole process.

In this sense, with the discovery of the Quantum Physics and the development of Quantum Mechanics, new doors for advances in metrology were open on the earlies decades of the 19th century. Later on, the Quantum Theory lead to the so-called field of Quantum Information which merges the notions of the theory of information and computer science, among others subfields, with the quantum mechanics. The role of the so-named entanglement, an exclusive feature of Quantum Mechanics, is essential in this context. Its complete understanding has integrated efforts of many researches world wide. Said this, the entanglement also is in the center of theoretical concepts in Quantum Metrology.

On the other hand and with the aim of interpreting raw data, there are the statistics, without which many descriptions of the actual and past physical findings would lack of the rigorous interpretation

Tree #	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18
Heights (m)	6	7	5	7	2	11	7	4	7	2	7	7	5	5	5	8	3	10

Table 2.1: A set of values for the heights of 18 trees. All measurements were rounded to integers for simplicity.

needed for the complexity of data samples.

2.1 Background on statistics and the theory of estimation

The main mathematical tools used by the metrology science belong to statistics. Moreover we are also interested on estimation theory. The statistics main characteristic is that makes the raw data under consideration comprehensible from the human perspective. The data can be anything, from a set of different heights of a basketball team, to the outcomes of a coin toss or the ages of a hundred students or even the outcomes of a thousand times repeated measurement of the electric field at some spatial point. The aim of this section is to give the reader sufficient material to follow this thesis and make it comprehensible from the beginning.

2.1.1 Data samples, average and variance

As we mentioned above, everything in statistics starts with a data sample. A data sample is defined as a set of values, we will restrict ourselves to quantitative values for simplicity, representing some magnitude. So, in our framework such a set can be written as $X = \{x_i\}_{i=1}^M$, where all M values are collected. An illustrative example with 18 heights of different trees is shown on the Table 2.1. It is worth to notice that this data sample represents a continuous variable, the height of some trees, while we have grouped more than a single instances into integer values. If we would use more decimals to describe the data probably all values would have been different among each other. This discussion is not so essential when discrete outcomes are analyzed

This picture can be extend to more outcomes coming from a single *event* or measure. For instance, a second value such as the width of the tree could be attached to each height on the previous example. This way we have opened our approach for higher dimensional data samples that can be described as a set of M outcomes with more than a single value each, $(X, Y, \ldots) = \{x_i, y_i, \ldots\}_{i=1}^M$.

Average

With this data at hand one of the first questions arises is whether the values of X are around

some mean value. If one tries to address this question one of the first approaches can be computed is the $arithmetic\ average$ of the set X, namely,

$$E[X] := \frac{1}{M} \sum_{i=1}^{M} x_i.$$
 (2.1)

Apart from the arithmetic average there are other types of averages such as the geometric mean, the root mean square or the harmonic mean, etc. For more information see those some of those fascinating works about the statistics [XXX]. For us and for the complete understanding of this thesis will be enough with the arithmetic average.

Let us take the previous example of the trees and compute the average. In this case the data sample is constituted by positive integer values for simplicity. The average of the data sample is straight forward $E[X] = \frac{1}{18} \sum_{i=1}^{18} x_i = 6 \text{m}$.

For the extension to the case of higher dimensional data samples one can say that each kind of outcome the average can be computed in a seamless way, $E[Y] = \frac{1}{M} \sum_{i=1}^{M} y_i$ and so on.

Variance

The second quantity one can compute is how spread the data is. This is usually done first subtracting the average to all values on the data sample, then squaring them, so the sign of the result is lost, and finally one averages over all the resulting set. This quantity is called the *variance* of the sample and it is written as follows,

$$V[X] := E[(X - E[X])^2],$$
 (2.2)

where X can be seen as a vector and when subtracting the scalar E[X] and X^2 stands for the elements wise squared of X, namely $X^2 := \{x_i^2\}_{i=1}^M$. The variance can also be written alternatively as $V[X] = E[X^2] - E[X]^2$. The definition of the *standard deviation* follows directly from the variance, $\sigma_X = \sqrt{V[X]}$. Many quantities on statistics require operations like the one described right before. This can be generalised again for more than one kind of outcome as it was done with the average.

Now let us see how result the variance for the example of the Table 2.1. In this case the variance would be the following, $V[X] = \frac{1}{18} \sum_{i=1}^{18} x_i^2 - 6^2 m^2 = 5.5 \overline{5} m^2$.

To summarise and using the standard deviation, one can say now about the original distribution that most values are around 6m with a deviation of 2.357m. Of course, some values are outside this range, but nevertheless the description is quite accurate, note that 12 values from 18 are inside the range $6m \pm 2.357m$.



Figure 2.1: (a) p_i for different values of i. All bars have width as 1, so it is drawn how many times each data value appears on the data sample. The width of the bars is called bin, and it can change so the bars would represent a wider range of values. (b) Yo can see the same data represented in this case by a histogram with the bin size equal to 2. To produce this histogram we have summed 2 adjacent p_i values starting from $p_1 + p_2$, $p_3 + p_4$ and so on. Those new bars we have chosen to represent a value in between, for instance (3 + 4)/2 = 3.5 and so on.

Histograms

At this point we introduce the *histograms* and with that, first, the *distribution function* of the data sample. Returning to the example of the Table 2.1, one may notice that the value 7m is repeated 6 times, as the value 5m is 4 times and so on. This is represented with the distribution function p_i , which in this case is for discrete values of the outcomes but which can be generalized for continuous variables as we will see later. This function takes the value of how many times the outcome i has appeared on the data sample. Let us plot the distribution function, see Figure 3.3.

Now some question arises immediately: How this is fully connected with the previous picture? How can one compute the average and other interesting quantities? The answer is simple but it has to be considered carefully. First of all, notice that p_i is defined for all the natural numbers including zero, see that in the example of the trees p_9 equals zero, so it can be extended for other heights too setting them to zero. This will depend on the physical property that our data sample represents but in this case it is the height of some trees, so the values cannot be negative. Second, notice that the sum of all the repetitions, all the values of p_i , is exactly 18 the number of data samples in the set. So we have that $M = \sum_{i=0}^{\infty} p_i$. Now we can formulate the ensemble average as $E[X] = \sum_{i=0}^{\infty} p_i i / \sum_{j=0}^{\infty} p_j$. The variance and with this the standard deviation immediately follow this approach. It is convenient to notice that the total measurement outcomes has not contribute anything but to normalize the quantities.

We can now without losing generality redefine the distribution function to be the number of

repetitions corresponding to the variable divided by the total outcomes, in this case M. It would have the same properties of a probability distribution function (PDF). For instance, now we have that the sum of all p_i -s equal to one, $\sum_{i=0}^{\infty} p_i = 1$, and the average is directly obtained by computing $E[X] = \sum_{i=0}^{\infty} p_i i$. This is the approach we will follow to represent data samples. The variance and other quantities also are simpler in this way.

2.1.2 Probabilities and frecuentist vs. bayesian approach

Let us talk now about probabilities. The notion of the probability comes, [XXX]. In our contest we will define the probability in the following way. Imagine we have an unknown data sample, let say X, from which we randomly choose one of the data values. The probability to obtain a random variable x is given by the parent distribution function divided by the population number M.

We can estimate the average height of the trees of that forest.

histogram

estimator

Explain how for bigger bin sizes, the error for higher statistical moments increases.

MLH

CLT

2.1.3 Estimators, Fisher information

Error propagation formula.

2.2 Quantum Mechanics from metrology perspective

The ubiquitous probabilistic nature of quantum mechanics forces all the community to know some basics on probability and statistics.

The quantum state, multi-particle state, entanglement

A system state in Quantum Mechanics lives on a Hilbert space, \mathcal{H} . The system state, ρ , has the following properties:

i) It is Hermitian, so it is invariant under the complex transposition, $\rho = \rho^{\dagger}$.

- ii) Its trace is equal to one, $tr(\rho) = 1$.
- iii) It is positive semi-definite, *i.e.*, all its eigenvalues are bigger or equal to zero, $\rho = \sum_{\lambda} p_{\lambda} \Pi_{\lambda}^{1}$ where $p_{\lambda} \geq 0$ are the scalar eigenvalues. It follows that $\sum_{\lambda} p_{\lambda} = 1$.
- iv) If all p_{λ} are zero except one, the state is a pure state, $\rho = \Pi_{\lambda}$.
- v) It follows that the quantum states form a convex set where the extremal points are pure states.

The composite system of N different parties lives on $\mathcal{H}=\mathcal{H}^{(1)}\otimes\mathcal{H}^{(2)}\otimes\cdots\otimes\mathcal{H}^{(N)}$ or for short $\mathcal{H}=\bigotimes_{i=1}^N\mathcal{H}^{(i)}$, where \otimes stands for a tensor product like construction. For instance, this composite Hilbert space could be used to represent a many-particle system, in this case N particles. A *separable* state on this Hilbert space can be described in the following way,

$$\rho = \sum_{i} p_{i} \rho_{i}^{(1)} \otimes \rho_{i}^{(2)} \otimes \cdots \otimes \rho_{i}^{(N)}, \tag{2.3}$$

where p_i are convex weights that sum to one and are equal or bigger than zero. If not the state is said to be *entangled*.

Angular momentum operators. The angular momentum algebra comes

Entanglement cannot be described classically.

Evolution

Unitary evolution

Markov

Limblad

Measurements (POVM)

Quantum Information

2.3 Quantum Metrology

The evolution of a quantum state can be used to infer on some hidden parameter. The estimation theory applied to the intrinsic statistical nature of a quantum states has lead to the the formulation of Quantum Metrology as an important subfield. Merging the probabilistic features of quantum mechanics and the estimation theory is not trivial. Nevertheless, starting from the pioneering works

 $^{^{1}}$ $ho = \sum_{\lambda} p_{\lambda} \Pi_{\lambda}$ is the eigen-representation of the state, where Π_{λ} are the eigenstates defined on \mathcal{H} too. They are as many as the dimension, d, of the Hilbert space. They are orthonormal under the product defined on such Hilbert space \mathcal{H} , i.e., $\mathrm{tr}(\Pi_{\lambda}\Pi_{\lambda'}) = \delta_{\lambda,\lambda'}$. Nevertheless, there is a extended discussion when Hilbert space is defined for continuous variables, see [XXX].

of Wotters, Braunstein and some other I probably forgot to mention [XXX] were the statistical distance of neighbor states is studied we have lead to amazing results which today drive many experiments and technology. From the works of Giovannetti et al and Paris on the middle of the last decade fundamental concepts arose, for example the figure of merit quantum Fisher information or the Heisenberg limit. In this section we will highlight the most important aspects of this field and with this we will conclude this introductory chapter.

The error propagation formula for an estimation of Θ based on the observable O.

$$(\Delta\Theta)^2 \ge \frac{(\Delta O)^2}{|\partial_{\Theta}\langle O \rangle|^2} \tag{2.4}$$

2.3.1 Quantum Magnetometry

One of the most basic tasks of Quantum Metrology is to address the precision of estimating the magnetic field strength, namely B, of an unknown external magnetic field. In this section we will assume that the magnetic field is homogeneous on the position space. To consider more advanced situations on which the magnetic field changes linearly with the position, see chapter [XXX]. With the aim of estimating the strength of the magnetic field, a prove state is used in order to interact with it, coupling the magnetic moment of the state and the field itself. After some time, the state has evolved. Finally, measuring how the state has changed one would be able to infer on the strength of the magnetic field, basically proportional to the change on the state.

In general, we will say that the magnetic moments of the states come exclusively from the spin angular momentum, neglecting any possible contribution from the orbital angular momentum. This way the physics is simpler. This is justified in the sense that most of the recent experiments on this context have been carried out with ion-traps, BECs or at most cold atomic ensembles, which have indeed a negligible orbital angular momentum.

Beside this considerations, the interaction Hamiltonian can be written as,

$$H = -\boldsymbol{\mu} \cdot \boldsymbol{B} \tag{2.5}$$

up to some constant factor. Now in the simplest case we will choose the magnetic field to be pointing to some fixed direction, for instance, the Oz direction. So the magnetic field vector can be written as B = Bk, where k is the unitary vector pointing to the Oz direction. This way estimation problem is much more simple, since one has not to determine the direction of the magnetic field.

The magnetic moment of the system is proportional to the spin angular momentum, $\mu = -\mu_{\rm B} g_{\rm s} \hbar^{-1} J$,

where μ_B and g_s are the Bohr magneton and anomalous gyromagnetic factor respectively. Finally, one can rewrite the interaction Hamiltonian as,

$$H = \gamma B J_z \tag{2.6}$$

where $\gamma = \mu_B g_s \hbar^{-1}$ and we have used the fact that $\mathbf{J} \cdot \mathbf{k} = J_z$. Finally, the unitary operator leading the evolution of the system can be written as,

$$U = \exp(-i\Theta J_z), \tag{2.7}$$

where the magnetic field strength is encoded into the variable $\Theta = -\mu_{\rm B} g_{\rm s} t B/\hbar$. Here $\mu_{\rm B}$ stand for the Magneton of Bhor and $g_{\rm s}$ for the giro-magnetic constant for the spin angular momentum, and t is the time spent on the evolution.

Metrology in the vicinity of Dicke states

In this chapter we will present recent results regarding the metrological usefulness of a family of unpolarised states. Such states can be used as prove states to estimate the homogeneous magnetic field strength, see Section 2.3.1 for refeneces about magnetometry. It turns out that those unpolarised states are the only ones that can bite the Heisenberg limit, as it was shown on the Section [XXX]. Hence, those states have attracted the interest of the community.

One of the figures of merit of this such unpolarized, but still useful states is the so-called unpolarized Dicke state [XXX], which consists of, on its *Oz*-axis representation, an equal number of qubits pointing up and pointing down while the whole state is at the same time symmetrised. It can be written as follows,

$$|D_N^0\rangle \equiv |D_N\rangle := \left(\frac{N}{N/2}\right)^{-\frac{1}{2}} \sum_{k \in \sigma_k} P_k \left(|1\rangle^{\otimes N/2}|0\rangle^{\otimes N/2}\right),\tag{3.1}$$

where k are elements of the set of all possible unique permutations of N elements of 2 kinds, σ_s . This state is also called twin-fock state on different contexts. Such state is known to be highly entangled [XXX] and to reach Heisenberg scaling when used in magnetometry. For these and other reasons unpolarised Dicke states have been created in photonic systems [XXX], in cold-gases [XXX]

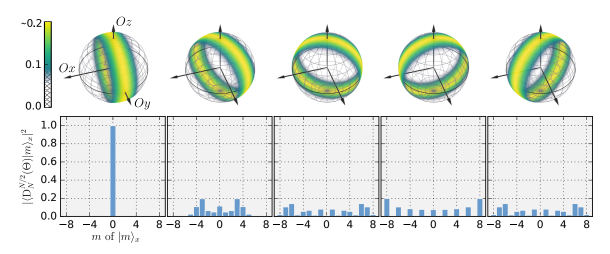


Figure 3.1: Secuence of the evolution of an unpolarized Dicke state of 16 qubits for $\Theta = \{i\pi/6\}_{i=0}^4$. Bloch spheres representing the Husimi quasi-probabilistic distribution of the state, and below PDF of the J_X POVM for each step of the secuence

and recently in trapped ions [XXX].

The symmetric subspace of N spin- $\frac{1}{2}$ particles is decomposed in N+1 orthonormal states, see Appendix B.2 for more details on how such subspaces behave even for different spins. Indeed, the subspace can be seen as a single spin- $\frac{N}{2}$ particle.

One of the most particular features this state has is that since it is a eigenstate of the collective operator J_z with corresponding eigenvalue equal to zero. At the same time, it lives on the subspace where the collective total spin is maximum, i.e., $\langle J^2 \rangle = N(N+2)/4$. Thus, with this together with the fact that the state is unpolarised, it turns that it must have a very large uncertainty for the collective spin operators perpendicular to J_z , namely J_x and J_y .

3.1 Unpolarised states for magnetometry

Having unpolarised states for magnetometry has been shown useful in Section XX. While the quantum Fisher information would give directly the performance one state would have, this is not feasible in general because a complete knowledge of the state is needed to compute such value. On the other hand, we can use the error propagation formula Equation (XX) to obtain a bound on the precision. As one can see on the Figure 4.6 a pure Dicke state is rotated along the Oz-axis. Say that in this case the initial state is an unpolarised Dicke state of aligned with the Ox-axis.

The state initially is an eigenstate of the J_x^k operators for $\forall k \in [1, \infty]$. Another feature of the sequence is that $\langle J_l \rangle = 0$ for $\forall l \in x, y, z$. It turns out that measuring the evolution of the second

statistical moment of J_x is one of the ways to go. It will start having value equal to zero for the pure unpolarized Dicke state, and rapidly will increase its value as it can be seen on the Figure 4.6. Another heuristic observation is that for $\Theta = \pi/2$ the value of $\langle J_x^2 \rangle$ will be at its maximum been it proportional to $\langle J^2 \rangle$ so to N^2 . Hence, the change on the second moment over the phase shift must be in this case proportional to N^2 . From this and for those states, we lead to the conclusion that one only needs to measure the second moment of the collective spin J_x to observe precisions that scale with the Heisenberg limit. In the following equation we show the error propagation formula that will give us the obtained precision,

$$(\Delta\Theta)^2 = \frac{(\Delta J_x^2)^2}{|\partial_{\Theta}\langle J_x \rangle|^2}.$$
 (3.2)

3.2 Evolution of the expectation values

With the aim of computing the precision, Equation (3.2), we will compute the dependence on Θ of the expectation value of the operator J_x and higher order moments. For that first of all we will move onto the Heisenberg picture where the operators evolve in time while the state remains the same. The operator J_x can be written a a function of Θ the following way,

$$J_x(\Theta) = e^{i\Theta J_z} J_x(0) e^{-i\Theta J_z} = J_x(0) c_{\Theta} - J_y(0) s_{\Theta},$$
 (3.3)

where $J_l(0)$ for $\forall l$ are the collective angular momentum operators at time equal zero, which we will write them simply J_l from now on, and c_{Θ} and s_{Θ} are the trigonometric functions introduced on the first chapter.

We need to compute the second and the fourth moments of J_x as it is required by the Equation (3.2). But before any calculation we will make a simplifying assumption which turn to be well supported on the most common situations. The assumption is that both expectation values are even functions on Θ ,

$$\langle J_x^2(\Theta) \rangle = \langle J_x^2(-\Theta) \rangle,$$

$$\langle J_x^4(\Theta) \rangle = \langle J_x^4(-\Theta) \rangle.$$
(3.4)

The square of J_x in the Heisenberg picture is written as follows,

$$J_x^2(\Theta) = J_x^2 c_{\Theta}^2 + J_y^2 s_{\Theta}^2 - (J_x J_y + J_y J_x) c_{\Theta} s_{\Theta}.$$
(3.5)

From the equation above and to fulfill the first constraint on the Equation (3.4) it turns out that the

expectation value over our yet to be shown initial state of the operator $(J_xJ_y + J_yJ_x)$ must vanish. Hence it is equivalent to the first assumption of the Equation (3.4) write that

$$\langle \{J_x, J_y\} \rangle = 0 \tag{3.6}$$

where { , } stands for the anticommutator. Apart for been simpler to compute the Equation (3.6) is based also on initial expectation values of the state. We will see that as we said before this is easily guarantied for most important cases.

As we have done recently with the square of J_x now we will do it for J_x^4 . This way one will be able to distinguish which other combination of operators must vanish in order to have Equation (3.4) quarantied. The fourth power of J_x can be written as follows in the Heisenberg picture,

$$J_{x}^{4}(\Theta) = J_{x}^{4}c_{\Theta}^{4} + J_{y}^{4}s_{\Theta}^{4} + (J_{x}^{2}J_{y}^{2} + J_{x}J_{y}J_{x}J_{y} + J_{x}J_{y}^{2}J_{x} + J_{y}J_{x}J_{y}J_{x} + J_{y}J_{x}^{2}J_{y} + J_{y}^{2}J_{x}^{2})c_{\Theta}^{2}s_{\Theta}^{2} - (J_{x}^{3}J_{y} + J_{x}^{2}J_{y}J_{x} + J_{x}J_{y}J_{x}^{2} + J_{y}J_{x}^{3})c_{\Theta}^{3}s_{\Theta} - (J_{x}J_{y}^{3} + J_{y}J_{x}J_{y}^{2} + J_{y}^{2}J_{x}J_{y} + J_{y}^{3}J_{x})c_{\Theta}s_{\Theta}^{3}.$$
(3.7)

And again assuming that its expectation value must be an even function on Θ it turns out that the second line must be equal to zero when the expectation value is considered. So $(J_x^3J_y + J_x^2J_yJ_x + J_xJ_yJ_x^2 + J_yJ_x^3)$ and $(J_xJ_y^3 + J_yJ_xJ_y^2 + J_y^2J_xJ_y + J_y^3J_x)$ must vanish again over any candidate state to be used as prove state. Hence, the second constraint of the Equation (3.4) can be rewritten as follows,

$$\langle \{J_x^2, \{J_x, J_y\}\} \rangle = 0,$$

$$\langle \{J_y^2, \{J_x, J_y\}\} \rangle = 0.$$
 (3.8)

Finally, we can write how the evolution of second and fourth moments of the J_x operator must look like,

$$\langle J_x^2(\Theta) \rangle = \langle J_x^2 \rangle c_{\Theta}^2 + \langle J_y^2 \rangle s_{\Theta}^2 \tag{3.9}$$

$$\langle J_x^4(\Theta) \rangle = \langle J_x^4 \rangle c_{\Theta}^4 + \langle J_y^4 \rangle s_{\Theta}^4 + \langle \{J_x^2, J_y^2\} + \{J_x, J_y\}^2 \rangle c_{\Theta}^2 s_{\Theta}^2.$$
(3.10)

From here we are able to write the evolution of the variance of the second moment when Equation (3.4)

must be obeyed. We obtain

$$(\Delta J_{x}^{2}(\Theta))^{2} = \langle J_{x}^{4}(\Theta) \rangle - \langle J_{x}^{2}(\Theta) \rangle^{2}$$

$$= \langle J_{x}^{4} \rangle c_{\Theta}^{4} + \langle J_{y}^{4} \rangle s_{\Theta}^{4} + \langle \{J_{x}^{2}, J_{y}^{2}\} + \{J_{x}, J_{y}\}^{2} \rangle c_{\Theta}^{2} s_{\Theta}^{2} - (\langle J_{x}^{2} \rangle c_{\Theta}^{2} + \langle J_{y}^{2} \rangle s_{\Theta}^{2})^{2}$$

$$= (\langle J_{x}^{4} \rangle - \langle J_{x}^{2} \rangle^{2}) c_{\Theta}^{4} + (\langle J_{y}^{4} \rangle - \langle J_{y}^{2} \rangle^{2}) s_{\Theta}^{4} + (\langle \{J_{x}^{2}, J_{y}^{2}\} + \{J_{x}, J_{y}\}^{2} \rangle - 2\langle J_{x}^{2} \rangle \langle J_{y}^{2} \rangle) c_{\Theta}^{2} s_{\Theta}^{2}$$

$$= (\Delta J_{x}^{2})^{2} c_{\Theta}^{4} + (\Delta J_{y}^{2})^{2} s_{\Theta}^{4} + (\langle \{J_{x}^{2}, J_{y}^{2}\} + \{J_{x}, J_{y}\}^{2} \rangle - 2\langle J_{x}^{2} \rangle \langle J_{y}^{2} \rangle) c_{\Theta}^{2} s_{\Theta}^{2}.$$
(3.11)

The remaining constituent of the Equation (3.2) on which we will base our result for the precision, is the modulus square of the derivative of the second moment of the J_x operator. Using Equation (3.9) for the expression of the evolution of the second moment, the denominator of Equation (3.2) follows

$$|\partial_{\Theta}\langle J_x^2(\Theta)\rangle|^2 = |-2\langle J_x^2\rangle c_{\Theta} s_{\Theta} + 2\langle J_y^2\rangle c_{\Theta} s_{\Theta}|^2$$

= $4\langle J_y^2 - J_x^2\rangle^2 c_{\Theta}^2 s_{\Theta}^2$. (3.12)

From the equations above directly follows expression for the precision of Θ ,

$$(\Delta\Theta)^{2} = \frac{(\Delta J_{x}^{2})^{2} c_{\Theta}^{4} + (\Delta J_{y}^{2})^{2} s_{\Theta}^{4} + (\langle \{J_{x}^{2}, J_{y}^{2}\} + \{J_{x}, J_{y}\}^{2} \rangle - 2\langle J_{x}^{2} \rangle \langle J_{y}^{2} \rangle) c_{\Theta}^{2} s_{\Theta}^{2}}{4\langle J_{y}^{2} - J_{x}^{2} \rangle^{2} c_{\Theta}^{2} s_{\Theta}^{2}}$$

$$= \frac{(\Delta J_{x}^{2})^{2} t_{\Theta}^{-2} + (\Delta J_{y}^{2})^{2} t_{\Theta}^{2} + \langle \{J_{x}^{2}, J_{y}^{2}\} + \{J_{x}, J_{y}\}^{2} \rangle - 2\langle J_{x}^{2} \rangle \langle J_{y}^{2} \rangle}{4\langle J_{y}^{2} - J_{x}^{2} \rangle^{2}}.$$
(3.13)

To this calculations further computations follows mainly regarding to the following expectation value $\langle \{J_x^2, J_y^2\} + \{J_x, J_y\}^2 \rangle$. This calculus is left for the Appendix A.1. Finally, the expression Equation (3.13) leads to the following,

$$(\Delta\Theta)^{2} = \frac{(\Delta J_{x}^{2})^{2} t_{\Theta}^{-2} + (\Delta J_{y}^{2})^{2} t_{\Theta}^{2} + 4\langle J_{y}^{2} \rangle - 3\langle J_{z}^{2} \rangle - 2\langle J_{x}^{2} \rangle (1 + \langle J_{y}^{2} \rangle) + 6\langle J_{x} J_{y}^{2} J_{x} \rangle}{4\langle J_{y}^{2} - J_{x}^{2} \rangle^{2}}$$
(3.14)

[WOW IN THE PAPER FINALLY IT IS CORRECT!!]

We have verified the correctness of our analytical formula with a direct numeric simulation of the Equation [ERROR PROP. FORM.] and the equation above. For that we, have used the ground-state of $H = J_x^2 + J_y$ for 6 qubits, $|GS\rangle$. This state in principle does not have extra symmetries where further simplifications would take part on the final expression of the formula, so is more adequate for testing. We have computed the evolution of the expectation values of the second and the fourth moments of the operator J_x in the range of half a cycle, *i.e.*, $\Theta \in [0, \pi]$, for thousand of equidistant points. We choose so high density of points on the range in order to compute a more accurate derivative of the



Figure 3.2: (a) Evolution of the precision for 6 qubits based on the simulation of the system and its expectation values. The agreement with the Equation [REF] is shown on the inset plot where the square of the difference between two approaches, the analytical and the simulation, is plotted. The difference is more or less two orders of magnitude below the actual value for the relevant points, which mainly comes because of the error coming from computing the derivative which has to be done approximately between two neighbor points. (b) Verification of the parity with respect to Θ of the expectation values of the second and the fourth moment, so to fulfill the constraint Equation [REF].

expectation value $\partial_{\Theta}\langle J_x^2 \rangle$ appearing on the Equation [EPF], see Figure [REF] (a). Finally, we have also checked that the constraints assumed at the beginning of this section are fulfilled. For that aim, the range of the evolution time has been $\Theta \in [-\pi, \pi]$ and we have computed the expectation values for five hundreds of equidistant points, see Figure [REF] (b). We can conclude saying that our formula exactly reproduces the evolution of the error propagation formula, Equation [REF].

3.2.1 The optimal precision

One can realize that the whole dependence on the phase shift is in the first two terms of the numerator. This way one can minimize the sum on the first two terms in order to find where the precision is best. So it follows that,

$$\tan^2(\Theta_{\text{opt}}) = \sqrt{\frac{(\Delta J_x^2)^2}{(\Delta J_y^2)^2}}$$
(3.15)

which inserted on the Equation (3.14) gives us the optimal precision when the second moment $\langle J_x^2 \rangle$ is measured based on the initial expectation values of the input state. The optimal precision is written in the following way,

$$(\Delta\Theta)_{\text{opt}}^{2} = \frac{\sqrt{(\Delta J_{x}^{2})^{2}(\Delta J_{y}^{2})^{2} + 4\langle J_{y}^{2} \rangle - 3\langle J_{z}^{2} \rangle - 2\langle J_{x}^{2} \rangle (1 + \langle J_{y}^{2} \rangle) + 6\langle J_{x}J_{y}^{2}J_{x} \rangle}}{4\langle J_{y}^{2} - J_{x}^{2} \rangle^{2}}.$$
 (3.16)

.

We conclude with this section checking our bound for the pure unpolarised Dicke state aligned with Ox, $|D_N\rangle_x$, whose precision bound is well known, Equation ([XXX]). With this aim we compute all the expectation values needed for the Equation (3.16) which almost all of them are trivial, $\langle J_x J_y^2 J_x \rangle = \langle J_x^4 \rangle = \langle J_x^2 \rangle = 0$. The rest are obtained in the following way,

$$\langle J_y^2 \rangle = \langle J_z^2 \rangle = \frac{N(N+2)}{8},$$
 (3.17)

$$\langle J_y^4 \rangle = \frac{N+2}{8} \left(\frac{3N(N+2)}{16} - \frac{1}{2} \right).$$
 (3.18)

The Equation (3.17) follows directly from the fact that the state is invariant under rotations on the Ox axis, so they are its expectation values, because the sum of all the second moments must give the value of the total angular momentum, in this case the maximum which is $\langle J^2 \rangle = \frac{N(N+2)}{4}$, and because $\langle J_x^2 \rangle = 0$. The proof of the Equation (3.18) needs more algebra and has been left for the Appendix ??.

From the equations above, one lead to the following expression for the precision of the phase shift for a pure unpolarised Dicke state,

$$(\Delta\Theta)^2 = \frac{2}{N(N+2)},\tag{3.19}$$

which coincides exactly with the inverse quantum Fisher information for such state.

3.3 Testing the formula against some known states

In this section we will compare our criteria based on few expectation values against the corresponding quantum Fisher information obtained for some known input states. Those input states will be first the family of states defined as the ground states of the following Hamiltonian, called the spin-squeezing Hamiltonian,

$$H_{\lambda} = J_{\nu}^2 - \lambda J_{\mu}. \tag{3.20}$$

For λ equal to zero we have the unpolarized Dicke state, Equation (3.1), and for λ large we recover the coherent totally polarized state pointing onto the Oy direction. In the meantime the state is also a spin-squeezed state, therefore the name of the Hamiltonian. The ground state is defined such that

$$\lambda_{\min} = \operatorname{tr}(H_{\lambda} \rho_{\lambda}), \tag{3.21}$$

where λ_{\min} is the lowest eigenvalue of H_{λ} . The state ρ_{λ} is invariant under permutations of the particles and it is also pure. Moreover the state ρ_{λ} is symmetric.

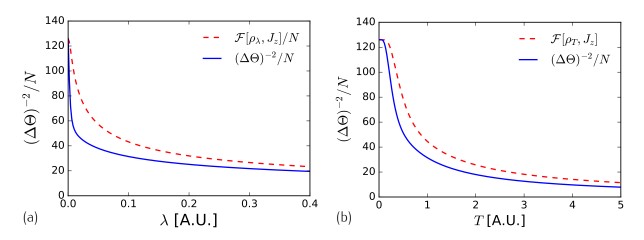


Figure 3.3: Comparison between our formula for the precision and the QFI for different states. (a) Comparison for ground states of H_{λ} . (b) Comparison with gaussian mixture of Dicke states.

The second family of input states we are going use are the Gaussian mixture of Dicke states around the unpolarized Dicke state, which have the following form as function of β ,

$$\rho_{(T)} \propto \sum_{m=-N/2}^{N/2} e^{-\frac{m^2}{T}} |\mathcal{D}_N^m\rangle_x \langle \mathcal{D}_N^m|_x. \tag{3.22}$$

After showing how the optimal precision formula behaves compared with the quantum Fisher information for those two families of states, we also have to proof that they indeed fulfill the constraints on Equation (3.4).

For the spin-squeezed state , ρ_{λ} , we have that it is invariant under

On the other hand for the thermal state, ρ_T , we have that it is invariant under rotations around the Ox axis

$$\rho_T = e^{-i\alpha J_x} \rho_T e^{i\alpha J_x}, \tag{3.23}$$

for arbitrary $\forall \alpha$.

$$\operatorname{tr}(e^{i\Theta J_z} J_y^m e^{-i\Theta J_z} \rho_T) = \operatorname{tr}(e^{i\Theta J_z} J_y^m e^{-i\Theta J_z} e^{-i\alpha J_x} \rho_T e^{i\alpha J_x})$$
(3.24)

If we choose $\alpha=\pi$ then we rotate all the orthogonal angular momentum operators such that $J_y\to -J_x$ and $J_z\to -J_z$. Therefore, $e^{i\pi J_x}e^{i\Theta J_z}J_y^me^{-i\Theta J_z}e^{-i\pi J_x}=e^{-i\Theta J_z}(-J_y)^me^{i\Theta J_z}$. Hence, for even m and particularly for m=2,4 we have that,

$$\operatorname{tr}(e^{i\Theta J_z} J_u^m e^{-i\Theta J_z} \rho_T) = \operatorname{tr}(e^{-i\Theta J_z} J_u^m e^{i\Theta J_z} \rho_T), \tag{3.25}$$

so the Equation (XXX) is granted.

3.4 Using our method with real experimental data

On reference [XXX], it is produced on the laboratory a state with the proper characteristics of an unpolarised Dicke state, small variance on one of the directions and very large one on the perpendicular directions to this. It is also invariant under rotations around Ox-axis. Effectively, the state has the following form,

$$\rho = \frac{1}{2\pi} \int e^{-i\alpha J_x} \rho_0 e^{i\alpha J_x},\tag{3.26}$$

where ρ_0 is what we would obtain if we would have access to the phase reference. Hence we have that at $\Theta=0$ all the statistical moments $\langle J_z^m \rangle = \langle J_y^m \rangle$ are equal.

Simplification of our precision formula on Equation (XXX). First, we simplify the expectation value $\langle J_x J_z^2 J_x \rangle$ in the following way,

$$\langle J_x J_y^2 J_x \rangle = \frac{\langle J_x (J_y^2 + J_z^2) J_x \rangle}{2} = \frac{\langle J_x (J_x^2 + J_y^2 + J_z^2) J_x \rangle - \langle J_x^4 \rangle}{2}$$

$$\leq \frac{N(N+2)}{8} \langle J_x^2 \rangle - \frac{\langle J_x^4 \rangle}{2}.$$
(3.27)

Notice that obtaining $\langle J_x J_z^2 J_x \rangle$ is hard experimentally. This simplification can only make our estimation of the precision worse while for symmetric states the equality holds. The bound from below for the precision can be written as,

$$(\Delta\Theta)_{\text{opt}}^{2} \le \frac{\sqrt{(\Delta J_{x}^{2})^{2}(\Delta J_{y}^{2})^{2}} + \langle J_{y}^{2} \rangle + \frac{3N(N+2)-8}{4} \langle J_{x}^{2} \rangle - 2\langle J_{x}^{2} \rangle \langle J_{y}^{2} \rangle - 3\langle J_{x}^{4} \rangle}{4\langle J_{y}^{2} - J_{x}^{2} \rangle^{2}}, \tag{3.28}$$

where some terms were reordered and further simplified.

It is worth to study this case and apply our methods such that we can extract conclusions about the metrological usefulness of the state. The system in consideration is a N=7900 particles system. The measured data for such system is

$$\langle J_x^2 \rangle = 112 \pm 31,$$
 $\langle J_y^2 \rangle = 6 \times 10^6 \pm 0.6 \times 10^6,$ $\langle J_x^4 \rangle = 40 \times 10^3 \pm 22 \times 10^3,$ $\langle J_y^4 \rangle = 6.2 \times 10^{13} \pm 0.8 \times 10^{13}.$ (3.29)



Figure 3.4: The (solid) line shows how the precision of Θ varies through the evolution. Notice that for the initial moment the precision is zero and that it reaches a maximum at $\Theta \approx 0.0057$ highlighted with the vertical (dashed) line. The gray area represent the region where the precision is below the shot-noise limit.

Hence we obtain the maximum precision,

$$\frac{(\Delta\Theta)_{\text{opt}}^{-2}}{N} \ge 3.7 \pm 1.5 \tag{3.30}$$

with boot straping methods. The direct substitution would yield to a 3.3 gain over the shot-noise limit.

Next we plot the value for the precision substituting directly the measured data into Equation (XXX), see Figure 3.4.

Further simplification of our method can be achieved for states of the kind of the one studied on this section. Based on $\langle AB \rangle \leq \lambda_{\max}(A) \langle B \rangle$ for two commuting positive-semidefinite observables,

$$\langle J_y^4 \rangle \le \frac{N^2}{4} \langle J_x^2 \rangle. \tag{3.31}$$

We can also approximate $\langle J_x^4 \rangle$ with $\langle J_x^2 \rangle$ in the sense that it is small and that mainly its value comes from technical noise,

$$\langle J_x^4 \rangle \approx \beta \langle J_x^2 \rangle^2.$$
 (3.32)

This approximation even if it is not a strict bound on the precision can be very useful in order to characterize the metrological usefulness of our input state based only on second statistical moments of only two angular momentum operators, namely $\langle J_z^2 \rangle$ and $\langle J_x^2 \rangle$. Those two expectation values are



Figure 3.5: Comparison between our formula for the precision and the QFI for different states. a) Comparison for ground states of H_{λ} . (b) Comparison with gaussian mixture of Dicke states.

related with the width of our state and also with how thin we can do it in one of the directions, for metrological purposes, perpendicular to the magnetic field. So in this case we use $\beta = 3$ assuming that the distribution function has Gaussian shape.

From these considerations we are able to write a second bound with fewer expectation values for the optimal precision such that,

$$(\Delta\Theta)_{\text{opt}}^{2} \leq \frac{\langle J_{y}^{2} \rangle + \frac{3N(N+2)-8}{4} \langle J_{x}^{2} \rangle + \left(\sqrt{\frac{N^{2}}{2\langle J_{y}^{2} \rangle} - 2} - 2\right) \langle J_{y}^{2} \rangle \langle J_{x}^{2} \rangle - 9\langle J_{x}^{2} \rangle^{2}}{4\langle J_{y}^{2} - J_{x}^{2} \rangle^{2}}.$$
(3.33)

We have used this formula to compute the bound on the optimal precision with the measured data shown on Equation (XXX), $(\Delta\Theta)_{opt}^{-2} \geq 2.9N$, see Figure 3.5.

Witnessing metrologically useful entanglement

TYPICALLY, one has not access to the density matrix of the system been used for metrology or for other quantum process. Moreover, for systems on which the particle number is very large, this is the case when one wants to do metrology with quantum states, the details of the density matrix are forbidden by practical reasons. Since the quantum Fisher information is based on the complete knowledge of the density matrix, shortcuts to avoid the complete tomography must be developed as we have had shown one practical case on the previous chapter. In this chapter, we obtain a general procedure to get an optimal bound for the quantum Fisher information based on as many expectation values of the initial state as one is ready to measure. Two main features are worth to mention again. First, in general this method gives us an optimal tight bound. Last but not least, the bound is based on the expectation values of the initial state only, so it is not necessary to perform an evolution of the state to estimate how well will it behave. This is in contrast to other approaches one can find in the literature, an it is a time saving concept.

The figure of merit of bounds from below for the quantum Fisher information based on expectation

values of the initial state is the following,

$$\mathcal{F}_{Q}[\rho, J_{z}] \ge \frac{\langle J_{x} \rangle^{2}}{(\Delta J_{y})^{2}}.$$
(4.1)

where the state is polarized along the Ox-axis and the variance of the J_y operator is smaller than the standard. On the previous chapter we also have shown one of these bound specifically designed for unpolarized Dicke states.

Homogeneous magnetometry, B = Bk where B is constant in time.

Section [REF], see magnetometry, generator J_z .

Quantum Fisher information $\mathcal{F}_{\mathbb{Q}}[\rho,J_z]$.

4.1 Minimum of a convex function of the state given some expectation values

Convex function of state, $q(\rho)$.

Which is the lowest possible value of $g(\rho)$ for a state where some expectation values of $\{W_i\}_{i=1}^M$ operators, $w_i = \text{tr}(\rho W_i)$ are known?

Such bound is defined in a way to fulfill the following inequality,

$$g(\rho) \ge \mathcal{B}_g(w_1, w_2, ...) := \left\{ \min_{\rho} g(\rho) \mid \{ w_i = \text{tr}(\rho W_i) \}_{i=1}^M \right\}.$$
 (4.2)

When computing this bound we will require to this to be as tight as possible.

For this purpose we simplify our discourse to one expectation value and therefore one observable, $w = \text{tr}(\rho W)$. Later we will extend our results to the multi-parameter case.

Notice that the tightest bound from below on of such a function $g(\rho)$ is convex too on the expectation values, *i.e.*, $\mathcal{B}_{q}(w)$ is convex if $g(\rho)$ is convex. This is shown in the following way,

Therefore, theory tells us that we can use the Legendre transform to get rid of the state preserving all the necessary information to reconstruct the \mathcal{B}_g bound from the expectation values, this can be found on the work of Prof. Gühne and Prof. Werner Reference [?] and for a brief description of the properties of the Legendre transform see Appendix ??. In this case the Legendre transform can be written in the following way,

$$\hat{g}(rW) = \sup_{\rho} \{ r \langle W \rangle_{\rho} - g(\rho) \}. \tag{4.3}$$

Now we have the Legendre transform, we recover the bound from below of $q(\rho)$ as

$$\mathcal{B}_{g}(w) = \sup_{r} \left\{ rw - \sup_{\rho} \left\{ r\langle W \rangle_{\rho} - g(\rho) \right\} \right\}. \tag{4.4}$$

To clarify, notice that the ρ appearing on the inner maximization is not related with the one appearing on $g(\rho)$ that is bounded by \mathcal{B}_g but a variable state for the maximization.

The Equation (4.4) has been applied on entanglement measures, when $g(\rho)$ is defined using the convex roof construction, *i.e.*, the function $g(\rho)$ is extended from the well defined region, namely the region of pure states, to the mixed states using the convex roof construction. For illustrative purposes the convex roof of a function defined for pure states is the following,

$$g(\rho) = \inf_{\{\rho_k, |\phi_k\rangle\}} \left\{ \sum_k p_k g(|\phi_k\rangle) \right\}$$
 (4.5)

for any decomposition $\rho = \sum_k p_k |\phi_k\rangle\langle\phi_k|$ In this case the Equation (4.4) can be further simplified. Let us focus by now on the inner maximization,

$$\hat{g}(rW) = \sup_{\rho} \{ r\langle W \rangle_{\rho} - g(\rho) \}
= \sup_{\rho} \{ r\langle W \rangle_{\rho} - \inf_{\{p_{k}, |\phi_{k}\rangle\}} \{ \sum_{k} p_{k} g(|\phi_{k}\rangle) \} \}
= \sup_{\{p_{k}, |\phi_{k}\rangle\}} \{ \sum_{k} p_{k} r\langle W \rangle_{|\phi_{k}\rangle} - \inf_{\{p_{k}, |\phi_{k}\rangle\}} \{ \sum_{k} p_{k} g(|\phi_{k}\rangle) \} \}
= \sup_{\{p_{k}, |\phi_{k}\rangle\}} \{ \sum_{k} p_{k} \{ r\langle W \rangle_{|\phi_{k}\rangle} - g(|\phi_{k}\rangle) \} \}
= \sup_{|\psi\rangle} \{ r\langle W \rangle_{|\psi\rangle} - g(|\psi\rangle) \}.$$
(4.6)

So now we have restricted the maximization to only pure states as it is shown on Reference [?].

One the other hand, one can find different definitions of the QFI in the literature. Among those definitions, there is one that defines the QFI as the convex roof of four times the variance of the generator of the unitary phase shift that suffers the state [?]. This can be written in the following way,

$$\mathcal{F}_{Q}[\rho, G] = \inf_{\{\rho_k, |\phi_k\rangle\}} 4 \sum_{k} p_k (\Delta G)_{|\phi_k\rangle}^2$$

$$\tag{4.7}$$

Even further simplification can be achieved for quantum Fisher information,

$$\hat{\mathcal{F}}_{Q}(rW) = \sup_{|\psi\rangle} \left\{ r \langle W \rangle_{|\psi\rangle} - 4(\Delta J_{z})_{|\psi\rangle}^{2} \right\}
= \sup_{|\psi\rangle} \left\{ r \langle W \rangle_{|\psi\rangle} - 4 \langle J_{z}^{2} \rangle_{|\psi\rangle} + 4 \langle J_{z} \rangle_{|\psi\rangle}^{2} \right\}
= \sup_{|\psi\rangle} \left\{ \langle rW - 4J_{z}^{2} \rangle_{|\psi\rangle} + \langle 2J_{z} \rangle_{|\psi\rangle}^{2} \right\},$$
(4.8)

which is clearly a second order maximization over pure states.

With that at hand we are able to prove the following observation: the Legendre transform of the highest lower bound of the quantum Fisher information when the expectation value of some operator W is known, can be written as the maximization over some parameter μ of the highest eigenvalue $\lambda_{\max}(rW-4(J_z-\mu)^2)$. This can be expressed as follows,

$$\hat{\mathcal{F}}_{Q}(rW) = \sup_{\mu} \{ \lambda_{\max}(rW - 4(J_z - \mu)^2) \}, \tag{4.9}$$

where at the maximum the derivative with respect to μ must be zero for this quadratic function on the state. Hence, the value of μ will coincide with the expectation value on the optimal state $\langle J_z \rangle_{\psi}$. Therefore, the finding of the optimal parameter μ is delimited to the range defined by $[\langle J_z \rangle_{\min}, \langle J_z \rangle_{\max}]$. For qubits this means that we have to perform a search for $\mu_{\rm opt}$ in between $-N/2 \le \mu \le N/2$.

Another way of getting the result of the maximization for the second order equation is parameterizing the eigenstate with the maximum eigenvalue of the operator $rW - 4(J_z - \mu)^2$, so we get $|\phi_{\text{max}}(\mu)\rangle$, and finally computing the expectation values of Equation (4.8) and maximizing it.

With this a tight bound from below for the quantum Fisher information can be found when the expectation value of an observable, say W, is known as follows,

$$\mathcal{B}_{\mathcal{F}}(w) = \sup_{r} \left\{ rw - \sup_{|\psi\rangle} \left\{ \langle rW - 4J_z^2 \rangle + \langle 2J_z \rangle^2 \right\} \right\}. \tag{4.10}$$

One may notice that the inner maximization is a second order maximization.

This can be extended for several observables as follows

$$\mathcal{B}_{\mathcal{F}}(w_1, w_2, ...) = \sup_{r} \left\{ r w - \sup_{|\psi\rangle} \left\{ \langle r W - 4J_z^2 \rangle + \langle 2J_z \rangle^2 \right\} \right\}, \tag{4.11}$$

where $\mathbf{r} = (r_1, \dots, r_m)$ is the vector form and the same applies to \mathbf{w} and the operators set \mathbf{W} . The contraction of two of those vectors must be seen as a scalar product, i.e., $\mathbf{r}\mathbf{W} = \sum r_i W_i$ where W_i is

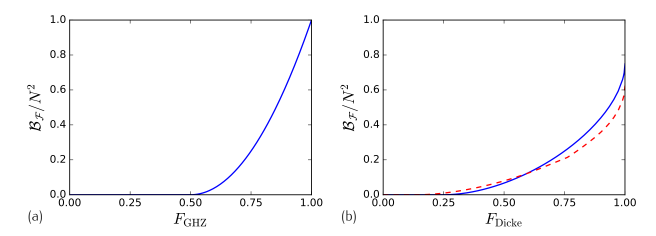


Figure 4.1: (a) Analytical solution of the bound $\mathcal{B}_{\mathcal{F}}$ for different values of the fidelity with respect to the GHZ state. (b) Numerical results for the minimum quantum Fisher information as a function of the fidelity with respect of unpolarise Dicke states perpendicular to the magnetic field, $|D_N^0\rangle$. (blue-line) For systems with 4 particles and (red-dashed) for system with 8 particles. One may notice that when the fidelity is at its maximum the bound approaches to 0.5 as it is the quantum Fisher information for large particle number.

the corresponding i^{th} operator of W.

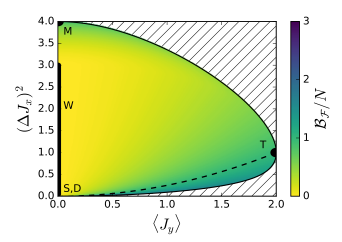


Figure 4.2: We show as a function of the expectation value, $\langle J_y \rangle$, and the variance in the perpendicular direction, $(\Delta J_x)^2$, the minimum sensitivity for a 4-qubit system. (hatched) The physically forbidden region is indicated. (M,T,S,D) Those points indicate where the mixed state (), the totally polarized state (), the single state and the unpolarized Dicke state would be located. (W) on this line sit any of the states which is a mixture between the completely mixed state of the symmetric subspace and the singlet state among others, for instance the completely mixed state of the whole Hilbert space. (dashed) It indicates the shot-noise threshold where below it non-classical sensitivities can be achieved.

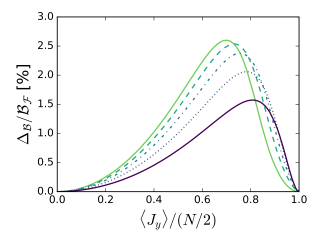


Figure 4.3: Difference between the bound of Pezze-Smerzi and the optimal bound for the quantum Fisher information normalized by the value of the optimal bound itself for the bosonic ground states of $H = J_x^2 - \lambda J_y$ for $\forall \lambda \in [0, \infty)$. From dark to lighter colors (line, point, dash-point, dashed, pointed, line), results for different particle numbers, N = 4, 6, 10, 20, 1000 respectively. Heuristically speaking, one can say that for large particle number the difference is biggest when the polarisation is around two thirds of the maximal polarisation and that this difference is about 2.6%.

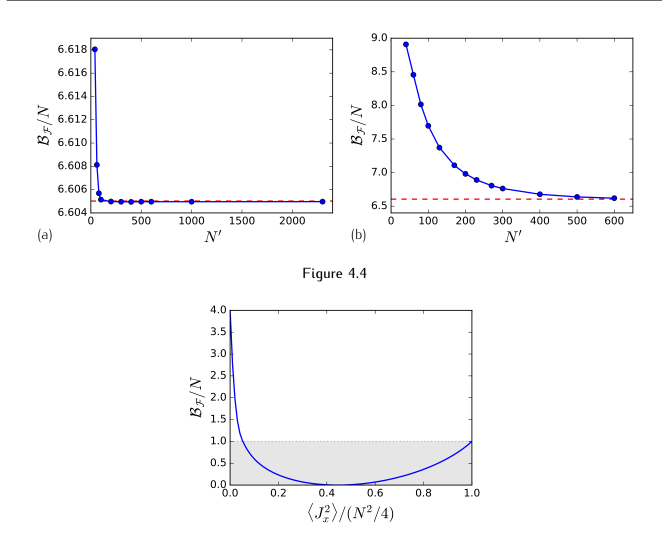


Figure 4.5: The numerics shows us a tiny region the symmetric system surpassing the shot-noise threshold.

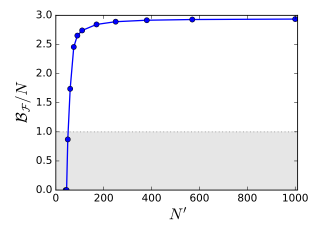


Figure 4.6: Secuence of the evolution of an unpolarized Dicke state of 16 qubits for $\Theta = \{i\pi/6\}_{i=0}^4$. Bloch spheres representing the Hirusi distribution of the state, and below PDF of the J_x POVM for each step of the secuence

4.1 Minimum of a convex function of the state given some expectation values

C

Accuracy bound for gradient field estimation with atomic ensembles

In this chapter, one of the most fundamental two-parameter estimation tasks in magnetometry is considered, namely gradient magnetometry. We will add the gradient of the magnetic field as the second parameter beside the constant (homogeneous) part of the field. While most works in magnetometry with a single ensemble focus only on the determination of the strength and direction of magnetic field, certain measurement schemes for the gradient have already been proposed and tested experimentally [XXX]. Some schemes use an imaging of the ensemble with a high spatial resolution, however, they do not count as single-ensemble methods in the sense we use this expression in our paper, since in this case not only collective observables are measured [1–3]. There is a method based on collective measurements of the spin length of a fully polarised ensemble [4]. Finally, there is a scheme where they use as a prove state a many-body singlet states, which is described in Reference [5]. This section provides precision bounds for that scheme, while expanding the quantum states considered to other interesting quantum states, including those that are not invariant under homogeneous fields.

The atoms will be distributed along the OX axis, so y=z=0, and in principle they will be able to feel differences on the magnetic field at different points of the axis. The magnetic field at the

atoms will be given by a linear function on the position x

$$B(x, 0, 0) = B_0 + xB_1 + O(x^2), (5.1)$$

where we will neglect the terms of order two or higher. We will consider the magnetic field pointing along the OZ direction direction only, $\mathbf{B}_0 = B_0 \mathbf{k}$ and $\mathbf{B}_1 = B_1 \mathbf{k}$, where \mathbf{k} is the unitary vector pointing on the OZ direction. For this configuration, due to the Maxwell equations, with no currents or changing electric fields, we have

$$\nabla \cdot \mathbf{B} = 0,$$

$$\nabla \times \mathbf{B} = 0.$$
(5.2)

This implies $\sum_{l=x,y,z} \partial_l B_l = 0$ and $\partial_m B_l - \partial_l B_m = 0$ for $\forall l \neq m$, where $\partial_m \equiv \partial/\partial_m$ stands for the partial derivative over the variable l. Thus, the spatial derivatives of the field components are not independent of each other. However, in the case of a linear arranged particle ensemble only the derivative along the OX axis has an influence on the quantum dynamics of the atoms.

We will determine the precision bounds for estimation of the magnetic field gradient B_1 based on the quantum Fisher information (QFI) [6–11]. We will show that for states insensitive to the homogeneous magnetic field, one can reduce the problem to a one-parameter estimation scenario. Such states can arise in a single-ensemble scheme, however, it will be shown that for single-ensembles insensitive to the homogeneous field the Heisenberg limit cannot be reached. Nevertheless, single-ensemble measurements have certain advantages since the spatial resolution can be higher and the experimental requirements are smaller since only a single ensemble must be prepared.

On the other hand, for states sensitive to the homogeneous field, the classical limit can be overcome only if the particle positions are highly correlated with each other. Our calculations are generally valid for any measurement, thus they are relevant to many recent experiments [1–4, 12–15]. We note that in the case of the singlet, our precision bounds are saturated by the metrological scheme presented in Reference [5].

We can also connect our results to entanglement theory [16–18]. We find that the shot-noise scaling cannot be surpassed with separable states, while the Heisenberg scaling can be reached with entangled states. However, the shot-noise scaling can be surpassed only if the particle positions are correlated, which is the case if the particles attract each other. Moreover, the case of two atomic ensembles, if the spin states of the ensembles are uncorrelated, can also be mapped to our model. This evidently shows how shot-noise scaling can be surpassed with the two-ensemble scenario.

Next, we will present characteristics of our setup. For simplicity, as well as following recent experiments (e.g., Ref. [3]), we will consider an ensemble of spin-j particles placed in a one-dimensional setup, x being the spatial coordinate. Furthermore, we assume that we have that we have particles that behave classically with respect to their spatial state. That is, they cannot be in a superposition of being in two different positions. On the other hand, they have internal degrees of freedom, their spin, that is quantum. This is a very good description to many of the cold gas experiments.

Based on these considerations, we assume that the state is factorizable into a spatial and a spin part as

$$\rho = \rho_{\mathsf{x}} \otimes \rho_{\mathsf{s}},\tag{5.3}$$

and that the spatial part can be characterized as an incoherent mixture of point-like particles that can be written as

$$\rho_{x} = \int d^{N}x \, P_{N}(x)|x\rangle\langle x|. \tag{5.4}$$

Note that the spatial part is diagonal on the position eigenbasis, where the entry x_n of $\mathbf{x} = (x_1, \dots, x_N)$ is the coordinate of the n^{th} particle, and $P_N(\mathbf{x})$ is the spatial probability distribution function of the atoms. This model can be seen as that the density matrix of the spatial part of the state is diagonal on the position eigenbasis and it simplifies much the problem. During the evolution of the state, correlations might arise between the two inner and spatial parts and the product form (5.3) might be not valid to describe the evolution of the system.

We note that our method could be easily extended to the case of Bose-Einstein condensates, not considered in this paper. In that case, the spatial state of the particles would be a pure state, and we would have $\rho_x = (|\Psi\rangle\langle\Psi|)^{\otimes N}$, where $|\Psi\rangle$ is a spatial single-particle state.

Although in our case the parameter to be estimated is B_1 , the time-evolution of the state is usually also affected by the second unknown parameter, the homogeneous field B_0 , which means that we generally have to consider a two-parameter estimation problem. The angular momentum of an individual atom is coupled to the magnetic field, yielding the following interaction term

$$h^{(n)} = \gamma B_z^{(n)} \otimes j_z^{(n)}, \tag{5.5}$$

where the operator $B_z^{(n)} = B_0 + B_1 x^{(n)}$ acts on the spatial part of the Hilbert space. The sum of all one particle interactions provide the total Hamiltonian

$$H = \gamma \sum_{n=1}^{N} B_{z}^{(n)} \otimes j_{z}^{(n)}, \tag{5.6}$$

which will generate the time evolution of the atomic ensemble.

We will estimate the precision of B_1 based on a measurement on the state after it passed through the dynamics expressed by the unitary evolution operator $U = \exp(-i\frac{H}{\hbar}t)$, where t is the time spent by the system under the influence of the magnetic field. Those type of unitary phase shift are the most easy ones to study with the quantum Fisher information, and this is the reason while we choose them to make the jump to the multi-parametric estimation problem. The unitary operator can be rewritten in the following way

$$U = e^{-i(b_0 H_0 + b_1 H_1)}, (5.7)$$

where the $b_i = \gamma B_i t/\hbar$. Here, the generator describing the effect of the homogeneous field is given as

$$H_0 = \sum_{n=1}^{N} j_z^{(n)} = J_z, (5.8)$$

while the generator describing the effect of the gradient is

$$H_1 = \sum_{n=1}^{N} x^{(n)} j_z^{(n)}. (5.9)$$

As in Eq. (5.9), we will usually omit \otimes for simplicity, and will use it only if it is necessary to make our explanation clearer.

Note that the operators H_0 and H_1 commute with each other. These two commuting dynamics are the two simplest in an atomic ensemble as they are based on collective operators not requiring an individual access to the particles.

Note also that it is not necessarily true that the operators we have to measure to estimate b_0 and b_1 commute with each other, which problem we will consider case by case. The reason for that is that both operators to be measured act on the same single atomic ensemble. On the other hand, in schemes in which the gradient is calculated based on measurements in two separate atomic ensembles or different atoms in a chain, the measuring operators can always commute with each other [12, 13, 19].

The paper is organized as follows. In Sec. 5.1, the basic concepts used in the paper are presented. In Sec. III, we restrict our calculations to single PI atomic ensembles and we develop some particular cases, such as the singlet spin state or the totally polarised state. In Sec. IV, we discuss our results.

5.1 Cramér-Rao precision bounds

In this section, we show how the Cramér-Rao bound and the QFI help us to obtain the precision bound that is valid for any measurement scenario. We will discuss gradient magnetometry using quantum states that are insensitive to homogeneous fields, which is a single-parameter estimation task. Then, we discuss the case of quantum states sensitive to homogeneous fields. We show that the precision bound obtained does not change under spatial translation. Since this is a two-parameter estimation task, we will introduce the two-parameter Cramér-Rao bound and the corresponding two-parameter QFI matrix, and we adapt those expressions to our problem.

For clarity we present our main tools inf subsequent paragraphs before going onto details for states that are not sensitive to the homogeneous field and states that they are. The expression for quantum Fisher information that we use along this paper will be suitable for the transition to the multi-parameter problem, *i.e.*, it is equivalent for the single parameter estimation problem still giving the chance to switch to the multi-parameter case. For two arbitrary operators A and B, it is written as follows

$$\mathcal{F}_{Q}[\rho, A, B] := 2 \sum_{k k'} \frac{(\lambda_{k} - \lambda_{k'})^{2}}{\lambda_{k} + \lambda_{k'}} A_{k, k'} B_{k', k}, \tag{5.10}$$

where the subscript for A and B stand for the matrix element on the eigenbasis of the initial state $\rho = \sum_k \lambda_k |k\rangle\langle k|$. If the two operators are the same, the usual form of the QFI on the literature can be recovered with only two arguments [6–11],

$$\mathcal{F}_{Q}[\rho, A, A] := \mathcal{F}_{Q}[\rho, A] = 2 \sum_{k, k'} \frac{(\lambda_{k} - \lambda_{k'})^{2}}{\lambda_{k} + \lambda_{k'}} |A_{k, k'}|^{2}.$$
 (5.11)

We mention that in our case the operators A and B will commute on all cases making the some computations easier. We also make use of the fact that the QFI as written as in Eq. (5.10) is linear on the second and last arguments,

$$\mathcal{F}_{Q}[rho, A, \sum_{i} b_{i}] = \sum_{i} \mathcal{F}_{Q}[\rho, A, b_{i}]$$
(5.12)

together with $\mathcal{F}_{Q}[\rho, A, B] = \mathcal{F}_{Q}[\rho, B, A]$ so the statement holds.

Some more useful properties of the Eq. (5.10). It can be rewritten as follows,

$$\mathcal{F}_{Q}[\rho, A, B] = 4\langle AB \rangle - 8 \sum_{k,k'} \frac{\lambda_k \lambda_{k'}}{\lambda_k + \lambda_{k'}} A_{k,k'} B_{k',k}. \tag{5.13}$$

This form leads to simple arguments on our derivations on following sections. For pure states it simplifies to,

$$\mathcal{F}_{Q}[|\psi\rangle, A, B] = 4\left(\langle AB\rangle_{\psi} - \langle A\rangle_{\psi}\langle B\rangle_{\psi}\right). \tag{5.14}$$

Notice that we recover $\mathcal{F}_Q[\rho,A]=4(\Delta A)^2$ as can be found in the literature for single parameter estimation with pure states [6, 20]. Another important feature of the qFI in Eq. (5.10) is that it is convex on the space of the states. This property takes the following form,

$$\mathcal{F}_{0}[q\rho_{1}+(1-p)\rho_{2}] \leq p\mathcal{F}_{0}[\rho_{1}]+(1-p)\mathcal{F}_{0}[\rho_{2}], \tag{5.15}$$

where we omit in writing the arguments for the operators of the qFI for simplicity.

In the following subsections we show the general form for the precision bounds for states insensitive to the homogeneous fields and for states sensitive to them. We also show that both bounds are invariant under spatial translation of the system which makes the computing for particular cases much easier.

5.1.1 Precision bound for states insensitive to homogeneous fields: Single-parameter dependence

Let us consider quantum states that are insensitive to the homogeneous field. For these states, $[\rho, H_0] = 0$ and hence the evolved state is a function of a single unknown parameter, b_1 .

For the unitary dynamics we consider, the QFI for single-parameter dependence can be expressed in terms of the eigendecomposition of the density matrix, $\rho = \sum_k \lambda_k |k\rangle\langle k|$, in the following way [6–11],

$$\mathcal{F}_{Q}[\rho, H_{1}] = 2 \sum_{k,k'} \frac{(\lambda_{k} - \lambda_{k'})^{2}}{\lambda_{k} + \lambda_{k'}} |(H_{1})_{k,k'}|^{2}.$$
 (5.16)

And due to the Cramér-Rao formula gives us an upper bound for the precision

$$(\Delta b_1)^{-2}|_{\text{max}} = \mathcal{F}_0[\rho, H_1]. \tag{5.17}$$

Note that it is always possible to find a measurement that saturates the above precision bound. Here, $\mathcal{F}_{Q}[\rho, H_{1}]$ denotes the QFI that depends, in the case of unitary transformation of the form Eq. (5.7), on the state ρ and on the generator of the evolution, H_{1} .

For the particular case on which the states has the form of Eqs. (??) and (??) the Equation (5.16) can be simplified in the following way, see that we have to compute the matrix elements of H_1 but it

is already diagonal on the spatial subspace. Therefore the following holds for the matrix elements,

$$(H_{1})_{x,k,y,k'} = \langle x, k | H_{1} | y, k' \rangle$$

$$= \langle x, k | \sum_{n=1}^{N} x^{(n)} j^{(n)} | y, k' \rangle$$

$$= \delta(x - y) \sum_{n=1}^{N} x_{n} \langle k | j^{(n)} | k' \rangle.$$
(5.18)

We will use the Dirac delta function to further simplify the Equation (5.16), for details of the simplification see the Appendix ??.

For the last part of the proof on which we have to show how translated systems have unchanged the sensitivity over the gradient estimation, we use the Heisenberg picture on which the operators must be reversely transformed instead of the states. Thus, we compute the transformation of H_1 in the following way,

$$U_{d}: H_{1} \to H_{1}(d) = U_{d}^{\dagger} H_{1} U_{d}$$

$$= \sum_{n=1}^{N} U_{d}^{\dagger} x^{(n)} U_{d} \otimes j_{z}^{(n)}$$

$$= \sum_{n=1}^{N} (x^{(n)} - d) j_{z}^{(n)}$$

$$= H_{1} - dH_{0}.$$
(5.19)

Hence, the new unitary evolution operator instead of Equation (5.7) will be

$$U = e^{-i(b_0 H_0 + b_1 H_1(d))} = e^{-i((b_0 - b_1 d)H_0 + b_1 H_1)},$$
(5.20)

which for states insensitive to the homogeneous field, $[\rho, H_0] = 0$, makes the transformation irrelevant.

With this, for states of the form Equations (5.3,??) the following bound for the precision of the estimation of the gradient parameter b_1 holds,

$$(\Delta b_1)^{-2}|_{\max} = \sum_{n,m}^{N} \int d^N \mathbf{x} P_N(\mathbf{x}) x_n x_m \mathcal{F}_{\mathbb{Q}}[\rho_s, j_z^{(n)}, j_z^{(m)}], \tag{5.21}$$

where the integral results onto a two-point correlation function of the spatial state.

Moreover, such bound is translationally invariant, i.e., it remains the same after an arbitrary

phase-shift of the form of $U_d = \exp(-i\boldsymbol{d} \cdot \boldsymbol{p}_x)$, where \cdot denotes scalar product, \boldsymbol{d} is a n-length vector of all elements equal to the phase-shift d and \boldsymbol{p}_x is the vector that collects all the single particle linear momentum operators, $\boldsymbol{p}_x^{(n)}$.

5.1.2 Precision bound for states sensitive to homogeneous fields: Two-parameter dependence

In order to obtain the precision bound for states sensitive to the homogeneous field, one has to consider the effect on the state of a second unknown parameter, in this case b_0 . The homogeneous field will rotate all the spins in the same way, while the field gradient rotates the spins differently depending on the position of the particles. Now, instead to the Cramér-Rao bound (5.17), we have a matrix inequality [6]. We have the QFI matrix \mathcal{F}_Q on one hand, which depends on ρ and the two generators H_0 and H_1 , and the covariance matrix from we are only interested on the variance of the gradient parameter, $(\Delta b_1)^2$ [6]. Just to record, H_0 and H_1 are Hermitian operators and commute with each other and thus the QFI matrix elements are computed as $\mathcal{F}_{ij} \equiv \mathcal{F}_Q[\rho, H_i, H_j]$, following the definition (5.10).

In the two-parameter estimation problem, \mathcal{F} is a 2 × 2 matrix and the precision bound for the estimation of the gradient is the following,

$$(\Delta b_1)^{-2} \le \mathcal{F}_{11} - \frac{\mathcal{F}_{01}\mathcal{F}_{10}}{\mathcal{F}_{00}},$$
 (5.22)

where \mathcal{F}_{ij} denotes the QFI matrix elements. One must estimate both parameters in most cases since the precision of one depends on the other. The bound is interpreted slightly different from the one-parameter Cramér-Rao bound. In order to be able to saturate the bound the measurements for estimating the two parameters must compatible [6].

To compute the bound (5.21) we need to consider the matrix elements of QFI one by one. First of all, we compute \mathcal{F}_{11} that without assuming anything else has the same form as Equation (5.21)

$$\mathcal{F}_{11} = \sum_{n,m}^{N} \int d^{N} \mathbf{x} P_{N}(\mathbf{x}) x_{n} x_{m} \mathcal{F}_{Q}[\rho_{s}, j_{z}^{(n)}, j_{z}^{(m)}].$$
 (5.23)

Second, the most trivial matrix element is \mathcal{F}_{00} which is a function of the internal state ρ_{s} . To compute

it one must notice that

$$(H_0)_{x,k,y,k'} = \langle x, k | H_0 | y, k' \rangle$$

$$= \langle x, k | J_z | y, k' \rangle$$

$$= \delta(x - y)(H_0)_{k,k'}.$$
(5.24)

And with this it is straightforward to obtain

$$\mathcal{F}_{00} = \mathcal{F}_{\mathcal{Q}}[\rho_{\mathsf{s}}, J_{\mathsf{z}}]. \tag{5.25}$$

Note that this is not a function of the whole state but only of the internal ρ_s state. Finally, we compute both \mathcal{F}_{01} and \mathcal{F}_{10} . To compute this, one may notice first that both matrix elements are equal, $\mathcal{F}_{01} = \mathcal{F}_{01}$. Therefore we have to compute only one of them. The computation follows,

$$\mathcal{F}_{01} = \sum_{n=1}^{N} \int d^{N} x P_{N}(x) x_{n} \mathcal{F}_{Q}[\rho_{s}, j_{z}^{(n)}, J_{z}].$$
 (5.26)

With those results the Equation (5.31) follows. For further details see Appendix ??.

For the last part of the proof, we need to show that his bound is invariant under translations as it is the Equation (5.21). With this aim, using the linearity of the last two arguments of $\mathcal{F}_{\mathbb{Q}}[\rho, A, B]$, Eq. (5.12), the fact that H_0 remains unchanged on the Heisenberg picture and using the shifted H_1 operator, Eq. (5.19), we have that

$$\mathcal{F}_{00}(d) = \mathcal{F}_{Q}[\rho, H_{0}(d)] = \mathcal{F}_{Q}[\rho, H_{0}],$$

$$\mathcal{F}_{01}(d) = \mathcal{F}_{Q}[\rho, H_{0}(d), H_{1}(d)]$$

$$= \mathcal{F}_{Q}[\rho, H_{0}, H_{1} - dH_{0}] = \mathcal{F}_{01} - d\mathcal{F}_{00},$$

$$\mathcal{F}_{11}(d) = \mathcal{F}_{Q}[\rho, H_{1}(d), H_{1}(d)]$$

$$= \mathcal{F}_{Q}[\rho, H_{1} - dH_{0}, H_{1} - dH_{0}]$$

$$= \mathcal{F}_{11} - 2d\mathcal{F}_{01} + d^{2}\mathcal{F}_{00}.$$
(5.27)
$$(5.28)$$

Simple algebra shows that the bound for a displaced system is the same bound as is it would not be

displaced,

$$(\Delta b_{1})^{-2} \leq \mathcal{F}_{11}(d) - \frac{(\mathcal{F}_{01}(d))^{2}}{\mathcal{F}_{00}}$$

$$= \mathcal{F}_{11} - 2d\mathcal{F}_{01} + d^{2}\mathcal{F}_{00}$$

$$- \frac{\mathcal{F}_{01}^{2} - 2d\mathcal{F}_{01}\mathcal{F}_{00} + d^{2}\mathcal{F}_{00}^{2}}{\mathcal{F}_{00}}$$

$$= \mathcal{F}_{11} - \frac{\mathcal{F}_{01}^{2}}{\mathcal{F}_{00}}.$$
(5.30)

Hence, the Observation XXX holds.

For states of the form Equations (5.3, 5.4), the expression to compute the precision precision bound takes the following form,

$$(\Delta b_{1})^{-2} \leq \sum_{n,m}^{N} \int d^{N} \mathbf{x} P_{n}(\mathbf{x}) x_{n} x_{m} \mathcal{F}_{Q}[\rho_{s}, j_{z}^{(n)}, j_{z}^{(m)}]$$

$$- \frac{\left(\sum_{n=1}^{N} \int d^{N} \mathbf{x} P_{n}(\mathbf{x}) x_{n} \mathcal{F}_{Q}[\rho_{s}, j_{z}^{(n)}, J_{z}]\right)^{2}}{\mathcal{F}_{Q}[\rho_{s}, J_{z}]}.$$
(5.31)

Despite it seems to be complicated, this equation is easily computed for the most common cases such as trapped-ions, cold atomic ensembles the like. This bound as well as Equation (5.21) is invariant under spatial translations of the system.

This observations make the computation of the next sections easier since the now we can place the system arbitrarily wherever we choose. It also allows us to place the origin of our coordinate system where the magnetic field is null. So, the linear magnetic field can be written as $\mathbf{B}(x) = xB_1\mathbf{k}$ where \mathbf{k} is the unitary vector pointing on the OZ direction perpendicular to OX and OY, making the estimation of the homogenous field irrelevant to compute the precision of the gradient parameter b_1 . Despite the redundant arguments used before, the discourse we have had on the preceding section has a vital importance to understand properly the gradient metrology on this context.

5.2 Ion chains and two separated ensembles for magnetometry

Despite the powerfulness of our tools it is always nice to start with simple but concise examples to see how the results developed on the previous chapter behave. For this goal we choose two state-of-the-art systems for the external states which a priori we know that behave well under the gradient interferometry.

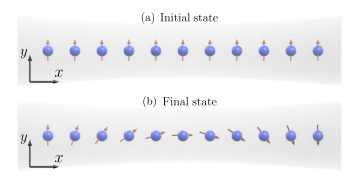


Figure 5.1: A one-dimensional chain of spin-j atoms (blue circles) confined in a potential (grey area). (a) The ensemble is initially totally polarised along a perpendicular direction of the magnetic field B_z and the direction of the chain OX. The internal state can be written as $|j\rangle_y^{\otimes N}$, the number represents m_y the eigenvalue of the one particle operator $j_y^{(n)}$. (b) One can see how the gradient field affects with a varying field strength the different spins when they are placed in different positions.

The first spatial state will be given by N distinguishable ions all placed to a constant distance from the neighbor constituents on a single dimensional configuration, *i.e.*, an ion-chain [Sanah's paper], see Figure 5.1. We have that for that matter the PDF describing the system is

$$P_N(\mathbf{x}) = \prod_{n=1}^N \delta(x_n - na). \tag{5.32}$$

With this at hand we compute the single point averages and the two point averages corresponding to the ion-chain. For the single point average we have that

$$\int d^N x P_N(x) x_n = na, (5.33)$$

and for the two point average we have the following,

$$\int d^N x P_N(x) x_n x_m = n m a^2. \tag{5.34}$$

We are going to compute the precision bound when the internal state is a product state of all particles pointing onto OY direction, $|j\rangle_{q}^{\otimes N}$, which is a state sensitive to the homogeneous field, so

we have

$$(\Delta b_{1})^{-2} \leq \sum_{n,m}^{N} nma^{2} \mathcal{F}_{Q}[|j\rangle_{y}^{\otimes N}, j_{z}^{(n)}, j_{z}^{(m)}]$$

$$-\frac{\left(\sum_{n=1}^{N} an \mathcal{F}_{Q}[|j\rangle_{y}^{\otimes N}, j_{z}^{(n)}, J_{z}]\right)^{2}}{\mathcal{F}_{Q}[|j\rangle_{y}^{\otimes N}, J_{z}, J_{z}]}$$

$$= a^{2} \left\{\sum_{n=1}^{N} 2jn^{2} - \frac{\left(\sum_{n=1}^{N} 2jn\right)^{2}}{2jN}\right\}$$

$$= 2a^{2}jN\frac{N^{2}-1}{12},$$
(5.35)

where we have used the $\mathcal{F}_{\mathbb{Q}}[|\psi\rangle$, A, $B]=4(\langle AB\rangle-\langle A\rangle\langle B\rangle)$ identity to compute the calculations.

Despite that Equation (5.35) is a third order function of the particle number N and that it seems to overcome the ultimate limit HL, on should notice that the length of the chain increases as we introduce more particles into the system. To solve this one can normalize the distance between the atoms or simply use some measure of spatial width of the complete system. In our case we decide to use on all results the standard deviation of the averaged particle position as the length measure. It is computed as follows,

$$\mu = \int d^N \mathbf{x} P_N(\mathbf{x}) \frac{\sum_{n=1}^N x_n}{N} \tag{5.36}$$

$$\sigma^{2} = \int d^{N} x P_{N}(x) \frac{\sum_{n=1}^{N} x_{n}^{2}}{N} - \mu^{2}$$
 (5.37)

$$\sigma_{\rm ch}^2 = a^2 \frac{N^2 - 1}{12}. ag{5.38}$$

It turns out that this exactly coincides with a factor we have on Equation (5.35). Substituting this onto the result the proof concludes.

For ion-chains where their constituents are separated by a constant distance and where the spinstate ρ_s is the totally polarised state along a perpendicular direction of both the field and the chain direction, in this case OY, the precision bound is given by the following formula,

$$(\Delta b_1)^{-2} \le 2\sigma_{\rm ch}^2 j N,\tag{5.39}$$

where σ_{ch} denotes the standard deviation of whole position spots on which the particles rest, j is the spin-number of each particle and N is the particle-number.

We continue this illustrative example with the state-of-the-art double-well spatial state. We also will use an internal state that with the maximal QFI so the reader gets familiar with our approach and sees how the best state to measure the gradient parameter looks like on our framework.

The spatial part is described by the following PDF where half of the particles are in one-well and the rest in the other, placed both positions at a distance of *a* from the origin,

$$P_N(\mathbf{x}) = \prod_{n=1}^{N} \delta(x_n + (-1)^n a), \tag{5.40}$$

where the "odd" particles go to the "right" and the "even" particles to the "left". With this we are able to compute the single-point and two-point correlation functions as,

$$\int d^{N} x P_{N}(x) x_{n} = (-1)^{n+1} a, \qquad (5.41)$$

$$\int d^{N} x P_{N}(x) x_{n} x_{m} = (-1)^{n+m} a^{2}.$$
 (5.42)

The state $|\psi\rangle$ is insensitive to the homogeneous field so we have that

$$(\Delta b_1)^{-2}|_{\text{max}} = \sum_{n,m}^{N} (-1)^{n+m} a^2 \mathcal{F}_{Q}[|\psi\rangle, j_z^{(n)}, j_z^{(m)}]$$

$$= a^2 \sum_{n,m}^{N} (-1)^{n+m} 4(-1)^{n+m} j^2$$

$$= 4a^2 j^2 N^2,$$
(5.43)

where we have used the definition for pure states of QFI, $4(\langle j_z^{(n)}j_z^{(m)}\rangle - \langle j_z^{(n)}\rangle\langle j_z^{(m)}\rangle)$.

On the other hand if we compute now the standard deviation as we did before for the case of the chain, Eqs. (5.36-5.38), we have that in this case $\mu = 0$ and the standard deviation

$$\sigma_{\rm dw}^2 = \int d^N \mathbf{x} P_N(\mathbf{x}) = a^2, \tag{5.44}$$

with which the proof follows. Before concluding the proof we want to show another more usual approach to the same problem.

It is as follows, given that the QFI is convex on states and having already fixed the external spatial state to be Equation (5.40), the inner state that maximises the QFI is the one that maximises $(\Delta H_1)^2$. In this case, taking into account the particle locations and that we have zero magnetic field at the

origin,

$$H_{1,\text{eff}} = b_1 a(\mathbb{I}_{\text{od}} \otimes J_{z,\text{ev}} - J_{z,\text{od}} \otimes \mathbb{I}_{\text{ev}}), \tag{5.45}$$

where we write the effective Hamiltonian that the particles feel and "od" and "ev" stand for odd and even particle index respectively. From here it is easy to show that the state maximising such variance is the following,

$$|\psi\rangle = \frac{1}{\sqrt{2}}(|0\cdots 0\rangle_{\text{od}} \otimes |1\cdots 1\rangle_{\text{ev}} + |1\cdots 1\rangle_{\text{od}} \otimes |0\cdots 0\rangle_{\text{ev}}), \tag{5.46}$$

which is equivalent to the state in Equation (5.47). This proofs that we have use the right state, and with this we can compute the QFI as in the literature as $\mathcal{F}_Q[|\psi\rangle, H_1] = 4(\Delta H_1)^2 = 4a^2j^2N^2$. With this we finally conclude the proof.

The maximally entangled internal state that maximized the QFI too is

$$|\psi\rangle = \frac{1}{\sqrt{2}}(|0101\cdots01\rangle + |1010\cdots10\rangle).$$
 (5.47)

Notice that this is a coherent state of all particles on the left "up" and all on the right "down" plus all on the left "down" and all on the right "up". We will choose without loosing of generality the particles to be spin-*j* particles. For this state insensitive to the homogeneous field and in the double-well spatial configuration, Eq. (5.40), the maximal achievable precision is

$$(\Delta b_1)^{-2}|_{\text{max}} = 4\sigma_{\text{dw}}^2 j^2 N^2 \tag{5.48}$$

In this section we have shown to the reader how one should handle the spatial width of the system for classifying it for gradient metrology as well as a state-of-the-art system on which the Heisenberg limit is achieved. Moreover, we have sown how to use the tools developed on previous section to compute simple bounds. In the next section we will focus on single cold-atoms ensembles since the play an important role on todays quantum technology and experiments and many groups are trying to realize them whit great success but with few theoretical support from our point of view.

5.3 Magnetometry with an atomic ensemble

In this section, we discuss magnetometry with a single atomic ensemble in a more deep way than in the previous section. For that, we present precision bounds for the estimation of the magnetic field gradient, for states that are insensitive to the homogeneous field. We also present precision bounds for states that are sensitive to the homogeneous field. We consider a one-dimensional cloud of spin-j

atoms placed in a one dimensional trap, which is elongated in the x-direction. The magnetic field points in the z-direction, and has a constant gradient along the x-direction. The setup is depicted in Fig. $\ref{fig:prop}$. In the last part of this section, we calculate precision bounds for the gradient estimation for some important multi-particle quantum states, for instance, Dicke states or Greenberger-Horne-Zeilinger (GHZ) states [21]. Note that all these states are permutationally invariant (PI), since we assume a PI procedure to prepare the states.

5.3.1 Precision bound for an atomic ensemble

An atomic ensemble is defined as a finite number of atoms where they cannot be labeled individually. Thus, the initial quantum state is assumed to be PI. Hence apart from ρ_s , the probability distribution function $P_N(x)$, appearing in Equation (??), must also be PI. The permutational invariance of $P_N(x)$ implies that

$$P_{N}(\mathbf{x}) = \frac{1}{N!} \sum_{k} \prod_{k} [P_{N}(\mathbf{x})], \tag{5.49}$$

where the summation is over all the possible permutations of the variables x_n denoted by Π_k .

As we have shown on Observations XXX and XXX, the precision bound is invariant under translations on the spatial Hilbert space. This allows us to place the "center of mass" of the system at the origin of the coordinates. With this simplifying assumptions the single-point average of the whole ensemble is

$$\mu = \int d^N \mathbf{x} P_N(\mathbf{x}) x_n = 0, \tag{5.50}$$

where we used the PI nature of $P_n(\mathbf{x})$ to eliminate the sum and the N from the Equation (5.36). We will do the same with the second moments appearing on the variance, Eq. (5.37). Because of this, the size of the system can be related with the single variable, x_n , variance

$$\sigma^2 = \int P_N(\mathbf{x}) x_n^2 d^N \mathbf{x}, \tag{5.51}$$

for any n, which is simplified due to the fact that the system is placed at the origin. In the same way, and due again to the PI nature of $P_N(x)$ we write the covariance of two particle positions as

$$\eta = \int P_N(\mathbf{x}) x_n x_m \, d^N \mathbf{x} \tag{5.52}$$

for any $n \neq m$. With this we have characterised all two-points correlations that potentially will appear on our calculations. An interesting property of the covariance of this type is that it is a value bounded from below and from above by the variance itself and the particle number N in the following

way,

$$\frac{-\sigma^2}{N-1} \le \eta \le \sigma^2. \tag{5.53}$$

So it cannot contribute for a better scaling on the precision than since it scales as much as σ^2 with the particle number. Notice that the lower bound scales worse.

First of all we show an important property of states insensitive to the homogeneous field and we do so using the fact that the QFI for the homogeneous field generator, i.e., J_z , on those states is zero, $\mathcal{F}_Q[\rho,J_z]=0$. The identity follows,

$$\mathcal{F}_{Q}[\rho, J_{z}] = 0$$

$$\sum_{n,m}^{N} \mathcal{F}_{Q}[\rho, j_{z}^{(n)}, j_{z}^{(m)}] = 0$$

$$\sum_{n=1}^{N} \mathcal{F}_{Q}[\rho, j_{z}^{(n)}] = -\sum_{n\neq m}^{N} \mathcal{F}_{Q}[\rho, j_{z}^{(n)}, j_{z}^{(m)}]$$
(5.54)

where we use the linearity on the second and third arguments of $\mathcal{F}_{\mathbb{Q}}[\rho,\cdot,\cdot]$ to jump to the second line and subsequently the last line follows.

From the definition of the QFI for states insensitive to the homogeneous field, Eq. (5.21), we compute the bound for single ensembles in the following way,

$$(\Delta b_{1})^{-2}|_{\text{max}} = \sum_{n,m}^{N} \int d^{N} \mathbf{x} P_{N}(\mathbf{x}) x_{n} x_{m} \mathcal{F}_{Q}[\rho_{s}, j_{z}^{(n)}, j_{z}^{(m)}]$$

$$= \sum_{n=1}^{N} \sigma^{2} \mathcal{F}_{Q}[\rho, j_{z}^{(n)}] + \sum_{n \neq m}^{N} \eta \mathcal{F}_{Q}[\rho, j_{z}^{(n)}, j_{z}^{(m)}].$$
(5.55)

Together with Equation (5.54) the Observation ?? follows.

The precision is bounded from below for a single atomic ensemble insensitive to homogeneous field with the following quantity

$$(\Delta b_1)^{-2}|_{\text{max}} = (\sigma^2 - \eta) \sum_{n=1}^{N} \mathcal{F}_{Q}[\rho_s, j_z^{(n)}].$$
 (5.56)

The bound in Eq. (5.56) can be saturated by an optimal measurement. Nevertheless, it is worth to notice that it cannot surpass the shot-noise scaling, $\sim N$, because $\mathcal{F}_Q[\rho_s, j_z^{(n)}]$, the QFI for the single-particle operator $j_z^{(n)}$, cannot be larger than j^2 .

First of all, notice that the second term appearing on Equation (5.31) is proportional to the single-point average $\int d^N x P_N(x) x_n$ which by definition is the same for any x_n and by decision is chosen to be zero, since we placed the system at the origin. So, we only have to compute the first term of the Equation (5.31),

$$(\Delta b_{1})^{-2} \leq \sum_{n,m}^{N} \int d^{N}x P_{n}(x) x_{n} x_{m} \mathcal{F}_{Q}[\rho_{s}, j_{z}^{(n)}, j_{z}^{(m)}]$$

$$= \sum_{n=1}^{N} \sigma^{2} \mathcal{F}_{Q}[\rho, j_{z}^{(n)}] + \sum_{n \neq m}^{N} \eta \mathcal{F}_{Q}[\rho, j_{z}^{(n)}, j_{z}^{(m)}]$$

$$= (\sigma^{2} - \eta) \sum_{n=1}^{N} \mathcal{F}_{Q}[\rho, j_{z}^{(n)}] + \eta \sum_{n,m}^{N} \mathcal{F}_{Q}[\rho, j_{z}^{(n)}, j_{z}^{(m)}],$$
(5.57)

where we add to the last term $\eta \sum_{n=1}^{N} \mathcal{F}_{Q}[\rho, j_{z}^{(n)}]$ and subtract it from the first term. From this, using the fact that the QFI is linear on the second and third arguments again, the proof holds.

For states sensitive to homogeneous fields, the precision of estimating the gradient is bounded from above as

$$(\Delta b_1)^{-2} \le (\sigma^2 - \eta) \sum_{n=1}^{N} \mathcal{F}_{Q}[\rho_s, j_z^{(n)}] + \eta \mathcal{F}_{Q}[\rho_s, J_z].$$
 (5.58)

The second term on the right-hand side of Eq. (5.58) is new in the sense that it did not appear on the bound for states insensitive to homogeneous fields. Note that the bound in Eq. (5.58) is not necessarily saturable if the optimal measurements to estimate the gradient parameter and the homogeneous parameter do not commute with each other. This question will be discussed in Appendix ??. Note that even if the first term cannot overcome the SL, in the second term the covariance is multiplied by QFI for estimating the homogeneous field and therefore this concrete term can make the bound, for extremely correlated particle positions, to scale as HL.

5.3.2 Precision limit for various spin-states

In this section, we present the precision limits for different classes of important quantum states such as the totally polarised state, the state having the largest precision among separable state, or the singlet state. We see how the precision bounds presented before, Eqs. (5.56, 5.58), are implemented. We show first the results for singlet that are insensitive to homogeneous fields. In this case, the bounds can be achieved by choosing the appropriate magnitude to measure. The rest of the results are for states sensitive to homogeneous fields which in general are not necessarily achievable bounds.

Singlet states

We consider now the singlet state, which is invariant under the influence of a homogeneous field along any direction. So we have to compute the formula for the bound of the precision Eq. (5.56), and we already know that it can be saturated for a certain optimal measurement.

A singlet state is an eigenstate of the collective J_z and J^2 operators, with an eigenvalue zero in both cases. Since this subspace is degenerate we have to take care in order to compute the precision bound. There are many different singlet states for an ensemble of N spin-j particles, and still a great amount of them are PI. Surprisingly the precision bound we compute is the same for any PI singlet. Atomic ensembles in a singlet state have been experimentally created with cold gases [22, 23].

In an N-particle system, there are several singlets pairwise orthogonal to each other. The number of such singlets, D_0 , depends on the particle spin j and the number of particles N.

The most general singlet state can be written in the total angular momentum basis, using D to label the degenerate states, $|J, M_z, D\rangle$, in the following diagonal way

$$\rho_{s} = \sum_{D=1}^{D_{0}} \lambda_{D} |0, 0, D\rangle \langle 0, 0, D|, \qquad (5.59)$$

where $\sum_{D} \lambda_{D} = 1$. In its complete form the eigenvalues of the spin density matrix are $\lambda_{J,M_z,D} = \delta_{0,J}\lambda_{D}$.

Looking at Eq. (5.56), we must compute the generalized QFI for the one-particle operator $j_z^{(n)}$ in order to compute the precision bound for PI singlet states. For that purpose we use the fact that when $j_z^{(n)}$ acts on a singlet state, it produces a state outside of the singlet subspace. This can be proved by noting that

$$e^{i\pi J_x} j_z^{(n)} e^{-i\pi J_x} = -j_z^{(n)}$$
 (5.60)

and that $e^{-i\pi J_x}|0,0,D\rangle=|0,0,D\rangle$ holds for any pure singlet state. Employing these equalities, we can arbitrarily flip the sign of $j_z^{(n)}$ so

$$\langle 0, 0, D | j_z^{(n)} | 0, 0, D' \rangle = -\langle 0, 0, D | j_z^{(n)} | 0, 0, D' \rangle,$$
 (5.61)

which implies

$$\langle 0, 0, D | j_z^{(n)} | 0, 0, D' \rangle = 0,$$
 (5.62)

for any pair of pure singlet singlet states.

In order to compute the QFI for the singlet state we use Eq. (5.13). Hence we can write the

following for the second term on Eq. (5.13),

$$8\sum_{D,D'} \frac{\lambda_D \lambda_{D'}}{\lambda_D + \lambda_{D'}} |\langle 0, 0, D | j_z^{(n)} | 0, 0, D' \rangle|^2 = 0.$$
 (5.63)

It follows that the QFI for any singlet is indeed simply

$$\mathcal{F}_{Q}[\rho_{s}, j_{z}^{(n)}] = 4 \operatorname{tr}(\rho_{s}(j_{z}^{(n)})^{2}). \tag{5.64}$$

For the last part of the proof, we must compute the expectation value of the operator $(j_z^{(n)})^2$. For that we have that

$$\operatorname{tr}(\rho_{s}(j_{k}^{(n)})^{2}) = \operatorname{tr}(\rho_{s}(j_{l}^{(n)})^{2}),$$
 (5.65)

for any pair $k, l \in x, y, z$ due to the rotationally invariance of the singlet, *i.e.*, all the singlets remain invariant under a SU(2) transformation of the kind $U = e^{i\phi J_{\vec{n}}}$, where \vec{n} is an unitary vector belonging to the positional space. We also have that

$$\langle (j_x^{(n)})^2 + (j_y^{(n)})^2 + (j_z^{(n)})^2 \rangle = j(j+1),$$
 (5.66)

for any state, since it represent the spin number of the particle, which is fixed. Hence, the expectation value of $(j_z^{(n)})^2$ on the singlet is

$$\operatorname{tr}(\rho_{s}(j_{z}^{(n)})^{2}) = \frac{j(j+1)}{3},$$
 (5.67)

for all the singlets. Inserting this into Eq. (5.64), we obtain $\mathcal{F}_{\mathbb{Q}}[\rho_s,j_z^{(n)}]=\frac{4j(j+1)}{3}$; and by Eq. (5.56) the statement is proved.

For PI spin states living in the singlet subspace, i.e., states composed of vectors that have zero eigenvalues for J_z and J^2 and all their possible statistical mixtures, the precision of the magnetic gradient parameter is bounded from above as

$$(\Delta b_1)_{\text{max}}^{-2} = \frac{4Nj(j+1)}{3} \left(\sigma^2 - \eta \right). \tag{5.68}$$

As mentioned earlier, singlet states are insensitive to homogeneous magnetic fields, hence determining the gradient leads to a single-parameter estimation problem. This implies that there is an optimal operator that saturates the precision bound given by Eq. (5.68). However, it is usually very hard to find this optimal measurement, although a formal procedure for this exists [6]. In Ref. [5], a particular set-up for determining the magnetic gradient with PI singlet states was suggested by the

measurement of the J_x^2 collective operator. For this scenario the precision is given by

$$(\Delta b_1)^{-2} = \frac{|\partial_{b_1} \langle J_x^2 \rangle|^2}{\langle J_x^4 \rangle - \langle J_x^2 \rangle^2}.$$
 (5.69)

In Appedix ??, we show that this measurement actually provides, in the short-time limit, an optimal precision for gradient metrology.

Totally polarised state

The totally polarised state can easily be prepared experimentally. It has already been used for gradient magnetometry with a single atomic ensemble [1, 3]. For the gradient measurement as for the measurement of the homogeneous field, the polarisation must be perpendicular to the field we want to measure in order to take advantage of the interaction between the particles and the field. Here we chose as before the totally polarized state along OY which is written as $|j\rangle_y^{\otimes N}$. Notice that this state is a state sensitive to the homogeneous field, hence, we must use the Equation (5.58) to compute the bound.

For the pure product states we have that $\mathcal{F}_{\mathbb{Q}}[|\psi\rangle,A]=4(\Delta A)^2$. Together with, $(\Delta j_z^{(n)})^2=j/2$ and $(\Delta J_z)^2=Nj/2$, when the polarisation is perpendicular to the OZ direction, the precision will be computed straightforward from Equation (5.58).

Before to do so, let us comment on the heuristics description of the evolution of the system. The homogeneous field rotates all spins by the same angle, while the gradient rotates the spin at different position by a different angle. Due to that, the homogeneous field rotates the collective spin, but does not change its absolute value. On the other hand, the field gradient decreases the absolute value of the spin, which has been used in Ref. [4] for gradient magnetometry, see Fig. 5.1.

Therefore, the Cramér-Rao bound fixes the highest value for the precision of the totaly polarised state as follows,

$$(\Delta b_1)^{-2} \le 2Nj\sigma^2. \tag{5.70}$$

Note that the precision bound for the totally polarised state is smaller than that of the optimal separable state we present later on. We can see clearly that the precision scales as $\mathcal{O}(N)$.

The best separable state

We will now turn our attention to the precision bound for all separable spin states. It is useful to obtain this value so we have a direct comparison of what is the best classically achievable precision.

It will turn out that for $j > \frac{1}{2}$, it is possible to achieve a precision higher than with the fully polarised state. We use the Equation (5.50) and substituting it onto the the low-level definitions of the precision bounds for the gradient magnetometry, Eqs. (5.21, 5.31). We see that if the state is sensitive to the homogeneous field only affects on the implications of the bound, one can be saturated for sure and on the other case it depends on the measurements compatibility as we discussed before. What we have is that the bound is the same $\mathcal{F}_Q[\rho, H_1]$ for both cases. Thus, it is easy to argue that the precision bound is a convex function on the states, even when the external state ρ_x is fixed. Therefore, the separable inner state that maximizes the precision must be a pure product state that maximises all possible $\mathcal{F}_Q[\rho_s,j_z^{(n)},j_z^{(m)}]$. For pure product states we have first that $\mathcal{F}_Q[\rho_s,j_z^{(n)},j_z^{(m)}]$ is four times the correlation between the single particle spin operators, which using the well known properties of the pure product states is zero when $n \neq m$. Finally, we have to maximise each $4(\Delta j_z^{(n)})^2$.

As we mentioned before, the best possible precision bound will be reached for a pure product state that maximises all $(\Delta j_z^{(n)})^2$. From the definition of the variance,

$$(\Delta j_z^{(n)})^2 = \langle (j_z^{(n)})^2 \rangle - \langle j_z^{(n)} \rangle^2. \tag{5.71}$$

Hence, We try a state that approaches to zero its polarisation on the OZ direction and maximises $\langle (j_z^{(n)})^2 \rangle$. We have that $|\psi\rangle = (|+j\rangle + |-j\rangle)/\sqrt{2}$ is ideal for this. The inner state for all particles is just the product state $\rho_s = |\psi\rangle\langle\psi|^{\otimes N}$. Notice that this state is permutationally invariant, hence it is a tight bound for what can be achieved with PI separable states. The variance of a single particle operator is $(\Delta j_z^{(n)})^2 = j^2$ so the proof holds.

After this discussion we make the following observation. The best achievable precision for separable states is written as

$$(\Delta b_1)^{-2} \le 4Nj^2\sigma^2,\tag{5.72}$$

where the state itself is sensitive to homogeneous fields. This bound coincides with the totally polarized state studied before when the spin number *j* is equal to half.

In the following we try to find a better precision bound making use of presumably better entangled states. Note that the bound for the singlet state, even if it is entangled, is above the bound for the totally polarised state but below of the bound defined for the best separable state. Nevertheless, when the singlet state is used effect of the homogeneous magnetic field has not to be compensated since it is insensitive to it and thus the bound can be saturated with an optimal estimator for the gradient field.

The unpolarised Dicke states $|Nj,0\rangle$ and $|Nj,0\rangle_x$

Unpolarised Dicke states play an important role in quantum optics and quantum information science. The Dicke state $|Nj,0\rangle_l$ with a maximal $\langle J_x^2 + J_y^2 + J_z^2 \rangle$ and $\langle J_l \rangle = 0$ for any $l \in x,y,z$ is particularly interesting due to its entanglement properties and its metrological usefulness. This state has been created in photonic experiments [24–26] and in cold atoms [27, 28], while a Dicke state with $\langle J_z \rangle > 0$ has been created with cold trapped ions [29].

The Dicke state $|Nj,0\rangle$ is an eigenstate of J_z so insensitive to homogeneous magnetic field pointing into the OZ direction, thus the precision can be saturate by some measurement. Whereas, The Dicke state $|Nj,0\rangle_x$ is sensitive to the homogeneous field. Moreover it is very useful for homogeneous magnetometry as it has been shown in Reference [XXX]. Here we consider large particle numbers, to make the results simpler.

Since both Dicke states are pure, we have that the QFI appearing on Equations (5.56, 5.58) are simply four times the following variances of $j_z^{(n)}$ and J_z . Since both Dicke states are unpolarized, all the first moments $\langle J_l \rangle$ are equal to zero and due to they are PI all $\langle j_l^{(n)} \rangle$ are also zero for all $l \in x, y, z$. Therefore, we only need to compute the second moments to compute the variances.

We will compute all the second moments on all directions of single particle operators $j_l^{(n)}$ as well as the global operators J_l for $|Nj,0\rangle$. Later on, we will map those result to the Dicke state for the OX direction. We have the following characteristic identities for $|Nj,0\rangle$, $\langle J_z^2\rangle=0$ and $\langle J_{\perp z}^2\rangle=\frac{Nj(Nj+1)}{2}$ where " $\perp z$ " can be seen as x or y. From the global second moments we can write for the single particle the following,

$$\langle (j_z^{(n)})^2 \rangle = -(N-1)\langle j_z^{(n)} j_z^{(m)} \rangle \tag{5.73}$$

$$\langle (j_{\perp z}^{(n)})^2 \rangle = \frac{j(Nj+1)}{2} - (N-1)\langle j_{\perp z}^{(n)} j_{\perp z}^{(m)} \rangle$$
 (5.74)

for all $\forall n \neq m$ due to the PI nature of the state. Together with $\langle (j_z^{(n)})^2 + 2(j_{\perp z}^{(n)})^2 \rangle = j(j+1)$ due to the rotational symmetry on OZ, we have three independent equations relating all the four moments. In Reference [5], a mapping between the Dicke states of spin-j ensembles and of spin- $\frac{1}{2}$ ensembles was provided, by which the expectation value

$$\langle (j_z^{(n)})^2 \rangle = \frac{(N-1)j^2}{2jN-1}$$
 (5.75)

was derived. With this we are able to obtain all the rest unknown values we are searching for. In the large N limit, this gives

$$\lim_{N \to \infty} \langle (j_z^{(n)})^2 \rangle = \frac{j}{2}. \tag{5.76}$$

From Eq. (5.56) we can conclude that $\mathcal{F}_{\mathbb{Q}}[\rho_s, j_z^{(1)}] = 2j$, hence the proof for homogeneous insensitive

 $|N_j, 0\rangle$ holds.

Now we have only to apply a mapping between the OX and OZ axis to obtain the respective second moments for $|N_j, 0\rangle_x$. We do that on the large N limit,

$$\lim_{N \to \infty} \langle (j_z^{(n)})^2 \rangle = \frac{j(2j+1)}{4}. \tag{5.77}$$

Finally, we have all the ingredients to go forward on writing the precision bound.

For large N, the precision bound for the Dicke $|N_i, 0\rangle$ state is

$$(\Delta b_1)_{\text{max}}^{-2} = 2Nj \left(\sigma^2 - \eta\right), \tag{5.78}$$

whereas for the homogeneous sensitive Dicke state $|N_j,0\rangle_x$ the precision is bounded from above by

$$(\Delta b_1)^{-2} \le Nj(2j+1)(\sigma^2 - \eta) + 2Nj(Nj+1)\eta, \tag{5.79}$$

which shows in principle a Heisenberg behavior in the second term on the right-hand side.

The Greenberger-Horne-Zeilinger (GHZ) states are also highly entangled states that play an important role in quantum physics [21]. They have been created experimentally in photonic systems [30–32] and trapped ions [33, 34].

The GHZ state is defined for qubits in the following way

$$|\mathsf{GHZ}\rangle = \frac{1}{\sqrt{2}}(|0\cdots 0\rangle + |1\cdots 1\rangle). \tag{5.80}$$

This state is very sensitive to the homogeneous field. On the other hand, as shown in Appendix \ref{Model} , for this state the optimal estimators for the homogeneous field and the gradient field are compatible. It means that both parameters can be estimated at once. Hence, in this case, the bound given in Eq. (5.58) can be saturated by some measurement set-up. In order to calculate this bound explicitly, let us recall that for pure states the QFI is simplified to Eq. (5.14). In the GHZ state the expectation value of $j_z^{(n)}$ and J_z is equal to zero, as it was for the Dicke state, and $\langle (j_z^{(n)})^2 \rangle = j^2$ and $\langle J_z^2 \rangle = N^2 j^2$, hence we obtain the following result

$$(\Delta b_1)_{\text{max}}^{-2} = 4Nj^2\sigma^2 + 4N(N-1)j^2\eta.$$
 (5.81)

This means that we can reach the Heisenberg-limit with such states, but only in cases where η is

Singlet states	$(\Delta b_1)^{-2} _{\max} = \frac{4Nj(j+1)}{3} (\sigma^2 - \eta)$
Totally polarised state	$(\Delta b_1)^{-2} \le 2N\sigma^2 j$
Best separable state	$(\Delta b_1)^{-2} \le 4N\sigma^2 j^2$
$ Nj,0\rangle$ Dicke state	$(\Delta b_1)^{-2} _{\max} = \frac{N_j}{2} \left(\sigma^2 - \eta\right)$
$ Nj,0\rangle_x$ Dicke state	$(\Delta b_1)^{-2} \le N(\sigma^2 - \eta)(2j^2 + j) + 2\eta Nj(Nj + 1)$
GHZ state	$(\Delta b_1)^{-2} _{\text{max}} = 4Nj^2(\sigma^2 + (N-1)\eta)$

Table 5.1: Precision bounds for differential magnetometry for various quantum states. For the definition of the states, see the text. If the bound are proved to be saturable then the maxisubscript is used instead of an inequality.

positive, i.e., that the particles stay spatially correlated.

Summary of results

Finally, we summarise the precision bounds obtained for various quantum states in Table 5.1. In Fig. 5.2, we show the mean values and variances of the collective angular momentum components for these states.



Figure 5.2: Angular momentum components and their variances for various quantum states. The mean value of the collective angular momentum J is represented by a red dot or arrow. The radius of the sphere is the maximal angular momentum, r = Nj. The green uncertainty ellipses describe the uncertainties of the spin components. In this case are represented different states for few particles due to the different scaling of the variances in terms of N, namely, singlet states (a), Dicke state (b), totally polarised state (c), best separable state (d), perpendicular Dicke state (e), and the GHZ (f).

6 Conclusions

ELLO it's me again

A. Long calculus appearing through the sections

In this appendix we will develop long calculations appeared throughout different chapters.

A.1 Simplification of $\langle \{J_x^2, J_y^2\} + \{J_x, J_y\}^2 \rangle$

The expectation value appearing on Equation (3.13) which we want to simplify has 6 different terms, all with two J_x and another two J_y ,

$$\langle J_x^2 J_y^2 \rangle + \langle J_x J_y J_x J_y \rangle + \langle J_x J_y^2 J_x \rangle + \langle J_y J_x^2 J_y \rangle + \langle J_y J_x J_y J_x \rangle + \langle J_y^2 J_x^2 \rangle. \tag{A.1}$$

From all those terms the third is somehow referent, since the pure unpolarised Dicke state used as reference is align with the Ox-axis, so $J_x|D_N^{N/2}\rangle_x=0$.

We use the commutation relations of the angular momentum operators $[J_k,J_l]=\epsilon_{klm}iJ_m$ to rearrange

all operators,

$$\langle J_x^2 J_y^2 \rangle = i \langle J_x J_z J_y \rangle + \langle J_x J_y J_x J_y \rangle, \tag{A.2a}$$

$$\langle J_x J_y J_x J_y \rangle = i \langle J_x J_y J_z \rangle + \langle J_x J_y^2 J_x \rangle, \tag{A.2b}$$

$$\langle J_x J_y^2 J_x \rangle = \langle J_x J_y^2 J_x \rangle,$$
 (A.2c)

$$\langle J_{u}J_{x}^{2}J_{u}\rangle = i\langle J_{u}J_{x}J_{z}\rangle + \langle J_{u}J_{x}J_{u}J_{x}\rangle, \tag{A.2d}$$

$$\langle J_y J_x J_y J_x \rangle = -i \langle J_z J_y J_x \rangle + \langle J_x J_y^2 J_x \rangle, \tag{A.2e}$$

$$\langle J_y^2 J_x^2 \rangle = -i \langle J_y J_z J_x \rangle + \langle J_y J_x J_y J_x \rangle. \tag{A.2f}$$

One may notice that with those relations is enough to see that we have $\operatorname{six} \langle J_x J_y^2 J_x \rangle$, for instance, Equation (A.2a) is $i \langle J_x J_z J_y \rangle$ plus Equation (A.2b) which at the same time is $i \langle J_x J_y J_z \rangle$ plus Equation (A.3c). So each equation has at the end one $\langle J_x J_y^2 J_x \rangle$ plus or minus some expectation value of the product of three operators.

For the three terms operators and again using the commutation relations we can further simplify this expression. Trying to get one $\langle J_x J_y J_z \rangle$ on each term, we obtain the following,

$$i\langle J_x J_z J_y \rangle = \langle J_x^2 \rangle + i\langle J_x J_y J_z \rangle,$$
 (A.3a)

$$2i\langle J_x J_y J_z \rangle = 2i\langle J_x J_y J_z \rangle, \tag{A.3b}$$

$$i\langle J_y J_x J_z \rangle = \langle J_z^2 \rangle + i\langle J_x J_y J_z \rangle,$$
 (A.3c)

$$-i\langle J_y J_z J_x \rangle = \langle J_y^2 \rangle - \langle J_z \rangle - i\langle J_x J_y J_z \rangle \tag{A.3d}$$

$$-3i\langle J_z J_y J_x \rangle = -3\langle J_x^2 \rangle - 3i\langle J_y J_z J_x \rangle$$

$$= -3\langle J_x^2 \rangle + 3\langle J_y^2 \rangle - 3i\langle J_y J_x J_z \rangle$$

$$= -3\langle J_x^2 \rangle + 3\langle J_y^2 \rangle - 3\langle J_z^2 \rangle - 3i\langle J_x J_y J_z \rangle.$$
(A.3e)

Now if we sum it all, note that the all 3 operators terms simplify, and if we take into account $6\langle J_x J_y^2 J_x \rangle$ the resulting expression is the following,

$$4\langle J_y^2 \rangle - 3\langle J_z^2 \rangle - 2\langle J_x^2 \rangle + 6\langle J_x J_y^2 J_x \rangle \tag{A.4}$$

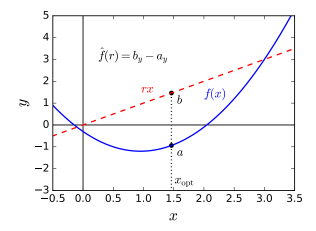


Figure A.1: Graphical representation of the Legendre transform. (blue-line) Convex function, $f(x) = x^2 - 1.9x - 0.3$, to be transformed. (red-dashed) Constant slope line passing by the coordinate system origin, rx. The Legendre transform is the maximal difference between rx and f(x) at the same x. In this case, the vertical distance between a and b.

A.2 Legendre transform

The Legendre transform of a convex function, say $f: x \to f(x)$, is defined as the maximum distance between the function the line rx and f(x) at same x. In can be written as follows,

$$\hat{f}(r) := \max_{x} \{ rx - f(x) \},$$
 (A.5)

where $\hat{f}(r)$ represents the transformed function. A geometric representation of the transform is given on the Figure A.1.

The inverse transformation is simply obtained by applying again the same technique. One fully recovers the

$$f(x) = \max_{r} \{rx - \hat{f}(r)\}.$$
 (A.6)

Let us develop the example shown in the Figure A.1, where the function is $f(x) = x^2 - 1.9x - 0.3$. In this case the problem is well defined on the complete real axis. Now, one has to find the maximum of g(r,x) = rx - f(x) for all $\forall r$. This maximum is easily obtained in this particular case with usual techniques. On has to solve for x the following equation $\partial_x g(r,x) = 0$. Thus, the maximum is at $x_{\text{opt}} = \frac{r+1.9}{2}$ and hence, the Legendre transform is the following,

$$\hat{f}(r) = \frac{r^2}{4} + 0.95r + 1.2025. \tag{A.7}$$

A.2 Legendre transform

If one applies again the transformation the resulting function is again the original one.

B. Miscellaneous mathematical tools

In this appendix we will illustrate basic mathematical tools used all through the thesis. They are shown here because without been figures of merit of the conceptual parts involving this thesis, they are nowadays sufficiently important for any whose intention is t expertise on this field of Quantum Metrology and Quantum Information.

B.1 Husimi Q-representation and the Bloch sphere

To represent states of total angular momentum bigger than $\frac{1}{2}$ we use the Husimi quasi-probability on the n unitary vector space defined as 2+2

$$Q(\alpha) = \langle |\varrho|\alpha\rangle \tag{B.1}$$

where ...

It is very common to express the particle density.

B.2 Angular momentum subspaces for different spins

Here I want to show how the whole Hilbert space of the spin angular-momentum of a multi-particle system splits. There are severas constituents such as the symmetric subspace, the PI subspace, the anti-symmetric subspace, etc.

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