On Quantum Statistical Inference

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Abstract

Interest in problems of statistical inference connected to measurements of quantum systems has recently increased substantially, in step with dramatic new developments in experimental techniques for studying small quantum systems. Furthermore, theoretical developments in the theory of quantum measurements have brought the basic mathematical framework for the probability calculations much closer to that of classical probability theory. The present paper reviews this field and proposes and interrelates a number of new concepts for an extension of classical statistical inference to the quantum context. (An earlier version of the paper containing material on further topics is available as quant-ph/0307189).

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Contents

1 Introduction ........................................ 3
   1.1 Overview ........................................ 4

2 States, Measurements and Instruments ............. 5
   2.1 The Basics ....................................... 5
   2.2 Spin-half ........................................ 9
   2.3 Superposition and Mixing ......................... 11
   2.4 The Schrödinger Equation ......................... 12
   2.5 Separability and Entanglement .................... 12
   2.6 Further Theory of Measurements .................. 13
   2.7 Further Theory of Instruments .................... 14

3 Parametric Quantum Models and Likelihood ........ 16

4 Quantum Exponential and Quantum Transformation Models ........ 17
   4.1 Quantum Exponential Models ....................... 17
   4.2 Quantum Transformation Models .................... 19

5 Quantum Exhaustivity, Sufficiency, and Quantum Cuts .... 19
   5.1 Quantum Exhaustivity ............................ 20
   5.2 Quantum Sufficiency ................................ 20
   5.3 Quantum Cuts and Likelihood Equivalence ....... 21

6 Quantum and Classical Fisher Information ............ 23
   6.1 Definition and First Properties ..................... 23
   6.2 Relation to Classical Expected Information ....... 24

7 Classical versus Quantum ................................ 27

8 Other Topics ........................................ 29
   8.1 Quantum Tomography ................................ 29
   8.2 Quantum Stochastic Processes and Continuous-Time Measurements ................... 30
   8.3 Quantum Tomography of Operations .................. 31
   8.4 Conclusion ....................................... 31
1 Introduction

Quantum mechanics has replaced classical (Newtonian) mechanics as the basic paradigm for physics. From there it pervades chemistry, molecular biology, astronomy, cosmology, . . . . The theory is fundamentally stochastic: the predictions of quantum mechanics are probabilistic. When used to derive properties of matter, the stochastic nature of the theory is typically swallowed up by the law of large numbers (very large numbers, like Avogadro’s, $10^{23}$). However, in some situations randomness does appear on the surface, most familiarly in the random times of clicks of a Geiger-counter. Present-day physicists, challenged by the fantastic theoretical promise of a quantum computer, are carrying out experiments in which half a dozen ions are held in an ion-trap and individually pushed into lower or higher energy states, and into quantum superpositions of such joint states. The existence of these wavelike superpositions of combinations of distinct states of distinct objects is a fundamentally quantum phenomenon called entanglement. Entanglement is of enormous importance in quantum computation and quantum communication. In other experiments, using supercooled electric circuits, billions of electrons behave as a single quantum particle which is brought into a wavelike superposition of macroscopically distinct states (clockwise and anti-clockwise current flow, for instance). This was recently achieved in Delft by [Mooij et al. (1993)] using a SQUID (semiconducting quantum interference device). [Hannemann et al. (2002)] recently implemented a Bayesian sequential adaptive design-and-estimation procedure to determine the state of 12 identically prepared two-level systems.

In these experiments, single quantum systems are individually manipulated and probed. The outcomes of measurements are random, with a probability distribution which depends on the one hand, on which quantum measurement (the experimental design) was carried out, and on the other hand, on the state of the quantum system being measured. If one does not know in advance the state of the quantum system, or wants to use the measurement results in order to prove that a certain state had been created, one is dealing with statistical estimation and testing problems for data from a probabilistic model with a rather elegant mathematical structure, as we shall see.

By the nature of quantum mechanics, measurement of a quantum system disturbs the system. The complete specification of a particular experiment tells us not only how the distribution of the data depends on the state of the quantum system being measured, but also how the state of the system after the measurement depends on its initial state and on the outcome which was observed. This complete specification is described mathematically by a quantum instrument. Measuring the system in one way precludes measuring it simultaneously in a different way. The total amount of information which can be obtained about an unknown parameter of the state of a quantum system is finite. Quantum physics delineates in a very precise way the class of all possible instruments. Thus, before looking at which experiments are practically feasible, one can already investigate mathematically the limits of the information which can be extracted from an unknown quantum system, leading to advice on various experimental strategies.

The field of quantum statistical inference studies these problems in a unified and systematic way. Established a quarter of a century ago in the classic monographs of Helstrom [1976] and Holevo [1982], it is currently under vigorous renewed development, stimulated by experimental efforts in nanotechnology, and the rapid theoretical development of quantum communication, quantum cryptography, quantum computation, and quantum information theory.
Though real laboratory experiments involve highly complex models and severe practical limitations, the basic theory and the basic statistical issues should be accessible to a general statistical audience. The most elementary models involve $2 \times 2$ complex matrices, some linear algebra and elementary probability. Such models already allow one to state problems of statistical design and inference which we are only just starting to understand, and which are relevant to experimentalists and theoreticians in quantum information.

The purpose of this paper is to introduce this problem area to the statistical community. We set up the basic statistical modelling in the simplest of settings, namely that of a small collection of two-dimensional quantum systems. Depending on the context, such quantum systems are called ‘spin-half systems’ (the spin of an electron, for instance), or ‘two-level atoms’ (ground state and first excited state for atoms in an ion trap, at very low temperature), or ‘qubits’ (the ‘bits’ of the RAM of a future quantum computer for which various technologies are being currently explored; one possibility being a supercooled aluminium ring in which an electric current might flow clockwise or anti-clockwise). Also covered is the polarisation of photons, leading us to phenomena studied in quantum optics such as violation of the Bell (1964) inequalities in the Aspect et al. (1982) experiment, of great current interest; see Wehrl et al. (1998), Gill (2003). Thus the same mathematical and statistical modelling covers a multitude of applications.

1.1 Overview

The paper is organised as follows. Section 2 describes the mathematical structure linking states of a quantum system, possible measurements on that system, and the resulting state of the system after measurement. Section 3 introduces quantum statistical models and notions of quantum score and quantum information, parallel to the score function and Fisher information of classical parametric statistical models. In Section 4 we introduce quantum exponential models and quantum transformation models, again forming a parallel with fundamental classes of models in classical statistics. In Section 5 we describe notions of quantum exhaustivity, quantum sufficiency and quantum cuts of a measurement, relating them to the classical notions of sufficiency and ancillarity. We next turn, in Section 6, to a study of the relation between quantum information and classical Fisher information, in particular through Cramér–Rao type information bounds. In Section 7 we discuss the interrelation between classical and quantum probability and statistics. Finally, in Section 8 we conclude with remarks on further topics of potential interest to probabilists and statisticians. Sections 4, 5, and 6 contain a considerable amount of new work.

This paper complements our more mathematical survey (Barndorff-Nielsen, Gill, and Jupp, 2001a) on quantum statistical information. A version of this paper with much further material (such as foundational questions, Bell inequalities, infinite dimensional spaces, continuous time observation of a quantum system) is available as Barndorff-Nielsen, Gill, and Jupp (2001b). Many further details can be found in Barndorff-Nielsen, Gill, and Jupp (2003). Gill (2001a) is a tutorial introduction to the basic modelling, while Gill (2001b) is an introduction to large sample quantum estimation theory. Some general references which we have found extremely useful are the books of Isham (1995), Peres (1995), Gilmor 4 (1994) and Holevo (1982, 2001a). The reader is also referred to the ‘bible of quantum information’ Nielsen and Chuang (2000), which contains excellent introductory material on the physics and the computer science, and to the basic probability and statistics text Williams (2001) which recognises (Chapter 10) quantum probability as a topic which should be in every statistician’s basic education. Fi-
nally, the former Los Alamos National Laboratory preprint service for quantum physics, now at Cornell, quant-ph at http://arXiv.org is an invaluable resource.

2 States, Measurements and Instruments

2.1 The Basics

The state of a finite-dimensional quantum system is described or represented by a $d \times d$ matrix $\rho$ of complex numbers, called the density matrix. The number $d$ is the dimension of the system and already the case $d = 2$ is rich both in mathematical structure and in applications, some of which were mentioned above. We shall write $\mathcal{H} = \mathbb{C}^d$ for the Hilbert space of $d$-dimensional complex vectors, also called the state space of the system. The inner product of vectors $\phi$ and $\psi$, written by physicists as $\langle \phi | \psi \rangle$ and by mathematicians as $\phi^* \psi$, equals $\sum \bar{\psi}_i \phi_i$ (the bar denotes complex conjugation). The length or norm of a vector is defined through $\| \phi \|^2 = \langle \phi | \phi \rangle$.

A density matrix $\rho$ has to be nonnegative and of trace 1, these properties being defined as follows. The trace of a square matrix is defined in the usual way as the sum of its diagonal elements. The definition of nonnegativity is a little more complicated. First, for an arbitrary complex matrix $X$ we define the adjoint $X^*$ of $X$ to be the matrix obtained from $X$ by taking its transpose and replacing each element by its complex conjugate. An element $\psi$ of the state space $\mathcal{H}$ is to be thought of as a column vector and hence $\psi^*$ is a row vector containing the complex conjugates of the elements of $\psi$. Since $\rho$ is a $d \times d$ matrix, the quadratic form $\psi^* \rho \psi$ is a complex scalar. The statement that $\rho$ is nonnegative means just that $\psi^* \rho \psi$ is real and nonnegative for every $\psi \in \mathcal{H}$.

Physicists would write $|\psi\rangle$ for the column vector $\psi$, $\langle \psi |$ for the row vector $\psi^*$, and $\langle \psi | \rho | \psi \rangle$ for the number $\psi^* \rho \psi$. In particular, $\langle \psi | \psi \rangle = \| \psi \|^2$ is a number, while if $\| \psi \| = 1$ then $| \psi \rangle \langle \psi |$ is the matrix which projects onto the one-dimensional subspace of $\mathcal{H}$ spanned by $\psi$. This bra-ket notation, due to Dirac, appears at first sight merely to require superfluous typing but it does gives a visual clue to the status of various objects and moreover provides a short-hand whereby the name of the bra or ket $\psi$ can be replaced by some identifying words or symbols, as in $|\varnothing\rangle$, $|\circ\rangle$.

It can be shown that a nonnegative matrix is automatically self-adjoint, i.e. $\rho = \rho^*$. Self-adjoint matrices share some familiar properties of symmetric real matrices: one can find an orthonormal basis of eigenvectors, and the eigenvalues are real numbers. The trace of a matrix equals the sum of the eigenvalues. A nonnegative matrix has nonnegative eigenvalues. Thus the eigenvalues of $\rho$ can be interpreted as a probability distribution over $\{1, \ldots, d\}$. As we shall see, this probability distribution has a physical meaning: the state $\rho$ can be thought of as a probability mixture over a collection of $d$ states, each associated with one of the eigenvectors and of a special type called a pure state. A probability mixture $\rho = \sum_i p_i \rho_i$ of density matrices is again a density matrix. This is the state obtained by taking the quantum system in state $\rho_i$ with probability $p_i$.

Example 1 (Pure state and mixed state). Let $|\phi_1\rangle, \ldots, |\phi_d\rangle$ denote an orthonormal basis of $\mathcal{H}$. If the basis is clear from the context, we can exploit the bra-ket notation and abbreviate these vectors to $|1\rangle, \ldots, |d\rangle$. Let $p_1, \ldots, p_d$, denote a probability distribution over $\{1, \ldots, d\}$. Write

$$\rho = \sum_i p_i |i\rangle \langle i|.$$
Note that $\rho_i = |i\rangle \langle i|$ is a $d \times d$ matrix of rank one. It represents the operator which projects an arbitrary vector into the one-dimensional subspace of $H$ consisting of all (complex) multiples of $|i\rangle$. One can easily check that it is a density matrix. Such a state, with density matrix being a rank-one projector and characterised by a unit length state vector in $H$, is a pure state. It follows that $\rho$, a probability mixture of density matrices, is also a density matrix. By the eigenvalue-eigenvector decomposition of self-adjoint matrices, any density matrix can be written in the form $|1\rangle$, with the vectors $|i\rangle$ orthonormal.

If a density matrix $\rho$ is of rank 1, one can write $\rho = |\phi\rangle \langle \phi|$ for some vector $|\phi\rangle$ with $\|\phi\|^2 = \langle \phi|\phi\rangle = 1$. The state is called a pure state and $|\phi\rangle$ is called the state vector; it is unique up to a complex factor of modulus 1. If the rank of a density matrix is greater than 1 then the state is called mixed. It can be written as a mixture of pure states in many different ways, especially if one does not insist that the state vectors of the pure states are orthogonal to one another.

The density matrix of a quantum system encapsulates in a very concise but rather abstract way all the predictions one can make about future observations on that system, or more generally, all results of interaction of the system with the real world.

So far we have been using the word ‘measurement’ in a rather loose way, but at this point it is important to make the technical distinction between mathematical models for a measurement when we do not care about the state of the system after the measurement, but only about the outcome, and models for a measurement including the state of the system after the measurement. The former is called a measurement and denoted by $M$; the latter, more complicated object, is called an instrument and denoted by $N$.

Let us start with the simpler object, a measurement. Consider a measurement with discrete outcome, i.e. the sample space of the outcome is at most countable. From quantum theory it follows that any measurement whatsoever, i.e. any experimental set-up, is described mathematically by a collection $M$ of $d \times d$ matrices $m(x)$ indexed by the outcomes $x$ of the experiment. The matrices have to be nonnegative (and hence also self-adjoint) and must add up to the identity matrix $1$. Let us write $p(x; \rho, M)$ for the probability that applying the measurement $M$ to the state $\rho$ produces the outcome $x$. Then we have the fundamental formula

$$p(x; \rho, M) = \text{trace}(\rho m(x)).$$  \hspace{1cm} (2)

One can see that this expression indeed defines a bona-fide probability distribution as follows. Writing $\rho = \sum p_i |\phi_i\rangle \langle \phi_i|$ and permuting cyclicly the elements in a trace of a product of matrices, one finds $\text{trace}(\rho m(x)) = \sum p_i \text{trace}(\langle \phi_i| m(x) |\phi_i\rangle)$. Thus, since $m(x)$ is a nonnegative matrix and the $p_i$ are probabilities, $p(x; \rho, M)$ is a nonnegative real number. Moreover, the sum over $x$ of these numbers is $\sum_x \text{trace}(\rho m(x)) = \text{trace}(\rho \sum_x m(x)) = \text{trace}(\rho 1) = \text{trace}(\rho) = 1$.

A quantum statistical model is a model for a partly or completely unknown state. This means that the state $\rho$ is supposed to depend on an unknown parameter $\theta$ in some parameter space $\Theta$. Write $\rho = (\rho(\theta) : \theta \in \Theta)$. When we apply a measurement $M$ to a quantum system from this model, the outcome has probability density

$$p(x; \theta, M) = \text{trace}(\rho(\theta) m(x)).$$  \hspace{1cm} (3)

Thus given the measurement and the quantum statistical model, a classical statistical inference problem is defined. Very important problems also arise when the measurement itself is indexed by an unknown parameter, but for reasons of space we do not address these here.
In principle, any measurement \( M \) whatever could be implemented as a laboratory experiment. Equation (3) tells us implicitly how much information about \( \theta \) can be obtained from a given experimental set-up \( M \). One may try to choose \( M \) in such a way as to maximise the information which the experiment will give about \( \theta \). Such experimental design problems are a main subject of this paper.

Often we are interested also in the state of the system after the measurement. In this case we need the more general notion of instrument. An instrument \( N \) (more precisely, a ‘completely positive instrument’) is represented by a family of collections of \( d \times d \) matrices \( n_i(x) \) satisfying \( \sum_x n_i(x)^*n_i(x) = 1 \) but being otherwise completely arbitrary. The index \( x \) refers to the observed outcome of the measurement, the index \( i \) could be thought of as ‘missing data’. Define \( m(x) = \sum_i n_i(x)^*n_i(x) \). It follows that the matrices \( m(x) \) are nonnegative (and self-adjoint) and add to the identity matrix, and thus represent a measurement (in the narrow or technical sense) \( M \). When we apply the instrument \( N \) to the quantum system, the outcome has the same probability density as (2), but we write it out in terms of \( N \) as

\[
p(x; \rho, N) = \sum_i \text{trace}(\rho m_i(x)^*n_i(x))
\] (4)

and the state of the system after applying the measurement, conditioned on observing the outcome \( x \), is

\[
\sigma(x; \rho, N) = \frac{\sum_i n_i(x)\rho m_i(x)^*}{\sum_i \text{trace}(\rho m_i(x)^*n_i(x))}.
\] (5)

The reader should check that the expression for \( \sigma(x; \rho, N) \) does define a bona-fide density matrix (nonnegative, trace 1). In some important practical problems the instrument itself depends on an unknown parameter, but here we suppose it is completely known.

It follows from quantum physics that whatever one can do to a quantum system has to have the form of a quantum instrument. Moreover, in principle, any quantum instrument whatsoever could be realised by some experimental set-up. Usually in the theory one starts by postulating some natural physical properties of the transformation from input or prior state to output or posterior state and data, and derives (4) and (5), which are then called the Kraus representation of the instrument, as a theorem. Here it is more convenient to start with (4) and (5). Further discussion and references are given in section 2.7.

One could consider applying two different quantum instruments, one after the other, to the same quantum system. One might even allow the choice of second instrument to depend on the outcome obtained from the first. The composition of two instruments in this way defines a new one; it is not difficult to express the matrices \( n_i(x) \) of the new instrument in terms of those of the old ones. Another way to get new instruments from old is by coarsening. Suppose one applies one instrument to a quantum system, then applies a many-to-one function of the outcome, and discards the original data. The new instrument can be written down in terms of the old by relabelling the matrices \( n_i(x) \) with new index \( j \) and variable \( y \) in obvious fashion.

In classical statistics, central notions such as sufficiency are connected to decomposing statistical models into parts (marginal and conditional distributions), and to reducing statistical models by reducing data. Starting with a quantum statistical model with density matrices \( \rho \) depending on a parameter \( \theta \), possibly with nuisance parameters too, it is now natural to ask whether notions akin to sufficiency and ancillarity can be developed for instruments. For instance, it might happen that the posterior state of a quantum system after applying a certain instrument no longer depends on the unknown parameter.
In the next subsection we shall work out many of these notions for the important special case of a two-dimensional quantum system. But first we present two special examples, connecting the notion of instrument to the classical notions (in quantum physics) of observables and unitary transformations.

**Example 2 (Simple instruments, simple measurements).** Let $x_1, \ldots, x_d$ denote $d$ distinct real numbers and let $|\psi_x\rangle, x \in \{x_1, \ldots, x_d\}$ be an orthonormal basis of $\mathcal{H}$ indexed by the numbers in $\mathcal{X} = \{x_1, \ldots, x_d\}$. We can now define an instrument $\mathcal{N}$ with outcomes in $\mathcal{X}$ by supposing that the index $i$ takes only one value, let us call it 0, and taking $n_0(x) = |\psi_x\rangle\langle\psi_x|$. This matrix is self-adjoint and idempotent (equal to its square). Therefore the corresponding matrices $m(x)$ are given by $m(x) = |\psi_x\rangle\langle\psi_x|$ too, and they sum to the identity matrix: the sum of projectors onto orthogonal one-dimensional subspaces spanning the whole space, is the identity. This shows that $\mathcal{N}$ is indeed an instrument, though of very special form indeed.

We can now compute the probability of observing the outcome $x$ and the posterior state of the quantum system, given the outcome is $x$, when the quantum system is originally in the state $\rho = \sum_i p_i |\phi_i\rangle\langle\phi_i|$. A straightforward calculation shows that they are given as follows:

$$p(x; \rho, \mathcal{N}) = \langle \psi_x | \rho | \psi_x \rangle = \sum_i p_i |\langle \psi_x | \phi_i \rangle|^2$$

$$(6)$$

$$\sigma(x; \rho, \mathcal{N}) = |\psi_x\rangle\langle\psi_x|.$$  

$$(7)$$

These formulae can be interpreted probabilistically as follows. The quantum system was initially in the pure state with density matrix $|\phi_i\rangle\langle\phi_i|$ with probability $p_i$. On being measured with the instrument $\mathcal{N}$, the system jumped to the pure state with density matrix $|\psi_x\rangle\langle\psi_x|$ producing the outcome $x$, with probability $|\langle \psi_x | \phi_i \rangle|^2$.

Let $X = \sum_x x |\psi_x\rangle\langle\psi_x|$. This is a self-adjoint matrix with eigenvalues $x_1, \ldots, x_d$ and eigenvectors $|\psi_1\rangle, \ldots, |\psi_d\rangle$. One says that the instrument $\mathcal{N}$ corresponds to the observable $X$. ‘Measuring the observable’ with this instrument produces one of the eigenvalues, and forces the system into the corresponding eigenstate. If the quantum system starts in a pure state with state vector $\phi$, then it jumps to the eigenstate $\psi_x$ with probability $|\langle \psi_x | \phi \rangle|^2$.

Suppose now $X$ is an arbitrary self-adjoint matrix. Let $\mathcal{X} = \{x_1, \ldots, x_d\}$ denote its distinct eigenvalues. Let $\Pi(x)$ denote the matrix which projects onto the eigenspace corresponding to eigenvalue $x$, not necessarily one-dimensional. Thus $X = \sum_x x \Pi(x)$. Define $n_0(x) = m(x) = \Pi(x)$. We see again that the matrices $n_0(x)$ define an instrument $\mathcal{N}$, and the matrices $m(x)$ define a corresponding measurement $M$. When this instrument is applied to the quantum system $\rho = \sum_i p_i |\phi_i\rangle\langle\phi_i|$, one obtains the outcome $x$ with probability

$$\sum_i p_i ||\Pi(x)|\phi_i\rangle||^2.$$ 

One may compute that the final state is the mixture, according to the posterior probabilities that the initial state was $|\phi_i\rangle$ given that the outcome is $x$, of the pure states with state vectors equal to the normalised projections $\Pi(x)|\phi_i\rangle/||\Pi(x)|\phi_i\rangle||$. Yet again we have the probabilistic interpretation, that with probability $p_i$ the quantum system started in the pure state with state vector $|\phi_i\rangle$. On measuring the observable $X$, the state vector is projected into one of the eigenspaces, with probabilities equal to the squared lengths of the projections. One gets to observe the corresponding eigenvalue. The posterior state is the mixture of these different pure states according to the posterior distribution of initial state given data $x$.

When one measures the observable $X = \sum_x x \Pi(x)$ with the corresponding simple instrument or simple measurement, the probability of each eigenvalue $x$ is trace($\rho \Pi(x)$). It follows
that the expected value of the outcome is trace(\rho X). More generally, let f be some real function. One may define the function f of the observable X by \( Y = \sum_x f(x) \Pi(x) \). This is the self-adjoint matrix with the same eigenspaces, and with eigenvalues equal to the function f of the eigenvalues of X. If the function f is many-to-one then some eigenspaces may have merged—consider the function ‘square’ for instance. It follows that the expected value of the function f of the outcome of measuring X is given by the elegant formula trace(\rho f(X)). We call this rule the law of the unconscious quantum physicist since it is analogous to the law of the unconscious statistician, according to which the expectation of a function \( Y = f(X) \) of a random variable X may be calculated by an integration (i) over the underlying probability space, (ii) over the outcome space of X, (iii) over the outcome space of Y. Note however that the simple instruments corresponding to X and to Y are different, and moreover neither is equal to the instrument ‘measure X, but record only \( y = f(x) \)’.

This calculus of expected values of (outcomes of measuring) observables is the basis of the mathematical theory called quantum probability; for some further remarks on this see Section 7.

Two observables P, Q are called compatible if as operators they commute: \( PQ = QP \). A celebrated result of von Neumann is that observables Q and P are compatible if and only if they are both functions \( f(R), g(R) \) of a third observable R. Taking R to have as coarse a collection of eigenspaces as possible, one can show that the results of the following three instruments are identical: the simple instrument for Q followed by the simple instrument for P, recording the values \( q \) of Q and \( p \) of P; the simple instrument for P followed by the simple instrument for Q, recording the values \( q = f(r) \) and \( p = g(r) \) where \( r \) is the observed value of R. It follows that the probability distribution of the outcome of measurement of an observable P is not altered when it is measured (simply, jointly) together with any other compatible observables.

An instrument such that the index \( i \) takes only one value, say 0, and such that all \( n_0(x) \) are projectors onto orthogonal subspaces of \( \mathcal{H} \), together spanning the whole space, is called a simple instrument. The corresponding measurement is called a simple measurement. Simple instruments and measurements stand in one-to-one correspondence with observables. The rule for the transformation of the state under a simple instrument is called the Lüders-von Neumann projection postulate.

Example 3 (Instrument with no data). It is possible that the quantum instrument \( \mathcal{N} \) transforms the quantum system \( \rho \) without actually producing any outcome \( x \); in the definition of an instrument, simply take the outcome space to consist of a single element, let us call it 0. Then the state \( \rho \) is transformed by the instrument into the state \( \sum_i n_i(0) \rho n_i(0)^* \) where the \( n_i(0) \) are matrices satisfying \( \sum_i n_i(0)^* n_i(0) = 1 \). A very special case is obtained when there is also only one value of the index \( i \) and the instrument is defined by a single matrix \( U = n_0(0) \) satisfying \( U^* U = U U^* = 1 \). State \( \rho \) is transformed into \( U \rho U^* \). Such a matrix U is called unitary and it corresponds to an orthogonal change of basis. A unitary transformation is invertible, and corresponds to the reversible time evolution of an isolated quantum system; see Subsection 2.4 below.

2.2 Spin-half

Our examples will concern mainly the spin of spin-half particles, where the dimension \( d \) of \( \mathcal{H} \) is 2. Unfortunately, it would take us too far afield to explain the significance of the word...
The classic example in this context is the 1922 experiment of Stern and Gerlach, see Brandt and Dahmen (1995, Section 1.4), to determine the size of the magnetic moment of the electron. The electron was conceived of as spinning around an axis and therefore behaving as a magnet pointing in some direction. Mathematically, each electron carries a vector ‘magnetic moment’. One might expect the sizes of the magnetic moment of all electrons to be the same, but the directions to be uniformly distributed in space. Stern and Gerlach made a beam of silver atoms move transversely through a steeply increasing vertical magnetic field. A silver atom has 47 electrons but it appears that the magnetic moments of the 46 inner electrons cancel and essentially only one electron determines the spin of the whole system. Classical physical reasoning predicts that the beam would emerge spread out vertically according to the component of the spin of each atom (or electron) in the direction of the gradient of the magnetic field. The spin itself would not be altered by passage through the magnet. However, amazingly, the emerging beam consisted of just two well separated components, as if the component of the spin vector in the vertical direction of each electron could take on only two different values, which in fact are $\pm \frac{1}{2}$ in appropriate units.

This example fits into the following mathematical framework. Take $d = 2$, then $\mathcal{H} = \mathbb{C}^2$ and $\rho$ is a $2 \times 2$ matrix

$$
\rho = \begin{pmatrix} \rho_{11} & \rho_{12} \\ \rho_{21} & \rho_{22} \end{pmatrix}
$$

with $\rho_{21} = \overline{\rho_{12}}$ and $\rho_{11}$ and $\rho_{22}$ real and nonnegative and adding to 1. The matrix has non-negative real eigenvalues $p_1$ and $p_2$ also adding to 1.

In this case the density matrices of pure states can be put into one-to-one correspondence with the unit sphere $S^2$, the surface of the unit ball in real, 3-dimensional space. Directions in the sphere correspond to directions of spin. This geometric representation is known in theoretical physics as the Poincaré sphere, in quantum optics as the Bloch sphere, and in complex analysis as the Riemann sphere. The mixed states, i.e. the convex combinations of pure states, correspond to points in the interior of the ball. The mapping from states (matrices) to points in the unit ball is affine, as we shall now show.

Any real linear combination of self-adjoint matrices is again self-adjoint. Since the 2 diagonal elements of a self-adjoint matrix must be real, and the 2 off-diagonal elements are one another’s complex conjugate, just 4 real parameters are needed to specify any such matrix. By inspection one discovers that the space of self-adjoint matrices is spanned by the identity matrix

$$
1 = \sigma_0 = \begin{pmatrix} 1 & 0 \\ 0 & 1 \end{pmatrix},
$$

together with the three Pauli matrices

$$
\sigma_x = \begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix}, \quad \sigma_y = \begin{pmatrix} 0 & -i \\ i & 0 \end{pmatrix}, \quad \sigma_z = \begin{pmatrix} 1 & 0 \\ 0 & -1 \end{pmatrix}.
$$

Note that $\sigma_x, \sigma_y$ and $\sigma_z$ satisfy the commutation relations

$$
[\sigma_x, \sigma_y] = 2i\sigma_z, \\
[\sigma_y, \sigma_z] = 2i\sigma_x, \\
[\sigma_z, \sigma_x] = 2i\sigma_y.
$$
where, for any operators $A$ and $B$, their commutator $[A, B]$ is defined as $AB - BA$. Note also that

$$\sigma_x^2 = \sigma_y^2 = \sigma_z^2 = 1.$$

Any pure state has the form $|\psi\rangle \langle \psi|$ for some unit vector $|\psi\rangle$ in $\mathbb{C}^2$. Up to a complex factor of modulus 1 (the phase, which does not influence the state), we can write $|\psi\rangle$ as

$$|\psi\rangle = \begin{pmatrix} e^{-i\phi/2} \cos(\theta/2) \\ e^{i\phi/2} \sin(\theta/2) \end{pmatrix}.$$

The corresponding pure state is

$$\rho = \begin{pmatrix} \cos^2(\theta/2) & e^{-i\phi} \cos(\theta/2) \sin(\theta/2) \\ e^{i\phi} \cos(\theta/2) \sin(\theta/2) & \sin^2(\theta/2) \end{pmatrix}.$$

A little algebra shows that $\rho$ can be written as $\rho = (1 + u_x \sigma_x + u_y \sigma_y + u_z \sigma_z) / 2 = \frac{1}{2}(1 + \vec{u} \cdot \vec{\sigma})$, where $\vec{\sigma} = (\sigma_x, \sigma_y, \sigma_z)$ are the three Pauli spin matrices and $\vec{u} = (u_x, u_y, u_z) = \vec{u}(\theta, \phi)$ is the point on the unit sphere with polar coordinates $(\theta, \phi)$.

An arbitrary mixed state is obtained by averaging pure states $\rho = \frac{1}{2}(1 + \vec{u} \cdot \vec{\sigma})$ with respect to any probability distribution over real unit vectors $\vec{u}$. The result is a density matrix of the form $\rho = \frac{1}{2}(1 + \vec{a} \cdot \vec{\sigma})$, where $\vec{a}$ is the centre of mass (a point in the unit ball) of the distribution of pure states seen as a distribution over the unit sphere. The coordinates of $\vec{a}$ are called the Stokes parameters when we are using this model to describe polarization of a photon, rather than spin of an electron.

### 2.3 Superposition and Mixing

Given two state vectors $|\phi_1\rangle$ and $|\phi_2\rangle$ and two complex numbers $c_1$, $c_2$, the state vector $(c_1|\phi_1\rangle + c_2|\phi_2\rangle) / ||c_1|\phi_1\rangle + c_2|\phi_2\rangle||$ is called the quantum superposition of the original two states, with complex weights $c_1$, $c_2$. This is a completely different way of combining two states from the mixture $p_1|\phi_1\rangle \langle \phi_1| + p_2|\phi_2\rangle \langle \phi_2|$. (Sometimes the latter is called an ‘uncoherent mixture’ and the former a ‘coherent mixture’.) For example, consider the case $d = 2$, let $|\phi_1\rangle$ and $|\phi_2\rangle$ form an orthonormal basis of $\mathcal{H} = \mathbb{C}^2$, and suppose $c_1 = c_2 = 1/\sqrt{2}$, $p_1 = p_2 = 1/2$.

We consider the equal weights superposition and the equal weights mixture of the pure states $|\phi_1\rangle$ and $|\phi_2\rangle$, showing how some measurements are able to distinguish between these states, whereas others do not.

The two matrices $m(1) = |\phi_1\rangle \langle \phi_1|$, $m(2) = |\phi_2\rangle \langle \phi_2|$ define a measurement $M^\phi$ with two possible outcomes 1 and 2, say. The probability distributions of the outcome under the superposition and under the mixture just defined are identical (probabilities 1/2 for each of the outcomes 1 and 2).

Define now a new orthonormal basis $|\psi_1\rangle = (|\phi_1\rangle + |\phi_2\rangle) / \sqrt{2}$, $|\psi_1\rangle = (|\phi_1\rangle - |\phi_2\rangle) / \sqrt{2}$. Corresponding to this basis, one can construct a measurement $M^\psi$ in the same way as before. When the superposition is measured with $M^\psi$, the outcome 1 is certain and the outcome 2 is impossible. However, when the mixture is measured with $M^\psi$, the two outcomes have equal probability 1/2.

It is very important to note that a pure state can be expressed as a superposition of others, and a mixed state as a mixture of others, in many different ways.
2.4 The Schrödinger Equation

Typically the state of a quantum system undergoes an evolution with time under the influence of an external field. The most basic type of evolution is that of an arbitrary initial state \( \rho_0 \) under the influence of a field with Hamiltonian \( H \). This takes the form

\[
\rho_t = e^{iH/t\hbar}\rho_0 e^{-iH/t\hbar},
\]

where \( \rho_t \) denotes the state at time \( t \), \( \hbar = 1.05 \times 10^{-34} \) J sec is Planck’s constant, and \( H \) is a self-adjoint operator on \( \mathcal{H} \). If \( \rho_0 \) is a pure state then \( \rho_t \) is pure for all \( t \) and we can choose unit vectors \( \psi_t \) such that \( \rho_t = |\psi_t\rangle\langle\psi_t| \) and

\[
\psi_t = e^{iH/t\hbar}\psi_0.
\]

Equation (8) is a solution of the celebrated Schrödinger equation \( i\hbar(d/dt)\psi = H\psi \) or equivalently \( i\hbar(d/dt)\rho = [H, \rho] \). The matrix \( e^{iH/t\hbar} \) is unitary. Conversely, every unitary matrix \( U \) can be written in the form \( e^{iH/t\hbar} \) for some self-adjoint matrix \( H \) and some time \( t \) and hence can be obtained by looking at some Schrödinger evolution at a suitable time.

2.5 Separability and Entanglement

When we study several quantum systems (with Hilbert spaces \( \mathcal{H}_1, \ldots, \mathcal{H}_m \)) interacting together, the natural model for the combined system has as its Hilbert space the tensor product \( \mathcal{H}_1 \otimes \cdots \otimes \mathcal{H}_m \). Then a state such as \( \rho_1 \otimes \cdots \otimes \rho_m \) represents ‘particle 1 in state \( \rho_1 \) and ... and particle \( m \) in state \( \rho_m \)’. Suppose the states \( \rho_i \) are pure with state vectors \( |\psi_i\rangle \). Then the product state we have just defined is also pure with state vector \( |\psi_1\rangle \otimes \cdots \otimes |\psi_m\rangle \).

A mixture of such states is called separable.

On the other hand, according to the superposition principle, a complex superposition of such state vectors is also a possible state vector of the interacting systems. Pure states whose state vectors cannot be written in the product form \( |\psi_1\rangle \otimes \cdots \otimes |\psi_m\rangle \) are called entangled. The same term is used for mixed states which cannot be written as a mixture of pure product states. A state which is not entangled is separable. The existence of entangled states is responsible for extraordinary quantum phenomena, which scientists are only just starting to harness (in quantum communication, computation, teleportation, etc.).

An important physical feature of unitary evolution in a tensor product space is that, in general, it does not preserve separability of states. Suppose that the state \( \rho_1 \otimes \rho_2 \) evolves according to the Schrödinger operator \( U_t = e^{iH/t\hbar} \) on \( \mathcal{H}_1 \otimes \mathcal{H}_2 \). In general, if \( H \) does not have the special form \( H_1 \otimes 1_2 + 1_1 \otimes H_2 \), the corresponding state at any non-zero time is entangled. The notorious Schrödinger Cat, is a consequence of this phenomenon of entanglement. For an illustrative discussion of this see, for instance, Isham (1993, Sect. 8.4.2).

Consider a product quantum system with density matrix \( \rho \). On its own, the first component has reduced density matrix \( \rho_1 \) obtained by “tracing out” the second component, \( (\rho_1)_{ij} = \sum_k (\rho)_{ik,jk} \). This procedure corresponds to computing a marginal from a joint probability distribution. Any mixed state can be considered as the reduction to the system of interest of a pure state on an enlarged, joint system. For instance, the completely mixed state \( 1/d \) is the result of tracing out the second component from the pure state \( \sum_j |j\rangle \otimes |j\rangle/\sqrt{d} \) on the product space formed from two copies of the original space.
2.6 Further Theory of Measurements

Example 4 (Spin-half, cont.). For any unit vector $\psi$ of $C^2$, let $\psi^\perp$ denote the unit vector orthogonal to it (unique up to a complex phase). The observable (self-adjoint matrix) $2|\psi\rangle\langle\psi| - 1 = |\psi\rangle\langle\psi| - |\psi^\perp\rangle\langle\psi^\perp|$ defines a simple instrument. It has eigenvalues 1 and $-1$ and one-dimensional eigenspaces spanned by $\psi$ and $\psi^\perp$. This observable corresponds to the spin of the particle in the direction (on the Poincaré sphere) defined by $\psi$. When ‘the spin is measured in this direction’ meaning, when this observable is measured, the result (in appropriate units) is either $+1$ or $-1$. Moreover, after the measurement has been carried out, the particle is in the pure state of spin in the corresponding direction. We mentioned two such measurements in Section 2.3 on mixing and superposition.

In particular, with outcome space $X = \{-1, 1\}$, the specification

$$
n_0(+1) = m(+1) = \frac{1}{2}(1 + \sigma_x)
$$

$$
n_0(-1) = m(-1) = \frac{1}{2}(1 - \sigma_x)
$$

defines a simple instrument (where the index takes only one value). It corresponds to the observable $\sigma_x$: spin in the $x$-direction.

We next discuss the notion of quantum randomisation, whereby adding an auxiliary quantum system to a system under study gives one further possibilities for probing the system of interest. This also connects to the important notion of realisation, i.e. representing a measurement by a simple measurement on a quantum randomised extension.

Suppose given a Hilbert space $H$, and a pair $(K, \rho_a)$, where $K$ is a Hilbert space and $\rho_a$ is a state on $K$. Any measurement $\tilde{M}$ on the product space $H \otimes K$ induces a measurement $M$ on $H$ by the defining relation

$$\text{trace } (\rho m(x)) = \text{trace } ((\rho \otimes \rho_a)\tilde{m}(x)) \quad \text{for all states } \rho \text{ on } H, \text{ all outcomes } x. \quad (9)$$

The pair $(K, \rho_a)$ is called an ancilla. The following theorem (Holevo’s extension of Naimark’s Theorem) states that any measurement $M$ on $H$ has the form $[\Box]$ for some ancilla $(K, \rho_a)$ and some simple measurement $\tilde{M}$ on $H \otimes K$. The triple $(K, \rho_a, M)$ is called a realisation of $M$ (the words extension or dilation are also used sometimes). Adding an ancilla before taking a simple measurement could be thought of as quantum randomisation.

**Theorem 1** [Holevo 1982]. For every measurement $M$ on $H$, there is an ancilla $(K, \rho_a)$ and a simple measurement $\tilde{M}$ on $H \otimes K$ which form a realisation of $M$.

We use the term ‘quantum randomisation’ because of its analogy with the mathematical representation of randomisation in classical statistics, whereby one replaces the original probability space with a product space, one of whose components is the original space of interest, while the other corresponds to an independent random experiment with probabilities under the control of the experimenter. Just as randomisation in classical statistics is sometimes needed to solve optimisation problems of statistical decision theory, so quantum randomisation sometimes allows for strictly better solutions than can be obtained without it.

Here is a simple spin-half example of a non-simple measurement which cannot be represented without quantum randomisation.

**Example 5 (The triad).** The triad, or Mercedes-Benz logo, has an outcome space consisting of just three outcomes: let us call them 1, 2 and 3. Let $\vec{v}_i$, $i = 1, 2, 3$, denote three unit vectors
in the same plane through the origin in \( \mathbb{R}^3 \), at angles of 120° to one another. Then the matrices
\[
m(i) = \frac{1}{2}(1 + \vec{u}_i \cdot \vec{\sigma})
\]
define a (non-simple) measurement \( M \) on the sample space \( \{1, 2, 3\} \). It turns up as the optimal solution to the decision problem: suppose a spin-half system is generated in one of the three states \( \rho_i = \frac{1}{2}(1 - \vec{v}_i \cdot \vec{\sigma}) \), \( i = 1, 2, 3 \), with equal probabilities. What decision rule gives the maximum probability of guessing the actual state correctly? There is no way to equal the success probability of \( M \) if one restricts attention to simple measurements, or to classically randomised procedures based on simple measurements. □

Finally, we introduce some further terminology concerning measurements. Given a measurement \( M \) and a function \( T \) from its outcome space \( \mathcal{X} \) to another space \( \mathcal{Y} \), one can define a new measurement \( M' = M \circ T^{-1} \) with outcome space \( \mathcal{Y} \). It corresponds to restricting attention to the function \( T \) of the outcome of the first measurement \( M \). We call it a coarsening of the original measurement, and we say that \( M \) is a refinement of \( M' \).

So far we have restricted attention to measurements with discrete outcome space. More generally, one considers measurements with outcomes in an arbitrary measure space \( (\mathcal{X}, \mathcal{A}) \) where \( \mathcal{A} \) is a sigma-algebra of measurable subsets of \( \mathcal{X} \). Such measurements are defined by a collection of matrices \( M(A) \) which are nonnegative, sigma-additive over \( \mathcal{A} \), and such that \( M(\mathcal{X}) = 1 \). The probability that the outcome lies in the set \( A \in \mathcal{A} \) is \( \text{trace}(\rho M(A)) \). A measurement \( M \) is called dominated by a (real, sigma-finite) measure \( \nu \) on the outcome space, if there exists a non-negative self-adjoint matrix-valued function \( m(x) \), called the density of \( M \), such that \( M(A) = \int_A m(x) \nu(dx) \) for all \( B \). When \( \mathcal{H} \) is finite dimensional, as in this paper, every measurement is dominated: take \( \nu(A) = \text{trace}(M(A)) \). In the dominated case, the outcome of the measurement has a probability distribution with density \( p(x; \rho, M) = \text{trace}(\rho m(x)) \) with respect to \( \nu \). If the outcome space is discrete and \( \nu \) is counting measure, then these notations are linked to our original setup by \( m(x) = M(\{x\}) \), \( M(A) = \sum_{x \in A} m(x) \).

To exemplify these notions, suppose for some dominated measurement \( M \) one can write \( m(x) = m_1(x) + m_2(x) \) for two non-negative self-adjoint matrix-valued functions \( m_1 \) and \( m_2 \). Define \( M' \) to be the measurement on the outcome space \( \mathcal{X}' = \mathcal{X} \times \{1, 2\} \) with density \( m(x, i) = m_i(x) \), \( (x, i) \in \mathcal{X}' \) with respect to the product of \( \nu \) with counting measure. Then \( M' \) is a refinement of \( M \).

We described earlier how one can form product spaces from separate quantum systems, leading to notions of product states, separable states, and entangled states. Given a measurement \( M \) on one component of a product space, one can naturally talk about ‘the same measurement’ on the product system. It has components \( M(A) \otimes 1 \). Given measurements \( M \) and \( M' \) defined on the two components of a product system, one can define in a natural way the measurement ‘apply \( M \) and \( M' \) simultaneously to the first and second component, respectively’: its outcome space is the product of the two outcome spaces, and it is defined using obvious notation by \( (M \otimes M')(A \times A') = M(A) \otimes M'(A') \).

A measurement \( M \) on a product space is called separable if it has a density \( m \) such that each \( m(x) \) can be written as a positive linear combination of tensor products of non-negative components. It can then be thought of as a coarsening of a measurement with density \( m' \) such that each \( m'(y) \) is a product of non-negative components.

### 2.7 Further Theory of Instruments

Just as we want to allow measurements also to take on continuous values, so we need instruments to do the same.
Consider an instrument $\mathcal{N}$ with outcomes $x$ in the measurable space $(\mathcal{X}, \mathcal{A})$. Let $\pi(dx; \rho, \mathcal{N})$ denote the probability distribution of the outcome of the measurement, and let $\sigma(x; \rho, \mathcal{N})$ denote the posterior state when the prior state is $\rho$ and the outcome of the measurement is $x$. It follows from the laws of quantum mechanics that the only physically feasible instruments have a special form, generalising in a natural way the definitions we gave for the discrete case. Namely, corresponding to $\mathcal{N}$ there must exist a $\sigma$-finite measure $\nu$ on $\mathcal{X}$ (which without loss of generality, can be taken to be a probability measure) and a collection of matrix-valued measurable functions $n_i$ of $x$ indexed by a finite or countable index $i$, such that

$$\sum_i \int_{\mathcal{X}} n_i(x)^* n_i(x) \nu(dx) = 1;$$

the posterior states for $\mathcal{N}$ are given by

$$\sigma(x; \rho, \mathcal{N}) = \frac{\sum_i n_i(x) \rho n_i(x)^*}{\sum_i \text{trace}(\rho n_i(x)^* n_i(x))} \quad (10)$$

and the distribution of the outcome is

$$\pi(dx; \rho, \mathcal{N}) = \sum_i \text{trace}(\rho n_i(x)^* n_i(x)) \nu(dx). \quad (11)$$

These formulae generalise naturally $[4]$ and $[5]$. In the physics literature, this kind of representation is often called the *Kraus representation of a completely positive instrument*. Space does not suffice to explain these terms, in particular ‘complete positivity’, further. The interested reader is referred to *Davies and Lewis* (1970), *Davies* (1976), *Kraus* (1983), *Ozawa* (1985), *Nielsen and Chuang* (2000), *Loubenets* (2001), and *Holevo* (2001a).

When the posterior state is disregarded, the instrument $\mathcal{N}$ gives rise to the measurement $M$ with density $m(x) = \sum_i n_i(x)^* n_i(x)$ with respect to the dominating measure $\nu$. Clearly, a measurement $M$ can be represented as the ‘data part’ of an instrument in very many different ways.

Further results of *Ozawa* (1985) generalise the realisability of measurements (Naimark, Holevo theorems) to the realisability of an arbitrary completely positive instrument. Namely, after forming a compound system by taking the tensor product with some ancilla, the instrument can be realised as a unitary (Schrödinger) evolution for some length of time, followed by the action of a simple instrument (measurement of an observable, with state transition according to the Lüders–von Neumann projection postulate). Therefore to say that the most general operation on a quantum system is a completely positive instrument comes down to saying: the only mechanisms known in quantum mechanics are Schrödinger evolution, von Neumann measurement, and forming compound systems. Combining these ingredients in arbitrary ways, one remains within the class of completely positive instruments; moreover, anything in that class can be realised in this way.

An instrument defined on one component of a product system can be extended in a natural way (similar to that described in Section 2.6 for measurements) to an instrument on the product system. Conversely, it is of great interest whether instruments on a product system can in some way be reduced to ‘separate instruments on the separate sub-systems’. There are two important notions in this context. The first (similar to the concept of separability of measurements) is the *mathematical* concept of separability of an instrument defined on a product system: this is that each $n_i(x)$ in the Kraus representation of an instrument
is a tensor product of separate matrices for each component. The second is the physical property which we shall call *multilocality*: an instrument is called multilocal, if it can be represented as a coarsening of a composition of separate instruments applied sequentially to separate components of the product system, where the choice of each instrument at each stage may depend on the outcomes of the instruments applied previously. Moreover, each component of the system may be measured several times (i.e. at different stages), and the choice of component measured at the $n$th stage may depend on the outcomes at previous stages. One should think of the different components of the quantum system as being localised at different locations in space. At each location separately, anything quantum is allowed, but all communication between locations is classical. It is a theorem of Bennett et al. (1999) that every multilocal instrument is separable, but that (surprisingly) not all separable instruments are multilocal. It is an open problem to find a physically meaningful characterisation of separability, and conversely to find a mathematically convenient characterisation of multilocality. (Note our terminology is not standard: the word ‘unentangled’ is used by some authors instead of ‘separable’, and ‘separable’ instead of ‘multilocal’).

Not all joint measurements (by which we just mean instruments on product systems), are separable, let alone multilocal. Just as quantum randomised measurements can give strictly more powerful ways to probe the state of a quantum system than (combinations of) simple measurements and classical randomisation, so non-separable measurements can do strictly better than separable measurements at extracting information from product systems, even if a priori there is no interaction of any kind between the subsystems.

### 3 Parametric Quantum Models and Likelihood

A measurement from a quantum statistical model $(\rho, m)$ results in an observation $x$ with density

$$p(x; \theta) = \text{trace}(\rho(\theta)m(x))$$

and log likelihood

$$l(\theta) = \log \text{trace}(\rho(\theta)m(x)) .$$

For simplicity, let us suppose that $\theta$ is one-dimensional. It is often useful to express the log likelihood derivative in terms of the *symmetric logarithmic derivative* or *quantum score* of $\rho$, denoted by $\rho_{\theta}$. This is defined implicitly as the self-adjoint solution of the equation

$$\rho_{\theta} = \rho \circ \rho_{\theta} ,$$

(12)

where $\circ$ denotes the Jordan product, i.e.

$$\rho \circ \rho_{\theta} = \frac{1}{2}(\rho \rho_{\theta} + \rho_{\theta} \rho) ,$$

$\rho_{\theta}$ denoting the ordinary derivative of $\rho$ with respect to $\theta$ (term-by-term differentiation in matrix representations of $\rho$). (We shall often suppress the argument $\theta$ in quantities like $\rho$, $\rho_{\theta}$, $\rho_{\theta}$, etc.) The quantum score exists and is essentially unique subject only to mild conditions. (For a discussion of this see, for example, p. 274 of Holevo 1982.)
The likelihood score \( l_\theta(\theta) = (d/d\theta)l(\theta) \) may be expressed in terms of the quantum score \( \rho_\theta(\theta) \) of \( \rho(\theta) \) as

\[
l_\theta(\theta) = p(x; \theta)^{-1} \text{trace}(\rho_\theta(\theta)m(x))
\]

\[
= p(x; \theta)^{-1} \frac{1}{2} \text{trace} \left( (\rho(\theta)\rho_{\bar{\theta}}(\theta) + \rho_{\bar{\theta}}(\theta)\rho(\theta))m(x) \right)
\]

where we have used the fact that for any self-adjoint matrices \( P, Q, R \) and any matrix \( T \) the trace operation satisfies \( \text{trace}(PQR) = \text{trace}(RQP) \) and \( \Re \text{trace}(T) = \frac{1}{2} \text{trace}(T + T^*) \). It follows that

\[
E_\theta(l_\theta(\theta)) = \text{trace}(\rho(\theta)\rho_{\bar{\theta}}(\theta)).
\]

Thus, since the mean value of \( l_\theta \) is 0, we find that

\[
\text{trace}(\rho(\theta)\rho_{\bar{\theta}}(\theta)) = 0.
\]  

The expected (Fisher) information \( i(\theta) = i(\theta; M) = E_\theta(l_\theta(\theta)^2) \) may be written as

\[
i(\theta; M) = \int p(x; \theta)^{-1} \left( \Re \text{trace}(\rho(\theta)\rho_{\bar{\theta}}(\theta)m(x)) \right)^2 \nu(dx).
\]  

(14)

It plays a key role in the quantum context, just as in classical statistics, and is discussed in Section 6. In particular, we shall discuss there its relation with the expected or Fisher quantum information

\[
I(\theta) = \text{trace}(\rho(\theta)\rho_{\bar{\theta}}(\theta)^2).
\]  

(15)

The quantum score is a self-adjoint operator, and therefore may be interpreted as an observable which one might measure on the quantum system. What we have just seen is that the outcome of a measurement of the quantum score has mean zero and variance equal to the quantum Fisher information.

## 4 Quantum Exponential and Quantum Transformation Models

In traditional statistics, the two major classes of parametric models are the exponential models (in which the log densities are affine functions of appropriate parameters) and the transformation (or group) models (in which a group acts in a consistent fashion on both the sample space and the parameter space); see Barndorff-Nielsen and Cox (1994). The intersection of these classes is the class of exponential transformation models, and its members have a particularly nice structure. There are quantum analogues of these classes, and they have useful properties. Below we outline some of these briefly. Considerably more discussion is given in our works Barndorff-Nielsen et al. (2001a,b, 2003).

### 4.1 Quantum Exponential Models

A quantum exponential model is a quantum statistical model for which the states \( \rho(\theta) \) can be represented in the form

\[
\rho(\theta) = e^{-\lambda}(\theta)e^{\frac{1}{2} \gamma^T(\theta)T_0} \rho_0 e^{\frac{1}{2} \gamma^T(\theta)T_0} \quad \theta \in \Theta,
\]
where \( \gamma = (\gamma^1, \ldots, \gamma^k) : \Theta \rightarrow \mathbb{C}^k \), \( T_1, \ldots, T_k \) are \( d \times d \) matrices, \( \rho_0 \) is self-adjoint and non-negative (but not necessarily a density matrix), the Einstein summation convention (of summing over any index which appears as both a subscript and a superscript) has been used, and \( \kappa(\theta) \) is a log norming constant, given by

\[
\kappa(\theta) = \log \text{trace}(e^{\frac{\imath}{2} \gamma^r(\theta) T_r^*} \rho_0 e^{\frac{\imath}{2} \gamma^r(\theta) T_r}).
\]

Three important special types of quantum exponential model are those in which \( T_1, \ldots, T_k \) are self-adjoint (and for the first type, \( T_0, T_1, \ldots, T_k \) all commute), and the quantum states have the forms

\[
\rho(\theta) = e^{-\kappa(\theta)} \exp \left( T_0 + \theta^r T_r \right),
\]

\[
\rho(\theta) = e^{-\kappa(\theta)} \exp \left( \frac{1}{2} \theta^r T_r \right) \rho_0 \exp \left( \frac{1}{2} \theta^r T_r \right),
\]

\[
\rho(\theta) = \exp \left( -i \frac{1}{2} \theta^r T_r \right) \rho_0 \exp \left( i \frac{1}{2} \theta^r T_r \right),
\]

respectively, where \( \theta = (\theta^1, \ldots, \theta^k) \in \mathbb{R}^k \) and \( \rho_0 \) is nonnegative (and self-adjoint), and the summation convention is in force.

We call these three types the quantum exponential models of **mechanical type**, **symmetric type**, and **unitary type** respectively. The mechanical type arises (at least, with \( k = 1 \)) in quantum statistical mechanics as a state of statistical equilibrium, see [Gardiner and Zoller (2000, Sect. 2.4.2)]. The symmetric type has theoretical statistical significance, as we shall see, connected among other things to the fact that the quantum score for this model is easy to compute explicitly. The unitary type has physical significance connected to the fact that it is also a transformation model (quantum transformation models are defined in the next subsection). The mechanical type is a special case of the symmetric type when \( T_0, T_1, \ldots, T_k \) all commute.

In general, the statistical model obtained by applying a measurement to a quantum exponential model is not an exponential model (in the classical sense). However, for a quantum exponential model of the form (17) in which

\[
T_j = t_j(X) \quad j = 1, \ldots, k
\]

for some self-adjoint \( X \), i.e. the \( T_j \) commute, the statistical model obtained by applying the measurement \( X \) is a full exponential model. Various pleasant properties of such quantum exponential models then follow from standard properties of the full exponential models.

The classical Cramér–Rao bound for the variance of an unbiased estimator \( t \) of \( \theta \) is

\[
\text{Var}(t) \geq i(\theta; M)^{-1}.
\]

Combining (20) with [Braunstein and Caves (1994)] (19) quantum information bound \( i(\theta; M) \leq I(\theta) \) (see (27) in Section 6.2) yields Helstrom’s (1976) quantum Cramér–Rao bound

\[
\text{Var}(t) \geq I(\theta)^{-1},
\]

whenever \( t \) is an unbiased estimator based on a quantum measurement. It is a classical result that, under certain regularity conditions, the following are equivalent: (i) equality holds in (20), (ii) the score is an affine function of \( t \), (iii) the model is exponential with \( t \) as canonical statistic (cf. pp. 254–255 of [Cox and Hinkley 1974]). This result has a quantum analogue, see
Theorems 3 and 4 and Corollary 1 below, which states that under certain regularity conditions, there is equivalence between (i) equality holds in (21) for some unbiased estimator \( t \) based on some measurement \( M \), (ii) the symmetric quantum score is an affine function of commuting \( T_1, \ldots, T_k \), and (iii) the quantum model is a quantum exponential model of type (17) where \( T_1, \ldots, T_k \) satisfy (19). The regularity conditions which we assume below are indubitably too strong: the result should be true under minimal smoothness assumptions.

4.2 Quantum Transformation Models

Consider a parametric quantum model \( (\rho, M) \) consisting of a parametrised family \( \rho = (\rho(\theta) : \theta \in \Theta) \) of states and a measurement \( M \) with outcome space \( (X, A) \). Suppose that there exists a group, \( G \), with elements \( g \), acting both on \( X \) and on \( \Theta \) in such a way that the following consistency condition (‘equivariance’) holds

\[
\text{trace}(\rho(\theta)M(A)) = \text{trace}(\rho(g\theta)M(gA))
\]

for all \( \theta, A \) and \( g \). If, moreover, \( G \) acts transitively on \( \Theta \) then we say that \( (\rho, M) \) is a quantum transformation model. In this case, the resulting statistical model for the outcome of a measurement of \( M \), i.e., \( (X, A, P) \), where \( P = \{\text{trace}(\rho(\theta)M) : \theta \in \Theta\} \), is a classical transformation model. Consequently, the Main Theorem for transformation models, see Barndorff-Nielsen and Cox (1994, pp. 56–57) and references given there, applies to \( (X, A, P) \).

Of special physical interest is the case in which the group acts on the states as a group of unitary matrices.

Example 6 (Spin-half: great circle model). Consider the spin-half model \( \rho(\theta) = U \frac{1}{2}(1 + \cos \theta \sigma_x + \sin \theta \sigma_y)U^* \) where \( U \) is a fixed \( 2 \times 2 \) unitary matrix, and \( \sigma_x \) and \( \sigma_y \) are two of the Pauli spin matrices, while the parameter \( \theta \) varies through \([0, 2\pi)\); see Subsection 2.2. The matrix \( U \) can always be written as \( \exp(-i\phi \vec{u} \cdot \vec{\sigma}) \) for some real three-dimensional unit vector \( \vec{u} \) and angle \( \phi \). Considered as a curve on the Poincaré sphere, the model forms a great circle. If \( U \) is the identity (or, equivalently, \( \phi = 0 \)) the curve just follows the equator; the presence of \( U \) corresponds to rotating the sphere carrying this curve about the direction \( \vec{u} \) through an angle \( \phi \). Thus our model describes an arbitrary great circle on the Poincaré sphere, parameterised in a natural way. Since we can write \( \rho(\theta) = UV_\theta U^* \rho(0)UV_\theta^* U^* \), where the unitary matrix \( V_\theta \) corresponds to rotation of the Poincaré sphere by an angle \( \theta \) about the z-axis, we can write this model as a transformation model. The model is clearly also an exponential model of unitary type. Perhaps surprisingly, it can be reparameterised so as also to be an exponential model of symmetric type. We leave the details (which depend on the algebraic properties of the Pauli spin matrices) to the reader, but just point out that a one-parameter pure-state exponential model of symmetric type has to be of the form \( \rho(\theta) = \exp(-\kappa(\theta))\exp(\frac{1}{2} \theta \vec{u} \cdot \vec{\sigma}) \frac{1}{2}(1 + \vec{v} \cdot \vec{\sigma})\exp(\frac{1}{2} \theta \vec{u} \cdot \vec{\sigma}) \) for some real unit vectors \( \vec{u} \) and \( \vec{v} \), since every self-adjoint \( 2 \times 2 \) matrix is an affine function of a spin matrix \( \vec{u} \cdot \vec{\sigma} \). Now write out the exponential of a matrix as its power series, and use the fact that the square of any spin matrix is the identity. This example is due to Fujiwara and Nagaoka (1995).

5 Quantum Exhaustivity, Sufficiency, and Quantum Cuts

This section proposes and interrelates a number of concepts that will constitute, we think, essential elements in the development of statistical inference for the quantum context. The
concepts are partly in the nature of quantum analogues of key ideas of classical statistical inference, such as sufficiency, the likelihood principle, etc.

5.1 Quantum Exhaustivity

Those quantum instruments for which no information on the unknown parameter of a quantum parametric model of states can be obtained from subsequent measurements on the given physical system deserve special attention. To simplify notation, we will write $\sigma(x; \rho(\theta), \mathcal{N})$ instead of $\sigma(x; \rho(\theta), \mathcal{N})$ for the posterior state. We propose the following definition of exhaustivity:

**Definition 1 (Exhaustive instrument).** A quantum instrument $\mathcal{N}$ is *exhaustive* for a parameterised set $\rho = (\rho(\theta) : \theta \in \Theta)$ of states if for all $\theta$ in $\Theta$ and for $\pi(\cdot; \theta)$-almost all $x$, the posterior state $\sigma(x; \theta, \mathcal{N})$ does not depend on $\theta$.

Thus the posterior states obtained from exhaustive quantum instruments are completely determined by the result of the measurement and do not depend on $\theta$.

A useful strong form of exhaustivity is defined as follows.

**Definition 2 (Completely exhaustive instrument).** A quantum instrument $\mathcal{N}$ is *completely exhaustive* if it is exhaustive for all parameterised sets of states. □

Recall that any completely positive instrument—in other words, any physically realisable instrument—has posterior states given by (10) and outcome distributed as (11). The following Proposition (which is a slight generalisation of a result of Wiseman 1999) shows one way of constructing completely exhaustive completely positive quantum instruments.

**Proposition 1.** Let the quantum instrument $\mathcal{N}$ be as above, with $n_i(x)$ of the form

$$n_i(x) = |\phi_{i,x}\rangle\langle\psi_x|,$$

for some functions $(i, x) \mapsto \phi_{i,x}$ and $x \mapsto \psi_x$. Then $\mathcal{N}$ is completely exhaustive.

**Proof.** By inspection we find that the posterior state is

$$\sigma(x; \rho, \mathcal{N}) = \frac{\sum_i |\phi_{i,x}\rangle\langle\phi_{i,x}|}{\sum_i \langle\phi_{i,x}|\psi_{i,x}|},$$

which does not depend on the prior state $\rho$. □

5.2 Quantum Sufficiency

Suppose that the measurement $M' = M \circ T^{-1}$ is a coarsening of the measurement $M$. In this situation we say that $M'$ is (classically) *sufficient* for $M$ with respect to a family of states $\rho = (\rho(\theta) : \theta \in \Theta)$ on $\mathcal{H}$ if the mapping $T$ is sufficient for the identity mapping on $(\mathcal{X}, \mathcal{A})$ with respect to the family $(P(\cdot; \theta; M : \theta \in \Theta)$ of probability measures on $(\mathcal{X}, \mathcal{A})$ induced by $M$ and $\rho$.

As a further step towards a definition of quantum sufficiency, we introduce a concept of inferential equivalence of parametric models of states.
Definition 3 (Inferential equivalence). Two parametric families of states $\rho = (\rho(\theta) : \theta \in \Theta)$ and $\sigma = (\sigma(\theta) : \theta \in \Theta)$ on Hilbert spaces $\mathcal{H}$ and $\mathcal{K}$ are said to be inferentially equivalent if for every measurement $M$ on $\mathcal{H}$ there exists a measurement $M'$ on $\mathcal{K}$ such that for all $\theta \in \Theta$

$$\text{trace}(M(\cdot)\rho(\theta)) = \text{trace}(M'(\cdot)\sigma(\theta))$$

and vice versa. (Note that, implicitly, the outcome spaces of $M$ and $M'$ are assumed to be identical.)

In other words, $\rho$ and $\sigma$ are equivalent if and only if they give rise to the same class of possible classical models for inference on the unknown parameter.

Remark 1. It is of interest to find characterisations of inferential equivalence. This is a nontrivial problem, even when the Hilbert spaces $\mathcal{H}$ and $\mathcal{K}$ are the same.

Next, let $\mathcal{N}$ denote a quantum instrument on a Hilbert space $\mathcal{H}$ and with outcome space $(\mathcal{X}, \mathcal{A})$ and let $\mathcal{N}' = \mathcal{N} \circ T^{-1}$ be a coarsening of $\mathcal{N}$ with outcome space $(\mathcal{Y}, \mathcal{B})$, generated by a mapping $T$ from $(\mathcal{X}, \mathcal{A})$ to $(\mathcal{Y}, \mathcal{B})$. It is easy to show that the posterior states for the two instruments are related by

$$\sigma(t; \theta, \mathcal{N}') = \int_{T^{-1}(t)} \sigma(x; \theta, \mathcal{N}) \pi(dx|t; \theta, \mathcal{N}),$$

where $\pi(dx|t; \theta, \mathcal{N})$ is the conditional distribution of $x$ given $T(x) = t$ computed from $\pi(dx; \theta, \mathcal{N})$.

Definition 4 (Quantum sufficiency of instruments). Let $\mathcal{N}'$ be a coarsening of an instrument $\mathcal{N}$ by $T : (\mathcal{X}, \mathcal{A}) \rightarrow (\mathcal{Y}, \mathcal{B})$. Then $\mathcal{N}'$ is said to be quantum sufficient with respect to a family of states $(\rho(\theta) : \theta \in \Theta)$ if

(i) the measurement $M'$ determined by $\text{trace}(M'(\cdot)\rho) = \pi(\cdot; \rho, \mathcal{N}')$ is sufficient for the measurement $M$ determined by $\text{trace}(M(\cdot)\rho) = \pi(\cdot; \rho, \mathcal{N})$, with respect to the family $(\rho(\theta) : \theta \in \Theta)$,

(ii) for any $x \in \mathcal{X}$, the posterior families $(\sigma(x; \theta, \mathcal{N}) : \theta \in \Theta)$ and $(\sigma(T(x); \theta, \mathcal{N}') : \theta \in \Theta)$ are inferentially equivalent.

5.3 Quantum Cuts and Likelihood Equivalence

In the theory of classical statistical inference, many important concepts (such as sufficiency, ancillarity and cuts) can be expressed in terms of the decomposition by a measurable function $T : (\mathcal{X}, \mathcal{A}) \rightarrow (\mathcal{Y}, \mathcal{B})$ of each probability distribution on $(\mathcal{X}, \mathcal{A})$ into the corresponding marginal distribution of $T(x)$ and the family of conditional distributions of $x$ given $T(x)$. We now define analogous concepts in quantum statistics based on the decomposition

$$\rho \mapsto (\pi(\cdot; \rho, \mathcal{N}), \sigma(\cdot; \rho, \mathcal{N}))$$

by a quantum instrument $\mathcal{N}$ of each state $\rho$ into a probability distribution on $(\mathcal{X}, \mathcal{A})$ and a family of posterior states; see Section 2.7.
The classical concept of a cut encompasses those of sufficiency and ancillarity and is therefore more basic. A measurable function $T$ is a cut for a set $\mathcal{P}$ of probability distributions on $\mathcal{X}$ if for all $p_1$ and $p_2$ in $\mathcal{P}$, the distribution on $\mathcal{X}$ obtained by combining the marginal distribution of $T(x)$ given by $p_1$ with the family of conditional distributions of $x$ given $T(x)$ given by $p_2$ is also in $\mathcal{P}$; see, e.g. Barndorff-Nielsen and Cox (1994, p. 38). Recent results on cuts for exponential models can be found in Barndorff-Nielsen and Koudou (1995), which also gives references to the useful role which cuts have played in graphical models. A generalisation to local cuts has become important in econometrics (Christensen and Kiefer, 1994, 2000).

Replacing the decomposition into marginal and conditional distributions in the definition of a cut by the decomposition (25) yields the following quantum analogue.

**Definition 5 (Quantum cut).** A quantum instrument $\mathcal{N}$ is said to be a quantum cut for a family $\rho$ of states if for all $\rho_1$ and $\rho_2$ in $\rho$

\[
\pi(\cdot; \rho_3, \mathcal{N}) = \pi(\cdot; \rho_1, \mathcal{N})
\]
\[
\sigma(\cdot; \rho_3, \mathcal{N}) = \sigma(\cdot; \rho_2, \mathcal{N}).
\]

for some $\rho_3$ in $\rho$. □

Thus, if $\mathcal{N}$ is a quantum cut for a family $\rho = (\rho(\theta) : \theta \in \Theta)$ with $\rho$ a one-to-one function then $\Theta$ has the product form $\Theta = \Psi \times \Phi$ and furthermore $\sigma(\cdot; \rho(\theta), \mathcal{N})$ depends on $\theta$ only through $\psi$, and $\pi(\cdot; \rho(\theta), \mathcal{N})$ depends on $\theta$ only through $\phi$.

Since a quantum instrument $\mathcal{N}$ is exhaustive for a parameterised set $\rho = (\rho(\theta) : \theta \in \Theta)$ of states if the family $\sigma(\cdot; \rho(\theta), \mathcal{N})$ of posterior states does not depend on $\theta$, exhaustive quantum instruments are quantum cuts of a special kind. They can be regarded as quantum analogues of sufficient statistics. At the other extreme are the quantum instruments for which the distributions $\pi(\cdot; \rho(\theta), \mathcal{N})$ do not depend on $\theta$. These can be regarded as quantum analogues of ancillary statistics.

Unlike exhaustivity, the concept of quantum sufficiency involves not only a quantum instrument but also a coarsening. The definition of quantum sufficiency can be extended to the following version involving parameters of interest.

**Definition 6 (Quantum sufficiency for interest parameters).** Let $\rho = (\rho(\theta) : \theta \in \Theta)$ be a family of states and let $\psi : \Theta \to \Psi$ map $\Theta$ to the space $\Psi$ of interest parameters. A coarsening $\mathcal{N}'$ of a quantum instrument $\mathcal{N}$ by a mapping $T$ is said to be quantum sufficient for $\psi$ on $\rho$ if

(i) the measurement $M'$ determined by $\text{trace}(M'(\cdot)\rho) = \pi(\cdot; \rho, \mathcal{N}')$ is sufficient for the measurement $M$ determined by $\text{trace}(M(\cdot)\rho) = \pi(\cdot; \rho, \mathcal{N})$, with respect to the family $\rho$,

(ii) for all $\theta_1$ and $\theta_2$ with $\psi(\theta_1) = \psi(\theta_2)$ and for all $x$ in $\mathcal{X}$, the sets $\sigma(x; \rho(\theta_1), \mathcal{N})$ and $\sigma(T(x); \rho(\theta_2), \mathcal{N}')$ of posterior states are inferentially equivalent.

□

Consideration of the likelihood function obtained by applying a measurement to a parameterised set of states suggests that the following weakening of the concept of inferential equivalence may be useful.
Definition 7 (Strong likelihood equivalence). Two parametric families of states $\rho = (\rho(\theta) : \theta \in \Theta)$ and $\sigma = (\sigma(\theta) : \theta \in \Theta)$ on Hilbert spaces $H$ and $K$ respectively are said to be strongly likelihood equivalent if for every measurement $M$ on $H$ there is a measurement $M'$ on $K$ with the same outcome space, such that

$$\frac{\text{trace}(M(dx)\rho(\theta))}{\text{trace}(M(dx)\rho(\theta'))} = \frac{\text{trace}(M'(dx)\sigma(\theta))}{\text{trace}(M'(dx)\sigma(\theta'))},$$

$\theta, \theta' \in \Theta$ (whenever these ratios are defined) and vice versa. □

Thus the likelihood function of the statistical model obtained by applying $M$ to $\rho$ is equivalent to that obtained by applying $M'$ to $\sigma$, for the same outcome of each instrument.

Consideration of the distribution of the likelihood ratio leads to the following definition.

Definition 8 (Weak likelihood equivalence). Two parametric families of states $\rho = (\rho(\theta) : \theta \in \Theta)$ and $\sigma = (\sigma(\theta) : \theta \in \Theta)$ on Hilbert spaces $H$ and $K$ respectively are said to be weakly likelihood equivalent if for every measurement $M$ on $H$ with outcome space $X$ there is a measurement $M'$ on $K$ with some outcome space $Y$ such that the likelihood ratios

$$\frac{\text{trace}(M(dx)\rho(\theta))}{\text{trace}(M(dx)\rho(\theta'))} \quad \text{and} \quad \frac{\text{trace}(M'(dy)\sigma(\theta))}{\text{trace}(M'(dy)\sigma(\theta'))}$$

have the same distribution for all $\theta, \theta'$ in $\Theta$, and vice versa. □

The precise connection between likelihood equivalence and inferential equivalence is not yet known but the following conjecture appears reasonable.

Conjecture. Two quantum models are strongly likelihood equivalent if and only if they are inferentially equivalent up to quantum randomisation.

6 Quantum and Classical Fisher Information

In Section 3 we showed how to express the Fisher information in the outcome of a measurement in terms of the quantum score. In this section we discuss quantum analogues of Fisher information and their relation to the classical concepts.

6.1 Definition and First Properties

Differentiating (13) with respect to $\theta$, writing $\rho_{\theta/\theta}$ for the derivative of the symmetric logarithmic derivative $\rho_{\theta}$ of $\rho$, and using the defining equation (12) for $\rho_{\theta}$, we obtain

$$0 = \text{trace}(\rho_{\theta/\theta}(\theta)\rho_{\theta/\theta}(\theta) + \rho(\theta)\rho_{\theta/\theta}(\theta))$$

$$= \text{trace} \left( \frac{1}{2} (\rho(\theta)\rho_{\theta/\theta}(\theta) + \rho_{\theta/\theta}(\theta)\rho(\theta)) \rho_{\theta}(\theta) \right) + \text{trace}(\rho(\theta)\rho_{\theta/\theta}(\theta))$$

$$= I(\theta) - \text{trace}(\rho(\theta)J(\theta)), \quad \text{where}$$

$$I(\theta) = \text{trace} (\rho(\theta)\rho_{\theta/\theta}(\theta)^2)$$

is the expected (or Fisher) quantum information, already mentioned in Sections 3 and 4, and

$$J(\theta) = -\rho_{\theta/\theta/\theta}(\theta),$$

23
which we shall call the observable quantum information. Thus

$$I(\theta) = \text{trace} \left( \rho(\theta) J(\theta) \right),$$

which is a quantum analogue of the classical relation $i(\theta) = E_{\theta}(j(\theta))$ between expected and observed information (where $j(\theta) = -l_{\theta}/\theta(\theta)$). Note that $J(\theta)$ is an observable, just as $j(\theta)$ is a random variable.

Neither $I(\theta)$ nor $J(\theta)$ depends on the choice of measurement, whereas $i(\theta) = i(\theta; M)$ does depend on the measurement $M$. Expected quantum information behaves additively, i.e. for parametric quantum models of states of the form $\rho : \theta \mapsto \rho_1(\theta) \otimes \cdots \otimes \rho_n(\theta)$ (which model ‘independent particles’), the associated expected quantum information satisfies

$$I_{\rho_1 \otimes \cdots \otimes \rho_n}(\theta) = \sum_{i=1}^n I_{\rho_i}(\theta),$$

which is analogous to the additivity property of Fisher information.

In the case of a multivariate parameter $\theta$, the expected quantum information matrix $I(\theta)$ is defined in terms of the quantum scores by

$$I_{\rho_1 \otimes \cdots \otimes \rho_n}(\theta) = \frac{1}{2} \text{trace} \left( \rho_{\theta_j}(\theta) \rho_{\theta_k}(\theta) + \rho_{\theta_k}(\theta) \rho_{\theta_j}(\theta) \right). \quad (26)$$

### 6.2 Relation to Classical Expected Information

Suppose that $\theta$ is one-dimensional. There is an important relationship between expected quantum information $I(\theta)$ and classical expected information $i(\theta; M)$, due to [Braunstein and Caves (1994)](1994), namely that for any measurement $M$ with density $m$ with respect to a $\sigma$-finite measure $\nu$ on $X$,

$$i(\theta; M) \leq I(\theta), \quad (27)$$

with equality if and only if, for $\nu$-almost all $x$,

$$m(x)^{1/2} \rho_{\theta_j}(\theta) \rho(\theta)^{1/2} = r(x)m(x)^{1/2} \rho(\theta)^{1/2}, \quad (28)$$

for some real number $r(x)$.

For each $\theta$, there are measurements which attain the bound in the quantum information inequality (27). For instance, we can choose $M$ such that each $m(x)$ is a projection onto an eigenspace of the quantum score $\rho_{\theta_j}(\theta)$. Note that this attaining measurement may depend on $\theta$.

**Example 7 (Information for spin-half).** Consider a spin-half particle in the pure state $\rho = \rho(\eta, \theta) = |\psi(\eta, \theta)\rangle \langle \psi(\eta, \theta)|$ given by

$$|\psi(\eta, \theta)\rangle = \begin{pmatrix} e^{-i\theta/2} \cos(\eta/2) \\ e^{i\theta/2} \sin(\eta/2) \end{pmatrix}.$$  

As we saw in Subsection 2.2, $\rho$ can be written as $\rho = (1 + u_x \sigma_x + u_y \sigma_y + u_z \sigma_z)/2 = \frac{1}{2}(1 + \vec{u} \cdot \vec{\sigma})$, where $\vec{\sigma} = (\sigma_x, \sigma_y, \sigma_z)$ are the three Pauli spin matrices and $\vec{u} = (u_x, u_y, u_z) = \vec{u}(\eta, \theta)$ is the point on the Poincaré sphere $S^2$ with polar coordinates $(\eta, \theta)$. Suppose that the colatitude $\eta$ is known and exclude the degenerate cases $\eta = 0$ or $\eta = \pi$; the longitude $\theta$ is the unknown parameter.
Since all the \( \rho(\theta) \) are pure, one can show that \( \rho_{\theta\theta}(\theta) = 2\rho_{\theta}(\theta) = \vec{u}(\theta) \cdot \vec{\sigma} = \sin(\eta) \vec{u}(\pi/2, \theta + \pi/2) \cdot \vec{\sigma} \). Using the properties of the Pauli matrices, one finds that the quantum information is
\[
I(\theta) = \text{trace}(\rho(\theta)\rho_{\theta\theta}(\theta)^2) = \sin^2 \eta.
\]
Summarising some results from Barndorff-Nielsen and Gill (2000), we now discuss a condition that a measurement must satisfy in order for it to achieve this information.

It follows from (28) that, for a pure spin-half state \( \rho = |\psi\rangle\langle\psi| \), a necessary and sufficient condition for a measurement to achieve the information bound is: for \( \nu \)-almost all \( x \), \( m(x) \) is proportional to a one-dimensional projector \( |\xi(x)\rangle\langle\xi(x)| \) satisfying
\[
\langle\xi(x)|2\rangle\langle2|a \rangle = r(x)|\xi(x)||1\rangle,
\]
where \( r(x) \) is real, \( |1\rangle = |\psi\rangle \), \( |2\rangle = |\psi\rangle^\perp \) (\( |\psi\rangle^\perp \) being a unit vector in \( \mathbb{C}^2 \) orthogonal to \( |\psi\rangle \)) and \( |a\rangle = 2|\psi\rangle_{\theta} \). It can be seen that geometrically this means that \( |\xi(x)\rangle \) corresponds to a point on \( S^2 \) in the plane spanned by \( \vec{u}(\theta) \) and \( \vec{u}_{\theta}(\theta) \).

If \( \eta \neq \pi/2 \), then distinct values of \( \theta \) give distinct planes, and all these planes intersect in the origin only. Thus no single measurement \( M \) can satisfy \( I(\theta) = i(\theta; M) \) for all \( \theta \). On the other hand, if \( \eta = \pi/2 \), so that the states \( \rho(\theta) \) lie on a great circle in the Poincaré sphere, then the planes defined for each \( \theta \) are all the same. In this case any measurement \( M \) with all components proportional to projector matrices for directions in the plane \( \eta = \pi/2 \) satisfies \( I(\theta) = i(\theta; M) \) for all \( \theta \in \Theta \). In particular, any simple measurement in that plane has this property.

More generally, a smooth one-parameter model of a spin-half pure state with everywhere positive quantum information admits a uniformly attaining measurement, i.e. such that \( I(\theta) = i(\theta; M) \) for all \( \theta \in \Theta \), if and only if the model is a great circle on the Poincaré sphere. This is actually a quantum exponential transformation model, see Example 4.

When the state \( \rho \) is strictly positive, and under further nondegeneracy conditions, essentially the only way to achieve the bound (27) is through measuring the quantum score. In the discussion below we first keep the value of \( \theta \) fixed. Since any nonnegative self-adjoint matrix can be written as a sum of rank-one matrices (using its eigenvalue-eigenvector decomposition), it follows that any dominated measurement can be refined to a measurement for which each \( m(x) \) is of rank 1, thus \( m(x) = r(x)|\xi(x)\rangle\langle\xi(x)| \) for some real \( r(x) \) and state vector \( |\xi(x)\rangle \), see the end of Section 2.6. If one measurement is the refinement of another, then the distributions of the outcomes are related in the same way. Therefore, under refinement of a measurement, Fisher expected information cannot decrease. Therefore if any measurement achieves (27), there is also a measurement with rank 1 components achieving the bound. Consider such a measurement. Suppose that \( \rho \) is positive and that all the eigenvalues of \( \rho_{\theta\theta} \) are different. The condition \( m(x)^{1/2}\rho_{\theta\theta}^{1/2} = r(x)m(x)^{1/2}\rho^{1/2} \) is then equivalent to \( |\xi(x)\rangle\langle\xi(x)|\rho_{\theta\theta} = r(x)|\xi(x)\rangle\langle\xi(x)| \), which states that \( \xi(x) \) is an eigenvector of \( \rho_{\theta\theta} \). Since we must have \( \int m(x)\mu(dx) = 1 \), it follows that all eigenvectors of \( \rho_{\theta\theta} \) occur in this way in components \( m(x) \) of \( M \). The measurement can therefore be reduced or coarsened to a simple measurement of the quantum score, and the reduction (at the level of the outcome) is sufficient.

Suppose now that the state \( \rho(\theta) \) is strictly positive for all \( \theta \), and that the quantum score has distinct eigenvalues for at least one value of \( \theta \). Suppose that a single measurement exists attaining (27) uniformly in \( \theta \). Any refinement of this measurement therefore also achieves the
bound uniformly; in particular, the refinement to components which are all proportional to projectors onto orthogonal one-dimensional eigenspaces of the quantum score at the value of \( \theta \) where the eigenvalues are distinct does so. Therefore the eigenvectors of the quantum score at this value of \( \theta \) are eigenvectors at all other values of \( \theta \). Therefore there is a self-adjoint operator \( X \) with distinct eigenvalues such that \( \rho_{\theta_0}(\theta) = f(X; \theta) \) for each \( \theta \). Fix \( \theta_0 \) and let \( F(X; \theta) = \int_{\theta_0}^{\theta} f(X; \theta) d\theta \). Let \( \rho_0 = \rho(\theta_0) \). If we consider the defining equation (12) as a differential equation for \( \rho(\theta) \) given the quantum score, and with initial condition \( \rho(\theta_0) = \rho_0 \), we see that a solution is \( \rho(\theta) = \exp\left(\frac{1}{2} F(X; \theta)\right) \rho_0 \exp\left(\frac{1}{2} F(X; \theta)\right) \). Under smoothness conditions the solution is unique. Rewriting the form of this solution, we come to the following theorem:

**Theorem 2 (Uniform attainability of quantum information bound).** Suppose that the state is everywhere positive, the quantum score has distinct eigenvalues for some value of \( \theta \), and is smooth. Suppose that a measurement \( M \) exists with \( i(\theta; M) = I(\theta) \) for all \( \theta \), thus attaining the Braunstein–Caves information bound (27) uniformly in \( \theta \). Then there is an observable \( X \) such that a simple measurement of \( X \) also achieves the bound uniformly, and the model is of the form

\[
\rho(\theta) = c(\theta) \exp\left(\frac{1}{2} F(X; \theta)\right) \rho_0 \exp\left(\frac{1}{2} F(X; \theta)\right)
\]  

(29)

for a function \( F \), indexed by \( \theta \), of an observable \( X \) where \( c(\theta) = 1/\text{trace}(\rho_0 \exp(F(X; \theta))) \), \( \rho_{\theta_0}(\theta) = f(X; \theta) - \text{trace}(\rho(\theta)f(X; \theta)) \), and \( f(X; \theta) = F_{\theta_0}(X; \theta) \). Conversely, for a model of this form, a measurement of \( X \) achieves the bound uniformly.

**Remark 2 (Spin-half case).** In the spin-half case, if the information is positive then the quantum score has distinct eigenvalues, since the outcome of a measurement of the quantum score always equals one of the eigenvalues, has mean zero, and positive variance.

**Theorem 3 (Uniform attainability of quantum Cramér–Rao bound).** Suppose that the positivity and nondegeneracy conditions of the previous theorem are satisfied, and suppose that for the outcome of some measurement \( M \) there is a statistic \( t \) such that, for all \( \theta \), \( t \) is an unbiased estimator of \( \theta \) achieving Helstrom’s quantum Cramér–Rao bound (21), \( \text{Var}(t) = I(\theta)^{-1} \). Then the model is a quantum exponential model of symmetric type (17),

\[
\rho(\theta) = c(\theta) \exp\left(\frac{1}{2} \theta T\right) \rho_0 \exp\left(\frac{1}{2} \theta T\right)
\]

for some observable \( T \), and simple measurement of \( T \) is equivalent to the coarsening of \( M \) by \( t \).

**Proof.** The coarsening \( M' = M \circ t^{-1} \) by \( t \) of the measurement \( M \) also achieves the quantum information bound (27) uniformly; i.e. \( i(\theta; M') = I(\theta) \). Applying Theorem 2 to this measurement, we discover that the model is of the form (29), while (if necessary refining the measurement to have rank one components) \( t \) can be considered as a function of the outcome of a measurement of the observable \( X \), and it achieves the classical Cramér–Rao bound for unbiased estimators of \( \theta \) based on this outcome. The density of the outcome (with respect to counting measure on the eigenvalues of \( X \)) is found to be \( c(\theta) \exp(F(x; \theta)) \text{trace}(\rho_0 \Pi_x) \) where \( \Pi(x) \) is the projector onto the eigenspace of \( X \) corresponding to eigenvalue \( x \). Hence, up to addition of functions of \( \theta \) or \( x \) alone, \( F(x; \theta) \) is of the form \( \theta \ell(x) \).
The basic inequality (27) holds also when the dimension of $\theta$ is greater than one. In that case, the quantum information matrix $I(\theta)$ is defined in (26) and the Fisher information matrix $i(\theta; M)$ is defined by
\[ i_{rs}(\theta; M) = E_\theta(l_r(\theta)l_s(\theta)), \]
where $l_r$ denotes $l/\theta$ etc. Then (27) holds in the sense that $I(\theta) - i(\theta; M)$ is positive semidefinite. The inequality is sharp in the sense that $I(\theta)$ is the smallest matrix dominating all $i(\theta; M)$. However it is typically not attainable, let alone uniformly attainable.

Theorem 2 can be generalised to the case of a vector parameter. This also leads to a generalisation of Theorem 3, which is the content of Corollary 1 below.

Theorem 4. Let $(\rho(\theta) : \theta \in \Theta)$ be a twice differentiable parametric quantum model. If

(i) there is a measurement $M$ with $i(\theta; M) = I(\theta)$ for all $\theta$,

(ii) $\rho(\theta)$ is positive for all $\theta$,

(iii) $\Theta$ is simply connected

then, for any $\theta_0$ in $\Theta$, there are an observable $X$ and a function $F$ (possibly depending on $\theta_0$) such that
\[ \rho(\theta) = \exp\left(\frac{1}{2}F(X; \theta)\right) \rho(\theta_0) \exp\left(\frac{1}{2}F(X; \theta)\right). \]

Corollary 1. If, under the conditions of Theorem 4, there exists an unbiased estimator $t$ of $\theta$ based on the measurement $M$ achieving (21), then the model is a quantum exponential family of symmetric type (17) with commuting $T_r$.

Versions of these results have been known for some time; see Young (1975), Fujiwara and Nagaoka (1995), Amari and Nagaoka (2000); compare especially our Corollary 1 to Amari and Nagaoka (2000, Theorem 7.6), and our Theorem 4 to parts (I)–(IV) of the subsequent outlined proof in Amari and Nagaoka (2000). Unfortunately, precise regularity conditions and detailed proofs seem to be available elsewhere only in some earlier publications in Japanese. Note that we have obtained the same conclusions, by a different proof, in the spin-half pure state case, Example 7. This indicates that a more general result is possible without the hypothesis of positivity of the state. See Matsumoto (2002) for important new work on the pure state case.

The symmetric logarithmic derivative is not the unique quantum analogue of the classical statistical concept of score. Other analogues include the right, left and balanced derivatives obtained from suitable variants of (12). Each of these gives a quantum information inequality and a quantum Cramér–Rao bound analogous to (27) and (21). See Belavkin (1976), and (as yet) unpublished new work by this author. There is no general relationship between the various quantum information inequalities when the dimension of $\theta$ is greater than one.

Asymptotic optimality theory for quantum estimation has only just started to be developed; see Gill and Massar (2001), Gill (2001a), and Keyl and Werner (2001); for an application see Hannemann et al. (2002).

7 Classical versus Quantum

This section makes some general comments on the relation between classical and quantum probability and statistics. This has been a matter of heated controversy ever since the discovery of quantum mechanics. It has mathematical, physical, and philosophical ingredients, and
much confusion, if not controversy, has been generated by problems of interdisciplinary communication between mathematicians, physicists, philosophers and more recently statisticians. Authorities from both physics and mathematics, perhaps starting with Feynman (1951), have promoted vigorously the standpoint that ‘quantum probability’ is something very different from ‘classical probability’. Malley and Hornstein (1993) conclude from a perceived conflict between classical and quantum probability that ‘quantum statistics’ should be set apart from classical statistics. Even Williams (2001) states that Nature chooses a different model for probability for the quantum world than for the classical world.

In our opinion, though important mathematical and physical facts lie at the root of these statements, they are misleading, since they seem to suggest that quantum probability and quantum statistics do not belong to the field of classical probability and statistics. However, quantum probabilities have the same meaning (whether you are a Bayesian or a frequentist) as classical probabilities, and statistical inference problems from quantum mechanics fall squarely in the framework of classical statistics. The statistical design problems are special to the field.

Our stance is that the predictions which quantum mechanics makes of the real world are stochastic in nature. A quantum physical model of a particular phenomenon allows one to compute probabilities of all possible outcomes of all possible measurements of the quantum system. The word ‘probability’ means here ‘relative frequency in many independent repetitions’. The word ‘measurement’ is meant in the broad sense of ‘macroscopic results of interactions of the quantum system under study with the outside world’. These predictions depend on a summary of the state of the quantum system. The word ‘state’ might suggest some fundamental property of a particular collection of particles, but for our purposes all we need to understand under the word is ‘a convenient mathematical encapsulation of the information needed to make any such predictions’. Some physicists argue that it is meaningless to talk of the state of a particular particle, one can only talk of the state of a large collection of particles prepared in identical circumstances; this is called a statistical ensemble. Others take the point of view that when one talks about the state of a particular quantum system one is really talking about a property of the mechanism which generated that system. Given that quantum mechanics predicts only probabilities, as far as real-world predictions are concerned the distinction between on the one hand a property of an ensemble of particles or of a procedure to prepare particles, and on the other hand a property of one particular particle, is a matter of semantics. However, if one would like to understand quantum mechanics by somehow finding a more classical (intuitive) physical theory in the background which would explain the observed phenomena, this becomes an important issue. It is also an issue for cosmology, when there is only one closed quantum system under study: the universe. At this level there is a remarkable difference between classical and quantum probabilities: according to the celebrated theorem of Bell (1964), it is impossible to derive the probabilities described in quantum mechanics by an underlying deterministic theory from which the probabilities arise ‘merely’ as the reflection of statistical variation in the initial conditions, unless one accepts grossly unphysical nonlocality in the ‘hidden variables’. Thus quantum probabilities are fundamentally irreducible, in contrast to every other physical manifestation of randomness known to us.

It follows from our standpoint that ‘quantum statistics’ is classical statistical inference about unknown parameters in models for data arising from measurements on a quantum system. However, just as in biostatistics, geostatistics, etc., many of these statistical problems have a common structure and it pays to study the core ideas and common features in detail.
As we have seen, this leads to the introduction of mathematical objects such as quantum score, quantum expected information, quantum exponential family, quantum transformation model, quantum cuts, and so on; the names are deliberately chosen because of analogy and connections with the existing notions from classical statistics.

Already at the level of probability (i.e. before statistical considerations arise) one can see a deep and fruitful analogy between the mathematics of quantum states and observables on the one hand, and classical probability measures and random variables on the other. Note that collections of random variables and collections of operators can both be endowed with algebraic structure (sums, products, . . . ). It is a fact that from an abstract point of view a basic structure in probability theory—a collection of random variables $X$ on a countably generated probability space, together with their expectations $\int XdP$ under a given probability measure $P$—can be represented by a (commuting) subset of the set of self-adjoint operators $Q$ on a separable Hilbert space, together with the expectations $\text{trace}(\rho Q)$ computed using the trace rule under a given state $\rho$. Thus a basic structure in classical probability theory is isomorphic to a special case of a basic structure in quantum probability. ‘Quantum probability’, or ‘noncommutative probability theory’ is the name of the branch of mathematics which takes as its starting point the mathematical structure of states and observables in quantum mechanics. From this mathematical point of view, one may claim that classical probability is a special case of quantum probability. The claim does entail, however, a rather narrow (functional analytic) view of classical probability. Moreover, many probabilists will feel that abandoning commutativity is throwing away the baby with the bathwater, since this broader mathematical structure has no analogue of the sample outcome $\omega$, and hence no opportunity for a probabilist’s beloved probabilistic arguments.

As statisticians, we would like to argue (tongue in cheek) that quantum probability is merely a special case of classical statistics. A quantum probability model is determined by specifying the expectations of every observable. This is equivalent to specifying a family of classical probability models: namely the joint probability distribution of the measurements of every commuting subset of observables. The basic structure of quantum probability is mathematically equivalent to a particular case of the basic structure of classical statistical inference—namely, an indexed family of probability models.

8 Other Topics

There are many further topics in quantum physics where more extensive use of knowledge and techniques from classical statistics and probability seems likely to lead to substantial scientific advances. However, classical concepts and results from the latter fields will often need considerable modification or recasting to be suitable and relevant for the quantum world, as is exemplified in parts of the previous Sections.

Here we shall indicate briefly a few of the topics. The selection of these is motivated mainly by our own current interests rather than by an aim to be in some way representative of the broad picture. However, the topics listed are all subject to considerable developments in the current literature. For more detailed accounts, with references to the physics and mathematics literature, see Barndorff-Nielsen, Gill, and Jupp (2001b, 2003).

8.1 Quantum Tomography

In its simplest form, the problem of quantum tomography is as follows.
The simple harmonic oscillator is the basic model for the motion of a quantum particle in a quadratic potential well on the real line. Precisely the same mathematical structure describes oscillations of a single mode of an electromagnetic field (a single frequency in one direction in space). In this type of structure one considers the quadrature observable at phase $\phi$, given by $X_\phi = Q \cos \phi + P \sin \phi$, where $Q$ and $P$ are the position and momentum operators. Here, the underlying Hilbert space $\mathcal{H}$ is infinite dimensional and the operators $Q$ and $P$ can be characterised abstractly by the commutation relation $[Q, P] = i\hbar 1$.

Given independent measurements of $X_\phi$, with $\phi$ drawn repeatedly at random from the uniform distribution on $(0, 2\pi]$, the aim is to reconstruct the unknown state $\rho$ of the quantum system. In statistical terms, we wish to do nonparametric estimation of $\rho$ from $n$ independent and identically distributed observations $(\phi_i, x_i)$, with $\phi_i$ as just described and $x_i$ from the measurement of $X_{\phi_i}$. In quantum optics, measuring a single mode of an electromagnetic field in what is called a quantum homodyne experiment, this would be the appropriate model with perfect photodetectors. In practice, independent Gaussian noise should be added.

Some key references are the book Leonhardt (1997) and the survey papers D’Ariano (1997a,b, 2001). Of special interest is a maximum likelihood based approach to the problem that has been taken in recent work by Banaszek et al. (2000). We think that it is a major open problem to work out the asymptotic theory of this method, taking account of data-driven truncation, and possibly alleviating the problem of the large parameter-space by using Bayesian methods. The method should be tuned to the estimation of various functionals of $\rho$ of interest, and should provide standard errors or confidence intervals.

8.2 Quantum Stochastic Processes and Continuous-Time Measurements

In this paper we have focussed directly on questions of quantum statistical inference. Of major related importance are the areas of quantum stochastic processes and continuous-time quantum measurements. These are currently undergoing rapid developments, and the concept of quantum instruments, discussed above, has a key role in parts of this. References to much of this work are available in Biane (1998) (see also the more extensive account by Meyer, 1993), Percival (1998), Holevo (2001a,b), Belavkin (2002), and Barndorff-Nielsen and Loubenets (2002).

There are quantum analogues of Brownian motion and Poisson processes, and more generally of Lévy processes, and a quantum stochastic analysis based on these. Interesting combinations of classical and quantum stochastic analysis occur in a variety of contexts, for instance in Monte Carlo simulation studies of the Markov quantum master equation; Mølmer and Castin (1996) is an important early reference. The Markov quantum master equation is important particularly in quantum optics which is one of the currently most active and exciting fields of quantum physics.

Other, mainly mathematically motivated, studies have strongly algebraic elements, such as in free probability. In this context a variety of ‘independence’ concepts have turned up, with associated Lévy processes, etc. See Barndorff-Nielsen and Thorbjørnsen (2002a,b) and Franz et al. (2001) and references given there. Note, moreover, that Biane and Speicher (2001) discuss a concept of free Fisher information.
8.3 Quantum Tomography of Operations

We have focussed on quantum statistical models where only the state depends on an unknown parameter. Of great interest is also the situation where an unknown operation acts on a known state.

Consider a quantum instrument $\mathcal{N}$ which produces no data but simply converts an input state $\rho$ into an output state $\sigma(\rho; \mathcal{N})$. By the general theory, $\sigma(\rho; \mathcal{N}) = \sum_i n_i \rho n_i^*$ for some collection of matrices $n_i$ satisfying $\sum n_i^* n_i = 1$. This representation is not unique but one can fix the $n_i$ by making some identifying restrictions. One could then proceed to estimate the $n_i$ by feeding the instrument with sufficiently many different input states $\rho$, many times, each time carrying out sufficiently many different measurements on the output state.

It has recently been discovered that there is an extremely effective short cut to this procedure. Consider two copies of the original quantum system, supposed here to be of dimension $d$. Consider the maximally entangled state $|\Psi\rangle = \sum_j |j\rangle \otimes |j\rangle/\sqrt{d}$ on the product system. Now allow the instrument $\mathcal{N}$ to act on the first component of the product system, while the second component is left unchanged. The output state, also on the product system, has density matrix $\sigma(|\Psi\rangle\langle\Psi|; \mathcal{N} \otimes \mathcal{I})$ where $\mathcal{I}$ denotes the identity instrument. It turns out that the output state completely characterizes $\mathcal{N}$. In fact, there is a one-to-one correspondence between on the one hand completely positive data-less instruments $\mathcal{N}$, and on the other hand density matrices on the product system such that the reduced density matrix of the second component is the completely mixed state $\sum_j |j\rangle\langle j|/d$ (i.e., the same as the reduced density matrix of the second component initially, which is left unchanged by the procedure).

Thus one does not need to probe the instrument with many different input states, but can effectively probe it with all inputs simultaneously, by exploiting quantum entanglement with an auxiliary system. The problem has been converted into a quantum statistical model $\sigma(|\Psi\rangle\langle\Psi|; \mathcal{N} \otimes \mathcal{I})$ with parameter being the unknown instrument $\mathcal{N}$.

This procedure has been pioneered by D’Ariano and Lo Presti (2001) and has already been exploited experimentally.

8.4 Conclusion

This paper has, in brief form, presented our present view of a role for statistical inference in quantum physics. We are keenly aware that many relevant parts of quantum physics and quantum stochastics have not been reviewed, or have only been touched upon.

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