Quantum Apices: Identifying Limits of Entanglement, Nonlocality, & Contextuality

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Quantum Apices: Identifying Limits of Entanglement, Nonlocality, & Contextuality

Elie Wolfe, Ph.D.
University of Connecticut, 2014

This work develops analytic methods to quantitatively demarcate quantum reality from its subset of classical phenomenon, as well as from the superset of general probabilistic theories. Regarding quantum nonlocality, we discuss how to determine the quantum limit of Bell-type linear inequalities. In contrast to semidefinite programming approaches, our method allows for the consideration of inequalities with abstract weights, by means of leveraging the Hermiticity of quantum states. Recognizing that classical correlations correspond to measurements made on separable states, we also introduce a practical method for obtaining sufficient separability criteria. We specifically vet the candidacy of driven and undriven superradiance as schema for entanglement generation. We conclude by reviewing current approaches to quantum contextuality, emphasizing the operational distinction between nonlocal and
contextual quantum statistics. We utilize our abstractly-weighted linear quantum bounds to explicitly demonstrate a set of conditional probability distributions which are simultaneously compatible with quantum contextuality while being incompatible with quantum nonlocality. It is noted that this novel statistical regime implies an experimentally-testable target for the Consistent Histories theory of quantum gravity.
Quantum Apices: Identifying Limits of
Entanglement, Nonlocality, & Contextuality

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B.A., Yeshiva University, 2006
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A Dissertation
Submitted in Partial Fullfilment of the
Requirements for the Degree of
Doctor of Philosophy
at the
University of Connecticut

2014
APPROVAL PAGE

Doctor of Philosophy Dissertation

Quantum Apices: Identifying Limits of Entanglement, Nonlocality, & Contextuality

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University of Connecticut

2014
to Noam and Eitan,

sweet and strong.
ACKNOWLEDGEMENTS

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Dr. Yelin opened up her own projects to me, and at the same time gave me complete freedom to pursue my own interests. She was able to be attentive and encouraging while also granting me flexibility and understanding to balance my family life. She was always available for questions, be they about research, writing, or tact. Every topic that I have looked into, every conference I have attended, every opportunity I have been afforded, are due to Dr. Yelin’s consistent support. Would that I might but emulate her in advising students of my own, then I’d be so very proud.

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Chapter 1

Nonlocality

[Quantum statistical limits are] something strange, neither mathematics nor physics, of very little interest. \(^1\)

Nonlocality is the most characteristic feature of quantum mechanics. \(^2\)

---

\(^1\) Anonymous Reviewer, 1980

*Letter to Boris Tsirelson* rejecting to publish Tsirelson’s derivation of his historic inequalities. [quoted by Tsirelson in private communication]

\(^2\) Sandu Popescu, 2014

*Nature Physics Insight, Quantum Foundations*, summarizing the modern perspective [1].
1.1 From Classical to General No-Signalling

It is now well understood that quantum measurements are contextual in the sense that the outcome of a measurement on a given subsystem can depend on the context of global measurements [2–6]. A context here means a tuple which uniquely maps all the subsystems to some measurements choices. A context can be nonlocal if the respective subsystems are spatially separated and the measurement choices for each party’s subsystem are selected nearly simultaneously. In such a scenario there isn’t enough time for a signal to propagate across the entire system to inform the subsystems of which global context has been selected. Nonetheless, the measurement outcomes manifest a striking awareness of the global context by yielding noticeably different statistics depending on the particular global context in play [7–17].

It is inevitably surprising to first learn that our universe does indeed exhibit such quantum nonlocality. How can such a thing be, if we accept that no signal can traverse any distance faster than light could? The answer is that the marginal statistics are independent of the global context. If we split the global system in two subsystems of any size, say subsystems A and B, then the probability of seeing some set out measurement outcomes in A upon selecting some choice of measurements for subsystem A, is guaranteed to be identical no matter what choice of measurements are selected for the space-like separated subsystem B. Thus, even Bayes’ Theorem
cannot offer any insight into which measurement choices were selected in subsystem $B$ given only the information accessible to system $A$. As such, the law of *No Signalling* is respected, even by quantum nonlocality.

So, we live in a nonlocal world: A world in which the systems we investigate demonstrate a sensitivity to global contexts but which nonetheless give no hint of what the global context actually is to the local investigators. Indeed, the investigators must collaborate through classical communication in order for them to even conclude that nonlocality has happened, as the evidence is concealed in the *correlation* (or anticorrelation) statistics. The specific classical assumptions which must be discarded upon witnessing such nonlocal correlations are discussed in Refs. [12–16].

If one accepts that the universe is nonlocal, one might presume that all forms of nonlocality which respect the no-signalling principle should be permitted. This, however, is not the case.

A careful analysis of the mathematical formalism underpinning quantum theory reveals that there are conceivable nonlocal no-signalling statistics which are nevertheless completely inaccessible through quantum mechanics [1, 17–23]. This no-go theorem is independent of the specific physical quantum system which might be utilized, or even the dimension of the Hilbert space: quantum mechanics excludes certain no-signalling statistics. And thus, the stage is set. The world is nonlocal, but
not maximally nonlocal. One can define the set of all local statistics [24], and one can define the set of all nonlocal no-signalling statistics [8, 25], but the quantum set is intermediate.

The first chapter of this thesis is dedicated to advancing and improving our understanding of the quantum set [26–32]. Indeed, while closed-form linear inequalities are known which tightly define local and nonlocal statistical polytopes [1, 17, 19, 20, 33], no closed-form description has been developed for the quantum set. The quantum boundary is certainly nonlinear [22, 34, 35], and there is presently a conjecture that it cannot even be defined by any guaranteed-terminating algorithm [36, Conjecture (8.3.3)]. Our contributions to this effort, reviewed herein, include the development of new inequalities which inscribe the quantum set [31, 37]; more broadly, this work supplies formal justification of the method used to derive the inequalities, correcting and explaining the pioneering work of Cirel’son [18]3, which was published without proof. Entirely unpublished prior to this thesis is our discussion explaining the role of Hermitian polynomials in justifying Tsirelson’s method, as well as the explicit contrast of our analytic quantum bounds with limits inferred from the principles of

3 “Cirel’son” is the Romanization from the Cyrillic “Цирельсон” employed in articles published before 1983. “Tsirel’son” was the form used this author from 1983 through 1991. Since Tsirelson emigrated to Israel from Russia in 1991 he has used only the form “Tsirelson”, which is the form we use in this thesis except when explicitly citing an early work. See http://www.math.tau.ac.il/~tsirel/faq1.html and Mathematical Reviews’ 1982 author index, appendix C, page C1.
Information Causality [38–40], Macroscopic Locality [41–43], and Local Orthogonality [36, 44–47].
1.2 Qubits are Sufficient for Binary and Dichotomic

A nonlocality scenario is described by the number of parties who are spatially separated, the number of measurement choices accessible to each party, and the (discrete) dimension of the possible measurement outcomes associated with each measurement. The set of quantum-compatible statistics are those which can be reproduced by associating every measurement with a set of quantum projectors, assigning some global quantum state to be shared among the parties, and ensuring that order of measurements is irrelevant by imposing commutation for any projector pair belonging to two distinct parties. In practice it is conveniently to implement this commutation structure by simply assigning each party to their own Hilbert space. It is not presently known if perhaps some generality is lost by using distinct Hilbert spaces for the parties, but to date there is no evidence for such a loss of generality. This question is known as Tsirelson’s Problem, see Refs. [48–51].

So, our goal is to decide if a given statistical “point” is quantum-compatible for some nonlocality scenario. A statistical point, here, refers to a tuple of statistics defining some experimental results. This is equivalently referred to as a “behavior” [52] or a “probabilistic model” [36]. There is no simple test to verify the quantum-ness of a statistical point, although the NPA hierarchy [35, 36, 52–54] provides an asymptotically converging algorithm which becomes computationally intractable very
quickly even for the most elementary of nonlocality scenarios.

If one restricts consideration to a particular shared quantum state, then it becomes relatively straightforward to determine all the nonlocal statistics (if any) that can be accessed by measurements on that state [55–58]. One very important result, for example, is that the shared quantum state must be entangled in order to exhibit any nonlocality [28, 59–62], and every entangled state can exhibit some form of nonlocality [63–66]. The goal in this thesis, however, is not to be tied down to any particular quantum state. Rather, we seek to quantify the set of nonlocal statistics accessible through quantum measurements upon any shared entangled state.

Although much progress has been made toward this goal [26, 28–30, 67], nevertheless, we find that increasing the dimension of the Hilbert spaces unlocks monotonically more nonlocal statistics. Thus, infinite dimensionality is required for true generality. One can even infer the dimension of the Hilbert spaces from the extent of some observed nonlocality [68–72].

There is an important exception [73], which can be leveraged to great effect [31]. What Masanes [73] proves (which Cirel’son [18] presumed) is that if the parties have access to two measurement choices each (“dichotomic”), and each measurement yields one of only two possible outcomes (“binary”), then shared qubits are completely sufficient to unlock all possible quantum nonlocal statistics; Using higher dimension
Hilbert spaces would not provide any advantage. This means that no matter how multipartite the scenario, it can be studied without loss of generality by considering nothing more than shared qubits, as long as it is binary and dichotomic.

For dichotomic and binary scenarios it is possible in principle to determine if some candidate statistics are quantum compatible by assessing whether or not they can be achieved using distributed qubits. To mathematically formalize what we mean, let us introduce a conventional notation for discussing binary and dichotomic multipartite scenarios. Firstly, it is convenient to think of the outcomes as equal to $\pm 1$ so that we can easily define both marginal and correlation expectation values,

$$
\langle A_x \rangle \equiv \text{Prob}(A_x = 1) - \text{Prob}(A_x \neq 1)
$$

$$
\langle A_x B_y \rangle \equiv \text{Prob}(A_x \times B_y = 1) - \text{Prob}(A_x \times B_y \neq 1)
$$

$$
\langle A_x B_y C_z \rangle \equiv \text{Prob}(A_x \times B_y \times C_z = 1) - \text{Prob}(A_x \times B_y \times C_z \neq 1)
$$

where the subscript indexes each party’s measurement choice, $x, y, z \ldots \in \{0, 1\}$, such that the total number of independent statistical parameters in given binary and dichotomic no-signalling nonlocality scenario is equal to $3^k - 1$ where $k$ is the number of parties, consistent with more general parameter counts derived by Pironio [74, Eqn. (9)] and Brunner et al. [17, Eqn. (8)].

Suppose we have some linear function of observables, such as a Bell inequality
[8, 25, 75, 76], and we are interested in determining the maximum of this linear function consistent with multipartite quantum mechanics. Quantum generalizations of Bell inequalities are known as Tsirelson inequalities [18, 31]. A Tsirelson inequality is a general statement about quantum-compatible statistics; a Tsirelson inequality can be used to reject a statistical point. If a point lies within a given Tsirelson inequality, however, this does not mean it is necessarily quantum compatible. For a point to be quantum compatible it must be within every possible Tsirelson inequality. Thus, the task of determining such quantum inequalities is a proxy task for assessing the quantum compatibility of a given point.4

As noted by Cirel’son [18, Theorem 1], the maximum value of a quantum measurement is equal to its largest eigenvalue. Since the measurement operator is independent of the state, we can determine its maximal eigenvalue by constructing the most general quantum measurement without having to also express a general quantum state.5 Following Wolfe and Yelin [31] we note that it is efficient to use a

4 To quote Ref. [77]: “A research program of ‘characterising quantum non-locality’ has arisen with two closely related goals: to provide a method of determining whether a given experimental behaviour could have been produced by an ordinary quantum model, and to discover physical or information-theoretic principles that result in constraints on possible behaviours.” Tsirelson inequalities are a tool for achieving the former goal.

5 The state-independent approach to determining quantum maxima only applies if the target function is being maximized over all quantum compatible points. If, on the other hand, some probabilistic degrees of freedom are pre-specified, then a quantum state must be introduced to perform constrained optimization. The best known example of quantum optimization over partially specified probability distributions is Hardy’s nonlocality [78–82]. Note that qubits are still sufficient to map out all binary and dichotomic Hardy nonlocality [83–86 and 2, Sec. IV.E].
reflection symmetry to parameterize the two measurements of each party. Therefore, the general quantum measurement operator lives in the $\mathbb{C}^{2 \otimes k}$ Hilbert space, and has only one free variable per party. Our task is to determine the largest possible value this operator’s largest eigenvalue can take. The Hermiticity of the quantum measurement is a distinct advantage in this process, as we now explain.
1.3 The Advantage of Hermitian Polynomials

A Hermitian matrix has only real eigenvalues. This has an important consequence for semidefinite programming. Suppose we have a matrix where some of the matrix elements are variables, and we would like to either maximize the largest eigenvalue or minimize the smallest eigenvalue over these variables. At first, this appears to be a problem of simultaneous optimization. The problem, however, can be recast more efficiently if we are guaranteed that the eigenvalues are real.

A degree $N$ polynomial has $N$ roots, by the fundamental theorem of algebra, however multiple roots can coincide if a root is also a critical point of the polynomial. Pursuant to the intermediate value theorem, we know there must be at least one critical point between any two real roots of a polynomial, as it must have changed direction in order to intersect the x-axis again. Note that this intermediacy of a critical point generalizes even to the case of a degenerate root, in the sense that we can define the critical point to be intermediate to two of the degenerate roots, effectively sandwiching it between two real roots separated by zero value. In other
Let \( p^{(0)}(x) \) be a polynomial in \( x \), with \( p^{(q)}(x) \equiv \frac{\partial^q p^{(0)}(x)}{\partial x^q} \)

where \( \forall q \geq 0, i < j : p^{(q)}(r_i^{(q)}) = 0 \) and \( r_i^{(q)} \in \mathbb{R} \) and \( r_i^{(q)} \leq r_j^{(q)} \) \hspace{1cm} (1.2)

then \( \exists_k : \) such that \( r_i^{(0)} \leq r_k^{(1)} \leq r_j^{(0)}, \ldots \)

or, alternatively,

**LEMMA:** A polynomial with \( N \) real roots (not necessarily distinct) has at least \( N - 1 \) critical points (not necessarily distinct) \hspace{1cm} (1.3)

in the region spanned by the real roots.

If the polynomial does have complex roots, then it is possible for there to be critical points smaller than the smallest real root or larger than the largest real root, for example \( p^{(0)}(x) = x^6 - 5x^4 + 7x^2 - 1 \). When the polynomial is the characteristic polynomial of a Hermitian matrix, however, all the critical points must be within the range of the real roots; the fundamental theorem of algebra dictates that there are more roots of a polynomial than roots of its derivative. The intermediate value
theorem demands that every pair of real roots sandwich some critical point.

Let $p^{(0)}(x)$ be a Hermitian polynomial in $x$

then $\forall k : \exists i < j$ such that $r^{(0)}_i \leq r^{(1)}_k \leq r^{(0)}_j$. \hfill (1.4)

Moreover because every root of the derivative now must be real in order to be intermediate to real roots, we can inductively realize the stronger statement

Let $p^{(0)}(x)$ be a Hermitian polynomial in $x$

then $\forall q \geq 1, k : \exists i < j$ such that $r^{(q+1)}_i \leq r^{(q)}_k \leq r^{(q+1)}_j$. \hfill (1.5)

In other words,

**LEMMA:** All the roots of a Hermitian polynomial’s $N$’th derivative are real, and lie between the $(N - 1)$’th derivative’s roots. \hfill (1.6)

If we define *change* as any reversal of the sign of the polynomial or the sign of any of its derivatives, then Lemma (1.6) tells us that a Hermitian polynomial\(^6\) has no change whatsoever in the region larger than its largest root or smaller than its smallest root.

This allows us to give a somewhat unconventional definition for the largest real root or the smallest real root of a Hermitian polynomial in terms of infimum and

\(^6\) Polynomials with exclusively real roots are also known as *hyperbolic* polynomials [87, 88].
supremum:

Let \( p^{(0)}(x) \) be a Hermitian polynomial in \( x \), then

\[
\begin{align*}
    r_{\text{max}}^{(0)} &\equiv \inf_{z \in \mathbb{R}} z, \text{ such that } \forall q \geq 0 : \sgn[p^{(q)}(z)] = \sgn\left[\lim_{x \to \infty} p^{(q)}(x)\right] \\
    r_{\text{min}}^{(0)} &\equiv \sup_{z \in \mathbb{R}} z, \text{ such that } \forall q \geq 0 : \sgn[p^{(q)}(z)] = \sgn\left[\lim_{x \to -\infty} p^{(q)}(x)\right].
\end{align*}
\]  

(1.7)

The limits at \( \pm \infty \) depend only on the leading coefficient of the polynomial, so the relevant concept can be simplified.

Let \( p^{(0)}(x) \) be a degree-\( N \) Hermitian polynomial in \( x \)

with leading term \( c \times x^N \), then taking \( \tilde{p}^{(0)}(x) \equiv p^{(0)}(x)/c \),

\[
\begin{align*}
    r_{\text{max}}^{(0)} &\equiv \inf_{z \in \mathbb{R}} z, \text{ such that } \forall q \geq 0 : \tilde{p}^{(q)}(z) > 0 \\
    r_{\text{min}}^{(0)} &\equiv \sup_{z \in \mathbb{R}} z, \text{ such that } \forall q \geq 0 : (-1)^{N-q} \times \tilde{p}^{(q)}(z) > 0.
\end{align*}
\]  

(1.8)

This transforms the nature of the task of maximizing the largest eigenvalue or minimizing the smallest eigenvalue or a rank \( N \) matrix. Instead of simultaneous optimization over multiple variables, the problem is recast now into optimization over the single dummy variable of the characteristic polynomial, but subject to \( N \) “for all”-type constraints which involve the matrix’s genuine variable elements. For
Let \( p^{(0)}(x) \) be the characteristic polynomial of a Hermitian matrix

with leading term \( c \times x^N \) and with “free” variables \( \{\text{vars}\} \), then

\[
\begin{align*}
\max \left\{ r^{(0)}_{\text{max}} \right\} & \equiv \inf_{z \in \mathbb{R}} z, \text{ such that } \forall q \geq 0 \forall \{\text{vars}\} : \tilde{p}^{(q)}(z) > 0 \\
\min \left\{ r^{(0)}_{\text{max}} \right\} & \equiv \inf_{z \in \mathbb{R}} z, \text{ such that } \forall q \geq 0 \exists \{\text{vars}\} : \tilde{p}^{(q)}(z) > 0 \\
\min \left\{ r^{(0)}_{\text{min}} \right\} & \equiv \sup_{z \in \mathbb{R}} z, \text{ such that } \forall q \geq 0 \forall \{\text{vars}\} : (-1)^{N-q} \tilde{p}^{(q)}(z) > 0 \\
\max \left\{ r^{(0)}_{\text{min}} \right\} & \equiv \sup_{z \in \mathbb{R}} z, \text{ such that } \forall q \geq 0 \exists \{\text{vars}\} : (-1)^{N-q} \tilde{p}^{(q)}(z) > 0.
\end{align*}
\]

Note that the final condition of Eq. (1.9) allows one to assess if a variable Hermitian matrix can - or cannot - be made positive semidefinite. The assessment can be rephrased as a question: “Can the smallest eigenvalue be made larger than or equal to zero?” Answering such a question requires only single-variate optimization, albeit constrained by a finite set of “exists” statements regarding the matrix’s free variables.

Note that for raw semidefinite programming the Hermitian matrix must be entirely numeric aside for the free variable. Our approach, however, allows one, in principle, to consider also non-numeric parameters in the Hermitian matrix aside from the free variable. We can now analytically compute the positive semidefinite domain of these parameters. That is, the condition \( \max \left\{ r^{(0)}_{\text{min}} \right\} \geq 0 \) effectively induces a
restriction on the parameters.

We note that the Navascués-Pironio-Acín (NPA) hierarchy \[35, 52, 54\] hinges on exactly this type of assessment. Their $\Gamma$ matrix, sometimes referred to as the certificate, is a Hermitian matrix parameterized by probabilities, namely both the marginal and correlation expectation values. The $\Gamma$ matrix also has extensive free variables, presumably corresponding to the expectation values of non-projective quantum operators, which are inherently unobservable. For probabilities consistent with a quantum bipartite experiment, \textit{Gamma} must be Hermitian and positive semidefinite. We can impose hermiticity on $\Gamma$ by construction, but the real test is demanding that $\Gamma$ be positive semidefinite. Do there exist free variable capable of making \textit{Gamma} positive semidefinite given the proposed experimental probabilities? Using Eq. (1.9) we can compute the largest possible value of the smallest eigenvalue. If this value is less than zero, then the given probabilities are exposed as incompatible with the outcome of a quantum bipartite experiment.

Because Eq. (1.9) permits non-numeric parameterization, this means that in principle one can obtain a restriction on the probabilities to be consistent with quantum from $\max \{ r_{\min}^{(0)} \} \geq 0$. Indeed, the first level of the hierarchy has been already been used to define such a restriction \[35, \text{Eq. (11)}, \text{see also 42, Eq. (11), 39, Eq. (3), and 31, Eq. (9)}\]. It would be of great interest to derive the stronger
restriction corresponding to the hierarchy level $Q^{1+AB}$ which has received extensive interest recently, as it is essential in Refs. [36, 44, 77, 89]. This special level of the hierarchy is discussed at length in Sec. 3.2 of this thesis.
1.4 Derivation of Linear Quantum Bounds

We have established that the calculation of a Tsirelson inequality \([26–32]\) is equivalent to a Hermitian matrix eigenvalue maximization problem, and that moreover, for binary and dichotomic scenarios, we have complete generality by considering each party as though in possession of only a single qubit. To demonstrate the general principle, we describe the calculation of a linear quantum bound for \(k\) parties.

We begin by defining a linear measurement operator to correspond to the general Tsirelson inequality with \(3^k - 1\) terms. For compactness, we illustrate explicitly only terms appearing in the bipartite scenario, which the multipartite generalization being fairly obvious. If the inequality is defined by seeking the quantum value of \(\gamma\), where

\[
\sum_{x=0}^{1} c^A_x \langle A_x \rangle + \sum_{y=0}^{1} c^B_y \langle B_y \rangle + \ldots + \sum_{x=0}^{1} \sum_{y=0}^{1} c^{AB}_{xy} \langle A_x B_y \rangle + \ldots \leq \gamma
\]  

\((1.10)\)

then the corresponding quantum linear operator can be defined as

\[
Z \equiv \sum_{x=0}^{1} c^A_x \hat{A}_x \otimes 1 \otimes \ldots + \sum_{y=0}^{1} c^B_y \hat{B}_y \otimes \ldots + \sum_{x,y=0}^{1} c^{AB}_{xy} \hat{A}_x \otimes \hat{B}_y \otimes \ldots + \ldots
\]  

\((1.11)\)

where without loss of generality we define each party’s two operators as measurements
in the x-y plane, with the second operator being the y-reflection of the first, such that

\[
\tilde{K}_\alpha \equiv v_K \hat{\sigma}_x + (-1)^\alpha \sqrt{1 - v_K} \hat{\sigma}_y = \begin{pmatrix} 0 & e^{-i(-1)^\alpha v_K} \\ e^{i(-1)^\alpha v_K} & 0 \end{pmatrix}
\] (1.12)

where \(v_A\) and \(v_B\) are the free variables in our matrix, they are arbitrary real numbers between zero and one. The operator \(Z\) is therefore a \(2^k \times 2^k\) Hermitian matrix. By using \(\hat{\sigma}_x\) and \(\hat{\sigma}_y\) as the spanning operators for each party, we are able to construct \(Z\) such that it has exclusively zeroes along its diagonal. Let the characteristic polynomial of \(Z\) be defined as \(p_Z(m)\). The leading coefficient of this polynomial is equal to one, by virtue of the diagonal zeroes.

The Tsirelson bound \(\gamma\) is equal to the largest possible value of the largest root of \(p_Z(m)\). Using the first definition in Eq. (1.9) we can express this quite formally now as

\[
\gamma \equiv \inf_{z \in \mathbb{R}} z, \text{ such that } \forall q \in \mathbb{N}, 0 \leq q \leq 2^k - 1, \forall 0 \leq v_A, v_B \leq 1 : 0 < \left. \frac{\partial^q p_Z(m)}{\partial m^q} \right|_{m=z} (1.13)
\]

or, since the last relevant derivative is always equal to \(2^k! \times m\),

\[
\gamma = \inf_{z \geq 0} z, \text{ such that } \forall q \in \mathbb{N}, 0 \leq q \leq 2^k - 2, \forall 0 \leq v_A, v_B \leq 1 : 0 < \left. \frac{\partial^q p_Z(m)}{\partial m^q} \right|_{m=z} (1.14)
\]
From here on out let us specialize further and consider only the bipartite case of a binary and dichotomic scenario; we refer to this bipartite binary and dichotomic scenario as the (2,2,2) scenario \cite{8}, although it is also often referred to as the CHSH scenario \cite{90}. It is the most fundamental scenario in which quantum nonlocality is observed. The explicit conditions to determine any Tsirelson inequality in the (2,2,2) scenario are

\[
\gamma = \inf_{z \geq 0} z, \text{ such that } \forall 0 \leq v_A, v_B \leq 1 : 0 < \min\{p_Z(m), p_Z'(m), p_Z''(m)\}.
\]

(1.15)

We calculated various novel marginal-involving Tsirelson inequality according to this procedure in Wolfe and Yelin \cite{31}. Our results are summarized here in Table 1.1.
Table 1.1: Linear Bounds (Number-Valued and Function-Valued): Note that TB\(^{(4)}\) is the well-known Tsirelson’s bound, and QB\(^{(4)}\) is its function-valued generalization. The bounds tabulated here should be read as 
\[
\sum_{x=0}^{1} c^A_x \langle A_x \rangle + \sum_{y=0}^{1} c^B_y \langle B_y \rangle + \sum_{x=0}^{1} \sum_{y=0}^{1} c^{AB}_{xy} \langle A_x B_y \rangle \leq (?) ,
\]
where the upper bound to the inequality can be determined for each class of general probabilistic theory, making the inequality a Bell Inequality, a Tsirelson Inequality, or a convex-hull description of the No-Signalling polytope. QB\(^{(8)}\) is invoked repeatedly in this thesis to construct constraints on the quantum region of various slices of the no-signalling polytope. See Eq. (1.31) and Eq. (3.9) in particular.

<table>
<thead>
<tr>
<th>Name</th>
<th>(c^A_0)</th>
<th>(c^A_1)</th>
<th>(c^B_0)</th>
<th>(c^B_1)</th>
<th>(c^{AB}_{00})</th>
<th>(c^{AB}_{01})</th>
<th>(c^{AB}_{10})</th>
<th>(c^{AB}_{11})</th>
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<th>QM(_{\text{Max}})</th>
<th>NOSIG(_{\text{Max}})</th>
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<tbody>
<tr>
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<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>1</td>
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1.5 *Comparison of Quantum Bounds*

Let us return to our primary task of trying to characterize the genuinely quantum *elliopote* of statistical points. The fundamental (2,2,2) general No-Signalling polytope \([1, 17, 19, 20, 33, 91]\) is the convex hull of the 16 local deterministic points and the 8 PR-box maximally nonlocal points. The local hidden variable polytope is the convex hull of only the 16 local boxes. The facets of the classical polytope are the 8 Bell inequalities. Every Bell inequality is saturated by six local points. At each facet, the quantum elliptope must touch the relevant 6 local points but extend “upward”, approaching (but not reaching) the maximally nonlocal point which lies above the facet.

We would like to map a quantum curve in this region. To do so we consider a plane spanned by three boxes: The PR box, the all-one fully-deterministic box, and the “origin”. In this thesis the concept of a probabilistic box is completely equivalent to a point in statistical space, or a “behavior” \([52]\) or a “probabilistic model” \([36]\). A box is nothing more than some multipartite conditional probability distribution. The plane spanned by these three points is exactly the “slice” studied in both Refs. \([39, \text{Fig. 3}]\) and \([44, \text{Fig. 4}]\).

The PR box \([1, 17, 19, 20, 33]\) corresponds to the extremal nonlocal probability point which perfectly correlates Alice and Bob for three out of four of their possible
pairwise measurement choices, but perfectly anticorrelates them should they happen to both choose measurements with index 1. It is named after Popescu and Rohrlich \[19\], and can be expressed as

\[
P_{PR}(ab|xy) = \frac{\delta_{(a \oplus b = xy)}}{2} \tag{1.16}
\]

where the \( \oplus \) in Eq. (1.16) indicates addition module two, i.e. the bitwise XOR function.

The all-one fully-deterministic box is an extremal local point which not only saturates the conventional Bell inequality \[21\] but also has no dependence whatsoever on the choice of Alice’s or Bob’s measurement choices. We indicate such measurement-invariant local boxes, even if they are not deterministic, by two numbers

\[
P_{m,n}(ab|xy) = \left( \begin{array}{c} m, \quad a = 1 \\ 1 - m, \quad a = 0 \end{array} \right) \times \left( \begin{array}{c} n, \quad b = 1 \\ 1 - n, \quad b = 0 \end{array} \right) \tag{1.17}
\]

noting that special cases include the fully-deterministic all-one box

\[
\text{All-one box: } P_{FD} \equiv P_{1,1}(ab|xy) = \begin{cases} 
1, & a = b = 1 \\
0, & \text{otherwise}
\end{cases} \tag{1.18}
\]
the semi-deterministic Bob-random box

\[
P_{SD} \equiv P_{1, \frac{1}{2}}(ab|xy) = \begin{cases} 
1/2, & a = 1 \\
0, & a = 0 
\end{cases} 
\] (1.19)

and the maximally random “white noise” box

\[
P_{\emptyset} \equiv P_{\frac{1}{2}, \frac{1}{2}}(ab|xy) = \frac{1}{4} 
\] (1.20)

The “origin” in statistical space refers to precisely the maximally random box, with no positive or negative bias along any expectation value, joint or marginal. We denote it here as \(P_{\emptyset}\), as per Eq. (1.20).

We introduce a mixed box spanned by \(P_{PR}\), \(P_{FD}\), and \(P_{\emptyset}\) such that

\[
P_{\xi\gamma}(ab|xy) = \xi P_{PR}(ab|xy) + \gamma P_{FD}(ab|xy) + (1 - \gamma - \xi) P_{\emptyset}(ab|xy) 
\] (1.21)

and proceed to assess the constraints on \(\gamma\) and \(\xi\) implied by presuming that the experimental outcomes are mediated through fundamentally quantum mechanisms.
The box $P_{\xi\gamma}$ defines the 2-dimensional statistical region

$$\langle A_0 \rangle = \langle A_1 \rangle = \langle B_0 \rangle = \langle B_1 \rangle = \gamma,$$

$$\langle A_0 B_0 \rangle = \langle A_1 B_0 \rangle = \langle A_0 B_1 \rangle = \langle A_1 B_1 \rangle = \gamma + \xi, \quad \langle A_1 B_1 \rangle = \gamma - \xi. \quad (1.22)$$

A well-known but relatively weak quantum bound is Uffink’s bound [22, 92], which also corresponds to the principle of Information Causality [38, 39].

$$((\langle A_0 B_0 \rangle - \langle A_1 B_1 \rangle)^2 + (\langle A_0 B_1 \rangle + \langle A_1 B_0 \rangle)^2 \leq 4 \quad (1.23)$$

For $P_{\xi\gamma}$, Uffink’s bound yields the restriction

$$\xi \leq \frac{\sqrt{2} - \gamma^2 - \gamma}{2} \quad (1.24)$$

A much stronger quantum bound is that corresponding to the first level of the NPA hierarchy [35, 52], which also corresponds to the principle of Macroscopic Locality [41–43]. It is the most commonly used bound in modern quantum foundations comparisons, because it nicely balances explicit analytical accessibility with strong
restrictive power. We denote this criterion by NPA$^1$, and it states that

$$\sum_{x,y=0}^{1} (-1)^{xy} \sin^{-1} \left( \frac{\langle A_x B_y \rangle - \langle A_x \rangle \langle B_y \rangle}{\sqrt{(1 - \langle A_x \rangle^2)(1 - \langle B_y \rangle^2)}} \right) \leq \pi$$

(1.25)

For $P_{\xi\gamma}$ the NPA$^1$ criterion it takes the explicit form

$$\sum_{x,y=0}^{1} \sin^{-1} \left( \frac{(-1)^{xy} \gamma (\gamma - 1) - \xi}{\gamma^2 - 1} \right) \leq \pi$$

(1.26)

One can invoke the identity

$$\sin \left( n \sin^{-1} (z) \right) = \sum_{k=0}^{\infty} \frac{nz^{2k+1}(2k+1+n)!!(2k-1-n)!!}{(2k+1)!(1+n)!!(-1-n)!!}$$

(1.27)

which, due to the $2k - 1 - n$ double factorial, terminates cleanly for odd $n$. Using $n = 3$ to consolidate like terms from the sum in Eq. (1.26) we obtain

$$\frac{(\gamma(\gamma - 1) - \xi)}{\gamma^2 - 1} \left( 4 \left( \frac{\gamma(\gamma - 1) - \xi}{\gamma^2 - 1} \right)^2 - 3 \right) \leq \frac{(\gamma - 1)\gamma + \xi}{\gamma^2 - 1}$$

(1.28)
which in terms of a bound on $\xi$ equivalent to

$$
\xi \leq \gamma (\gamma - 1) + \frac{(\gamma^2 - 1)^2}{h(\gamma)} + \frac{h(\gamma)}{6} \quad \text{with}
$$

$$
h(\gamma) \equiv \sqrt[3]{6(1 - \gamma)^3(\gamma + 1)^2 \left( 9\gamma + \sqrt{3\gamma(25\gamma - 4) - 6} \right)}
$$

(1.29)

We also would like to consider $QB_{3}^{(8)}$ from our earlier work seeking out new Tsirelson inequalities [31], reproduced here in Table 1.1. We specifically select $QB_{3}^{(8)}$ in order to illustrate its superior restrictive power relative to NPA on this slice of the no signalling polytope. It reads

$$
\forall c^2 \leq 1:
\begin{align*}
\langle A_0 \rangle + \langle A_1 \rangle - \langle B_0 \rangle - \langle A_1 B_1 \rangle &\leq \frac{c^2 - \sqrt{(3c^2 - 4)(c^2 - 2)}}{c^2 - 1} \\
+ \langle A_0 B_0 \rangle + \langle A_1 B_0 \rangle + \langle A_0 B_1 \rangle
\end{align*}
$$

(1.30)

For $P_{\xi\gamma}$ the left hand side of $QB_{3}^{(8)}$ is $(2 + c)\gamma + 4\xi$ such that we are effectively bounding $\xi$ by

$$
\xi \leq \frac{\min_{c^2 \leq 1}\left\{ \frac{c^2 - \sqrt{(3c^2 - 4)(c^2 - 2)}}{c^2 - 1} - c\gamma \right\} - 2\gamma}{4}
$$

(1.31)

or, in Mathematica, $\xi \leq \min\left\{ \frac{c^2 - \sqrt{(3c^2 - 4)(c^2 - 2)}}{c^2 - 1} - c\gamma, 0 < c < 1 \right\}$.

which resolves to the explicit envelope defining the entire set of linear bounds, namely
via the minimal root of an order 8 (non-Hermitian) polynomial,

\[ \xi \leq \frac{\lambda_{\min}(\gamma) - 2\gamma}{4} , \quad \text{where} \]

\[ \lambda_{\min}(\gamma) \equiv \min_{\lambda_{1} \cdots \lambda_{8}} \lambda \quad \text{such that} \]

\[ 3 \left( \lambda^{2} - 8 \right) \left( \lambda^{2} + 2\lambda - 2 \right)^{3} + \]

\[ 8\gamma^{8} - 2\gamma^{6} \left( 13\lambda^{2} - 56\lambda + 220 \right) + \gamma^{4} \left( 31\lambda^{4} - 104\lambda^{3} - 470\lambda^{2} + 960\lambda + 4080 \right) \]

\[ - \gamma^{2} \left( 16\lambda^{6} + 10\lambda^{5} - 449\lambda^{4} - 1060\lambda^{3} + 1360\lambda^{2} + 7264\lambda + 8032 \right) = 0 \]

We additionally consider a quantum bound due to Local Orthogonality [36, 44–47]. In particular, we invoke the 10-term LO^2 inequality \(^7\)

\[ P(1111|1100) + P(1110|1100) + P(1101|1001) + P(1100|0001) \]

\[ + P(1011|1101) + P(1001|1001) + P(0111|1100) \]

\[ + P(0011|1111) + P(0010|0111) + P(0000|1010) \leq 1 \]

where the four-partite boxes in Eq. (1.33) are to be understood as a wiring of two bipartite boxes [47], such that

\[ P(abcd|xyzw) = P(ab|xy) \times P(cd|zw) . \]

\(^7\)This clique was conveyed in private communication from the authors of Refs. [44].
If both the bipartite boxes that comprise \( P(abcd|xyzw) \) are identical copies of \( P_{\xi\gamma} \) then

\[
\frac{\xi (3\xi - 2(\gamma - 1)) - 3(\gamma - 2)\gamma + 5}{8} \leq 1
\]  
(1.35)

or, as a bound on \( \xi \):

\[
\xi \leq \frac{\sqrt{10} - 1}{3} (1 - \gamma)
\]  
(1.36)

which is precisely what is plotted in Figure 4 of Ref. [44].

We know that the quantum elliptope includes the deterministic box as well as the so-called Tsirelson box. By convexity, all points below the line connected these two points must be with the quantum elliptope. We therefore are particularly interested in the quantum region which lies outside this plane, if any. This line corresponds to boundary

\[
\xi \leq \frac{1}{\sqrt{2}} (1 - \gamma)
\]  
(1.37)

which we subtract from all of the relevant nontrivial quantum bounds to obtain the plot of Fig. 1.1.

To illustrate that even all these quantum bounds are still inadequate, we have numerically calculated the maximum possible \( \xi \) permitted by the \( 1 + AB \) level of the NPA hierarchy, \( NPA^{1+AB} \). The \( NPA^1 \) criterion is a relaxation of \( NPA^{1+AB} \), which
itself is a relaxation of the genuine bipartite quantum boundary. This is discussed further in Sec. 3.2 of this thesis.

Fig. 1.1: (Color online) A comparison of various quantum bounds and their implications for the quantum elliptope. Prior to the determination of the LO\(^2\) bounds first visualized in Ref. [44] in 2013, our own \(\text{QB}_3^{(8)}\) was the best-known analytic boundary in this slice of the no-signalling polytope. The grey region bounded numerically by \(\text{NPA}^{1+AB}\) corresponds to quantum contextuality, which is known to be a relaxation to quantum nonlocality even for the (2,2,2) scenario [89]. The black dots are the numeric points for which we calculated the maximal \(\xi\) per \(\text{NPA}^{1+AB}\).

We note here that on this slice of the polytope the condition of \(\text{QB}_3^{(8)}\) is grossly inadequate at characterizing the true quantum elliptope, as evidenced by both \(\text{LO}^2\) as well as \(\text{NPA}^{1+AB}\). This inadequacy was not known at the time Ref. [31] was published.
In that reference a volume analysis was presented in an effort to compare the relative sizes of the statistical region consistent with local, quantum, and no-signalling models. The assumption was that the genuine quantum boundary was well approximated by the collection of linear quantum bounds dominated by $\text{QB}^{(8)}_3$. This assumption was in retrospect not valid, and therefore we do not reproduce the results of that volume analysis in this thesis.

We conclude this chapter by noting that the study of characterizing quantum nonlocality has become an entire burgeoning field in quantum information theory. Quantum nonlocality has enjoyed a meteoric rise to prominence, as indicated by this chapter’s opening epigraphs, and is presently at the forefront of many current active research areas. Nonlocality is the device-independent resource which powers many novel applications in quantum information theory, such as the device-independent variants of quantum cryptography [61, 91, 93–97] and state tomography [98, 99], among other extraordinary uses.
Entanglement

Entanglement is a trick that quantum magicians use to produce phenomena that cannot be imitated by classical magicians. 

\footnote{Asher Peres
Quoted by Dagmar Bruß, providing a favored definition of entanglement Bruß [100].}
2.1 Entanglement as a Resource

Entanglement is well-understood to be the fundamental resource requisite for quantum nonlocality. While there are possible indications that entanglement is not fundamental for quantum computational speedup [101–105], and that the amount of entanglement in a system might not even directly translate into the strength of nonlocality a system can exhibit [106–114], there is no doubt that entanglement represents a firm qualitative demarcation in the nature of correlations that a quantum system can be utilized to generate. Firstly, separable states give rise only to classical correlations [64]. The converse, i.e. the statement that every entangled state violates a Bell inequality, is known as Gisin’s Theorem [115–118]. Gisin’s theorem was proven true for all pure states in 2012 [116], two decades after its conjecture. Long believed to hold for mixed states as well [23, 119], the general proof extending Gisin’s theorem to mixed states appears to have been established very recently as well, albeit with some qualifications [66, 117, 120]. Thus, determining if a state is separable or entangled is fundamentally critical to our understanding of quantum nonlocality.

There is extensive interest in schema for generating entanglement [121–125] due to the plethora of nonclassical tasks which entanglement makes possible. Examples include Quantum Key Distribution [61, 93, 95, 96, 126], Quantum Computation [102, 127–132], and Precision Measurement [133–139]. Various measures of entanglement
have been developed to quantify the resource value of entangled systems for the different tasks. These various entanglement measures reveal the presence of different classes of entanglement [100, 155–159], but they do not, however, tightly characterize the set of entangled states. Entanglement measures are effectively necessary separability criteria, in that for a quantum state to be separable it must have a measure of zero in the relevant entanglement metric, but they are not sufficient. A mixed quantum state may yield zero on some entanglement measure and yet nevertheless not be a separable state. All readily-evaluatable separability criteria are of the necessary-but-not-sufficient variety [65, 160–163]. Such is the essential nature of entanglement witnesses [64, 106, 164]. It is worth noting that an asymptotically sufficient separability criterion has been developed [165, 166] in a manner that parallels the semidefinite hierarchy for characterizing quantum correlations [54].

To certify that a system is separable, one must show that the density matrix can be decomposed as a convex mixture of separable states. There is no general rule for how to go about seeking such a decomposition, even if one is convinced that such a decomposition exists. Kraus et al. [167], Karnas and Lewenstein [168] describe a limited algorithm to certify biseparability [169], but these presume that the decomposition can be represented as a finite mixture, which is not always the case [170]. Korbicz et al. [171] delineate an in-principle approach suitable for symmetric
states, but it is not amenable for practical application. Refs. [172, 173] provide sufficient criteria for $\mathbb{C}^d \otimes \mathbb{C}^d$ mixed states based on mutually unbiased bases [174–176] and symmetric informationally-complete POVMs [177, 178], but these criteria are limited to system with a particular bipartite structure. There is, therefore, an resolved desire for a general method capable of practically certifying separability; such a method is needed, for example, to delineate truly entangled phenomenon as distinct from merely superficially cooperative-behaving systems [170, 179–181].

As established in Sec. 1.2, for multipartite scenarios which are binary and dichotomic, the full range of quantum nonlocality can be achieved using qubits [73]. Additionally, we note that no single party holds a privileged position with regards to nonlocality. That is, the quantum nonlocality elliptope in statistical space must be invariant with respect to permutations of the parties, maintaining this natural symmetry of the local and no-signalling polytopes [8, 21, 28, 30, 33].

This suggests that we should prioritize permutationally-invariant multi-qubit states in our study of entanglement and separability certification, because the quantum statistical boundary for binary and dichotomic scenarios corresponds physically to measurements on such quantum states. This direct correspondence between nonlocality and entanglement only holds for permutationally-invariant multi-qubit states. See Refs. [28, 55, 182] for a survey of recent works advancing the quantification
of nonlocality in multi-qubit states. It is especially interesting that all entangled permutationally-invariant multi-qubit states are necessarily genuinely multipartite entangled \cite{163}, but it should also be noted that genuine multipartite entanglement does not necessarily imply genuinely multipartite nonlocality \cite{114}.

The second chapter of this thesis is dedicated to studying entanglement in permutationally-invariant multi-qubit states. Our contributions include the development of an explicit method for certifying separability for diagonally-symmetric multi-qubit mixed states \cite{170}, the counter-intuitive finding that uniquely-quantum effects in Dicke model superradiance take place absent any entanglement \cite{170}, and the analytic validation of driven Dicke model superradiance as a scheme for generating spin-squeezed entangled states \cite{183}. Entirely unpublished prior to this thesis is our idea to utilize the structure of independently evolving systems in order to develop sufficient separability criteria for cooperatively evolving systems. The analysis of driven superradiance in terms of normalized Dicke states, and the closed form expressions for the $N$-particle SDS jacobian and diagonally symmetric PPT conditions, are also unique to this thesis.
2.2 Separability of Diagonally Symmetric States

Any permutationally invariant pure state can be expressed in terms of symmetric\(^2\) Dicke basis states. The Dicke states are the superposition of equal-energy states; each is essentially a normalized sum-over-all-permutations of some (separable) computational-basis state, such as

\[
|D_1^4\rangle = \frac{|0001\rangle + |0010\rangle + |0100\rangle + |1000\rangle}{\sqrt{4}}.
\]

(2.1)

or in general

\[
|D_N^{n_1}\rangle = \sqrt{\frac{n_0!n_1!}{N!}} \sum_{\text{perms.}} \sum_{\{\{0,1\}\}} \frac{|0...0, 1...1\rangle}{n_0, n_1}
\]

(2.2)

where \(n_i\) is number of instances of \(|i\rangle\) in the computational-basis state being permuted, and we have introduced the shorthand \(n_0 = N - n_1\), where \(N\) the total number of qubits. Indexing Dicke states by \(N\) and \(n_1\) is the convention used in Refs. [182, 184], whereas in our earlier work in Ref. [170] we used \(n_0\) and \(n_1\).

We begin by studying the separability properties of mixed states which are diagonal in the Dicke basis. To indicate such states we use the subscript GDS for

\(^2\) Our use of the term “symmetric” here is equivalent to permutationally invariant, which is to say, a bosonic symmetry of indistinguishable particles. This is in contrast to the more reserved use of the terminology “symmetric” such as is considered by Tóth and Gühne [148].
“General Diagonally Symmetric”. The general mixed state which is diagonal in the Dicke basis can be parameterized as

$$\rho_{GDS} = \sum_{n_{1}=0}^{N} \chi_{n_{1}} |D_{n_{1}}^{N}\rangle \langle D_{n_{1}}^{N}|$$  \hspace{1cm} (2.3)

where the $\chi_{n_{1}}$ represent the eigenvalues in the eigendecomposition of $\rho_{GDS}$, which, in the convention of quantum optics, we refer to as the populations of $\rho_{GDS}$.

Certifying separability amounts proving the existence of a decomposition of target mixed state into some convex combination of separable states; determining the existence of such a decomposition is “hard”. We show that it is effective to instead ask if the target mixed state “fits” some preconstructed separable form. For GDS states, we take as our preconstructed separable form a given mixture of symmetric separable states. First, the fully-general separable multi-qubit pure state which is permutationally invariant is just the $N$-fold tensor product of a fully-general two-level pure state, that is,

$$|\psi[y,\phi]\rangle = \left(\sqrt{y}|0\rangle + \sqrt{1-y}e^{i\phi}|1\rangle\right)^{\otimes N} \text{ or, in operator form } \rho^{N} \equiv |\psi\rangle \langle \psi|$$  \hspace{1cm} (2.4)

ie. $\rho^{N}[y,\phi] = \left(y|0\rangle\langle 0| + (1-y)|1\rangle\langle 1| + \sqrt{y(1-y)} \left(e^{-i\phi}|0\rangle\langle 1| + e^{i\phi}|1\rangle\langle 0|\right)\right)^{\otimes N}$

which we would like to express more explicitly. By multinomial expansion we can
expand $\rho^N[y, \phi]$ in terms of a sum over four exponents, $\gamma_{00}, \gamma_{10}, \gamma_{01}, \gamma_{11}$, which are to be understood as ranging over nonnegative integers $\gamma \in \mathbb{Z}^+$ in such a manner that the sum of the exponents total $N$, $\gamma_{00} + \gamma_{10} + \gamma_{01} + \gamma_{11} = N$. In this manner we are also taking tensor-exponents of each of the four operator-basis product states, summing over all permutations of the operators as well. That is, $\rho^N[y, \phi] =$

$$= \sum_{\{all \gamma\}} y^{(\gamma_{00}+\gamma_{01}/2)} (1-y)^{(\gamma_{11}+\gamma_{10}/2)} e^{i\phi(\gamma_{10}-\gamma_{01})} \sum_{\text{operator permutations}} \rho \left[ \begin{array}{cc} \gamma_{00} & \gamma_{01} \\ \gamma_{10} & \gamma_{11} \end{array} \right]$$

where $\rho \left[ \begin{array}{cc} \gamma_{00} & \gamma_{01} \\ \gamma_{10} & \gamma_{11} \end{array} \right] \equiv (|0\rangle\langle0|)^{\otimes \gamma_{00}} (|1\rangle\langle1|)^{\otimes \gamma_{11}} (|0\rangle\langle1|)^{\otimes \gamma_{01}} (|1\rangle\langle0|)^{\otimes \gamma_{11}}$ (2.5)

is the natural generalization of computational basis states to product states. Note that the sum over operator permutations is intentionally not normalized as each permutation of each has equal weight in the expansion of $\rho^1[y, \phi] \otimes^N$.

We elect to consider only such states which are mixed uniformly over all $\phi$, namely $\rho^N[y] \equiv (2\pi)^{-1} \int_0^{2\pi} \rho^N[y, \phi] \ d\phi$. While this does induce a loss of generality, this is desirable in order that $\rho^N[y] \in \mathcal{Q}_{\text{GDS}}$. Note that

$$\int_0^{2\pi} e^{i\phi(\gamma_{10}-\gamma_{01})} \ d\phi = \begin{cases} 1, & \gamma_{10} = \gamma_{01} \\ 0, & \gamma_{10} \neq \gamma_{01} \end{cases} \quad (2.6)$$

which allows us to perform a four-to-three change-of-variable, such that $\gamma_{10} = \gamma_{01} \rightarrow \kappa$, $\gamma_{00} \rightarrow n_0 - \kappa$, $\gamma_{11} \rightarrow n_1 - \kappa$. Note that in these three new variables, the earlier
implicit condition $\gamma_{00} + \gamma_{10} + \gamma_{01} + \gamma_{11} = N$ is *automatically satisfied* once we define $n_0 = N - n_1$, but to preserve the positivity of both $\gamma_{00}$ and $\gamma_{11}$ we must be careful to upper bound $\kappa \leq \min[n_0, n_1]$. Thus the uniform mixing over all phases results in

$$
\rho^N [y] = \sum_{n_1=0}^{N} \min[n_0, n_1] \sum_{\kappa=0}^{y^{n_0}(1-y)^{n_1}} \sum_{\text{operator permutations}} \rho \left[ \frac{(n_0 - \kappa)}{\kappa} \frac{\kappa}{(n_1 - \kappa)} \right] \quad (2.7)
$$

To translate explicitly into the Dicke basis states we rearrange the order of summation and make use of a binomial argument, namely

$$
\sum_{\text{perms.}} \frac{1}{4 \times \{j\langle k|\}} \sum_{\kappa} \rho \left[ \frac{(n_0 - \kappa)}{\kappa} \frac{\kappa}{(n_1 - \kappa)} \right] = \left( \sum_{\text{perms.}} \frac{1}{\{0\langle 1|\}} \frac{1}{\{0\langle 1|\}} \right) \left( \sum_{\text{perms.}} \frac{1}{\{0\langle 1|\}} \frac{1}{\{0\langle 1|\}} \right) \quad (2.8)
$$

The left hand side of Eq. (2.8) is a double sum, over permutations of the four operators as well as over all possible partition schemes indexed by $k$. This is equivalent to the right hand side of Eq. (2.8), namely taking the product of unpaired permutation summations. This counting scheme follows from

$$
\sum_{\kappa} \frac{N!}{\kappa! (n_0 - \kappa)! (n_1 - \kappa)! \kappa!} = \left( \frac{N!}{n_0! n_1!} \right)^2.
$$

To clarify what is meant in Eq. (2.8) let us consider the explicit example where $n_0 = 3$ and $n_1 = 1$ as in Eq. (2.1). We’ll start with the right hand side of Eq. (2.1)
and work backward to the left hand side.

\[
\left( \sum_{\text{perms.}} |0\ldots01\ldots1\rangle \right) \left( \sum_{\text{perms.}} \langle 0\ldots01\ldots1| \right) \\
= \left( |0001\rangle + |0010\rangle + |0100\rangle + |1000\rangle \right) \left( \langle 0001| + \langle 0010| + \langle 0100| + \langle 1000| \right) \\
= \left( |00\ldots00| + |00\ldots10| + |01\ldots00| + |01\ldots10| \right) \\
+ \left( |00\ldots01| + |00\ldots11| + |01\ldots01| + |01\ldots11| \right) \\
+ \left( |01\ldots00| + |01\ldots10| + |10\ldots00| + |10\ldots10| \right) \\
+ \left( |10\ldots01| + |10\ldots11| \right)
\]

\[
= \sum_{\text{perms.}} \rho \begin{bmatrix} 3 & 0 \\ 0 & 1 \end{bmatrix} + \sum_{\text{perms.}} \rho \begin{bmatrix} 2 & 1 \\ 1 & 0 \end{bmatrix} = \sum_{\text{perms.}} \sum_{\kappa=0}^{\min[3,1]} \rho \begin{bmatrix} (3 - \kappa) & \kappa \\ \kappa & (1 - \kappa) \end{bmatrix}
\]

Of course by inspection of Eq. (2.2) it is clear that
\[
\sum_{\text{perms.}} |0\ldots01\ldots1\rangle \langle 0\ldots01\ldots1| = \sqrt{\frac{N!}{n_0!n_1!}} |D_{n_1}^N \rangle \langle D_{n_1}^N|
\]
such that we can conveniently now express Eq. (2.7) in a manner which makes it clear that \( \rho^N \langle y | \in \mathcal{E}_{\text{GDS}}, \) namely

\[
\rho^N \langle y | = \sum_{n_1=0}^{N} \frac{N!}{n_0!n_1!} y^{n_0} (1 - y)^{n_1} |D_{n_1}^N \rangle \langle D_{n_1}^N|.
\]  

(2.10)

This suggests a suitably-generic parameterization of separable diagonally symmetric
states, namely

\[ \rho_{SDS} = \sum_{n_0=0}^{N} \sum_{j=1}^{j_{\text{max}}} \frac{N! x_j y_j^{n_0} (1 - y_j)^{n_1}}{n_0! n_1!} |D_{n_1}^N\rangle \langle D_{n_1}^N| \]  

(2.11)

which implies a sufficient criterion for separability of diagonally symmetric states:

Does the state \( \rho_{GDS} \) fit for the form of \( \rho_{GDS} \)? Formally, a comparison of Eqs. (2.3) and (2.11) indicates that

\[ \rho_{GDS} \in \mathcal{Q}_{SDS} \text{ iff } \exists x_1 \ldots x_{j_{\text{max}}} , y_1 \ldots y_{j_{\text{max}}} \]

satisfying \( \forall n_1 \chi_{n_1} = N! \sum_{j=1}^{j_{\text{max}}} \frac{x_j y_j^{n_0} (1 - y_j)^{n_1}}{n_0! n_1!} \)  

(2.12)

such that \( \forall j : 0 \leq x_j , y_j \leq 1 \).
2.3 Volume Analysis of Separability Criteria

It is possible to show that not only is the separability criterion implied by conditions (2.12) sufficient to certify separability, but that it is furthermore also apparently necessary! That is to say, we can show that every fully separable symmetric $N$-qubit state is of the form of $\rho_{\text{GDS}}$. We establish the universality of $\rho_{\text{GDS}}$ by showing that the volume of states which it parameterizes is equal to the volume of diagonally-symmetric states which satisfy the necessary separability criterion of positivity under partial transpose (PPT) [160, 161].

The property of PPT is generally necessary but insufficient for separability [65, 106, 119, 169], although for symmetric states it known to be is sufficient for $N = 2, 3$, but still insufficient for $N \geq 4$ [150, 162, 163, 179]. Our finding of $\text{SDSVol}_N = P\text{PTDSVol}_N$ implies that the PPT criterion apparently is sufficient for $N \geq 4$ for diagonally symmetric states. We provide a complete numeric proof for $N = 4$, and additional numeric evidence to suggest that the correspondence is not broken for larger $N$.

To define a volume of a set of quantum mixes states we must establish a metric on the spaces of density matrices; the metric can be arbitrary but must be consistent. We choose the populations of $\rho_{\text{GDS}}$ as our integration coordinates, ie. $\chi_0 \ldots \chi_N$. Note that this is in contrast to the more conventional metrics established by Życzkowski
et al. [185–189].

When computing the volume of SDSVol$_N$ it is natural to integrate not using the populations $\chi_{n1}$ but rather the variables $x_j, y_j$. This requires that we introduce not only a volume element to compensate for the change of variable of the integration basis, but that we furthermore take care to establish a one-to-one mapping between $x_1 \ldots x_{j_{\text{max}}}, y_1 \ldots y_{j_{\text{max}}}$ and $\chi_0 \ldots \chi_N$.

This implies that $j_{\text{max}}$ must be chosen such that the central system of equations comprising criterion (2.12) should be well behaved, i.e. that there should be exactly $N + 1$ variables $x_j, y_j$ total appearing in the $N + 1$ equations. Considering that $x_j$ and $y_j$ always come in pairs, this poses an obstacle for those instances when $N + 1$ is odd. Our solution is to take $j_{\text{max}} = \lceil (N + 1)/2 \rceil$ and to mediate the extraneous variable by manually adjusting $y_{(N+2)/2} = 0$. This ansatz is validated when we find that SDSVol$_N = \text{PPTDSVol}_N$ still holds when $N$ is an even number. We summarize by defining

$$j_{x_{\text{max}}} = \left\lceil \frac{N + 1}{2} \right\rceil$$
$$j_{y_{\text{max}}} = \left\lfloor \frac{N + 1}{2} \right\rfloor$$

(2.13)

$$y_{j_{x_{\text{max}}}} = 0 \text{ if } j_{x_{\text{max}}} \neq j_{y_{\text{max}}}$$

We must also break the symmetry of the decomposition inherent to conditions
(2.12), in the sense that as it stands, any solution for \( y_1 \) in terms of \( \chi_0 \ldots \chi_N \) can be exchanged with a solution for \( y_2 \) by also exchanging \( x_1 \) and \( x_2 \), implying a degeneracy of solutions. We can solve this by imposing the ordering \( \forall 1 \leq i < j \leq j_{\text{max}} \ y_i \geq y_j \) which fits nicely with the sometimes-relevant constraint \( y_{(N+2)/2} = 0 \). In practice it is easiest to integrate without restriction, and then to compensate by dividing by \( jy_{\text{max}}! \) to account for the degeneracy.

The volume element we mentioned is of course the absolute value of the determinant of Jacobian matrix for the change-of-variable. The \( N + 1 \) columns of this matrix corresponds to the populations \( \chi_{n_1} \). The rows of this matrix corresponds to differentiation of the populations with respect to the \( N + 1 \) new variables \( x_j, y_1 \ldots y_j \), as the populations are with regards to these new variable in Eq. (2.12). One can show that this determinant has the form

\[
\text{Jac}_N = Z_N \prod_{k=0}^{\lfloor N+1/2 \rfloor} x_k \prod_{j=0, j \neq k}^{\lfloor N+1/2 \rfloor} (y_j - y_k)^2 \quad \text{where} \quad Z_N \equiv \prod_{n_1=0}^{N} \frac{N!}{n_0!n_1!} \quad (2.14)
\]

which is valuable as it allows us to readily factor the integral into one integral over the \( x_j \) and another integral over the \( y_j \). Note that the index of \( x \) in Eq. (2.14) goes up only to \( jy_{\text{max}} \), whereas the index of \( y \) goes up to \( jx_{\text{max}} \). This is a result of \( x_{jx_{\text{max}}} \) not appearing at all in the Jacobian if \( jx_{\text{max}} \neq jy_{\text{max}} \). Recall also that when \( N \) is
even then $y_{jx_{\text{max}}} = 0$ as per Eq. (2.13).

The integral over the $x_j$ is not entirely trivial, as we must include some condition of normalization when considering a volume of states, $1 = \text{Tr} [\rho_{\text{GDS}}] = \sum_{n_1=0}^{N} \chi_{n_1} = \sum_{j=1}^{j_{\text{max}}} x_j$. We could define integration limits that are inherently normalized, but it is easier to simply introduce a Dirac delta function in the integrand to enforce normalization. One can verify that

$$
\int_0^1 \cdots \int_0^1 \delta \left( 1 - \sum_{j=1}^{\left\lfloor \frac{N+1}{2} \right\rfloor} x_j \right) \prod_{j=0}^{\left\lfloor \frac{N+1}{2} \right\rfloor} x_k \, dx_1 \cdots dx_{\left\lfloor \frac{N+1}{2} \right\rfloor} = \frac{1}{N!}
$$

(2.15)

which means that

$$
\text{SDSVol}_N = \frac{Z_N}{\left\lfloor \frac{N+1}{2} \right\rfloor! N!} \int_0^1 \cdots \int_0^1 \prod_{j=0, j \neq k}^{\left\lfloor \frac{N+1}{2} \right\rfloor} (y_j - y_k)^2 \, dy_1 \cdots dy_{\left\lfloor \frac{N+1}{2} \right\rfloor}
$$

$$
= \prod_{z=1}^{N} z^{(z-1)} \left( \frac{z-1}{(2z-1)!} \right) \left( \frac{z}{(2z-1)!} \right) \quad (\text{proof not shown})
$$

(2.16)

which we explicitly give for $N = 4$, namely

$$
\text{SDSVol}_4 = \frac{Z_4}{\left\lfloor \frac{4+1}{2} \right\rfloor! 4!} \int_0^1 \int_0^1 y_1^2 y_2^2 (y_1 - y_2)^4 \, dy_1 dy_2
$$

$$
= \frac{96}{2 \times 24} \times 525^{-1} \approx (3809.5) \times 10^{-6}.
$$

(2.17)

To calculate the volume of the positive-under-partial-transpositions diagonally
symmetric states, PPTDSVol\(_N\), we perform the integration directly in the basis of the populations, so we must introduce indicator functions to enforce that we only count PPT states. Here the PPT conditions mean that all eigenvalues are nonnegative for all bipartitions of the qubits for partial transposition. The permutation symmetry of \(\rho_{GDS}\) means we need only consider \([N/2]\) bipartitions: partial transposition of the first qubit \(\rho^{\text{PT}1;N}\) or of the first two qubits \(\rho^{\text{PT}2;N-1}\), etc, akin to the considerations in Refs. [162, 190].

For diagonally symmetric states the positivity conditions associated with each bipartition has a clean form, namely

\[
\rho^{\text{PT}q;N-q} \geq 0 \quad \text{iff} \quad \forall_{0 \leq m \leq N-2q} : \text{Det}[\tilde{\rho}_m^q] \geq 0
\]  

(2.18)

where \(\tilde{\rho}_m^q\) is a \((q+1) \times (q+1)\) matrix with elements

\[
\tilde{\rho}_m^q[x,y] \equiv \chi_{m+x+y} \frac{\binom{q}{x} \binom{q}{y} \binom{N-q}{m+x} \binom{N-q}{m+y}}{\binom{N}{m+x+y}^2}
\]  

(2.19)

which implies the special case PPT condition for one-qubit bipartition as a result of
the closed-form expression

\[
\text{Det}\left[\hat{\rho}_m^{-1}\right] = \frac{(m + 2)(N - m)\chi_m\chi_{m+2} - (m + 1)(N - m - 1)\chi_{m+1}}{N^2}
\]  

(2.20)

corresponding to the PPT condition used by Quesada and Sanpera [184, Eq. (14)].

Per Eq. (2.18), we set up our integral for the volume of PPT diagonally-symmetric states in terms of Heaviside Theta functions, such that

\[
PPTDSVol_N = \int_0^1 \cdots \int_0^1 \delta \left( 1 - \sum_{n_1=0}^{N} \chi_{n_1} \right) \prod_{q=1}^{\lfloor N/2 \rfloor - 2q} \prod_{m=0}^{N-2q} \Theta \left( \text{Det}\left[\hat{\rho}_m^q\right] \right) \, d\chi_0 \cdots d\chi_N
\]

(2.21)

which is difficult to evaluate in general. For \(N = 4\) we have the explicit integral

\[
PPTDSVol_4 = \int_{0}^{1} \int_{0}^{1} \int_{0}^{1} \int_{0}^{1} \Theta \left( 8\chi_0\chi_2 - 3\chi_1^2 \right) \Theta \left( 9\chi_1\chi_3 - 4\chi_2^2 \right) \Theta \left( 8\chi_2\chi_4 - 3\chi_3^2 \right) \\
\times \Theta \left( 9(\chi_1\chi_3 + 8\chi_0\chi_4)\chi_2 - 2\chi_2^3 - 27(\chi_4\chi_1^2 + \chi_0\chi_3^2) \right) \\
\times \delta(1 - \chi_0 - \chi_1 - \chi_2 - \chi_3 - \chi_4) \, d\chi_0 d\chi_1 d\chi_2 d\chi_3 d\chi_4
\]

(2.22)

\[= (3808 \pm 2) \times 10^{-6}\]

evaluated numerically. Because we \textit{must} have \(PPTDSVol \geq SDSVol\) we are forced to revise \(PPTDSVol_{N=4}\) to the upper limit of its numerical uncertainty, which indicates convincingly that \(PPTDSVol_4 = SDSVol_4\).
For larger $N$ we have numerical evidence that $\text{PPTDVol}_N = \text{SDSVol}_N$ by a Monte Carlo argument. We programmed a numerical survey of billions and billions of random states which were positive under all partial transpositions, and without exception, ever such state also satisfied the necessary separability criterion (2.12). This would not be expected if $\text{PPTDVol}_N > \text{SDSVol}_N$ and is as such a proof by contraposition of apparent equivalence between $\text{PPTDVol}_N$ and $\text{SDSVol}_N$. 
2.4 Separability of Superradiance

An example of a diagonally symmetric system of physical consequence is the Dicke model of superradiance [191–194]. Superradiance is a phenomenon in which excited atoms spaced together very closely radiate in an induced cascade. This can occur if the volume of the system is smaller than the wavelength of the emitted radiation. The Dicke model of superradiance is the maximally idealized phenomenological model. The idealization employed in the Dicke model is that of perfect indistinguishability of the particles, such that we treat the system as existing entirely in only highest symmetry of the Hilbert space. Experimentally it corresponds to the small-volume limit and an absence of dipole-dipole induced dephasing. A thorough treatment of the volume-dependent many-body effects not considered in the Dicke model can be found in Refs. [193, 194].

The Dicke model describe the spontaneous decay of the (open) system with decay rate $\Gamma$ by means of Lindblad operators. The Liouville master equation [129, 195–197] which governs the time evolution of driven Dicke model superradiance is

$$\frac{\partial \rho}{\partial t} = \Gamma \left( A^- \rho A^+ \rho A^+ A^- A^- \right)$$

(2.23)
where

\[ A^+ = \sum_{n=1}^{N} \mathbb{1}_{n-1} \otimes |1\rangle \langle 0| \otimes \mathbb{1}_{N-n} \]  

(2.24)

with the annihilation operator being the adjoint of the creation operator, \( A^- = (A^+)^\dagger \).

The high symmetry of the Lindblad operators lead to a time evolution entirely within the manifold of the diagonally symmetric states, such that if initially diagonally symmetric then it remains diagonally symmetric, of the form \( \rho_{GDS} \) per Eq. (2.3).

Thus, the rate equation for Dicke model superradiance Gross and Haroche [192] can be expressed in the form of a recursive relation between the populations, namely

\[ \frac{\partial \chi_{n_1}(t)}{\partial t} = -\Gamma (n_0 + 1) n_1 \chi_{n_1}(t) + \Gamma n_0 (n_1 + 1) \chi_{n_1+1}(t) \]  

(2.25)

as per Gross and Haroche [192]. We choose to consider initially separable states and ask if superradiance time evolution leads to entanglement. The only diagonally symmetric separable state which superradiates is the maximally excited state, and thus we take as our initial conditions

\[ \chi_{n_1}(t \to 0) = \begin{cases} 1, & n_1 = N, n_0 = 0 \\ 0, & n_1 < N, n_0 > 0 \end{cases} \]  

(2.26)
When we solve for the $\rho_{SDS}$ weights and amplitudes per Eq. (2.12) we find that all the $x$ and $y$ are real number between zero and one throughout the entire time evolution, that is $\rho_{\text{SuperRad}} \in \mathcal{Q}_{SDS}$. Indeed, we numerically verified that for pure Dicke Model superradiance, conditions (2.12) are satisfied for all $\tau > 0$, thereby certifying full separability throughout the time evolution, for $N \leq 8$. This is demonstrated graphically in Fig. 2.2.
Fig. 2.1: (Color online) Superradiance populations as a function of time, per Eq. (2.25). The system starts out entirely in the maximally excited state and evolves towards existing entirely in the ground state. At intermediate times the system is a mixture of populations.

Fig. 2.2: (Color online) This plot illustrates the permanent full separability of a superradiating system, for $N = 4$. The various decomposition elements are plotted as a function of time; they have been solved-for from the system of polynomial equations defined in Eq. (2.12). For $N = 4$ there are five decomposition elements, as $y_3$ has been manually set to zero in accordance with Eq. (2.13).
2.5 Independently Radiating Model

It is valuable to consider, by an apophatic contrasting approach, a system which has no quantum enhanced effects and radiates according to a constant-hazard decay principle, essentially the large volume limit. To understand statistical independence we must first recognize the relationship between populations and probability functions. For example, if we define a probability distribution PDF\(t\) giving the likelihood that a single two-level atom will decay at any instant in time, then the ground state occupancy of that atom over time is equivalent to the cumulative probability distribution, CDF\(t\). The excited states occupancy, being one minus the ground state occupancy, corresponds therefore to the survival function, also known as the reliability function, REL\(t\). The failure rate, ie. the emission intensity, is given by the statistical hazard function, HAZ\(t\). The hazard function is defined as

\[
\text{HAZ}[t] = \frac{\text{PDF}[t]}{1 - \text{CDF}[t]} = -\frac{\text{REL}'[t]}{\text{REL}[t]} \tag{2.27}
\]

such that if the atoms behave statistically independently then each has a constant hazard function HAZ\(t\) = \(\Gamma\) leading to the occupancy differential equation REL\(t\) = -\(\Gamma\) REL\(t\).

Suppose now we have an infinite number of two-level atoms radiating according
to constant hazard \( \Gamma \), which we shall refer to as standardradiance as a foil to superradiance. Let us mentally group these infinite atoms into sets of size \( N \). Any given set has \( n_1 \) excited atoms and \( n_0 \) ground-state atoms, where the ratio changes in time. At any intermediate time some sets will have more or less excited atoms than others. We define the populations to mean the relative frequency of having sets with \( n_1 \) excited atoms. A set of \( n_1 \) atoms has an \( n_1 \)-fold hazard for the possibility that any one of its atoms might decay. On the other hand, the collection of \( n_1 \)-excited sets is constantly being replenished by the decay of any atom from the sets with \( n_1 + 1 \) excited atoms, which have a hazard rate to transition equal to \( \Gamma(n_1 + 1) \). Thus the independently-radiating rate equation for standardradiance populations is

\[
\frac{\partial X_{n_1}(t)}{\partial t} = -\Gamma n_1 X_{n_1}(t) + \Gamma (n_1 + 1) X_{n_1+1}(t)
\]  

(2.28)

where we use \( X \) for standardradiance to distinguish it from \( \chi \) used for superradiance in Eq. (2.25).

From a set of populations evolving in time it is possible to compute the per-particle florescence rate, ie. the per-particle probability density function to see a photon coming out of the system. For standardradiance the per-particle florescence rate is presumed to be an exponential distribution per the premise of constant
hazard. By contrast, the superradiance per-particle florescence rate has a higher peak emissivity along with a narrower distribution width, from whence comes its name. The per-particle florescence rate for both forms of radiation is given simply by

\[
\text{Superradiance PDF} = \sum_{n_1} \frac{n_1}{N} \frac{\partial \chi_{n_1}(t)}{\partial t} = \sum_{n_1} \frac{(n_0 + 1)}{N} n_1 \chi_{n_1}(t) \\
\text{Standardradiance PDF} = \sum_{n_1} \frac{n_1}{N} \frac{\partial X_{n_1}(t)}{\partial t} = \sum_{n_1} \frac{n_1}{N} \chi_{n_1}[t] = \Gamma e^{-\Gamma t}.
\]

The relatives rates of florescence are for \( N = 4 \) are contrasted in Fig. 2.4.

Let us pause for a moment to prove what is already physically obvious, namely that standardradiance is always described by a separable system, ie. \( \rho_{\text{StandardRad}} \in \varrho_{\text{SDS}} \). As we proceed in this proof, we shall find that standardradiance corresponds to a very special case.

We seek a closed-form solution to the standardradiance populations. To this end we note that the independent radiation model is fully equivalent to the well-studied physical system of nuclear decay. We analogize the radiative evolution of set of \( N \)-atoms through various excitation levels with a decay chain consisting of \( N \) nuclides. Specifically, we recognize that the concentration of a given nuclide in a sample parallels the relative frequency of finding \( N \)-atom sets containing a particular \( n_1 \) count of excited atoms. Decay chains in nuclear systems are described by Bateman equations [198]. Bateman equations give the concentration of the \( k \)'th nuclide at time
Concentration\(_k\) [\(t\)] = \[
\sum_{j=1}^{k} \left( \frac{e^{-\lambda_j t}}{\prod_{p=1, p\neq j}^{k} \lambda_j - \lambda_p} \right)^{k-1} \prod_{p=1}^{\lambda_p} \right)
\]  (2.30)

where \(\lambda_j\) is the decay constant for nuclide \(j\). For our model of independently radiating atoms, the decay constant is nothing more than the multiplicity of excited atoms in a given population\(^3\) times \(\Gamma\). Note the Bateman equations index the initial nuclide by \(k = 1\), whereas for standard radiance, however, we index the initial state \(\chi_{n_1=N}\) by \(n_0 = 0\). Therefore to translate the Bateman equations for our purposes we set \(k \rightarrow n_0 + 1\) and \(\lambda_{n_0} \rightarrow \Gamma n_1\) or \(\lambda_k \rightarrow \Gamma (N + 1 - k)\) such that we have

\[
X_{n_1} (t) = \sum_{j=1}^{n_0+1} \left( \frac{e^{-\Gamma(N+1-j)t}}{\prod_{p=1, p\neq j}^{\Gamma(N+1-j)} \Gamma(N+p-1)} \right)^{n_0} \right)
\]  (2.31)

This variant of the Bateman equations, with integer-values decay constants, can be readily simplified, to yield

\[
X_{n_1} (t) = \frac{N!}{n_0!n_1!} \left( 1 - e^{-\Gamma t} \right)^{n_0} \left( e^{-\Gamma t} \right)^{n_1}.
\]  (2.32)

\(^3\) In principle Bateman’s equations can be adapted to give a closed-form solution to superradiance populations as well. The outer sum is not analytically amenable however. The best simplification we have been able to identify is \(\chi_{n_1} (t) = \sum_{j=0}^{n_0} \frac{(-1)^j e^{-(N+1-j)\Gamma t} n_0! N!(N-2j-1)(n_1-j-2)!}{j! n_1!(n_0-j)! (N-j-1)!} \).
It is valuable at this point to introduce a dimensionless and rescaled time parameter, namely

\[ y = 1 - e^{-\Gamma t} \quad \text{ie.} \quad t = -\frac{\ln(1 - y)}{\Gamma} \quad (2.33) \]

so that \( y \) ranging between zero and one maps to \( t \) ranging from zero to infinity. With this rescaled time we see that

\[ X_{n_1}[y] = N! \frac{y^{n_0}(1 - y)^{n_1}}{n_0!n_1!} \]

ie. \[ \rho_{\text{StandardRad}}[y] = \sum_{n_1=0}^{N} N! \frac{y^{n_0}(1 - y)^{n_1}}{n_0!n_1!} |D_{n_1}^N \rangle \langle D_{n_1}^N| = \rho^N[y] \quad (2.34) \]

and therefore not only \( \rho_{\text{StandardRad}} \in \mathcal{E}_{\text{SDS}} \), but furthermore the standard radiating systems are themselves the basis states of the SDS parameterization!

This has two important physical ramifications. Firstly, whereas the SDS states were originally constructed purely for their mathematical form, this correspondence with standard radiating system provides a physical interpretation of those states. Secondly, this tells us that we could have defined a sufficient separability criteria capable of certifying the full separability of superradiance without requiring a priori separable states with similar form. By presuming that the phenomenological model of independently radiating atoms must be absent entanglement, one can define a
separability criteria by testing whether or not it is possible to express the superradiant state $\rho_{\text{SuperRad}}[t^\ast]$ in terms of a mixture of fundamentally-independently-radiating systems at various snapshots in time, $\rho_{\text{SuperRad}}[t_i]$. This amounts to the test of possible decomposition $\rho_{\text{SuperRad}}[t^\ast] \Rightarrow \sum_{j=1}^? x_j \rho_{\text{SuperRad}}[t_i]$. Just as per Eq. (2.12), the possibility of decomposition into particular separable states implies a sufficient separability criterion directly in terms of the matrix elements, namely

$$\rho_{\text{SuperRad}}[t^\ast] \in \mathcal{Q}_{\text{StandardRad}} \text{ iff } \exists x_1 \ldots x_{j_{\text{max}}}, t_1 \ldots t_{j_{\text{max}}}$$

satisfying $\forall n_1 \chi_{n_1} = \sum_{j=1}^{j_{\text{max}}} x_j X_{n_1}$ \hspace{1cm} (2.35)

such that $\forall j : 0 \leq x_j \leq 1$ and $0 \leq t_j \leq \infty$

which, if we change times variables such that $t_j \rightarrow -\ln(1-y_j)$ and we substitute the closed form expression for the standard radiance populations given by Eq. (2.32), then the criterion (2.35) is transformed identically into the criterion (2.12).

This alternative method of separability certification, namely by forcing a comparison with a statistically-independent analog of the target model, is of particularly practical value. We hope that similar comparisons, and the sufficient separability criteria which follow, may be of use to other researchers attempting to certify separability as well.
**Fig. 2.3:** (Color online) Standardradiance populations as a function of time, per Eq. (2.28) and Eq. (2.32). The system starts out entirely in the maximally excited state and evolves towards existing entirely in the ground state. At intermediate times the system is a mixture of populations. Note that the evolution is “slower” than that of superradiance; note the much shorter timescale plotted in Fig. 2.1.

**Fig. 2.4:** (Color online) This is a plot of the per-particle fluorescence rates for both superradiance and standardradiance, per Eq. (2.29). See how the superradiance per-particle florescence rate has a higher peak emissivity (along with a narrower distribution width) from whence comes its name.
2.6 Entanglement via Driven Superradiance

Although we have established, perhaps counter to expectations that there is no entanglement in Dicke model superradiance, it is interesting to explore a variant of this model in which we can show that entanglement is indeed generated. The model which we consider now is that of driven Dicke model superradiance, which has been considered repeatedly [195, 199–203], and which González-Tudela and Porras [203] have shown to lead to a spin-squeezed steady state. Here we analytically affirm the numerical results of González-Tudela and Porras [203], however using a spin-squeezing parameter more sensitive at detecting entangled states. This section of the thesis summarizes the salient results discussed in our work in Ref. [183]. Please note that in Ref. [183] we parameterized the density matrix using unnormalized Dicke states in the $j, m$ spin notation. To be consistent with the body of this thesis, however, we have translated our results into the notation of established by the Dicke basis of Eq. (2.2).

Driven superradiance is a generalization of Dicke model superradiance when the system is additionally driven by some external field; we take the external driving frequency in our model to be $\omega$ and use the Rotating Wave Approximation [196, 204, 205]. Thus the Liouville master equation [195, 197] which governs driven superradiance
\[ \frac{\partial \rho}{\partial t} = -i \left[ \frac{\omega}{2} (A^+ + A^-), \rho \right] + \Gamma \left( A^- \rho A^+ - \frac{A^+ A^- \rho + \rho A^+ A^-}{2} \right), \] (2.36)

is a simple extension of Eq. (2.23), and where the raising and lowering operators are still defined by Eq. (2.24).

To solve Eq. (2.36) we need not consider a fully-general density matrix \( \rho \). Firstly, the equation is symmetric with respect to permutation of the individual qubit Hilbert spaces, so we can take our density matrix to be symmetric, that is, expandable in symmetric basis states of the Dicke states of Eq. (2.2), although no longer diagonal in that basis. Second, the raising and lowering nature of the driving potential allows us to infer which matrix elements must be real and which must be (entirely) imaginary, and therefore we can define a sufficiently-general \( N \)-particle density matrix

\[ \rho_N = \sum_{n_a=0}^{N} \sum_{n_b=0}^{N} X(N)_{n_a}^{n_b} i^{(n_a-n_b)} |D_{n_a}^N \rangle \langle D_{n_b}^N| \] (2.37)

with real symmetric \( X(N)_{n_a}^{n_b} = X(N)_{n_b}^{n_a} \in \mathbb{R} \)

for shorthand, we shall refer to the various \( X(N)_{n_a}^{n_b} \) as the matrix elements of \( \rho_N \), although technically we ought to account for the phase as well.

It is possible to infer a rate equation in terms of the matrix elements from
Eq. (2.37) in the same manner that Eq. (2.25) follows from Eq. (2.23). To do so we need only recall the effects of the ladder operators on the Dicke states, namely

\[ A^+ |D^N_n \rangle = \begin{cases} \sqrt{(n+1)(N-n)} |D^N_{n+1} \rangle \equiv f_+(n) |D^N_{n+1} \rangle , & n + 1 \leq N \\ 0 , & n + 1 > N \end{cases} \tag{2.38} \]

\[ A^- |D^N_n \rangle = \begin{cases} \sqrt{(n)(N-n+1)} |D^N_{n-1} \rangle \equiv f_-(n) |D^N_{n-1} \rangle , & n - 1 \geq 0 \\ 0 , & n - 1 < 0 \end{cases} \]

which allows us to apply the operators of Eq. (2.36) inside the summations in Eq. (2.37). If we then re-index the dummy variables of summation so as to have a common index in the Dicke basis |D^N_{na} \rangle \langle D^N_{nb}|, as opposed to a common index in X(N)_{nb}^{na}, the we obtain a set of coupled first-order differential equations defined by

\[ \frac{\partial X(N)^{n_b}_{na}(t)}{\partial t} = -\frac{\omega}{2} \left( f_+(n_a) X(N)^{n_b}_{na+1}(t) + f_+(n_b) X(N)^{n_b+1}_{na}(t) \right) \\
-\frac{\omega}{2} \left( f_-(n_a) X(N)^{n_b}_{na-1}(t) + f_-(n_b) X(N)^{n_b-1}_{na}(t) \right) \\
+ \Gamma \left( f_+(n_a)f_+(n_b) X(N)^{n_b+1}_{na+1}(t) + \frac{f_-(n_a)^2 + f_-(n_b)^2}{2} X(N)^{n_b}_{na}(t) \right) \tag{2.39} \]

which have no imaginary elements, hence justifying the manual choice of phases in Eq. (2.37). Setting the left hand side of Eq. (2.39) to zero defines the steady state
condition, along with

$$\text{tr}[\rho_N] = \sum_{n=0}^{n} X(N)^n = 1$$

(2.40)

to account for normalization.

To obtain, practically, the steady-state matrix elements from Eq. (2.39) we need to iterate it over all possible $0 \leq n_a, n_b \leq N$, amounting to $(N + 1)^2$ equations. Without loss of generality we can invoke the symmetry of the matrix elements to consider only $0 \leq n_a \leq n_b \leq N$, which reduces the set of equations by about a factor of two. Even leveraging the symmetry, however, the set of linear equations scales like $O(N^2)$, and thus has quadratic computational complexity.

Spin Squeezing provides a valuable metric of entanglement [133–135, 171, 206, 207], with extensive immediate application in precision metrology [133–139]. We use the explicit form of the spin squeezing parameter of Ma et al. [135, Eq. (57)] and Lee and Chan [208, Eq. (45)], as follows:

$$\xi^2 = \frac{\langle \vec{J}_1^2 + \vec{J}_2^2 \rangle - \sqrt{\langle \vec{J}_1^2 - \vec{J}_2^2 \rangle^2 + \langle \vec{J}_1 \vec{J}_2 + \vec{J}_2 \vec{J}_1 \rangle^2}}{2/N}$$

(2.41)
where

\[ \tilde{J}_1 = \tilde{J}_y \cos \phi - \tilde{J}_x \sin \phi \]

\[ \tilde{J}_2 = \tilde{J}_x \cos \theta \cos \phi + \tilde{J}_y \cos \theta \sin \phi - \tilde{J}_z \sin \theta \]

and

\[ \theta = \cos^{-1} \left( \frac{\langle \tilde{J}_z \rangle}{\sqrt{\langle \tilde{J}_x \rangle^2 + \langle \tilde{J}_y \rangle^2 + \langle \tilde{J}_z \rangle^2}} \right) \]

\[ \phi = \tan^{-1} \left( \frac{\langle \tilde{J}_y \rangle}{\langle \tilde{J}_x \rangle} \right) , \text{ sensitive to quadrant,} \]

and where

\[ \tilde{J}_{x/y/z} = \frac{1}{N} \sum_{n=1}^{N} \frac{1}{n} \cdots \frac{1}{n-1} \otimes \sigma_{x/y/z} \otimes \frac{1}{n} \cdots \frac{1}{N-n} . \]

This particular measure of spin squeezing is denoted with a subscript \( S \) in Ref. [135, Table 1], where it is credited to Kitagawa and Ueda [209].

The calculation of \( \xi^2 \) can be immensely simplified by recognizing that the entire system’s spin is encoded in the \( \rho \)’s one or two particle reduced states. For states with real and imaginary parts à la Eq. (2.37) we show in the Supplementary Online
Materials of Ref. [183] that

\[ \xi_N^2 = 1 + (N - 1) \left< \sigma_x \otimes \sigma_x \rho_N^{(2)} \right>, \tag{2.45} \]

for driven superradiance, where subscript \(N\) indicates this special-case form. We have also introduced here \(\rho_N^{(d)}\) to indicate the reduced state of \(d\) particles, and we elect to explicitly specify the reduced-state \(\rho\) in the expectation value purely for pedagogical clarity. Note that Eq. (2.45) is also derived for symmetric states in Ref. [210, Eq. (7)].

Spin-squeezing is defined by \(\xi^2 < 1\), which is also a sufficient criterion for the presence of entanglement. With no loss of generality we therefore have certification of nonzero entanglement [100, 146–148, 179] via

\[ \forall N : \rho_N \in \{ \mathcal{Q}_{\text{entangled}} \} \text{ if } \left< \sigma_x \otimes \sigma_x \rho_N^{(2)} \right> < 0 \tag{2.46} \]

which, since \(\forall N : \left< \sigma_x \rho_N^{(1)} \right> = 0\), means that Eq. (2.46) is just a special case of the general entanglement criteria of [207, 210–212, Eq. (33)], which recognizes that all separable symmetric states satisfy \(\langle A \otimes A \rangle - \langle A \otimes 1 \rangle \geq 0\) for all Hermitian operators \(A\).

The essential contribution of our work in Ref. [183] is to provide an explicit
expression for $\langle \sigma_x \otimes \sigma_x \rho^{(2)}_N \rangle$ directly in terms of the unreduced matrix elements $X(N)_{n_a+1}^{n_b}$ of $\rho_N$ per Eq. (2.37). We found that $\rho^{(d)}_N$ can be expressed as linear map acting on $\rho_d$. We briefly summarize the argument there, translating it to the notation of this thesis. We recognize that the Dicke states can be expressed in an arbitrary bipartitioned basis through the Clebsch-Gordan coefficients [213–221], such that

$$
\rho_N = \sum_{n_a=0}^d \sum_{n_b=0}^{N-d} \sum_{n_a'=0}^d \sum_{n_b'=0}^{N-d} \left( X(N)_{n_a'+n_b'+n_a''+n_b''} \left( n_a'+n_a''-n_b'-n_b'' \right) \times CG(n_a',n_a'') CG(n_b',n_b'') \right.

\times |D_{n_a'}^d\rangle \otimes |D_{n_a''}^{N-d}\rangle \langle D_{n_b'}^d| \otimes \langle D_{n_b''}^{N-d}| \right)

(2.47)
$$

where $CG(n',n'')$ is shorthand for the full-spin Clebsch-Gordan coefficient. Explicitly,

$$
CG(n',n'') \equiv \frac{\binom{d}{n'}\binom{N-d}{n''}}{\binom{N}{n'+n''}} = \sqrt{\frac{d!(N-d)!(n'+n'')!(N-n'-n'')!}{n'!(d-n')!n''!(N-d-n'')!N!}}

(2.48)
$$

per Wolfe [221, Eq. (7)] and O’Hara [217, Eq. (10)]. Now the definition of the reduced state requires tracing out the second “half” of the partitioned Hilbert space, namely

$$
\rho^{(d)}_N = \sum_{k=0}^{N-d} \left( 1 \otimes \langle D_{k}^{N-d}| \right) \rho_N \left( 1 \otimes |D_{k}^{N-d}| \right)

(2.49)
$$

which acts to isolate only those elements which are diagonal in the Dicke basis of
that Hilbert space, such that we have

\[
\rho_N^{(d)} = \sum_{k=0}^{N-d} \sum_{n_a=0}^d \sum_{n_b=0}^d \left( \text{CG}(n_a^\dagger, k) \text{CG}(n_b^\dagger, k) X(N)^{n_a+k}_{n_b+k} |D^{n_a}_{n_b}\rangle \langle D^{n_b}_{n_a}| \right) \tag{2.50}
\]

which we choose to express as a linear map acting on \(\rho_d\), such that

\[
\rho_N^{(d)} = \text{Map}_N(\rho_d), \quad \text{where}
\]

\[
\text{Map}_N \left( X(d)^{n_b}_{n_a} \right) \rightarrow \sum_{k=0}^{N-d} \text{CG}(n_a^\dagger, k) \text{CG}(n_b^\dagger, k) X(N)^{n_a+k}_{n_b+k}.
\tag{2.51}
\]

Note that this mapping can readily be generalized for all symmetric states, not just those of the ansatz of Eq. (2.37).

Now, since one can readily verify that

\[
\langle \sigma_x \otimes \sigma_x \rho_2 \rangle = \sum_{s=0}^{1} (-1)^s \binom{2}{s} X(2)^{1+s}_{1-s} \quad \text{and since}
\]

\[
\langle \sigma_x \otimes \sigma_x \rho_N^{(2)} \rangle = \text{Map}_N \left( \langle \sigma_x \otimes \sigma_x \rho_2 \rangle \right) \tag{2.53}
\]

\[
= \sum_{k=0}^{N-2} \sum_{s=0}^{1} (-1)^s \binom{2}{s} \text{CG}(1+s, k) \text{CG}(1-s, k) X(N)^{1+s+k}_{1-s+k}
\]

we are now able to explicitly define the spin squeezing parameter of Eq. (2.45) directly in term of the matrix elements. Furthermore, we can simplify Eq. (2.53) by noting
that, for the relevant parameter range,

\[
\left(\begin{array}{c}2 \\ s \end{array}\right) \left\{ C \left(1 + s', k\right) C \left(1 - s', k\right) \right. \\
= \frac{2}{N(N-1)} \sqrt{(i+1)(i+1+s)(N-i-1)(N-i-1+s)}.
\]  

(2.54)

Finally, reindexing the summation and inserting our result into Eq. (2.45), we conclude that

\[
\xi_N^2 = 1 + \frac{2}{N} \sum_{q=1}^{N-1} \sum_{s=0}^{1} (-1)^s \sqrt{(q)(q+s)(N-q)(N-q+s)} X(N)^{q+s}_{q-s},
\]  

(2.55)

a simple expression that only draws upon the diagonal and the one-off-diagonal matrix elements of \( \rho_N \).

Our question now is can we find some \( \omega \) for a given \( \Gamma \) such that we can drive the system into an entangled state characterized by \( \xi^2 < 1 \)? Yes! We quantify the entanglement of the steady state in terms of \( \Omega \equiv \omega/\Gamma \), defined as the ratio of the two experimental parameters. We find the steady state to be spin squeezed, i.e. with measure \( \xi^2 < 1 \), for sufficiently small \( \Omega \); see for example Fig. 2.5. To make a general statement, we note that for all \( N \), when \( |\Omega| \lesssim 0.475N \) the resulting steady state is always at least somewhat spin-squeezed state, see Fig. 2.7.

One would like to know how to tune \( \Omega \) so as to maximize the entanglement in
the resulting steady state. To this end, see Fig. 2.6 where it appears that the optimal
\( \Omega \) scales like \( (\Omega/N)^2 \sim a \ln N + b \) for large \( N \). It is also desirable to quantify the
maximal extent of the spin squeezing that can achieved in the model of driven Dicke
superradiance. Per Fig. 2.6, the squeezing extent rapidly strengthens for large systems.
Indeed, the value of the best-possible \( \xi^2 \) almost appears to drop off logarithmically as
a function of \( N \), descending below 0.5 at the right edge of Fig. 2.6 with no sign yet of
tapering off. This suggest that by increasing the size of the system, \( \xi^2 \) can perhaps
be made arbitrarily small in the steady state of this model. With the usual caveats
that genuine superradiance suffers from volume-dependent effect not accounted for
in the Dicke model [193, 194], this result nevertheless further suggest that driven
superradiance may be a viable scheme for generating large tightly squeezed states.

It is worth noting that the spin squeezing parameter is related to to the
entanglement monotone \textit{Negativity}[141, 142, 145, 146]. The Negativity is equal to the
combined magnitude of all negative eigenvalues in the partial transpose of \( \rho \), ie.

\[
\mathcal{N}(\rho) \equiv \frac{\left(\sum_i |\lambda_i|\right) - 1}{2}. \tag{2.56}
\]

The Negativity is a common benchmark of a state’s distillability and resource value
for nonlocality [55, 56].
For a $2 \times 2$ system, such as $\rho_{N}^{(2)}$, it is known that the partial transpose is always full rank and has at most one negative eigenvalue \cite{141}, in which case the Negativity is the magnitude of that single negative eigenvalue. By direct computation we find that
\[
\langle \sigma_x \otimes \sigma_x \rho_2 \rangle / 2
\]
is one of the eigenvalues of $\rho_{2}^{PT}$, and thus via the mapping of Eq. (2.51) we also have that $\langle \sigma_x \otimes \sigma_x \rho_{N}^{(2)} \rangle / 2$ must be an eigenvalue of general $\rho_{N}^{(2),PT}$. What we see is that the spin-squeezing parameter is effectively a linear function of the reduced state Negativity, such that Eq. (2.45) has the corollary

If $\xi_{N}^{2} < 0$, then

\[
\xi_{N}^{2} = 1 - 2(N - 1)\mathcal{N} \left( \rho_{N}^{(2)} \right).
\]

(2.57)

See Refs. \cite{141, 150} for a translation between the Negativity and Concurrence entanglement monotones, as the Concurrence has in some sense become a conventional standard metric for multiparticle entanglement \cite{149}, such as in Refs. \cite{124, 151}. Spin squeezing is directly related to the two-particle Concurrence in Ref. \cite[Eq. (5)]{211} and to the CCNR criteria in Ref. \cite[Obs. 2]{148}. 
Fig. 2.5: (Color online) We graph the spin-squeezing parameter $\xi^2$ for a steady state driven superradiant system as a function of $\Omega/N$ for various small $N$. Recall that $\Omega$ is the ratio of the driving frequency to relaxation frequency, $\Omega \equiv \omega/\Gamma$. The state is spin-squeezed whenever $\xi^2 < 1$; shown as shaded regions in this graph. The minima of the curves descends further with increasing $N$.

Fig. 2.6: (Color online) We plot the best-case scenario values for entanglement generation as a function of system size $N$. Note the dual meaning of the $Y$ axis: The upper curve indicates the minimal possible $\xi^2$, it is shaded upward to indicate that all larger values are also achievable. The lower curve indicates the optimal choice of $(\Omega/N)^2$ to achieve the corresponding minimal $\xi^2$. The lower shading indicates the complete parameter region where the steady state is spin squeezed. Note the logarithmic scaling of the $X$ axis.
Fig. 2.7: (Color online) This contour plot shows the spin-squeezing parameter $\xi^2$ for a steady state driven superradiant system as a function of $\Omega = \omega/\Gamma$ over a dense set of $N$, among which are the discrete $N$ plotted in Fig. 2.5. We plot only the region where $\xi^2 < 1$. Red indicates strongest spin squeezing, ie. minimal $\xi^2 < 1$. Although hard to see, the $\xi^2 = 1$ boundary is not monotonically decreasing; rather, it’s minimized at $\Omega/N = 36$. 

\[ \xi^2 = \begin{array}{c}
0.9 \\
0.8 \\
0.7 \\
0.6 \\
0.5 \\
\end{array} \]
The bipartite entanglement detected by spin squeezing is an indication of the nonlocal capacity of the state, and vice versa. Tóth and Gühne [222] discuss how one cannot infer the degree of entanglement and the extent of spin squeezing from the observation of bipartite statistical correlations. Tura et al. [182] discuss how general nonlocality of a quantum state can be inferred by Bell-testing it using exclusively bipartite measures such as that of Eq. (2.46), see also Refs. [223, 224]. Bera [225] goes so far as to demonstrate that the well-known enhancement of precision measurement due to entangled states [133–139] is actually more a consequence of the nonlocality capacity of the state than its entanglement.

We conclude that while entanglement may not be a direct proxy for nonlocal potential [106–112] it is nevertheless an extraordinarily valuable resource, in that it allows for a plethora of non-classical phenomena [61, 93, 95, 96, 102, 126–139].

Entangled qubits represent a uniquely valuable subtype of quantum resource in that they are a direct proxy to multipartite binary and dichotomic quantum nonlocality [73], see also Refs. [55, 56, 159]. More broadly, symmetric quantum states are valuable in that they mirror the symmetry structure of multipartite no-signalling polytopes [8, 21, 28, 30, 33]. The relationship between entanglement and nonlocality remains an open question; we hope that the techniques for assessing and generating entanglement that we have developed in this chapter may prove useful in furthering
quantum information theory, both conceptually and practically.
Chapter 3

Contextuality

The fundamental theorem of quantum mechanics [is...] if you have several questions, and you can answer any two of them, then you can also answer all of them. \(^1\)

\(^1\) **Ernst Specker**, 2009

*Recorded by Adán Cabello*, reflecting on the Kochen-Specker Theorem [226].
3.1 The Graph-Theoretic Formalism

As mentioned in Sec. 1.1, the outcomes of quantum measurements are contextual. Contextuality is broader than merely nonlocality, in the sense that quantum contextuality can be exhibited without requiring spatially separated subsystems. Contextuality, tautologically, is evident whenever measurement outcomes for a single-partite experiment are inconsistent with noncontextually (albeit probabilistically) assigning outcomes to the various measurements [227, 228]. For explicit examples of quantum contextuality distinct from quantum nonlocality, see Refs. [2–6, 9, 229, 230]. It is precisely the inability to pre-assign probabilities to even a single-partite scenario (“noncomposite” in the language of Man’ko and Markovich [231]) scenario which makes contextuality more broadly applicable than nonlocality [9, 10].

Just as with nonlocality, contextuality is valued as a resource for information theoretic tasks [232, 233]. Further similarly, the extent of quantum contextuality is intermediate between absolute noncontextuality and maximal contextuality conceivable without violating probability normalization. The distinction between classes has recently been formalized with the aid of graph-theoretic approach due to Cabello, Severini, and Winter [234]. This formalism is also reviewed in Refs. [6, 36, 235].

One can express the contextuality scenario as an exclusivity graph. This is done by assigning each vertex of the graph a label indicating some particular
measurement outcome tuple (output) for some particular apparatus choice tuple (input). Noncontextuality is the notion that each individual output depends only on the individual input queried, whereas contextuality is when the output depends differently on the input depending on *which tuple* of inputs it is drawn from. To complete the exclusivity graph we draw edges between vertexes that are incompatible. By incompatible we mean that at least one input is common between both vertexes such that the output for that input is not the same at both vertexes. We refer to incompatible vertexes as orthogonal, adjacent, or exclusive.

In this work, exclusivity between vertexes is represented through edges, thus in our notation exclusivity=orthogonality=adjacency. This is the notation used in Refs. [6, 44, 47, 76, 234–236]. By contrast, in Refs. [36, 45] the authors work within the framework of nonorthogonality graphs, electing to represent exclusivity by non-adjacent vertices. \(^2\) Our work accommodates both conventions by explicitly specifying which graphical representation is meant in each statement. To do this, we refer to graphs where exclusivity is represented by adjacency as orthogonality graphs - O \((G)\). Graphs where exclusive vertexes are not adjacent are referred to as nonorthogonality graphs - NO \((G)\).

Firstly, note that the total probability assigned to two adjacent vertexes in

---

\(^2\) Note that different conventions for the graphical representation of exclusivity are used in different works even by the same authors. Fritz *et al.* [36] are also authors of Refs. [44, 47].
$O(G)$ must not exceed 1. This is a simple consequence of probability normalization; summing the probability of seeing all various different outputs for a given constant input equals one, so any two different outputs for a given input have total probability no greater than one. We now proceed to assign probabilities to the various vertexes per various classes of statistical behaviors.

A deterministic behavior is one which assigns zero-or-one probabilities to the various vertexes. Note that the total probability of any deterministic behavior on the graph is bounded by the Independence Number - $\alpha(O(G))$, which is the vertex count of the largest possible set of vertexes in $G$ which do not mutually conflict, that is, there are no edges linking vertexes in an independent subset. A probabilistic noncontextual behavior essentially presumes that for every experimental query the experimenter is probabilistic exposed to some different possible deterministic underlying behavior, with the weights of the likelihood of the different behavior being normalized probabilities. Note that because the total probability of every deterministic behavior is bounded by $\alpha$, so to any convex combination of behaviors will also be bounded by $\alpha$. As such, having total probability less than or equal to the orthogonality graph’s independence number is a universal feature of all noncontextual behaviors. Such behaviors are
called NCHV, or noncontextual hidden variable models.

\[
\text{If } \vec{p}(v) \in \text{NCHV, then } \sum_v p(v) \leq \alpha(O(G))
\] (3.1)

Note that Eq. (3.1) is a necessary condition for classicality, but not a sufficient one. A necessary and sufficient condition is given by Fritz et al. [36, Prop. 4.3.1].

A quantum contextual behavior requires modelling the probability with quantum measurement operators, i.e., projectors, acting on a quantum state. As thoroughly discussed in Refs. [6, 36, 234–236], the total probability of any quantum behavior on a graph is bounded by the graph’s Lovász Theta Number - \( \vartheta \). Following Refs. [6, 234] we have that

\[
\vartheta(O(G)) = \max_{|\Psi\rangle, |\psi_v\rangle} \sum_{v \in V(G)} |\langle \Psi | \psi_v \rangle|^2
\]

where \(|\Psi\rangle, |\psi_v\rangle\) range over all unit vectors

\[
\text{such that } u \perp v \text{ and } u \neq v \text{ implies } |\psi_u\rangle \perp |\psi_v\rangle.
\] (3.2)

where the orthogonal representation may be defined for \( O(G) \) if one insists that orthogonality \((u \perp v)\) be represented by non-adjacency \((u \not\sim v)\) such as in Refs. 3

\[3\] There is a conflict in the definition of the Lovász theta number of a graph, depending on one’s preference for representing orthogonality by adjacency or non-adjacency. We use the modern convention per Knuth [237]. This is the convention used by both Cabello et al. [6] and Fritz et al. [36].
As such we have

\[
\text{If } \vec{p}(v) \in Q_{\text{contextual}}, \text{ then } \sum_v p(v) \leq \vartheta(O(G))
\]

which, analogously to Eq. (3.1), is a necessary condition for statistics to be quantum-contextual-achievable, but not a sufficient one. A necessary and sufficient condition for quantum contextuality in terms of the Lovász Theta Number is given by Fritz et al. [36, Prop. 6.3.2]. In the language of graph theory, the quantum measurement projectors are called “ribs” and the quantum state being measured is called the “handle”, after an archetypal five-vertex exclusivity graph with a three dimensional orthogonal representation that looks like remarkably like an umbrella [238, Theorem 2; see also 106, 236].

We note with some interest that we are not aware of a graph invariant which captures all general contextuality scenario beyond quantum contextuality. Such highly-general scenarios, however, are defined by only the restriction that adjacent vertexes have total probability less than one, see Liang et al. [2] for illustrative examples. Every probabilistic model of Fritz et al. [36], by definition there, satisfies this general criterion.

Recently there has been intense interest in trying to derive quantum contextual
statistics purely from a minimal set of informational principles [239–242]. Effort has shifted now to searching for a *single* principle capable of recovering quantum nonlocality and contextuality. Landmark candidates included Macroscopic Locality and Information Causality [38–43]. The most recent and celebrated ideas, however, are the Exclusivity Principle [235, 243–249] and Local Orthogonality [36, 44–47].

Expressed in the graphical formalism, both these new principles posit imposing total probability less than one for any set of mutually exclusive vertices. Bounding clique total probability is a more restrictive condition than merely bounding pairwise probability [2]. Indeed Refs. [6, 36, 44–47, 235, 247, 249] have exploited this simple idea to tremendous effect.

A third approach has also been pioneered independently by Sorkin [250]. Sorkin’s Quantum Measure Theory was uniquely motivated by efforts to formulate a quantum theory of gravity, and serves as motivation for the Consistent Histories interpretation of quantum mechanics [251, 252]. Quantum Measure Theory operationally posits “Sorkin’s Sum Rule” which forbids third-order interference phenomena. Recently it has been noted that Sorkin’s Sum Rule recovers various properties of contextuality and nonlocality [253, 254]. In this thesis we treat Sorkin’s principle as merely a special case of the Exclusivity Principle since Henson [255] has shown that
Sorkin’s Principle implies Consistent Exclusivity.\footnote{We note that the sum rule is effectively equivalent to requiring zero tripartite Interaction Information \cite[Eq. (6)]{256} if one understands third-order interference as merely a special physical case of tripartite Interaction Information. Interaction Information is the generalization of Mutual Information, which expresses the amount information (redundancy or synergy) bound up in a set of variables \textit{beyond} that which is present in any subset of those variables. Phrased as such, it becomes clear that Sorkin is concurring with Specker’s sentiment \cite[226, 243, 257, 258, and 36, Sec. 7.1]{226, 243, 257, 258, 36} that “A collection of propositions about a quantum mechanical system is precisely simultaneously decidable when they are pairwise simultaneously decidable.” It is not clear to the authors why this correspondence between the principles of Sorkin and Specker has not been noted in earlier literature.}

There are two important limitations which we must keep in mind to modulate any expectations that principles such as the aforementioned might fully recover quantum nonlocality:

Firstly, the statistical constraints due to the Exclusivity Principle (EP) and Local Orthogonality (LO) have not been shown to recover quantum contextual statistics. Fritz \textit{et al.} \cite[Sec. 7.3]{36} has shown that ‘Consistent Exclusivity” (CE) is related to the Shannon Capacity of nonorthogonality graphs, $\Theta(\text{NO}(G))$. CE is a \textit{gedankenexperiment} based on applying EP or LO to an infinite number of copies of a contextuality scenario, in which each copy is identical and independent. The existence of graphs for which the Shannon Capacity diverges from the Lovász Theta Number shows that CE fails, in general, to recover quantum contextual statistics \cite[Theorem 7.4.3]{36}. Admittedly, EP/LO may have ramifications beyond those which can be gleaned from the CE thought experiment, such as demonstrated recently by Cabello \cite{249}. It is therefore open that EP/LO might perhaps recover quantum contextual statistics.
An extension of EP due to Yan [246] is known to exactly correspond to quantum
contextuality; see [36, Theorem 7.7.2].

Secondly, and especially germane for this thesis, is that the statistical constraints
due quantum contextuality fail to tightly define the constraints of quantum nonlocality.
Although quantum nonlocality is innately a demonstration of quantum contextuality,
the statistical boundaries of single-partite quantum contextuality per Eq. (3.3) are
not sufficient to define the set of nonlocal quantum statistics. As a consequence,
EP/LO are not candidates for recovering quantum nonlocal statistics, only quantum
contextual statistics.

This is not particular surprising, given that the Lovász theta number is defined
in terms of general quantum projective measurements without reference to any kind
of product structure. In a genuine multipartite quantum experiment, although the
shared state $|\Psi\rangle$ may be entangled, the measurement projectors $|\psi_v\rangle$ must represent
tuples, ie. tensor products, of individual projectors for the different parties in different
Hilbert spaces; at the very least the projectors must commute. It is the product
structure (mandated commutativity) in the measurement projectors which gives rise
to the “gap” between the quantum-contextual and quantum-nonlocal elliptopes in
conditional probability space. Contrast Fritz et al. [36]'s Proposition 5.2.1, where the
projectors commute, with Corollary 6.4.2 there, where the projectors only appear to
commute, in that they are permutation invariant with respect to a particular state, ie. with respect to the handle associated with the orthogonal representation of the exclusivity graph. See also Refs. [259, and 89, Lemma 3].

The formal mathematical distinction between quantum nonlocality and quantum contextuality is the commutation of the parties’ projectors which is “imposed” in quantum nonlocality. This is such a critical point that it bears repeating explicitly.

A nonlocality scenario can be “relaxed” to a contextuality scenario by maintaining the same exclusivity (hyper)graph, but generating the probabilities which live on the graph via noncommuting projectors. The measurements of Alice and Bob are now just considered different dimensions in a larger sample space; in general, nonlocality scenarios can readily be viewed as contextuality scenarios by identifying each party with a different degree of freedom of a single physical system. The fact that the temporal order of Alice and Bob’s measurements is irrelevant can be implemented by imposing permutation invariance of the projectors with respect to whatever quantum state is being measured.

As expressed by Fritz et al. [36] and Navascués et al. [89], this idea can be extended to distinguish N-partite quantum nonlocality from a quantum contextuality scenario involving N simultaneous measurements as follows: Let there be a normalized state $|\phi\rangle \in \mathcal{H}$ and normalized projector operators $\{E_{k}^{a,x}\} \subset B(\mathcal{H})$ such that $\sum_{a} E_{k}^{a,x} =$
\(P(a_1, \ldots, a_N | x_1, \ldots, x_N) = \langle \phi | \bigotimes_{k=1}^N E_{a_k}^{x_k} | \phi \rangle\). Now the condition on the projectors which distinguishes nonlocality from contextuality is as follows:

**Contextuality:** The projectors commute with respect to the order of the simultaneous measurements: 

\[
E_{a_1}^{x_1} \ldots E_{a_N}^{x_N} | \phi \rangle = E_{\pi(1)}^{a_{\pi(1)}} \ldots E_{\pi(N)}^{a_{\pi(N)}} | \phi \rangle,
\]

where \(\pi \in S_N\) is an arbitrary permutation of the “parties” \(\{1, \ldots, N\}\).

**Nonlocality:** The projectors for each party are assigned the own Hilbert spaces:

\(\{E_k^{a,x}\} \subset B(H_k)\), implying that the global Hilbert space is effectively defined as

\(H = \bigotimes_{k=1}^N H_k\).

Note that projectors with satisfy the nonlocality condition inherently also satisfy the contextuality condition, but not vice versa.

In this thesis we refer to this relaxation of nonlocality as precisely contextuality because it allows for the joint probability distribution to be generated by noncomposite measurements. Given a nonlocality scenario, one would naturally define the corresponding quantum contextual correlations as those which share the same exclusivity graph as the nonlocality scenario but which can be generated by noncomposite quantum measurements [36, Proposition 6.3.1.d]; indeed this is equivalent the the operational definition above in terms of “almost” commuting projectors [36, Corollary...
6.4.2. This perspective on quantum nonlocality is also expressed by Rabelo et al. [236], to whom we are grateful for sharing a preprint of Ref. [236].

The third chapter of this thesis is wholly dedicated to exploring the statistical discrepancy between correlations consistent with quantum nonlocality and those consistent with quantum contextuality, and the implications of this distinction. Our contribution to this effort is the novel identification of a region in conditional probability space where the correlations which are impossible to implement in a quantum nonlocality scenario can nevertheless be achieved through a quantum contextuality scenario. Our results in this chapter have never before been published prior to this thesis.
3.2 Quantum Contextuality is the NPA\textsuperscript{1+AB} Set

Quantum contextuality, as just defined, corresponds to what we refer to here as the NPA\textsuperscript{1+AB} level of the NPA hierarchy [35, 36, 52], although it is also referred to as \(Q^{1+AB} [77]\), or \(Q_1 [36]\), or \(\tilde{Q} [89]\). The subtle distinction between quantum correlations in the sense of contextuality vs. nonlocality has received increasing attention in recent years [36, 89, 236 and 259, Eq. (20)], although very few explicit statistical discrepancies have been identified. We note that other references rarely refer to such statistics as quantum contextuality statistics, even though an unlabelled exclusivity graph, even if motivated by a multipartite scenario, tautologically defines an abstract contextuality scenario; Ref. [236], however, does use this language.

Some physicists have even suggested that perhaps quantum nonlocality is only an approximation to quantum contextuality, and that the universe might in fact permit NPA\textsuperscript{1+AB}-saturating correlations in a genuine bipartite scenario; Popescu [1] has questioned “...whether nature is in fact more nonlocal than expected from quantum theory.” Dowker et al. [77] have further explicitly shown that the existence of correlations consistent with NPA\textsuperscript{1+AB} is the (strongest possible) consequence of the Consistent Histories interpretation due to Griffiths, Omnès, Gell-Mann, and Hartle, [252, 260, 261, and references therein], specifically the convex and closed-under-wirings variant of the Strongly Positive Joint Quantum Measure formalism,
Therefore, it is a matter of urgent importance to improve our understanding of the different experimental predictions of conventional multipartite quantum mechanics versus NPA$^{1+AB}$.

As mentioned, there are a few known instances where NPA$^{1+AB}$ gives a non-trivially weaker restriction of the allowed probabilities than does genuine bipartite nonlocality as implemented through quantum mechanics. The first examples were the I3322 scenario [262–264] and the CGLMP scenario for $d = 3$ [265, 266], and now more recently a proof has been provided for the fundamental (2,2,2) scenario [89]. While Navascués et al. [89] provide a singular counterexample point, we here demonstrate an entire region where NPA$^{1+AB}$ lies outside of the genuine bipartite quantum elliptope.

To demonstrate the inadequacy of NPA$^{1+AB}$ we require another (stronger) quantum bound which we can show that NPA$^{1+AB}$ is relatively looser than. This is somewhat challenging as the default criterion of NPA\textsuperscript{1} Eq.(1.25) is already a relaxation of by NPA$^{1+AB}$. To get around this, we use our own QB\textsuperscript{3}(8) from Ref. [31], reproduced here in Table 1.1 and Eq.(1.30), as our contrasting bound. Recall that

\footnote{To quote Dowker et al. [77]: “Thus, if $Q^{1+AB}$ is strictly larger than $Q$, as indicated by the computational evidence, then non-local correlations beyond those achievable in ordinary quantum mechanics are achievable within strongly positive quantum measure theory.” $Q^{1+AB}$ refers to the set of all points in conditional probability space consistent with the NPA$^{1+AB}$ criterion, and $Q$ refers to the set of nonlocal correlations achievable by assigning Alice and Bob to separate Hilbert spaces, see Ref. [89]. The sentiment that $Q \neq Q^{1+AB}$ is merely “indicated by the computational evidence” is deprecated by the stronger analytical results graphed in Fig.3.1 of this thesis.}
the NPA$^{1+AB}$ certifies points as non-quantum if the corresponding $\Gamma$ matrix cannot be made positive semidefinite. We reproduce the $\Gamma$ matrix corresponding to binary and dichotomic NPA$^{1+AB}$ in the $9 \times 9$ explicit matrix in Table 3.1.
Table 3.1: We reproduce the $\Gamma$ matrix corresponding to NPA$^{1+AB}$ level of Navascués-Pironio-Acín hierarchy [35, 52] for the bipartite binary and dichotomic scenario. The coordinates in conditional probability space define the eight “given” parameters: $\langle A_0 \rangle, \langle A_1 \rangle, \langle B_0 \rangle, \langle B_1 \rangle, \langle A_0 B_0 \rangle, \langle A_1 B_0 \rangle, \langle A_0 B_1 \rangle, \langle A_1 B_1 \rangle$. There are also eight “free” variables in the matrix which have no experimentally-determinable values, namely: $\langle A_0 A_1 \rangle, \langle B_0 B_1 \rangle, \langle A_0 A_1 B_0 \rangle, \langle A_0 A_1 B_1 \rangle, \langle A_0 B_0 B_1 \rangle, \langle A_0 A_1 B_0 B_1 \rangle, \langle A_0 A_1 B_1 B_0 \rangle, \langle A_0 A_1 B_1 B_1 \rangle$. In the matrix below, the free variables are colored purple and are not wrapped by BraKet. A conditional probability distribution is inside (outside) the NPA$^{1+AB}$ elliptope iff values exist (do not exist) for the free variables such that the matrix $\Gamma$ can be made positive semidefinite.

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In Sec. 1.5 we considered the parametric probabilistic box $P_{\xi\gamma} = \xi P_{\text{PR}} + \gamma P_{\text{FD}} + (1 - \gamma - \xi) P_{\emptyset}$ spanned by the PR box, the fully-deterministic box, and the totally random box, per Eq. (1.21). We found that, on this facet, the criterion NPA$^{1+AB}$ was far and away the most restrictive quantum bound known. In other words, there did not appear to be any instance in which quantum nonlocality was distinct from quantum contextuality along that particular slice of the no-signalling polytope.

We have found another slice, however, for which we can show that NPA$^{1+AB}$ is outside of, ie. not tight against, the elliptope of bipartite quantum correlations. The three boxes which span this new slice are the PR box, the totally random box, and the semi-deterministic Bob-random box of Eq. (1.19). Analogous to Eq. (1.21) we define here a box mixture which lives in this span, namely

$$P_{\xi\beta}(ab|xy) = \xi P_{\text{PR}}(ab|xy) + \beta P_{\text{SD}}(ab|xy) + (1 - \beta - \xi) P_{\emptyset}(ab|xy)$$ (3.4)

Recall that $P_{\text{SD}}$ is a local box which returns 1 for Alice (independent of her measurement choice), and is completely random for Bob (independent of his measurement choice).

The 2-dimensional statistical region we are considering with this box span, is
therefore

\[ \langle A_0 \rangle = \langle A_1 \rangle = \beta, \quad \langle B_0 \rangle = \langle B_1 \rangle = 0 \]

\[ \langle A_0 B_0 \rangle = \langle A_1 B_0 \rangle = \langle A_0 B_1 \rangle = -\langle A_1 B_1 \rangle = \xi \]
Fig. 3.1: (Color online) A comparison of quantum bounds along a slice of the no-signalling polytope which illustrates the inadequacy of NPA\(^{1+AB}\) for characterizing multipartite quantum correlations. The grey shaded area represents the cumulative best known approximation of the genuine quantum boundary. Note that unlike in Fig. 1.1, the contextuality quantum bound corresponding to NPA\(^{1+AB}\) is not tight along this boundary. Specifically, for \(\frac{1}{4} \lesssim \beta < \frac{1}{2}\), it is inadequate. The region of the quantum contextuality elliptope which is known to lie outside of the quantum nonlocality elliptope is shaded in red. The lower figure is a closeup view highlighting this region of interest, with a \(\beta\)-dependent rescaling of the y-axis for visual clarity. For \(\beta > \frac{1}{2}\) the quantum elliptope exactly coincides with the no-signalling polytope, and thus the quantum boundary is simply \(\xi \leq 1 - \beta\) for \(\beta > \frac{1}{2}\). It should be noted that the LO\(^2\) and NPA\(^1\) bounds bump into the no-signalling polytope earlier, at \(\beta = \frac{1}{3}\). The black dots are the numeric points for which we calculated the maximal \(\xi\) per NPA\(^{1+AB}\).
Using $P_{\xi\beta}$ per Eq. (3.4), and recognizing that normalization of probabilities demands $\xi \leq 1 - \beta$, it is easy to recalculate the familiar quantum bounds we first considered in Sec. 1.5. We define the bounds piecewise along $\beta$, switching to $\xi \leq 1 - \beta$ when that becomes the more restrictive condition on $\xi$. It is easy to see that Uffink’s bound of Eq. (1.23) becomes

$$\xi \leq \begin{cases} 2^{-1/2} & \text{for } 0 \leq \beta \leq 1 - 2^{-1/2} \\ 1 - \beta & \text{for } 1 - 2^{-1/2} \leq \beta \leq 1 \end{cases}$$

(3.6)

and the bound corresponding to the first level of the NPA hierarchy per Eq. (1.25) is simply

$$\xi \leq \begin{cases} 2^{-1/2} \sqrt{1 - \beta^2} & \text{for } 0 \leq \beta \leq 1/3 \\ 1 - \beta & \text{for } 1/3 \leq \beta \leq 1 \end{cases}$$

(3.7)

, an obvious improvement over Uffink’s bound in Eq. (3.6).

Using the clique of mutually-exclusive events tallied in Eq. (1.33) we can show that $L_{\text{O2}}$ implies for the $P_{\xi\beta}$ box

$$\xi \leq \begin{cases} \frac{\sqrt{2}(1-\beta)(\beta+5)+\beta-1}{3} & \text{for } 0 \leq \beta \leq 1/3 \\ 1 - \beta & \text{for } 1/3 \leq \beta \leq 1 \end{cases}$$

(3.8)

Lastly, we can recycle the envelope of linear bounds given in Eq. (1.32) in
order to characterize the constraint implied by $QB_{3}^{(8)}$ of Eq. (1.30) for the $P_{\xi\beta}$ box of Eq. (3.4). By direct substitution of Eq. (3.5) into the function-valued bound Eq. (1.30), we see that

$$\xi \leq \min_{c^{2} \leq 1} \left\{ \frac{c^{2} - \sqrt{(3c^{2} - 4)(c^{2} - 2)}}{c^{2} - 1} - c(2\beta) \right\},$$

(3.9)

Note that Eq. (3.9) shares the same minimization term as Eq. (1.31), such that we can identify the envelope of bounds on $P_{\xi\beta}$ due to $QB_{3}^{(8)}$ as

$$\xi \leq \begin{cases} \lambda_{\min} \left( \frac{2\beta}{4} \right) & \text{for } 0 \leq \beta \leq \frac{1}{2} \\ 1 - \beta & \text{for } \frac{1}{2} \leq \beta \leq 1 \end{cases}$$

(3.10)

where $\lambda_{\min}(\gamma)$ is defined in Eq. (1.32).
3.3 Concluding Thoughts

When we plot the quantum boundary due to $QB_3^{(8)}$ per Eq. (3.9) in Fig. 3.1 we observe a region in which this boundary is more restrictive than even $NPA^{1+AB}$. This affirms that finding that quantum contextuality allows for statistics not accessible through quantum nonlocality [36, 89].

Note that any computational error in the numeric determination of the $NPA^{1+AB}$ boundary would result in an underestimation of $\xi$ for every discrete $\beta$, such that the plotted $NPA^{1+AB}$ curve represents a lower bound, and consequently there is no ambiguity that $NPA^{1+AB}$ is plainly more permissive than $QB_3^{(8)}$ in the region highlighted in Fig. 3.1. The underestimation of $\xi$ results from the positivity constraint of the $\Gamma$ matrix of Table 3.1; the computer may falsely reject a permitted $\xi$ if it is unable to instances of free variables to make $\Gamma$ positive semidefinite in a finite number of iterations.

Thus Fig. 3.1 reiterates the fundamental limitation inherent to both the Exclusivity Principle (EP) [235, 243, 244, 246–249] and Local Orthogonality (LO) [36, 44–47] as means to recover quantum nonlocal correlations. Since those principles are expressed in terms of the graph-theoretic formalism of quantum contextuality they are innately blind to the distinction between quantum contextual and quantum nonlocal correlations.
Our identification of a broad region in condition probability space where $\text{NPA}^{1+AB}$ diverges from the predictions of quantum multipartite nonlocality opens the door to experimental testing of the true nature of quantum reality. A bipartite experiment which could trace out the $\text{NPA}^{1+AB}$ curve in Fig. 3.1, such as a generalization of that performed recently by Vermeyden et al. [37], would violate the predictions of conventional quantum mechanics. It would, instead, provide long-sought-after experimental evidence for relativistic quantum theories, such as Consistent Histories [77].

It is inspiring to recognize that the pursuit of a physical principle capable of recovering quantum nonlocality - as opposed to contextuality - is still a wide open question in quantum foundations. Nonlocality is the dominant feature exploited in the quantum information theory. Yet, it remains an unexplained consequence of the mathematics underpinning quantum mechanics. Although much is known about quantum nonlocality, its fundamental raison d’être still escapes us. The quest for a more satisfying explanation will surely lead to numerous applications in the future. Once a strong conceptual foundation is established, who can fathom the potential of the “second quantum revolution” [267, 268]?
References


Glossary

Box

See Point.

Clique

In graph theory, a clique is a set of vertices each of which is adjacent to all the others in that set. In the context of Orthogonality Graphs, a clique refers to a set of measurement outcomes (non-necessarily of the same measurement tuple) which are mutually exclusive, or contradictory. An example of a clique of incompatible measurement outcomes is given by the ten terms in the inequality (1.33). See Ref. [235] for a quick introduction to the relevant concept.

Conditional Probability Space

The abstract space of possible experimental statistical parameters, also known as statistical space. A given experimental parameter, for example, could be the probability that Alice and Bob obtain correlated measurement outputs given
that Alice uses measurement apparatus $A_1$ and Bob using apparatus $B_2$. A fixed experimental setup yields a single points in conditional probability space in the limit of infinite experimental trials. (A Finite trial count results in an uncertainty of the coordinates of the experiment in statistical space, the study of which is generally subsumed in Martingale analysis [269–272].) See Polytope.

**Consistent Exclusivity**

Consistent Exclusivity is a special case of the Exclusivity Principle when inferences on the correlations of a contextuality scenario are made by considering the possibility that independent copies of the scenario could exist simultaneously [245 and 36, Sec. 7.1]. Consistent Exclusivity is characterized by various integer levels, corresponding the number of independent copies from which inferences are being drawn. For correlations to be unambiguously consistently exclusive, they must be compatible with Consistent Exclusivity in the limit of infinite simultaneous copies. Consistent Exclusivity is directly related to the Shannon Capacity of nonorthogonality graphs [36, Sec. 7.3]. Extended Consistent Exclusivity requires additionally that the set of allowed correlations be convex and closed under wirings. Extended Consistent Exclusivity is known to perfectly recover quantum contextual - but not quantum nonlocal - correlations, in that it is equivalent to the $1 + AB$ level of the NPA hierarchy [36, Sec. 7.7]. See
**Exclusivity Principle, NPA.**

**Elliptope**

An elliptope is a higher-dimensional analog of a convex curved ellipsoid. Of special interest is the quantum elliptope in conditional probability space. The quantum boundary in conditional probability space is known to be nonlinear, but does not presently have a closed form characterization [22, 34, 35]. One can use linear Tsirelson inequalities to generate a pseudo-quantum polytope which contains the genuine quantum elliptope within it; in the infinite limit the region bounded by the linear equalities converge to the quantum elliptope per the supporting hyperplane theorem [31]. The quantum elliptope can, alternatively, be inner characterized by the convex hull of extremal quantum-achievable conditional probability distributions. *See Polytope.*

**Exclusivity Principle**

A graph-theoretic principle which states that any exclusivity graph must have vertex probabilities less than or equal to one in every orthogonality clique [235, 243–249]. Quantum mechanics innately obeys this principle, because if a set of quantum projectors are all pairwise orthogonal, than any string of such projectors, or any length, is equal to the null vector. One can, however,
imagine probabilistic theories which violate the Exclusivity Principle [2]; such correlations are not possible in a quantum reality, however. The Exclusivity Principle differs from Local Orthogonality in that it applies to contextuality scenarios as well as nonlocality scenarios. The Exclusivity Principles differs from Consistent Exclusivity in that it considers all possible extensions of the target scenario [245, 249]. See Clique.

Local Orthogonality

A first principle which partially recovers quantum correlations. Nearly identical to Consistent Exclusivity, Local Orthogonality applies only to multipartite nonlocality scenarios. It was developed by Fritz et al. [44], independently from Cabello’s Exclusivity Principle [235]. See Refs. [36, 44–47]. Local Orthogonality constructs composite scenarios through a hypergraph formalism [36], but then reduces to standard orthogonality graphs when its primary principle is applied. See Exclusivity Principle, Consistent Exclusivity.

NPA

An acronym for the three authors of the Navascués-Pironio-Acín (NPA) hierarchy [35, 52, 54]. The hierarchy converges asymptotically to the genuine set of quantum nonlocal correlations. Finite levels of the hierarchy, however, permit
additional correlations which are not achievable in a quantum nonlocality scenario. Of special interests in the $1 + AB$ level of the NPA hierarchy, as this corresponds the the correlations achievable in a single-partite quantum contextuality scenario [36, 236] as well the set of correlations recovered by the Consistent Histories theory of quantum gravity [77], the convex and closed-under-wirings variant of Consistent Exclusivity [246 and 36, Sec. 7.7], as well as various other principles [89].

**Point**

In this thesis, we reserve the use of “point” to indicate a particulate set of coordinates in *Conditional Probability Space*. Thus, a point refers to a particular set of statistics which might characterize a nonlocality or contextuality scenario. Completely equivalent is the notion of a *behavior*, or a probabilistic model, and a bipartite *box* such as the famous Popescu and Rohlich - PR - box [1, 17, 19, 20, 33]. In this thesis we consider composite boxes, that is to say, a normalized mixture of boxes where the the weights are variable parameters. Such variable boxes defines curves or surfaces in *conditional probability space*.
Polytope

A polytope is a higher-dimensional analog of a polyhedron. Of special interest are statistical polytopes in conditional probability space. For any nonlocality scenario, both the set of no-signalling conditions and the set of Bell inequalities form polytopes in conditional probability space which bound the set correlations consistent with general probabilistic theories and with local hidden variable models, respectively [1, 17, 19, 20, 33]. Polytopes are described by either the convex hull of extremal points, or equivalently, by linear equalities. The quantum boundary in conditional probability space is nonlinear, and therefore quantum correlations are described by a convex elliptope instead of a polytope. See Conditional Probability Space.

QB

An initialism of “quantum bound”. In this thesis we make reference primarily to QB$^3_8$ featured in Ref. [31] and reproduced in Table 1.1. The subscript 3 indicates that it belongs to the third class of quantum bounds in the table, whereas the superscript 8 indicates that QB$^3_{8}$ represents a boundary in the full 8-dimensional conditional probability space of the (2,2,2) scenario [31]. QB$^3_{8}$ is a function-valued quantum bound, in that it describes a contiguous collection
(or envelope) of linear quantum bounds. Linear quantum bounds receive the special designation “Tsirelson Bounds” [18], or “TB”, in this thesis.