Experimental adaptive quantum tomography of two-qubit states

I. INTRODUCTION

Quantum state tomography (QST) is a procedure allowing one to estimate an unknown state of a quantum system from the outcomes of measurements on an ensemble of identically prepared copies of this system. Since this primitive is essential for experimental quantum information studies, much effort was devoted to the optimization of state tomography. An optimal tomographic protocol should demonstrate the best possible accuracy of reconstruction for a given number of performed measurements. It is clear that adaptive measurement strategies, where the choice of the next measurement depends on the previous outcomes, may be advantageous [1]. First implementations were carried out in a Bayesian framework and used precomputed decision trees to choose optimal measurement strategies [2]. Obviously, such an approach is applicable only for very limited sample sizes. Recent advances in Bayesian methods, inspired by machine-learning applications, led to the development of fast algorithms allowing for an online determination of optimal measurements [3,4]. On the other hand, the frequentist approach makes use of the maximum-likelihood estimation and suggests determination of the optimal measurements by minimizing the variance of locally unbiased estimators [5]. The latter ideas were implemented experimentally for a single-parameter estimation [6,7]. Two recent experimental implementations [8,9] have demonstrated the quadratic enhancement in the accuracy of reconstruction, measured as the infidelity with the “true” state, for the adaptive estimation of qubit states. The experiment in [8] relied on a single adaptive step: when half of the data are collected, the most likely state is estimated and the following measurements are performed in the eigenbasis of the estimated state. The work in [9] used the fully adaptive Bayesian protocol based on maximizing the information gain for each measurement for a given Bayesian posterior distribution [3]. Other utility functions besides information gain were considered for optimization; for example, the infidelity-optimized Bayesian protocol for pure states was considered in [10]. The Bayesian framework for state estimation may be advantageous for several reasons. First of all, the posterior distribution provides the most complete statistical description of the knowledge about the state inferred from the measurements and never yields unphysical or unjustified estimates [11]. Even more importantly, Bayesian experimental design techniques offer rich possibilities for construction and optimization of adaptive tomographic protocols.

Adaptive tomography was previously implemented for single-qubit states only. The estimation of higher-dimensional states introduces new features, which should be addressed. For example, available measurements are usually restricted in a specific way, e.g., the experimental system is split into two parts and projective measurements on each part can be implemented easily, while performing the most general measurement on the whole system is experimentally challenging [12]. In this case, the optimization of measurements in the adaptive protocol should be performed over some subset. Besides that, there are purely computational difficulties arising from the exponentially increasing dimensionality of the parameter space. In this work, we address these issues and present the experimental realization of the fully adaptive Bayesian quantum state estimation protocol for two-qubit states. Our software is implemented for arbitrary dimensions and the experiment is performed for the polarization state of entangled photon pairs. Two types of measurements are considered:

1. general projective measurements,
2. factorized measurements that can be represented as a tensor product of projective measurements on subsystems.

Another issue in real experiments is the influence of the instrumental noise. We study the performance of adaptive tomography under noisy conditions and find it advantageous over nonadaptive protocols. The paper is organized as follows: first, we review the inference algorithm and design of our experimental realization in Sec. II. Then, in Sec. III, we present...
experimental results and the results of numerical simulations. Finally, the conclusions are given in Sec. IV.

II. TOMOGRAPHY ALGORITHM

Each tomographic protocol consists of two essential parts: a measurement strategy and an estimator. The measurement is characterized by a set of positive operator-valued measures (POVMs) \( \mathcal{M} = \{ \mathbb{M}_\alpha \} \), with index \( \alpha \in \mathcal{A} \) numbering different configurations of the experimental apparatus. Having the measured system in the state \( \rho \) and given the configuration \( \alpha \), the probability of observing an outcome \( y \) is given via Born’s rule:

\[
P(y | \alpha, \rho) = \text{Tr}(\mathbb{M}_\alpha \rho),
\]

where \( \mathbb{M}_\alpha \in \mathbb{M}_\alpha \) are POVM elements. We will assume that each POVM is complete, \( \sum_\alpha \mathbb{M}_\alpha = \mathbb{I} \). The estimator maps the set of all observed outcomes, \( \mathcal{D}_N = \{ y_n \}_{n=1}^N \), onto an estimation of the state \( \hat{\rho} \).

A. Bayesian approach

The Bayesian approach to quantum tomography deals with a probability distribution over the space of density matrices. First, a prior distribution \( p(\rho) \) is specified. Then, after data acquisition, a posterior probability \( p(\rho | \mathcal{D}) \) is obtained via Bayes’ rule:

\[
p(\rho | \mathcal{D}) \propto \mathcal{L}(\mathcal{D}; \rho)p(\rho),
\]

where \( \mathcal{L}(\mathcal{D}; \rho) = \prod_\alpha P(y_n | \alpha, \rho) \) is a likelihood function, which incorporates a statistical model. It is equal to the probability of observing data \( \mathcal{D} \) given the state \( \rho \). The posterior distribution provides the most complete description of our knowledge about the quantum state, inferred from data \( \mathcal{D} \).

The posterior distribution can be utilized for both statistical assessment and constructing a measurement strategy. Point estimates of desired quantities are found as expectations over the posterior distribution. Their error bars are given by variances. For example, we may obtain the Bayesian mean estimate (BME) of the state \( \hat{\rho} = \mathbb{E}_{p(\rho | \mathcal{D})}[\rho] \). The measure of its uncertainty is the distribution size \( \bar{x} = \mathbb{E}_{p(\rho | \mathcal{D})}[d(\rho, \hat{\rho})] \), where \( d(\cdot, \cdot) \) is a distance between two states. In our analysis, we used Bures metric \( d_B \), which is closely related to the fidelity \( F(\rho, \hat{\rho}) = \text{Tr}^2 \sqrt{\rho^{1/2} \hat{\rho} \rho^{1/2}} \) [13]: \( d_B^2(\rho, \hat{\rho}) = 2 - 2\sqrt{F(\rho, \hat{\rho})} \approx 1 - F(\rho, \hat{\rho}) \) for \( 1 - F \ll 1 \).

Prior. For the purposes of quantum tomography, the prior distribution over the state space should be noninformative or uniform. It is natural to identify uniformity with maximal symmetry. In the case of pure states, such prior distribution is unique and is determined by the requirement of unitary invariance [14]. Therefore, the Haar measure is used. But for mixed states, unitary invariance only implies the Haar measure over eigenvectors of density matrices, leaving the distribution in the simplex of eigenvalues \( \lambda_i \) undetermined (each density matrix has a unit trace, so a set of eigenvalues forms a simplex \( \{ \lambda_i : \sum \lambda_i = 1 \} \)).

There are several approaches to specify a probability distribution of eigenvalues [15]. The first one is based on a purification procedure: the \( D \)-dimensional system of interest is supplemented with a \( K \)-dimensional environment, and the composite \( D \times K \) system is assumed to be in a pure state. The measure on the composite system is assumed to be known (Haar measure) and after tracing out the environment, it induces a certain measure \( d\mu_{DK} \) in the state space of the initial system. The second approach relies on the fact that each metric \( d \) generates a measure \( d\mu_{ij} \): any ball with a fixed radius has the same measure. Interestingly, the measure \( d\mu_{DK} \) induced by partial tracing for \( D = K \) coincides with the measure induced by the Hilbert-Schmidt distance \( d_{HS}^2(\rho, \hat{\rho}) = \text{Tr}((\rho - \hat{\rho})^2) \); \( d\mu_{HS} = d\mu_{DD} \).

We shall consider the following prior distributions [16]:

(i) induced by Bures distance,

\[
p_B(\rho) \propto \prod_{i=1}^D \frac{1}{\sqrt{\lambda_i}} \prod_{i<j} \frac{1}{\lambda_i + \lambda_j},
\]

(ii) induced by Hilbert-Schmidt distance,

\[
p_{HS}(\rho) \propto 1,
\]

(iii) uniform in the eigenvalues simplex,

\[
p_\Delta(\rho) \propto \prod_{i<j} \frac{1}{(\lambda_i - \lambda_j)^2}.
\]

After integrating over eigenvectors, one obtains the probability distributions \( p(\lambda_1, \ldots, \lambda_D) \) in the eigenvalues simplex. The expressions are formally similar to the right-hand sides of (3)–(5), except for the appearance of a geometric factor \( \prod_{i<j} (\lambda_i - \lambda_j)^2 \). For instance,

\[
p_{HS}(\lambda_1, \ldots, \lambda_D) \propto 1 \times \prod_{i<j} (\lambda_i - \lambda_j)^2.
\]

It is clear that \( p_\Delta(\lambda_1, \ldots, \lambda_D) \propto 1 \) is indeed a uniform distribution in the eigenvalues simplex.

This geometric factor causes zero probability for the states with degenerate eigenvalues in both Hilbert-Schmidt and Bures distributions. At the same time, degenerate states are important, as they appear when a pure state \( |\psi\rangle \) passes through a depolarizing channel \( |\psi\rangle \langle \psi | \rightarrow (1 - \varepsilon)|\psi\rangle \langle \psi | + \varepsilon|\phi\rangle \langle \phi | \) for \( \varepsilon I / D \), where \( I / D \) is a \( D \)-dimensional completely mixed state. Thus, the third, “unnatural” prior is introduced essentially to avoid such gaps at degenerate states. In Appendix A, it is shown that the accuracy of a degenerate state tomography depends rather strongly on the prior choice and a potential explanation of this behavior is given.

B. Adaptivity criterion

The Bayesian approach is a natural framework for constructing an adaptive measurement strategy. Indeed, the posterior can be updated as soon as a new outcome is observed, and given this new knowledge about the state one may find the next optimal measurement. There are several criteria for optimal experiment design known in the literature (see Ref. [17] for a review). One possible way is to choose an experimental configuration \( \alpha \) that is expected to reduce Shannon entropy \( \mathbb{H} \) of the posterior distribution the most. Usage of Shannon entropy as an objective function allows one to reformulate this criterion as the following optimization procedure (which is
simpler to work with in practice) [3]:
\[ \alpha = \text{argmax}_\gamma \{ \mathbb{H}[\mathbb{P}(y|\alpha, D)] - \mathbb{E}_{p(\rho|D)}[\mathbb{H}[\mathbb{P}(y|\alpha, \rho)]] \}, \tag{7} \]
where \( \mathbb{P}(y|\alpha, D) = \int \mathbb{P}(y|\alpha, \rho)p(\rho|D)d\rho = \mathbb{P}(y|\alpha, \hat{\rho}) \) is the expected probability of outcome \( y \). In our experimental realizations, the maximization is performed among all POVMs \( \gamma \) of the approximation drops down. To monitor the situation, the algorithm proceeds, the weights tend to collapse, i.e., all but a small number of them become essentially zero. Consequently, the quality of the approximation drops down. To monitor the situation, the effective sample size \( S_{\text{eff}} = (\sum_{i=1}^{N} w_i^2)^{-1} \) is calculated. If it is below a certain threshold, then resampling is performed and the weights are equalized. A detailed description of the resampling procedure is provided in Appendix C.

At the start of the protocol, samples are generated based on the chosen prior. For Hilbert-Schmidt \( p_{\text{HS}}(\rho) \) and Bures \( p_B(\rho) \) induced priors, we utilized algorithms from [20, 21], while the prior uniform in the eigenvalues simplex \( p_\Delta(\rho) \) was constructed in two steps: first, samples are generated according to \( p_{\text{HS}}(\rho) \) and, second, samples are moved by the resampling routine with the probability distribution (5).

**Block measurements.** Achieving the best performance implies application of the adaptivity criterion after each successful measurement. However, this may be impractical due to the excessive burden of changing the measurement configuration. Instead, one can use block measurements by keeping the same configuration for \( b \) consecutive measurement outcomes. Instead, we provide a direct connection between the number of independent measurements \( m \) made so far and the number of accumulated outcomes \( N \). Based on the asymptotics \( b(N) \approx dN/dm \), a useful expression applicable for \( N \geq 100 \) is \( m \approx 50 \ln N - 160 \).

There is some evidence that a block size may be as large as \( O(N) \). Let us denote, for simplicity, \( d_{\text{true}}(N) \equiv d_B^0(\hat{\rho}(N), \rho_0) \), which is the distance between the estimator \( \hat{\rho}(N) \) and the true state \( \rho_0 \). It was shown in [9] that the measurement basis is optimal if it is aligned according to the true state with the error \( d_{\text{align}}^2 \) smaller than \( d_{\text{true}}^2(N) \): \( d_{\text{align}}^2 = kd_{\text{true}}^2(N), \ 0 < k < 1 \). After measuring the block of maximal size \( b \), the basis ceases to be optimal: \( d_{\text{align}}^2 = d_{\text{true}}^2(N + b) \). With these two equations at hand, we obtain \( d_{\text{true}}^2(N + b) = kd_{\text{true}}^2(N) \). Assuming the behavior \( d_{\text{true}}^2(N) = cN^a \) (see Sec. III), finally, we find that the block size scales linearly with \( N \):
\[ b = (k^{1/a} - 1)N = O(N). \]

Interestingly, our simulations show that the amount of information extracted during one block \( b \propto N \) [the entropy gain under maximization in (7) times the block size \( b \)] holds approximately constant, while the tomography proceeds. This can be used to construct a rule for selecting a block size on the fly: the block lasts until the given amount of information is not gathered. Another valid approach, where changes of the measurement configuration are based on its deviation from the optimal one, was used in [10]. The authors have shown that on average, the number of configuration changes \( m \) grows logarithmically with \( N \), and therefore the block size \( b \approx dN/dm \) again scales linearly with \( N \).

**III. IMPLEMENTATION**

**A. Experiment**

The described protocols are implemented experimentally using polarization degrees of freedom of photon pairs produced by spontaneous parametric down-conversion (SPDC). The sketch of the experimental setup is shown in Fig. 1. State preparation is performed by pumping two nonlinear 2-mm-thick BBO crystals with linearly polarized light.
A Littrow configuration [22] external cavity-diode laser (408 nm) is used as a pump. The pump power just before the crystals is 7.0 mW. The Faraday isolator OI serves for both blocking parasitic light, reflected backwards, and preparing the initial linear polarization. This polarization is then rotated by means of two wave plates WP5-6. The crystals are cut for a slightly noncollinear frequency-degenerate type-I phase matching. Such configuration allows us to prepare polarization-entangled states of variable degree of entanglement and purity [23]. The purity of the generated state can be altered by changing the size of the collecting iris A1-2 or by inserting the birefringent compensators C1-2 with high wavelength-dependent phase shifts after the crystals [24,25]. A degree of polarization entanglement is controlled by rotating the linear polarization of the pump. A typical coincidence count rate was 130 s$^{-1}$. Achieving better state accuracy required smaller collecting apertures, so the best obtained fidelity of the Bell state, $F = 0.958$ (purity Tr $\rho^2 = 0.931$), was observed at a count rate of 20 s$^{-1}$.

The long pass filter F (450 nm) and the interference filter IF (820±5 nm) are used to block the pump and to select a frequency-degenerate range. The two components of generated photon pairs propagate through similar channels, which consist of wave plates WP1-4 (zero-order half- and quarter-wave plate in each channel) mounted on computer-controlled rotation mounts and Wollaston prisms WL1-2. Finally, they are coupled to multimode fibers connected to single-photon counting modules (SPCMs) D1-4. This setup allows us to make any factorized four-dimensional projective measurement (i.e., realize projectors onto $|\psi\rangle = |\psi_1\rangle \otimes |\psi_2\rangle$, where $|\psi_{1,2}\rangle$ are the polarization states of photons in channels 1 and 2).

We have compared the performance of the adaptive measurement strategy with the random measurements strategy that is known to be optimal in a nonadaptive case [26], Theorem 3.1]. A comparison of protocols was carried universally within the Bayesian framework. The Bayesian mean state $\hat{\rho} = E_{\rho|D}|D\rangle\langle D|$ was used as an estimate of the true state. A natural figure of merit for any estimation technique is the distance $d_{B}^2(\hat{\rho},\rho_0)$ between the estimation $\hat{\rho}$ and the true state $\rho_0$. However, in a practical tomography, the true state is usually unknown and the tomography provides the best estimation at hand: e.g., one may perform a very long series of measurements and use the final result as the “true” state. Its accuracy, though, is still limited by the experimental imperfections. Therefore, instead of using distance to the true state $d_{B}^2(\hat{\rho},\rho_0)$ as a measure of accuracy, we turned to its assessment—the posterior distribution size $d_B^2 = E_{\rho|D}[d_{B}^2(\hat{\rho},\rho_0)]$. In Appendix B, it is shown that the distribution size $d_B^2$ is, in fact, connected with $d_{B}^2(\hat{\rho},\rho_0)$ and thus provides a reasonable estimation.

Figure 2 shows the dependence of the posterior distribution size $d_B^2(N)$ on the number $N$ of the detected pairs for the factorized random (FR) and factorized adaptive (FA) projective measurements. The data are a result of averaging over many (typically 15) single full runs of the tomography protocol. These results clearly demonstrate an advantage of the adaptive strategy over the random one for both factorized [Fig. 2(a)] and maximally entangled [Fig. 2(b)] state tomography. The results of fitting the data with a power law in the form $cN^n$ are presented in Table I. The adaptive protocol clearly demonstrates an improved distribution size scaling in the Bures metric, although it is not exactly the expected $N^{-1}$ dependence. We attribute this reduction in performance to two main reasons: the inaccuracies in the wave-plate positioning (on the order of 0.2°) and deviations of the measured states from truly pure. The former was previously observed (see supplemental information of [8]) and is known to reduce the performance of the adaptive protocol, while the latter, which is studied below in the simulations section, is due to the degraded behavior of the restricted factorized adaptive protocol for mixed states compared to the ideal adaptive tomography; thus, the observed reduction of the convergence rate is also expected. There is an apparent saturation of the FA protocol curve for a factorized state [Fig. 2(a)]. The most probable explanation is that a state under the tomography has happened to cause a plateau in the dependence $d_{B}^2(N)$ (see Appendix A).

We have experimentally observed that the adaptive tomography of two qubits is less sensitive to instrumental errors than a nonadaptive one. Previously, the same was shown for a single-qubit tomography by means of numerical simulations [8]. The following noise model was used: each time the wave plates WP1-4 are rotated, their positions are intentionally set to $\theta_i + \delta\theta_i$, rather than to the exact angles $\theta_i$ used by the estimation algorithm. Here, $\delta\theta_i$ is a random variable uniformly distributed in the range $[-\Delta\theta, \Delta\theta]$. Again, due to the absence of knowledge about the true state, the quantity $d_{B}^2$, which we will refer to as spread over different runs of tomography, was used. Let $\hat{\rho}_k(N)$ be the Bayesian mean estimate in the $k$th experimental run. Then the

![FIG. 1. Experimental setup. The entangled photon pairs are generated in two BBO crystals with perpendicular axes, pumped by linearly polarized light from a 408 nm diode laser. Projective measurements are performed separately for each photon of the pair by means of half- and quarter-wave plates followed by Wollaston prisms.](image-url)
stands for factorized random measurements and FA stands for factorized adaptive measurements. The shaded areas correspond to one standard deviation for the instrumental noise in the experimental run: no additional noise (blue), adaptive tomography. Color encodes different amounts of additional "software" noise is added, and the noise floor is determined by the intrinsic errors in positioning introduced by wave-plate rotators ($\approx 0.2\degree$). For moderate statistics $N \lesssim 10^3$, there is no visible difference between adaptive and random protocols tend to converge. In our case, the difference becomes negligible for $\Delta \theta = 5\degree$. It is worthwhile to mention that even for $N = 10^5$, there is no visible saturation for the adaptive protocol without additional noise ($\Delta \theta = 0\degree$), while the random protocol reaches its noise floor at $N \approx 10^4$.

Reduced sensitivity to errors in wave-plate positioning for the adaptive tomography can be intuitively understood in the following way. If the exact probabilities of a tomographically complete set of measurements are known, then one can easily reconstruct the true state. In real-world applications, probabilities can only be determined with some finite accuracy. The higher the accuracy, the better the assessment of the true state. Therefore, the goal is to provide the best estimation of probabilities. Suppose we perform a tomography of a state $\rho_0$, and POVM elements $M_{\alpha \gamma} = M_{\alpha}(\theta_i)$ are parameterized by parameters $\theta_i$ (wave-plate angles in our case). The adaptive tomography tends to select parameters $\theta_i$, which give low probabilities of outcomes $p_A(\theta_i)$ (see Sec. II B). Therefore, $p_A(\theta_i)$ take extremal values, and error in probability estimation $\delta p_A$ depends quadratically on the uncertainties in parameters $\delta \theta_i$: $\delta p_A = O(\delta \theta_i^2)$. This is not the case for nonadaptive protocols, unless the POVM is aligned with the true state [8]. For nonadaptive tomography, typically $\delta p_R = O(\delta \theta_i) \gg \delta p_A$.

B. Simulations

The chosen experimental approach allows factorized projective measurements only, which we refer to as F-class measurements. Projective measurements of a general form (G class), without factorization constraints, could also be implemented, although with a significantly larger experimental effort [12]. Since G class is much wider, it is reasonable to expect higher accuracy of the state estimation. In this section, we focus on a question of whether or not an extension $F \rightarrow G$ dominates over adaptivity. Several results for pure states have been previously known from the literature. It
was shown [12] that nonadaptive measurements in mutually unbiased bases (which belong to G class because they require projections onto entangled states) can improve precision for some states. On average, general measurements are not worse than factorized ones (symbolically, one can write GR ≥ FR, where “larger” means more accurate). However, the situation is different for adaptive protocols [3]: on average, factorized adaptive measurements outperform general nonadaptive ones (FA > GR). We have performed numerical simulations of four protocols with different types of measurements: factorized random (FR), factorized adaptive (FA), general random (GR), and general adaptive (GA). Our goal was to compare the average performance of Bayesian tomography for these four types.

Let us start with the pure states. For each protocol, we have averaged over 1000 runs with different pure true states, uniformly distributed with respect to the Haar measure. The simulation results are shown in Fig. 4. We have tried various priors, defined according to Eqs. (3)–(5), and found that the prior choice only changes the accuracy by a multiplicative constant, thus preserving a relationship between different protocol performances (see Appendix A for details). Here we present the results for the prior uniform in the eigenvalues simplex $p_A(\rho)$. As a figure of merit, the distance $d_B^2(\hat{\rho}(N), \rho_0)$ between a current estimate $\hat{\rho}(N)$ and the true state $\rho_0$ is used [Fig. 4(a)]. It is also instructive to show histograms for the values of $N$ for which a fixed accuracy $d_B^2 = 10^{-3}$ is reached in different runs [Fig. 4(b)]. The results indicate that the determining factor is adaptivity because the performance is independent of the measurement class F or G. There is an obvious enhancement in the estimation accuracy for adaptive protocols. Power-law fits of the dependencies $d_B^2(\hat{\rho}(N), \rho_0)$ of the form $cN^a$ give convergence rates $a = -0.958 \pm 0.008$ for adaptive protocols and $a = -0.588 \pm 0.011$ for the random ones, which is close to the theoretically expected scalings $N^{-1}$ and $N^{-1/2}$, respectively. To summarize, for pure states we have a relation $GA = FA > GR = FR$—on average, there is no advantage in using general measurements.

As genuine pure states are experimentally unfeasible, it is important to study how the protocol performs for the
general form mixed states. The results of simulations averaged 
over mixed states are shown in Fig. 5. The dependencies 
represent the average behavior over 1000 true states, randomly 
distributed with respect to the Bures-induced measure. The 
results are different from those for the pure states: GA \(>\) FA = 
GR \(>\) FR. Only the GA protocol provides solid enhancement 
with \(N^{-1}\) scaling (\(\alpha = -1.00 \pm 0.02\)) for nonpure states, 
while the other three protocols behave nearly similarly and 
perform at the \(N^{-3/4}\) level (\(\alpha = -0.77 \pm 0.03\)). Previously, 
the average convergence \(N^{-3/4}\) was predicted for nonadaptive 
mixed state tomography of qubits [27]. The accuracy of the GA 
protocol remains unaffected as expected, but the performance 
of the FA protocol is worse for mixed states as compared to 
pure ones. This is rather surprising because, in the single-qubit 
tomography, it is known that mixed states are in some sense 
easier to estimate than pure [8]. Therefore, naively extending 
this result to the high-dimensional tomography, one may 
expect that accuracy will at least not be reduced for mixed 
states. However, this is not the case for the FA protocol. At the 
same time, the accuracy of nonadaptive protocols for mixed 
states is improved.

From continuity considerations, it is clear that for pure 

enough states, the pure state tomography behavior GA = 

FA > GR = FR should still hold. To understand the 
transition between the two, we have performed averaging over 
several layers of states with different purity, \(\text{Tr} \rho^2 = \text{const}\). 
To construct each layer, we effectively cut 10 appropriate 
true states from the Bures distribution. Then the distribution 
size dependence \(d_{GR}^2(N)\) is averaged over these states. Value of \(N\) required to reach the specified precision, measured by the 
distance to the true state of \(d_{GR}^2(N) = d_{err}^2 = 5 \times 10^{-3}\), is 
plotted against the purity in Fig. 6. For FA and FR protocols, 
we also plot the experimentally obtained points. They are in 
reasonable agreement with the simulated data. For purities 
above \(p_{\text{pure}} \approx 0.94\), the advantage of general measurements 
is negligible and the results for pure states hold (GA = FA 

> GR = FR). For purities below \(p_{\text{pure}}\), general measurements 
are preferable for either adaptive (GA > FA) or nonadaptive 
protocols (GR > FR). On the other hand, the adaptivity 
is still dominating the accuracy (FA > GR) for purities 
above \(p_{\text{adapt}} \approx 0.7\). However, the boundaries \(p_{\text{adapt}}\) and \(p_{\text{pure}}\) 

themselves depend on the selected error level \(d_{err}^2\). For lower 
error levels, \(p_{\text{adapt}}\) shifts towards smaller values, due to the 
different scaling for adaptive and nonadaptive protocols. On 
the contrary, \(p_{\text{pure}}\) approaches unity because nearly pure states 
become tomographically distinguishable from truly pure ones 
for smaller \(d_{err}^2\) (for example, \(p_{\text{pure}} \approx 0.98\) for \(d_{err}^2 = 10^{-3}\)).

IV. CONCLUSION

We have demonstrated an experimental realization of a 
fully adaptive Bayesian state estimation protocol for two-qubit 
states. Even with the restriction of the experimentally attractive 
factorized measurements performed separately on each qubit, 
the protocol shows significant improvement of the estimation 
accuracy compared to the optimal nonadaptive protocol. This 
improvement is most significant for pure and nearly pure states. 
Achieving the advantage for mixed states, though, requires 
implementation of the joint entangling measurements on both 
qubits. This is somewhat unexpected because mixed states are 
known to be easier to estimate with nonadaptive strategies 
[8]. However, the most interesting area for quantum informa-
tion applications—nearly pure states—still shows very good 
performance, so the advantage of the general measurements 
probably does not justify the experimental efforts for their 
realization. Thus, the factorized adaptive protocol seems to be 
the optimal choice, considering the trade-off between accuracy 
and challenging implementation.

A natural question is whether it is possible to scale the 
protocol to higher dimensions. The straightforward answer 
will be that it is probably not too far, since the computation 
complexity grows exponentially with the dimensionality of the 
Hilbert space of the reconstructed system. This is, however, 
the case for any tomographic protocol, so the only extension here 
is the search for an optimal measurement, which is by itself 
not more demanding than the estimation of the posterior. All 
of the computations, performed for the experiment presented 
here, were done online during the experimental run with a 
standard desktop computer, so the required computational 
power for two qubits (15-dimensional parameter space) is 
reasonable. Some modifications may be introduced as well to 
allow running the algorithm for several qubits with tolerable 
performance. For example, the time-consuming resampling 
procedure may be simplified by relaxing the requirement for 
exact sampling from the likelihood function, and substituted 
by some approximation such as the Liu-West resampling 
algorithm [28], which only preserves the first and the second 
moment of the sampled distribution, as it was done for 
Hamiltonian learning in [4]. This will push the tractable 
dimension boundary a little bit higher; however, “the curse 
of dimensionality” is inevitable, and one should not expect 
that the full tomography of a several-dozen-qubit state will 
be feasible. Other avenues of research may be pursued here, 
such as restricting the tomography to some physically relevant 
subspace, e.g., matrix-product states [29], using compressed 
sensing techniques for low-rank states [30] or permutationally
invariant tomography of multipartite systems [31]. It will be interesting to investigate the advantages of self-learning techniques, like those presented here, offered in these settings. In general, we expect to see new interesting applications for machine-learning and Bayesian methods in quantum state and process tomography and in other related areas.

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APPENDIX A: PRIOR INFLUENCE

Though tomography accuracy, measured as a posterior average Bures distance to the true state averaged over some ensemble of states, varies by a (small) multiplicative constant for different priors, a pointwise behavior can change dramatically. The reason is that a neighborhood of a true state may include states which are rare or exceptional in the given prior, i.e., have low probability. Among the considered priors (3)–(5), Bures and Hilbert-Schmidt distributions assign zero probability to states with degenerate eigenvalues [see Eq. (6)]. This behavior is related to the geometry of the density matrix space [32]. One can cancel out the geometric factor, obtaining, for example, a uniform in the eigenvalues simplex prior \( p_\Delta(\lambda) \propto 1 \), which assigns nonzero probability for all states.

The dependence of the average distance to the true state \( d_B^2(\hat{\rho}(N), \rho_0) \) on \( N \) is compared for the Bures and simplex priors for the FA protocol in Fig. 7. The Hilbert-Schmidt prior gives rise to a similar behavior as the Bures one. The true state is chosen to have a diagonal density matrix with a triply degenerate eigenvalue: \( \lambda_i = \{0.9925, 0.0025, 0.0025, 0.0025\} \) (this state has a purity of \( \approx 0.985 \)). For the FA protocol with the Bures prior, the scaling is close to \( N^{-1} \) in the beginning but then a plateau occurs. When a sufficient amount of data is obtained, the normal behavior \( N^{-1} \) restores; however, the prefactor \( c \) becomes larger by two orders of magnitude. As a consequence of the plateau existence, accuracy does not increase during a long interval \( 2 \times 10^5 \lesssim N \lesssim 4 \times 10^6 \), and it looks like tomography is useless in this range. The simplex prior recovers the situation: the scaling \( N^{-1/2} \) and the initial prefactor remain the same during the whole tomography run.

As a reference, we have utilized the maximum likelihood estimator (MLE; numerical algorithm is the one from [33]), which does not involve any priors inherent to the Bayesian inference (though we used an optimal measurement set found by the FA protocol with the simplex prior). As compared to the BME with the Bures prior, the MLE demonstrates a \( N^{-1/2} \) scaling instead of the plateau; nevertheless, the accuracy is worse. Asymptotically, at large \( N \), the MLE behaves similarly to the BME within error bars. Altogether, the MLE is not better than the BME.

We have presented the results for an extreme case of a triply degenerate density matrix. For the states which lie farther from degenerate ones, the plateau becomes shorter. A position of the plateau \( d_{\text{th}}^2 \) is proportional to a distance between the true state \( \rho_0 \) and the border of the space of physical density matrices, formed by rank-3 states: \( d_{\text{th}}^2 = \min_{\rho_\text{rank3}} d_B^2(\rho_0, \rho) \approx \lambda_{\text{min}} \) for \( \lambda_{\text{min}} \ll 1 \), where \( \lambda_{\text{min}} \) is the minimal eigenvalue of the true state. It is worthwhile to mention that the plateau shape is protocol dependent: for GA, it is significantly shorter.

For critical applications, to achieve the best performance, we suggest construction of an informative prior from physical considerations, e.g., based on a particular decoherence model for a given system. For a recent recipe of constructing useful informative priors, see [34].

APPENDIX B: DISTRIBUTION SIZE

In order to verify that the distribution size \( \bar{d}_B^2 \) can be considered as an appropriate estimation of the tomography accuracy \( d_B^2(\hat{\rho}, \rho_0) \), we studied how the ratio \( R_{\text{dd}} = \bar{d}_B^2(\hat{\rho}, \rho_0)/\bar{d}_B^2 \) of the distance to the true state and the distribution size evolves with \( N \) in numerical simulations. The ratios \( R_{\text{dd}}(N) \), found for single tomography runs with different true states, randomly chosen from the Bures distribution, have been averaged over 1000 runs. The behavior of the averaged ratio \( R_{\text{dd}} \) for \( 10^3 \) and \( 10^4 \) samples used in the Monte Carlo calculations is compared in Fig. 8. For \( 10^3 \) samples, the ratio holds almost constant, not larger than 2, until \( N \approx 10^5 \), so the accuracy \( d_B^2(\hat{\rho}, \rho_0) \) in an experiment can be estimated by means of the distribution size \( \bar{d}_B^2 \). For larger \( N \), the sampling technique should be refined by increasing the number of samples to guarantee that the ratio remains constant. Interestingly, the behavior of \( R_{\text{dd}}(N) \) is protocol dependent: for the GA protocol, \( 10^3 \) samples are sufficient within the whole interval \( 1 \leq N \leq 10^5 \), while for the FA protocol, this is not the case because the ratio starts to increase heavily from \( N \approx 10^5 \).

The distance to the true state \( d_B^2(\hat{\rho}, \rho_0) \) is a coarse estimate of the tomography accuracy because it gives only one number to characterize a region in the state space, where the true state is likely situated; it does not provide any information about the shape of the region. A more rigorous approach in
Bayesian tomography is to use region estimates, e.g., posterior covariance ellipsoids [35], to specify the credible region.

**APPENDIX C: RESAMPLING**

The resampling procedure is used to redraw samples \( \{ \rho_i \}_{i=1}^S \) from the posterior distribution \( p(\rho|D) \) when the effective sample size \( S_{\text{eff}} = (\sum_{i=1}^S w_i^2)^{-1} \) is below a certain threshold \( S_{\text{eff}} < S_{\text{thr}} \). In our inference, we typically use \( S = 1000 \) samples and a threshold of \( S_{\text{thr}} = 0.1S \). The resampling algorithm consists of several steps:

1. Pick a particle \( \rho_i \) from the sample distribution with a probability \( w_i \), repeat \( S \) times. At this point, particles with tiny weights are eliminated and the ones with high weights are duplicated;

2. equalize all weights \( w_i := 1/S \);

3. move each particle according to the Metropolis-Hastings (MH) algorithm [36] to sample the posterior distribution correctly.

The MH procedure requires a random step rule \( \rho_0 \rightarrow \rho \) [also called a proposal distribution \( Q(\rho|\rho_0) \)], according to which a candidate particle \( \rho^\prime \) is generated. This particle is then accepted or rejected based on the acceptance ratio

\[
r = \frac{p(\rho'|D) Q(\rho|\rho')} {p(\rho_0|D) Q(\rho'|\rho_0)}. \tag{C1}
\]

Here the posterior probability \( p(\rho|D) \) has to be calculated using a full-likelihood function (2). This is a time-consuming task and therefore a trade-off between resampling accuracy and calculation speed exists. In our implementation, 50 MH accept or reject iterations were found to be sufficient.

The random walk of the MH algorithm relies on the purification procedure [11]. Every density matrix \( \rho_0 \) can be purified in a space of extended dimension \( D \times D \), i.e., \( \rho_0 = Tr_2 |\psi_0\rangle \langle \psi_0| \), where \( Tr_2 \) is a partial trace over the auxiliary system and \( |\psi_0\rangle \) is a pure state in the extended system. “Cholesky-like” matrix decomposition \( \rho_0 = AA^\dagger \) for the density matrix is useful to perform the purification in practice. Indeed, one can easily check that \( |\psi_0\rangle = \sum_i A_i |i\rangle |j\rangle \), with \( |i\rangle |j\rangle \) being the standard computational basis. Once the purification \( |\psi_0\rangle \) is found, the random step \( W \), followed by partial tracing, is applied,

\[
|\psi\rangle = W(|\psi_0\rangle) = a|\psi_0\rangle + b \frac{|g\rangle - |\psi_0\rangle \langle \psi_0|g\rangle} {\|g\rangle - |\psi_0\rangle \langle \psi_0|g\rangle\|}, \tag{C2}
\]

where \( a, b \) are the parameters controlling the step size (\( a^2 + b^2 = 1 \) and \( |g\rangle \sim N(0,1) \) is a vector with real and imaginary parts of its elements being independent and identically distributed random variables with normal distribution \( N(0,1) \)). We choose a Gaussian step size in Bures metric by setting \( a = 1 - d^2/2 \) with \( d \sim N(0,\sigma) \). To achieve a nearly constant fraction of accepted steps, the standard deviation \( \sigma \) was set proportional to the posterior distribution size \( d_B \).

Let us multiply the random operator \( W \) by an arbitrary unitary matrix \( U \). Then the generated probability distribution does not depend on the order of \( W \) and \( U \) in the product (however, the commutator \( [W,U] \neq 0 \)):

\[
UW(|\psi_0\rangle) \sim W(U|\psi_0\rangle). \tag{C3}
\]

This can be proved by a substitution of the explicit form of \( VW \) into (C3). The left-hand and the right-hand sides will be equal up to the substitution \( |g\rangle \rightarrow U^\dagger |g\rangle \). However, \( U^\dagger |g\rangle \sim N(0,1) \) is again from the same ensemble of Gaussian vectors as \( |g\rangle \) is. Let us consider \( U \) in (C3) as an arbitrary unitary operator with a fixed point: \( U|\psi_0\rangle = |\psi_0\rangle \). Then we find that the distribution of vectors \( |\psi\rangle = W(|\psi_0\rangle) \) is invariant under such unitary transformations. Thus, the expression (C2) generates a “spherical” neighborhood of the center \( |\psi_0\rangle \) with a radial probability distribution determined by \( a \). This property is desired for the MH algorithm because it allows one to explore the space more isotropically, reducing the required number of iterations.

In our case, the random step procedure is based on the purification procedure and hence induces a uniform Hilbert-Schmidt probability distribution \( p_{HS}(\rho) \propto 1 \). As a result, the second multiplier in (C1) vanishes: \( Q(\rho|\rho')/Q(\rho'|\rho) = 1 \). Indeed, if one sets a uniform probability distribution \( p_{MH}(\rho) \equiv 1 \) as a desired probability in the MH routine \( p(\rho|D) \propto p_{MH}(\rho) \), then \( r = 1 \), i.e., all steps are accepted. So the algorithm will converge to the distribution induced by a random walk \( p_{HS}(\rho) \), which coincides with the desired distribution \( p_{MH}(\rho) \).

For other types of a random walk, the second multiplier in (C1) may not be equal to 1. Nevertheless, we believe that one can force the multiplier to unity and use a random step that inherently generates, for example, the Bures uniform distribution. The development of such random step rule would be an eligible task.

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