Wang, J., Paesani, S., Santagati, R., Knauer, S., Gentile, A. A., Wiebe, N., Petruzella, M., O’Brien, J. L., Rarity, J. G., Laing, A., & Thompson, M. G. (2017). Experimental quantum Hamiltonian learning. Nature Physics, 13(6), 551-555. https://doi.org/10.1038/nphys4074
Efficiently characterising quantum systems [1–3], verifying operations of quantum devices [4–6] and validating underpinning physical models [7–9], are central challenges for the development of quantum technologies [10–12] and for our continued understanding of foundational physics [13, 14]. Machine-learning enhanced by quantum simulators has been proposed as a route to improve the computational cost of performing these studies [15, 16]. Here we interface two different quantum systems through a classical channel — a silicon-photonics quantum simulator and an electron spin of performing these studies [15, 16]. Here we interface two different quantum systems through a classical channel — a silicon-photonics quantum simulator and an electron spin — and use the former to learn the latter’s Hamiltonian via Bayesian inference. We learn the salient Hamiltonian parameter with an uncertainty of approximately $10^{-3}$. Furthermore, an observed saturation in the learning algorithm suggests deficiencies in the underlying Hamiltonian model, which we exploit to further improve the model itself. We go on to implement an interactive version of the protocol and experimentally show its ability to characterise the operation of the quantum photonic device. This work demonstrates powerful new quantum-enhanced techniques for investigating foundational physical models and characterising quantum technologies.

In science and engineering [17, 18], physical systems are approximated by simplified models to allow the comprehension of their essential features. The utility of the model hinges upon the fidelity of the approximation to the actual physical system, and can be measured by the consistency of the model predictions with the real experimental data. However, predicting behaviour in the exponentially large configuration space of quantum systems is known to be intractable to classical computing machines [19, 20]. A fundamental question therefore naturally arises: How can underpinning theoretical models of quantum systems be validated?

To address this question, quantum Hamiltonian learning (QHL) was recently proposed [15, 16] as a technique that exploits classical machine learning with quantum simulations to efficiently validate Hamiltonian models and verify the predictions of quantum systems or devices. QHL is tractable in cases in which other known methods fail because quantum simulation is exponentially faster than existing techniques [19–21] for simulating broad classes of complex quantum systems [22–26].

Our experimental demonstration of QHL uses a digital quantum simulator [20] on a programmable silicon photonic circuit, shown in Figs. 1a–c, to learn the electron spin dynamics of a negatively charged nitrogen-vacancy (NV) centre in bulk diamond, shown in Figs. 1d,e. We further demonstrate an interactive QHL protocol that allows us to characterise and verify single-qubit gates using other trusted gates on the same quantum photonic device.

Silicon quantum photonics is a promising platform for the realisation of manufacturable quantum technologies [27–30]. Our silicon device integrates entangled photon generation, projective measurements, single-qubit and two-qubit operations onto a single chip, as shown in Fig. 1c. This device implements the quantum circuit in Fig. 1b. Photons are generated and entangled in the path-encoded state $(|0\rangle_0|0\rangle_1 + |1\rangle_0|1\rangle_1)/\sqrt{2}$, with $s$ and $i$ indicating signal and idler photons [29]. Then the idler photon is prepared in the state $|\psi_i\rangle$ and undergoes an arbitrary unitary evolution, $U$ or $V$, conditional upon the logical state of the signal photon [31]. This entangled state $(|0\rangle_0U|\psi_i\rangle + |1\rangle_0V|\psi_i\rangle)/\sqrt{2}$ is realised upon the coincidental detection of the idler photon indicated by the blue dots, and the signal photon indicated by the red dots in Fig. 1c. The overlap between $U|\psi_i\rangle$ and $V|\psi_i\rangle$ is evaluated measuring the control qubit, enabling the estimation of the likelihoods for our QHL implementations. More details on the silicon photonic device are provided in Methods and Suppl. Info. 1.

The solid-state spin-qubit dynamics [32–36] under test are between the $m_s = 0$ and $m_s = -1$ states of the electron ground-state triplet (Fig. 1f) in the NV$^-$ centre. Optical addressing, read-out, and microwave (MW) manipulation of the electron spin are performed with a bespoke confocal microscope arrangement. At the transition frequency of 2.742 GHz, the electron spin is optically initialised into the $m_s = 0$ state. The electron spin is then coherently driven in a single Rabi sequence (Fig. 1g), for a given evolution time $t$, by applying MW pulses of a fixed but arbitrarily chosen power. The photoluminescence (PL) indicating the spin state is detected and used to obtain the output probability. For more details on NV$^-$ spin see Methods and Suppl. Info. 2.

The general aim of QHL is to find the parameters $x_0$ that best describe the dynamical Hamiltonian evolution of the system via $\hat{H}_0 = \hat{H}(x_0)$. Learning the Hamiltonian relies on an estimation of likelihoods, which can be exponentially hard to compute on classical machines. However, a quantum simulator can be programmed for a parametrised Hamiltonian $\hat{H}(\vec{x})$ such that the observed data allows the efficient estimation of its associated likelihoods. The first QHL protocol we implemented is called quantum likelihood estimation (QLE). The initial state $|\psi\rangle$ of the target system is evolved for a time $t$ and measured in a basis $\{\{|D\rangle\}\}$, as shown in Fig. 2a. The observed data $D$ is fed to the quantum simulator which simulates state evolution and measurement assuming $\hat{H}(\vec{x})$ as the true Hamiltonian. Given $\vec{x}$, the probability $\Pr(D|\vec{x}) = |\langle D|e^{-i\hat{H}(\vec{x})t}|\psi\rangle|^2$ of obtaining $D$ is known as the likelihood function for QLE. We then use $\Pr(D|\vec{x})$ in combination with an approximate form of Bayesian inference known as Sequential Monte-Carlo algorithms (SMC) to learn $\vec{x}$ and estimate its uncertainty. In this approximation, a finite set of points in the parameters space $\{\vec{x}_i\}$, called particles, is used to describe the probability distri-
Si quantum photonic simulator

Magnet
Si chip
Gate control
Si quantum photonic simulator

Laser
Initialise
Photon
sources
Gate controller
Laser
ˆM
ˆUˆV
ˆV
MMI
ˆU ˆV
| i
|+i ˆM
Si-WG
TO-gate
Crosser
c

Figure 1 Quantum photonic simulator and diamond nitrogen-vacancy centre. The silicon quantum photonic simulator: a, experimental setup, b, quantum circuit, and c, device schematic. The operation, either ˆU or ˆV, on the qubit |ψ⟩, is entangled with the states, |0⟩ or |1⟩ respectively, of the control qubit. The device in (c) implements the logical circuit. A 10 mW continuous-wave pump laser with near 1550 nm wavelength is coupled into the chip through optical fibres. In (c), black lines are silicon nano-photonic waveguides (Si-WG), and gold wires represent thermo-optical (TO) phase-shifters and their transmission lines. A pair of idler (blue) and signal (red) photons with different wavelengths are generated via spontaneous four-wave mixing in the spiral waveguide sources. These photons are split equally via multi-mode interferometer (MMI) beam-splitters, producing a post-selected maximally entangled state. The operation ˆU or ˆV performed on idler qubit is coherently controlled by the state of the signal qubit. Performing measurements ˆM on the signal qubit allows to estimate the likelihood function for the chosen configuration of the device. Photons are detected off-chip by fibre-coupled superconducting nanowire single-photon detectors (SNSPD). d, Confocal setup with diamond (inset) containing NV − centres. e, Structure and f, energy level diagram of an NV − centre in diamond. The ground-state electron spin Hamiltonian, describing the coherent dynamics between m s = 0 and −1, is to be characterised and learned using the quantum simulator in (a), g. A single Rabi sequence for the initialisation, manipulation and read-out of the electron spin state. A laser pulse at 532 nm is used to initialise the spin into m s = 0. Two microwave (MW) π/2-pulses with a time t delay are then used to coherently drive the spin. The spin state is measured by detecting photo-luminescence (PL) with an avalanche photodiode (APD). These two different physical systems are interfaced through a classical computer.

bution (See Methods for details).

Our silicon-photonics device and the NV − centre were interfaced with a classical computer, such that experimental data directly enabled QLE. Rabi oscillations of the NV − centre’s electron spin can be modelled by a Hamiltonian of the form ˆH(f) = ˆσ f/2 acting on the initial state, defining ˆσ as the quantisation axis (this definition is equivalent to the conventional model ˆσ f/2 up to a rotation of reference frame). The silicon-photonics chip simulated the model ˆH(f) to learn the Rabi frequency f and to enable the calculation of the likelihood function for each particle. At each step of the QLE implementation, the evolution time t was chosen adaptively for the NV − electron spin performing a single Rabi sequence. PL results were calculated from 3 million iterations for each sequence. The likelihoods obtained were then used to update the prior distribution via the classical computer, before proceeding to the next step. The prior distribution Pr(f) of the particles was initialised to be uniform between 0 and an arbitrary value Δf, where we chose Δf = 100/2π MHz. For clarity, we consider the rescaled quantity ω = f/Δf distributed in the interval ω ∈ [0, 1].

We performed QLE with 50 steps using a 20 particles SMC approximation to learn the electron spin dynamics of the system. Figures 2b,c show the particle’s distribution converging to the correct value ω0. The final learned value corresponds to the Rabi frequency f QLE = (6.93 ± 0.09) MHz, given by the mean and standard deviation of the distribution, which is consistent with the referenced value f0 = 6.90 MHz obtained with the fit of the Rabi oscillations measurements (see Suppl. Info. 2). Thus the simulator successfully learns the parameter that best represents this Hamiltonian, without prior knowledge of the Rabi frequency. We note that the total number of measurements on the NV − system required for QLE is smaller than those for the fit (≃ 200). The fast experimental convergence of the algorithm to ω0 is observed through the evolution of the quadratic losses (here equal to the mean-squared errors) of the particles distribution achieving a final value of approximately 10−5, as shown in Fig. 2d.

Figure 2e reports the evolution of the distribution variance and shows an exponential decay in the first 35 steps. The stepping of data points, e.g. near steps 15 and 24, arises from the probabilistic nature of the learning algorithm. We note that the variance c2 saturates at approximately 4.2 × 10−5. This saturation indicates that the algorithm stops learning within this model (Model I). Such saturations are easy to spot within a Bayesian framework, because c2 can be directly computed from the posterior distribution. This strikingly illustrates that QHL can estimate the limitations of the physical model used to describe the dynamics of the system.

Knowing when a model has failed affords us the opportunity to improve upon it. The present model was improved by introducing chirping, described by a time-dependent Hamiltonian ˆH′(f, a; t) = ˆσ Δ f (f + at)/2 (Model II), where a is a chirping constant. Including chirping allows the algorithm to con-
Within 50 steps, the distribution converges to the correct value of the two parameters are evolved for a time $t$ and measured. The simulator mimics the system dynamics according to the model and obtains an estimate of the likelihood function using the outcomes from the system. The likelihoods are then used to infer the posterior distribution of the parameters $\hat{\omega}$ (dashed red line). Insets: the distribution of particles after 15, 30 and 50 steps. The points represent experimental data and the shaded areas are un-normalised Gaussian fittings. c, Evolution of the mean and standard deviation of the distribution. Error bars are ±1 s.d. of the distribution. d, Evolution of the quadratic losses, here equivalent to the mean-squared error. Circles are experimental data and the line represents theoretical simulation results with a 67.5% credible interval (shaded area). The theoretical simulation was averaged over 500 runs of QLE. e, Model validation and improvement. The presence of other physical effects in the system that are not describable by the model $\hat{H}(\vec{x})$ (Model I) limits the amount of extractable information, as manifested by a saturation of the variance in $\Sigma^2(\omega) \approx 2.9 \times 10^{-3}$ after approximately 35 steps. The adoption of a new two-parameters model $\hat{H}(\omega, a)$ (Model II), which includes the presence of chirping, allows to achieve a covariance below $\|\Sigma\|^2 = 7.5 \times 10^{-6}$ (the shaded area). Inset: covariance norm evolution of the Model II.

Figure 2 Learning the electron spin dynamics on the quantum photonic simulator using QLE. a, Schematic of QLE. The system Hamiltonian $\hat{H}(\vec{x}_0)$ (shaded green) is to be learned by a quantum simulator (shaded red) that embeds an abstract model $\hat{H}(\vec{x})$ of the target system. We here choose an electron spin in NV$^-$ centre as the target system and use a silicon photonic device as the simulator. In the system, the initial state $|\psi\rangle$ is evolved for a time $t$ and measured. The simulator mimics the system dynamics according to the model and obtains an estimate of the likelihood function using the outcomes from the system. The likelihoods are then used to infer the posterior distribution of the parameters $\vec{x}$ via Bayes’ rule and to calculate the next step time $t$. b, The QLE progressive learning of the electron spin Hamiltonian parametrised by a rescaled Rabi frequency $\omega = f/\Delta f$. The probability distribution over Hamiltonian parameter $\omega$ is described by a discrete approximation using 20 particles. The logarithmic (rather than linear) scale of probability Log$\{P(\omega) + 1\}$ is used to better elucidate the key information including the convergence. Within 50 steps, the distribution converges to the correct value $\hat{\omega}_0$ (dashed red line). Inset: the distribution of particles after 15, 30 and 50 steps. The points represent experimental data and the shaded areas are un-normalised Gaussian fittings. c, Evolution of the mean and standard deviation of the distribution. Error bars are ±1 s.d. of the distribution. d, Evolution of the quadratic losses, here equivalent to the mean-squared errors. Circles are experimental data and the line represents theoretical simulation results with a 67.5% credible interval (shaded area). The theoretical simulation was averaged over 500 runs of QLE. e, Model validation and improvement. The presence of other physical effects in the system that are not describable by the model $\hat{H}(\omega)$ (Model I) limits the amount of extractable information, as manifested by a saturation of the distribution variance at $\sigma^2(\omega) \approx 2.9 \times 10^{-3}$ after approximately 35 steps. The adoption of a new two-parameters model $\hat{H}(\omega, a)$ (Model II), which includes the presence of chirping, allows to achieve a covariance below $\|\Sigma\|^2 = 7.5 \times 10^{-6}$ (the shaded area). Inset: covariance norm evolution of the Model II.

Though QLE is scalable, it often requires short evolution times to ensure the likelihood evaluation is tractable, which can preclude exponential reductions in the number of experiments needed, and makes the SMC approximation more error prone. Yet if it is possible to couple two quantum devices via a quantum (rather than a classical) channel, such as photon-NV spin coupling systems [33] or different gates on a single photonic chip [10], an interactive quantum likelihood estimation (IQLE) algorithm can be adopted to overcome these problems [15, 16].

Similar to QLE, in IQLE the state initially evolves forward in time with the Hamiltonian of the system $\hat{H}(\vec{x}_0)$. However, the transformation is then inverted by the time-reversed Hamiltonian evolution $\hat{H}_- = \hat{H}(\vec{x}_-)$, with $\vec{x}_-$ sampled from the prior distribution (Fig. 3a). To ensure the backwards transformation via $H_-$, the state must be transferred from the system to the simulator. Thus IQLE requires the presence of a coherent quantum channel between them. IQLE enables a number of significant features. It has been shown that the likelihood function for the two-outcome experiments, which involves computing $|\langle \psi | e^{\hat{H}_- t} e^{-i\hat{H}(\vec{x}) t} | \psi \rangle|^2$, is efficient for $\hat{H} \approx \hat{H}_-$ even if...
now respectively represent the trusted hardware and the untrusted system to be validated. This application illustrates how IQLE could be implemented to use small trusted quantum circuits to characterise and verify large quantum circuits, improving the scalability in many-qubit systems in which characterisation will be a key challenge.

We implement IQLE entirely on the single photonic chip, showing its ability to characterise single-qubit operations of quantum devices. In our experiment the photonic device plays the role of both the untrusted system and the trusted hardware, which is relevantly equivalent to the case that integrates two of the devices on a single chip. This further allows a natural implementation of a quantum device self-verification by demonstrating the algorithm, widening the context of quantum characterisation and verification. The operation to be characterised here is $e^{-i0\Delta f/2}$, where $f_0$ matches the value of the fitted Rabi frequency, chosen for consistency with the previous QLE demonstration. Thus characterising this $\hat{\sigma}_z$-rotation operation is equivalent to learning the Rabi frequency. Similar to the QLE demonstration, the Hamiltonian $\hat{H}(f)$ of Model I was simulated to learn the parameter $\omega = f / \Delta f$. In each step of IQLE, the experiment was implemented twice: once for measuring the outcomes from the untrusted $\hat{\sigma}_z$-rotation (top part in Fig. 3a), and once for estimating the likelihoods (bottom part in Fig. 3a). See Methods for more details. Figure 3b shows the experimental results for the estimated $\omega$ as given by the posterior mean and standard deviation at each step of IQLE. The particle distribution converges quickly to the correct value $\omega_0$. After 50 algorithm steps we obtain $f_{\text{IQLE}} = (6.92 \pm 0.08)$ MHz, which is within 1 s.d. of the implemented Rabi frequency $f_0 = 6.90$ MHz. The evolution of the quadratic losses (Fig. 3c) indicates that the parameter is learned exponentially fast, with a final quadratic loss value of approximately $10^{-9}$. The convergence of the algorithm to the implemented value $\omega_0$ indicates the successful self-verification of the quantum device.

We report the first demonstration of QHL showing the capability of validating Hamiltonian models and verifying quantum devices. While these experiments use a digital quantum photonic simulator for the demonstration, QHL is universal and can be implemented on any quantum computing platform (e.g. [10–12]). Furthermore, this learning protocol applies to non-digital simulators, which is particularly of interest when certain classes of analogue quantum simulations are likely to approach a regime beyond that available to classical supercomputers in the medium term [7, 8]. With anticipated future developments in quantum hardware, the QHL protocol can be scaled up to learn more complex Hamiltonians, and promises the early delivery of quantum-enhanced computational techniques to efficiently characterise and verify quantum systems and technologies, and to investigate foundational physics.

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Corresponding authors:
Jianwei Wang (jianwei.wang@bristol.ac.uk)
Nathan Wiebe (nawiebe@microsoft.com)
Mark G. Thompson (mark.thompson@bristol.ac.uk)

Author contributions
J.W., S.P. and R.S. contributed equally to this work. J.W., S.P., S.K. and N.W. conceived and designed the experiments. N.W. provided theoretical support. S.P. and M.P. performed the simulations. J.W., S.P., R.S., S.K. and A.A.G built the setups, carried out the experiment, analysed the data and wrote the manuscript with feedback from all authors. N.W., A.L., J.G.R., J.L.O’B. and M.G.T. managed the project.
Methods

Diamond NV$^-$ centre sample and setup. The bulk diamond hosting the negative charged NV$^-$ centre is a chemical vapour deposition (CVD) grown sample (electronic grade) with a natural abundance of 1 ppb nitrogen impurities, see Inset in Fig. 1d. The NV$^-$ centre was positioned in the static magnetic field at room temperature. All the measurements were performed on a home-built scanning confocal microscope, as shown in Supplementary Fig. 2. With the help of optical detected magnetic resonance (ODMR), we perfectly aligned a small external magnetic field of 5mT in the direction of the NV$^-$ centre’s axis, lifting the degenerated $m_s = \pm 1$ spin states. Supplementary Fig. 3a shows the ODMR of the NV$^-$ centre used in the experiment, which was scanned under continuous optical laser and microwave (MW) excitation, indicating the transition from $m_s = 0$ to $m_s = -1$ at the frequency of 2.742 GHz. More details on the confocal setup and spin measurement are reported in Suppl. Info. 2.

Silicon quantum photonic device and setup. The quantum device was manufactured on the standard Silicon-on-Insulator (SOI) wafer using 248 nm-ultraviolet photolithography and reactive-ion etching. Single photons were generated and guided in silicon waveguides with a cross-section of $20 \times 220 \text{ nm}^2$. The single photon sources were designed with a 1.2 cm length. The relative phase between different paths was manipulated using TO phase-shifter used by metal deposition of titanium upon the silicon waveguides. MMs couplers with a size of 2.8 $\mu \text{m} \times 28 \mu \text{m}$ were used as beam-splitters with near a 0.5 reflectivity. The crossers in the device showed a ~40 dB crosstalk between the two intersected waveguides. The schematics of components are shown in Fig. 1c.

The chip was optically accessed via single-mode optical fibers using spot-size converters. The chip was wired-bounded on a PCB and each phase-shifter was individually controlled by an electronic driver with 12 bits resolution. Photons were detected using superconducting nanowire single-photon detectors. A classical computer was used to process the photon statistics obtained through a time interval analyser from the quantum device, and perform the Bayesian inference to update the Hamiltonian model. The detailed experimental setup for the quantum chip is provided in Supplementary Fig. 1.

State evolution on the quantum photonic device. The schematic of the photonic device is provided in Fig. 1c. The state generated by the SFWM sources [37], is given by $(|20⟩ + |02⟩)/\sqrt{2}$ in the Fock basis [27]. After the first pair of photons which here works as a probabilistic filter, the state becomes $(|1100⟩ + |0011⟩)/\sqrt{2}$ by post-selecting photons [29], where the number indicates the photon number occupying in different spatial modes. We re-label the top and last mode as the first two, i.e. applying the following transformation $|a,b,c,d⟩ → |a,b,c⟩$, which is physically realized using waveguide crossers. Then, the state evolves into a maximally two-qubits entangled state.

$$
\begin{align*}
|1010⟩ + |0101⟩ & \quad \sqrt{2} \\
\end{align*}
$$

If we convert the Fock state to the logic state via $|0⟩_{\text{logic}} → |0⟩_{\text{Fock}}$ $(|1⟩_{\text{logic}} ↔ |1⟩_{\text{Fock}})$, we obtain $|0⟩_{D} |0⟩_{1} |0⟩_{2} |1⟩_{3} / \sqrt{2}$, where the subscripts denote the qubit 1 and qubit 3. By adding two additional modes in the bottom paths (expanding into a 4 dimensional space) which can represent one of the additional photon qubit 2, we can obtain a state equivalent to $|0⟩_{D} |0⟩_{1} |0⟩_{2} |1⟩_{3} / \sqrt{2}$, where the input state $|\psi⟩_{2}$ can be prepared in the higher-dimensional space. Evolving the state $|\psi⟩_{2}$, using two different unitaries $U$ in the upper path and $V$ in the lower one, we have the state as

$$
|0⟩_{D} |0⟩_{1} |\tilde{U}|\psi⟩_{2}⟩ |0⟩_{3} + |0⟩_{D} |0⟩_{1} |\tilde{V}|\psi⟩_{2}⟩ |1⟩_{3} / \sqrt{2}
$$

Then, to obtain two-new crossers and interfering photons at the last two MMs, it allows us to erase the path information between the upper and the lower path, i.e. whether the photon went through the $\tilde{U}$ or the $\tilde{V}$ operation. This is equivalent to apply a Hadamard gate to the third qubit. The state emerging by this evolution can be described as

$$
\begin{align*}
& \frac{1}{\sqrt{2}} \left( |0⟩_{D} |0⟩_{1} |\tilde{U}|\psi⟩_{2}⟩ |0⟩_{3} + |1⟩_{D} |0⟩_{1} |\tilde{V}|\psi⟩_{2}⟩ |1⟩_{3} \right) \\
& \frac{1}{\sqrt{2}} \left( |0⟩_{D} |1⟩_{1} |\tilde{U}|\psi⟩_{2}⟩ |0⟩_{3} + |0⟩_{D} |1⟩_{1} |\tilde{V}|\psi⟩_{2}⟩ |1⟩_{3} \right) \\
& \frac{1}{\sqrt{2}} \left( |0⟩_{D} |0⟩_{1} |\tilde{U}|\tilde{U} |\psi⟩_{2}⟩ |0⟩_{3} + |1⟩_{D} |0⟩_{1} |\tilde{V}|\tilde{V} |\psi⟩_{2}⟩ |1⟩_{3} \right) \\
& \frac{1}{\sqrt{2}} \left( |0⟩_{D} |1⟩_{1} |\tilde{U}|\tilde{U} |\psi⟩_{2}⟩ |0⟩_{3} + |0⟩_{D} |1⟩_{1} |\tilde{V}|\tilde{V} |\psi⟩_{2}⟩ |1⟩_{3} \right)
\end{align*}
$$

Applying a post-selection for those cases where the second photon emerges from one of the upper modes, i.e. projecting the third qubit into the state $|0⟩_{3}$, it is possible to achieve the desired state

$$
|0⟩_{D} |0⟩_{1} |\tilde{U} |\psi⟩_{2}⟩ |0⟩_{3} + |1⟩_{D} |0⟩_{1} |\tilde{V} |\psi⟩_{2}⟩ |1⟩_{3} / \sqrt{2}
$$

In our QHL experiments, we choose $|\psi⟩_{2}$ as $|0⟩$ which can be naturally realised after the stage of generating entangled photon state, with no complication between the operation of unitaries and the preparation of $|\psi⟩_{2}$. Thus, the operations $\tilde{U}$ and $\tilde{V}$ are solely used to represent the Hamiltonian evolution.

We remark that the operation, either $\tilde{U}$ or $\tilde{V}$, performed on the second qubit on the initial state $|\psi⟩_{2}$ is determined by the state of the first qubit $|\tilde{p}⟩_{1}$ and $|\tilde{q}⟩_{2}$ projective basis, we can directly estimate the overlap between the states evolved according to the $\tilde{U}$ and $\tilde{V}$ operations. This method can be seen as an entanglement based scheme for calculating the overlap [38, 39]. Unfortunately we point out that the probabilistic and post-selected nature of this approach makes it not scalable, but it allows flexibility to perform likelihoods estimation in our demonstration. The details on likelihood estimation are discussed in the last section in Methods.

Bayesian inference and Hamiltonian learning. Bayesian inference is a commonly used method in physics and statistics for model estimation. The method is fundamentally based on the Bayes’ theorem, which gives the proper way to update a probability distribution given evidence. The fundamental object in Bayesian inference is called a prior distribution $Pr(\tilde{x})$. From a Bayesian perspective, the prior distribution describes the subjective beliefs that an experimenter may have about the model in question which is parameterized by $\tilde{x} \in \mathbb{R}^N$. For example, if we want to learn a Rabi frequency we may know from prior knowledge that $f = 6.9 \pm 0.01 \text{ MHz}$. If that is the case then a reasonable prior distribution for describing this is to take $\tilde{x} = [f] \text{ and}$

$$
Pr(\tilde{x}) = e^{-\left(f - 6.9\ f \text{ MHz}\right)^2 / 0.0002} / \sqrt{0.01 \pi} \text{ s} \text{ MHz}.
$$

A major advantage of this approach is a posterior distribution not only gives an estimate of the most likely outcome, argmax $\left(Pr(\tilde{x})|D\right)$, but also gives an estimate in the uncertainty of that estimate in the form of the covariance matrix of $Pr(x)$. We use this feature in the main body for model selection.

While the subjectivity of the initial prior distribution is a hotly debated subject among statisticians, the use of priors does an excellent job here, addressing the fact that we almost always start experiments with prior understanding of the system in question. The Bayesian formalism gives us a language to articulate it. Also the Bernstein von-Mises theorem shows that, under relatively generic assumptions, a poor choice of the prior distribution does not affect the ultimate conclusions reached by Bayesian inference. Rather it only affects the time required to learn the model in question.

The next most important object in Bayesian inference is the likelihood function. The likelihood function takes an experimental datum, $D$, and parameters $\tilde{x}$ and outputs the probability that $\tilde{x}$ generates the observed data. This is expressed as $Pr(D|\tilde{x})$ and is essential for most applications of Bayesian inference because it allows Bayes’ theorem to update the prior distribution given an observed datum:

$$
P(\tilde{x}|D) = \frac{Pr(D|\tilde{x})Pr(\tilde{x})}{\int Pr(D|\tilde{x})Pr(\tilde{x})d\tilde{x}}
$$

The output of Bayes’ theorem, $Pr(\tilde{x}|D)$, is known as the posterior distribution. It represents how the experimentalists’ beliefs about the model should be distributed after observing the evidence given their prior beliefs.

This update process is seldom efficient in Bayesian inference. This is because, unless there is a special relationship between the prior and the likelihood function (i.e. conjugate priors), there will not be a simple analytic form for the posterior distribution. This means that the entire function must be stored in memory, which is not possible for a continuous models. Such problems are often solved by using approximate inference methods.

Sequential Monte-Carlo (SMC) algorithms are a class of approximate inference algorithms that are increasingly popular for approximating Bayesian inference. The idea behind these methods is to ap-
proximate the probability density by a discrete sum of points, known as particles. In particular, we wish to find a set of weights $w_i$ and positions $x_i$ such that $\Pr(\vec{x}) \approx \sum x_i w_i (\vec{x} - x_i)$ and $\sum w_i = 1$. This allows us to replace the integrals with a sum and further allow the density to be represented in a finite amount of memory. Formally this approximation is not known to be efficient, in that the number of particles needed can be shown to scale at most sub-exponentially (rather than polynomially) with the number of model parameters. In practice, the dependence on the number of model parameters is often quite weak and often depends more strongly on the properties of the likelihood function than the size of the model.

An important technical issue about SMC algorithms is that the particles often need to move as the inference algorithm proceeds. To see this, consider the Rabi model. If we assume that the system has no noise then the Rabi frequency can be learned with infinite precision. This corresponds to a probability density of $\Pr(\vec{x}) = \delta(\omega - \omega_{true})$. This density can only be described exactly inside the SMC formalism if there is a particle $\vec{x}_i = \omega_{true}$. If a finite number of particles is chosen then the probability of this is zero. Therefore in order to model very narrow distributions that will occur in Bayesian inference we need to move particles.

One procedure for doing this is known as resampling. The goal of the resampler is to effectively move the particles by redrawing them from a distribution that resembles the current distribution. These "resampled" particles are then assigned equal weight and the learning process is resumed, now with more particles concentrated in the important parts of the posterior distribution. This resampling is triggered when the number of "effective" particles in the approximation becomes too small. In particular, we resample if and only if the inverse participation ratio, $\frac{1}{\sum w_i^2}$, becomes too small (usually $\frac{1}{\sum w_i^2} < N_{part}/2$ is taken to be the threshold). The resampler we used was given by Liu and West [40] and it has the property of preserving the first two moments of $\Pr(\vec{x})$ while approximately maintaining the structure of the particle cloud.

If we specialize to the problem of QLE and IQLE experiments then our likelihood function is always of the form [15, 16]

$$\Pr(D | \vec{x}, t) = |\langle e^{-i\hat{H}(\vec{x})t}| \psi \rangle|^2, \quad (7)$$

for QLE or

$$\Pr(D | \vec{x}, t, \vec{x}') = |\langle e^{-i\hat{H}(\vec{x}, \vec{x}')t}| D \rangle|^2, \quad (8)$$

for IQLE.

An important question to ask is: "how do we choose $t$ to best estimate $\hat{H}(\vec{x})$?" We employ an adaptive strategy known as the particle guess heuristic to provide near-optimal experiments [41, 42]. This strategy takes the evolution time to scale as the reciprocal of the current uncertainty. For the application using QLE to estimate Rabi frequencies, we adopt a near-optimal choice of the evolution time $t = 1.26/\sigma$, where $\sigma$ is the standard deviation (s.d.) of $\Pr(\vec{x})$. For IQLE, we adopt the following heuristic $t = 1/|\langle \hat{H}(\vec{x}) - \hat{H}(\vec{x'}) | 2 = 1/|\omega_1 - \omega_2|$, with both $\vec{x}_1$ and $\vec{x}_2$ sampled from the prior distribution $\Pr(\vec{x})$ [15].

For both QLE and IQLE algorithms are reported in Supplementary Algorithms 1-5.

**Estimation of the likelihoods on the photonic chip.** The likelihood estimation for QLE requires the inner product between the evolved state $e^{-i\hat{H}(\vec{x})t}| \psi \rangle$ and the state $| D \rangle$. In this work we have used $| D \rangle = | \psi \rangle = | 0 \rangle$, in the chosen computational basis state of both NV centre and photonic device. The NV spin’s reference frame we use, defining the $\hat{e}_z$ as the quantisation axis, is obtained by a rotation of basis from the standard reference frame, where the quantisation axis is the $\hat{e}_z$ axis.

The choice of this reference frame is due to the ease of preparing the input state $| \psi \rangle$ as $| 0 \rangle$ in the photonic chip, with no need of compiliation between state preparation and unitary operations (see the state evolution section in Methods).

In order to calculate the inner product in our photonic device we exploit an entanglement-based technique [38, 39]. The scheme realised in the integrated device allow us to produce entangled states $| \psi \rangle = | 0 \rangle\langle \psi | + | 1 \rangle\langle \psi | \sqrt{2}$, where the subscripts $s$ and $t$ now represent the qubits 1 and 2 (see Eq. 4) encoded in the signal and idler photons, respectively. When performing a projective measurement on the $\hat{e}_z$ eigenbasis $| \pm \rangle = (| 0 \rangle \mp | 1 \rangle)/\sqrt{2}$, we obtain

$$\Re\langle \langle \psi | 0 | \psi \rangle \rangle = 2p_+ - 1 \quad (9)$$

where $p_+$ is the probability to get the outcome $| + \rangle$. Similarly, when performing a projective measurement on the $\hat{e}_x$ eigenbasis $| \pm \rangle = (| 0 \rangle \pm | 1 \rangle)/\sqrt{2}$, we obtain

$$\Im\langle \langle \psi | 0 | \psi \rangle \rangle = 2p_{+1} - 1 \quad (10)$$

where $p_{+1}$ is the probability to get the outcome $| + \rangle$. The likelihood for QLE is given by $L_{QLE} = |\langle \psi | e^{-iH(\vec{e}_x)t} | \psi \rangle|^2$ and particularly $|\langle 0 | e^{-iH(\vec{e}_x)t} | 0 \rangle|^2$ in this experimental demonstration, which can be easily obtained setting

$$\hat{U} = I, \quad \hat{V} = e^{-i\hat{H}(\vec{e}_z)t} / \sqrt{2} \quad (11)$$

The schematic in Fig. 2a shows the QLE implementations on the photonic device. Using Eq. 9 and Eq. 10, we have

$$L_{QLE} = (2p_+ - 1)^2 + (2p_{+1} - 1)^2 \quad (12)$$

The values of $p_+$ and $p_{+1}$ are calculated performing the single-qubit operations on the control qubit and measuring photon coincidences.

For IQLE the likelihood $L_{IQLE} = |\langle \psi | e^{-i\hat{H}(\vec{e}_x,t)\hat{H}(\vec{e}_x,t)} | \psi \rangle|^2$ is obtained similarly using Eq. 12 but setting

$$\hat{U} = e^{-i\hat{H}(\vec{e}_x,t)}, \quad \hat{V} = e^{-i\hat{H}(\vec{e}_x,t)} \quad (13)$$

given the same format of Hamiltonians and evolved state $| \psi \rangle$. The schematic in Fig. 5a shows the IQLE implementations on the photonic device. The quantum channel required for IQLE in our case is thus given by the entanglement generated in the sources. We remark that using this scheme to implement IQLE all evolutions are forward in time, in contrast with the original approach where the time reversal $e^{-i\hat{H}(\vec{e}_x,t)}$ is performed by a backwards evolution in time [15]. This can make our entanglement-based approach amenable for analogue quantum simulators. It however comes at the cost of additional entanglement between the system and an ancillary qubit.

**Data availability.** The data that support the plots within this paper and other findings of this study are available from the corresponding author upon reasonable request.

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