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## Tuning methods for semiconductor spin-qubits

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We present efficient methods to reliably characterize and tune gate-defined semiconductor spin qubits. Our methods were developed for double quantum dots in GaAs heterostructures, but can easily be adapted to other quantum-dot-based qubit systems. These tuning procedures include the characterization of the inter-dot tunnel coupling, the tunnel coupling to the surrounding leads and the identification of various fast initialization points for the operation of the qubit. Since semiconductor-based spin qubits are compatible with standard semiconductor process technology and hence promise good prospects of scalability, the challenge of efficiently tuning the dot's parameters will only grow in the near future, once the multi-qubit stage is reached. With the anticipation of being used as the basis for future automated tuning protocols, all measurements presented here are fast-to-execute and easy-to-analyze characterization methods. They result in quantitative measures of the relevant qubit parameters within a couple of seconds, and require almost no human interference.

#### I. INTRODUCTION

The recent developments in semiconductor based spin qubits show their great potential as building blocks of a quantum computer and demonstrate their promise for scalable architectures.<sup>1–9</sup> However, with increasing number of physical qubits, challenges like device architecture,<sup>10–12</sup> long-range coupling, <sup>13–17</sup> error correction, <sup>18,19</sup> decoherence due to charge noise, 20,21 and scalable implementation 22,23 of the control electronics<sup>24,25</sup> will play an increasingly important role. One further obstacle, which has not received much attention to date, is the tuning of the qubit devices. Especially in the case of gate-defined quantum dots, even tuning a double quantum dot is a non-trivial task, as each quantum dot comprises at least three electrostatic gate-electrodes, each of which influences the number of electrons in the dot, the tunnel coupling to the adjacent lead, and the inter-dot tunnel coupling. The current practice of manually tuning the qubits is a relatively time-consuming procedure. While it can be simplified with improved gate designs that feature little cross talk between different target parameters $^{26}$ , manual tuning is inherently impractical for scale-up and applications.

In this work we present tuning and characterization methods for double-quantum dots which have evolved over the course of the experiments on two-electron spin qubits presented in Refs.<sup>1</sup> and<sup>21,27–32</sup>. These procedures are used to tune up one and two two-electron spin qubits but they also involve aspects needed for multi-qubit devices.

Complementary to Ref.<sup>33</sup>, which shows a computerautomated scheme for the coarse-tuning of quantum dots into the single-electron regime, we focus here on the fine-tuning of the spin qubit once the single-electron regime is reached. In addition to the tuning of the interdot tunnel couplings<sup>34,35</sup>, the fine-tuning includes the adjustment and the characterization of the tunnel couplings to the adjacent leads and the identification of the energy transitions relevant for the qubit functionality. We exploit high-bandwidth readout by radiofrequency (RF)-reflectometry<sup>36,37</sup> and present fast, easyto-analyze, quantitative measurements to characterize semiconductor spin qubits. Contrary to the relatively slow tuning based on direct current (DC) electron transport through the dot,<sup>38</sup> all scans necessary for characterising a device in our scheme can be performed within a few seconds by using pulsed gate measurements and charge sensing with RF readout. As the tuning parameters of interest are obtained directly as fit parameters and require no human intervention, these analysis methods are well suited as a basis for the full automation of the complete tuning procedure. Such an automation will be crucial for the scalability of any qubit that requires tuning.

Importantly, while all measurements presented here were performed on GaAs double quantum dots operated as two-electron spin qubits, the procedures can easily be adapted to other quantum-dot-like qubit systems. In particular, most aspects are not specific to GaAs or twoelectron spin qubits, as devices containing two exchange coupled single-spin qubits<sup>39,40</sup> are subject to very similar requirements. Moreover, our procedures are also adaptable to devices with a larger number of dots or qubits, which will also require the adjustment of inter-dot and dot-lead tunnel couplings.

The outline of this paper is as follows: In Sec. II we introduce the device layout of the two-electron spin qubit in GaAs, and explain the basics of the experimental setup including the RF-reflectometry circuit. In Sec. III we present our methods to quantitatively characterize and fine-tune the qubit. We first motivate the use of virtual gates, a linear combination of several gates that allows changing specific quantum dot parameters individually. We continue by describing the characterization of the inter-dot tunnel coupling  $t_c$  and the tunnel couplings to the electron reservoirs, which must be tuned to certain values for the proper operation of the qubit.

Additionally, we provide routines for locating fast–reload points used to initialize the qubit in different states, and the location of the energy transitions that allow setting up a hardware feedback-loop to polarize and stabilize the nuclear spin bath in the GaAs host material.<sup>28</sup>

# II. DEVICE LAYOUT AND EXPERIMENTAL SETUP

All data shown in this paper were obtained from the qubit in Refs.<sup>1and32</sup>, depicted in Fig. 1. This is a so-called singlet-triplet spin qubit (ST<sub>0</sub>-qubit), embedded in a GaAs double quantum-dot formed by electrostatic gates (gray and blue features in Fig. 1) on top of a two-dimensional electron gas (2DEG). The ST<sub>0</sub>qubit is encoded in the  $m_z = 0$  subspace of the regime where each of dot is occupied by a single electron, i.e. to the subspace spanned by  $S = (|\uparrow\downarrow\rangle - |\downarrow\uparrow\rangle)/\sqrt{2}$  and  $T_0 = (|\uparrow\downarrow\rangle + |\downarrow\uparrow\rangle)/\sqrt{2}$ ,<sup>41</sup> where  $\uparrow$  or  $\downarrow$  describes the spin state of the electron in one of the dots. This type of qubit can be fully manipulated using only electric pulses.

In more detail, the gates depicted in gray in Fig. 1 represent DC static gates used to define the quantum dots and to tune them into the single-electron regime. They are heavily filtered and fed with voltages of order 1 V. The broad side gates S1 and S2 adjust the number of electrons in quantum dot 1 and 2. The barrier gates B1 and B2 control the tunnel coupling to the leads and the inter-dot coupling is controlled by the "nose" and "tail" gates, N12 and T12.

Two additional control–gates (named RFX and RFY and depicted in blue in Fig. 1) are used for qubit manipulation by applying mV-scale signals. They are DCcoupled to an arbitrary waveform generator Tektronix AWG5014C operated at 1 GS/s, and attenuated by 33 dB at various cryogenic stages to reduce thermal noise from room temperature. Using dedicated static– and control– gates eliminates the need for bias tees and the resulting pulse imperfections<sup>28</sup>, and ensures a nearly flat frequency response of the control gates from DC to a few hundred



Figure 1. False color SEM image of a device similar to the one used in this work, including contacting scheme. Applying static voltages to the grey gates confines two electrons in a double dot potential in the 2DEG of a GaAs/Al<sub>0.31</sub>Ga<sub>0.69</sub>As heterostructure. The blue gates, RFX and RFY, are used exclusively for fast manipulation. The dot on the left is used for charge sensing of the double dot, and is embedded in an impedance-matching circuit as the resistive element. The crossed boxes represent ohmic contacts to the leads.

MHz, at the cost of one additional gate electrode.

The double dot is capacitively coupled to a third dot (named sensing dot), which is used as a charge detector. Its conductance depends on the local electrostatic landscape, allowing to read out the charge state of the double dot.<sup>42</sup> The spin-state of the double dot can be probed through the sensing dot by spin-to-charge conversion based on Pauli-spin-blockade.<sup>43,44</sup> The sensing dot is embedded as a resistive component in an impedance matching circuit, so that the conductance through the dot can be monitored using RF-reflectometry  $^{36,37,\overline{4}5}$  at a local oscillator frequency of approximately 230 MHz and a bandwidth of 20 MHz. We employ a setup similar to Ref. <sup>37</sup>, with the addition of a cryogenic circulator at base temperature. The demodulated signal  $V_{\rm rf}$  is a function of the conductance of the sensing dot, and is recorded using an Alazar ATS9440 digitizer board.

We typically use a hardware sample rate of 100 MS/s, which we downsample on-the-fly at full data rate to 250 kHz using a multithreaded, high throughput C++based driver for the Alazar card. This downsampled rate arises from a typical length of 4  $\mu$ s for experiments, which usually comprises a 2.5  $\mu$ s long measurement window during which we power the RF-circuit. Effects of 1/f-like noise are eliminated from the data by changing the sweep-pulse parameter after each cycle, and then averaging over many repetitions of the parameter sweep to elude slow drifts in the sensor or gate voltage configuration. For a typical tuning dataset, the sweep comprises 100 parameter values and it is repeated 1536 times for a total measurement time of  $1536 \times 100 \times 4 \,\mu s \approx 600 \,\mathrm{ms}$ , and then averaged again over 1-5 repetitions, if necessary. These acquisition parameters are not yet optimized for speed, and we expect that a speed-up of at least a factor of 10 is possible while still maintaining an adequate accuracy of the extracted parameters.

#### III. FINE TUNING OF THE QUBIT

This section describes in detail the fine-tuning of a  $ST_0$ -qubit after the double dot has been tuned in the two-electron regime, either around (2,0)-(1,1) or the (0,2)-(1,1)) charge transition (see Fig. 2 (a))). The procedure for the coarse-tuning to this charge regime is described in the Appendix. All measurements presented in this section are performed using RF-reflectometry on the sensing dot.



Figure 2. (a) Charge stability diagram of the double dot measured using RF-reflectometry. Different values of  $V_{\rm rf}$  correspond to different charge ground states. (b) Fit of the stability diagram using the model described in Sec. III A. Circles and lines mark the automatically detected triple points and the positions of the charge transition. The gray arrow represents the direction of the detuning  $\epsilon$ . In both figures we subtracted a background value due to the direct influence of the charge sensor extracted from the fit.

#### A. Locating the triple points

The operation of a qubit requires the accurate characterization of the charge stability diagram in the RFX– RFY plane (here and in the following RFX and RFY refer to the voltage applied to the respective gate). In other words, it is necessary to know the exact position of the triple points – the points in the charge stability diagram where three charge states are energetically degenerate (e.g. (1,0),(1,1) and (2,0)) – and of the so– called lead transitions – the transitions between different charge–states of the double dot that involve electron exchange with one of the reservoirs (e.g. the transition  $(1,0) \leftrightarrow (1,1)$ ). We present instead an automated routine, based on a measurement of the charge stability diagram near the (2,0)-(1,1) transition, followed by the fit to The fitting model consists of two parts. The first part is a two-site Hubbard Hamiltonian without spin:

$$H = \begin{pmatrix} E_{1,0} + v_1 & 0 & 0 & 0\\ 0 & E_{1,1} + v_1 + v_2 & t_c & 0\\ 0 & t_c & E_{2,0} + 2v_1 & 0\\ 0 & 0 & 0 & E_{2,1} + 2v_1 + v_2 \end{pmatrix}$$

in the charge basis  $j \in \{(1,0), (1,1), (2,0), (2,1)\}$ . Here,  $E_j$  are the basis state energies at RFX = RFY = 0,  $t_c$  is the inter-dot tunnel coupling and  $v_i$  the on-site potential, which can be calculated knowing the voltages applied to the RF-gates  $V_i$  and their respective lever arms, including cross-capacitances. The index *i* indicates RFX and RFY, respectively ( $V_1$  = RFX and  $V_2$  = RFY). The spectrum of this Hamiltonian can be calculated analytically to find the charge configuration of each eigenstate at each point in the RFX-RFY plane. Assuming that the occupation probability of each state corresponds to thermal equilibrium, we obtain a vector **p**, describing the occupation probabilities of the various charge basis states.

Since measurements like those presented in Fig. 2 are slow compared to the system dynamics, we can use the ground-state charge population vector  $\mathbf{p}$  as input in a linear fitting model for the charge sensor:

$$S = \mathbf{p} \cdot \mathbf{s} + s_{\text{ct},1} V_1 + s_{\text{ct},2} V_2 + S_0, \tag{1}$$

where S is the charge sensor output,  $\mathbf{s}$  is a vector that contains the sensor output for each charge eigenstate,  $s_{\text{ct},i}$  account for direct crosstalk between the RF-gates and the sensor, and  $S_0$  is an offset. The components of **s**, as well as  $s_{\text{ct},i}$  and  $S_0$ , the lever arms, the crosscapacitances, the energies  $E_i$ , and the inter-dot tunnelcoupling  $t_c$  are treated as fitting parameters, while the input parameters for the fit are the 2D sensor output data and the voltages  $V_i$  applied to the RF-gates. A typical measurement and a fit to the data is presented in Fig. 2. From the fit parameters the position of the triple points as well as the location of the leads transitions in the RFX-RFY plane are extracted. These values are used as reference points in all the following tuning procedures, and to recalibrate the set-up after a charge rearrangement. Furthermore, the direction orthogonal to the segment where the charge states (2,0) and (1,1) are energetically degenerate, defines the so-called detuning axis  $\epsilon$  (gray arrow in Fig. 2b). In the following, we fix  $\epsilon = 0$  to correspond to the measurement point defined in Sec. III E.

#### B. Setting up virtual gates

The first step of the tuning procedure is the coarse tuning of the double dot in the two electron regime, i.e. the identification of the (2,0)-(1,1) (or the (0,2)-(1,1)) charge transition. This step can be based on standard quantum transport measurements (see Appendix) or charge sensing and was automated in Ref.<sup>33</sup>. Once the double dot has been tuned in the appropriate charge regime, the fine-tuning of the qubit can start. For doing so, it is convenient to switch to virtual gates. These virtual gates are given by a linear combination of three physical gates (see Tab. I) that allow tuning the parameters of the double dot while leaving the charge stability diagram in the RFX and RFY plane unaffected. Virtual gates are chosen such that each of them affects primarily one specific dot parameter: gates LeadY and LeadX change the tunnel coupling to the respective lead, while the tunnel coupling between the dots is manipulated by the virtual gates T and N. In each case, in addition to changing the physical gate that mostly influences the desired parameter, a compensating voltage is applied to the S1 and S2 gates to cancel out any cross-capacitance effect. Virtual gates X and Y depend only on S1 and S2 and are used to readjust the position of features in the charge stability diagram in case of imperfect compensation from the virtual gates or charge rearrangements.

To obtain the virtual gate coefficients shown in Tab. I, we focus on the lead transitions and measure how their position in the RFX–RFY plane is shifted by the potential applied on a certain gate. To do so, we apply two different voltages (differing typically by 2-6 mV) to each of the DC gates in turn. For each set of voltages, we measure the dot's response while sweeping RFX or RFY across both lead transitions, as shown in Fig. 3(a) for case of the Y-lead. In these curves, the two plateaux correspond to two different charge states of the double dot. The labels close to each curve indicate the gate on which the potential is changed. To obtain the influence of that gate, the value of RFX (or RFY) at which the transition between the two plateaus occurs, is extracted using a fit-model corresponding to a Fermi distribution<sup>46</sup>,

$$V_{\rm rf}(v) = V_{\rm rf,0} + \delta V_{\rm rf} v - \frac{1}{2} A \left( 1 + \tanh\left(\frac{v - v_{\rm lead}}{w}\right) \right).$$
(2)

Here, v is the voltage on either the RFX or RFY sweeping gate,  $V_{\rm rf,0}$  represents the background value of the charge sensing signal  $V_{\rm rf}$ , the linear term  $\delta V_{\rm rf} v$  accounts for the direct influence of the sweeping gate on the conductance through the sensor (assumed to be linear), the third term accounts for the excess charge once an electron tunnels into or out of the quantum dot and includes a finite electron temperature and lever arm via w, while  $v_{\text{lead}}$  defines the position of the lead transition, and finally A is the contrast of the transition. We use  $V_{\rm rf,0}, \delta V_{\rm rf}, A, w$  and  $v_{\rm lead}$  as fit parameters. The values of  $v_{\text{lead}}$  extracted from these fits depend on the voltage applied to all the DC-gates, and are used to construct a  $2 \times 6$  cross-capacitance matrix. Virtual-gate coefficients are then extracted by inverting the appropriate sub-matrices of the cross-capacitance matrix. Typical values are given in Tab. I. The virtual gate coefficients can be further fine-tuned by applying the same principle

to study the influence of the DC–gates on the location of the triple points of the (2,0)-(1,1) charge transition or on the position of the ST<sub>+</sub> anti-crossing.

A similar concept is used in Ref.<sup>2</sup> to perform orthogonal charge stability diagrams in a three-electron quantum dot, and in Refs.<sup>29and47</sup>.

Table I. Typical values for the coefficients of the virtual gates. The values correspond to the ratio of change in physical gate voltages to that of the virtual gate.

Virtual gate								
			$\operatorname{LeadY}$	LeadX	Т	Ν	Х	Y
$\mathbf{Physical}$	gate	B1	1	0	0	0	0	0
		S1	-0.76	0.5	-0.52	-0.5	1.5	-2.1
		T12	0	0	1	0	0	0
		N12	0	0	0	1	0	0
		B2	0	1	0	0	0	0
		S2	0.26	-1.1	-1.25	-0.5	-3.68	1.03

#### C. Tunnel coupling to the leads

The next step is the tuning of the tunnel coupling to leads X and Y (Ohmic contacts next to the RF-gates, see Fig. 1), which act as electron reservoirs. The coupling to these leads is controlled by the virtual gates LeadY and LeadX, and it must be weak enough to prevent excess  $T_1$ relaxation due to cotunneling or thermal activation and, at the same time, be strong enough to allow fast qubit initialization within tens of nanoseconds.

To extract the tunneling time to the X lead, we apply 25 MHz square-wave pulses that force the system to switch between the charge states  $(1,0) \leftrightarrow (1,1)$  (or, equivalently, between  $(2,0) \leftrightarrow (2,1)$ , and use the sensing dot to measure the time-dependent occupation of the double dot. For this purpose, we average the signal over approximately 1500 periods, recorded at a hardware sampling rate of  $100 \,\mathrm{MS/s}$ . A typical time trace is shown in Fig. 3(b). Applying the square–wave pulses to regions of charge stability (i.e. where no charge transition is possible) allows us to subtract the background due to direct sensor coupling. The tunnelling time to the lead can be extracted from the rise times of the response to the square pulses, Fig. 3(b), with a lower sensitivity bound of about 25 ns determined by the bandwidth of the tank circuit attached to the sensing dot (faster tunneling times can be resolved with the reload sweep discussed in Sec. III E). To fit these data, we use the model

$$V_{\rm rf}(t,t_0) = \begin{cases} V_{\rm rf,0} + \frac{1}{2}A \frac{\cosh\left(\frac{t_0}{2t_{1,1}}\right) - \exp\left(\frac{t_0-2t}{2t_{1,1}}\right)}{\sinh\left(\frac{t_0}{2t_{1,2}}\right)} & \text{for } t < t_0 \\ V_{\rm rf,0} - \frac{1}{2}A \frac{\cosh\left(\frac{t_0}{2t_{1,2}}\right) - \exp\left(\frac{t_0-2t}{2t_{1,2}}\right)}{\sinh\left(\frac{t_0}{2t_{1,2}}\right)} & \text{for } t \ge t_0. \end{cases}$$
(3)



Figure 3. (a) To determine the influence of the various DCgates on the position of a lead transition, we apply two different voltages to each one of the DC-gates in turn. For each set of voltages, we measure the occupation of the double dot as we sweep across two regions in the charge stability diagram. (b,c) To measure the tunnelling times to the leads, we apply MHz-frequency square voltage-pulses across the respective lead transition, forcing an electron to be exchanged with the respective reservoir. The tunnelling time can be extracted from the rise time of the response signal. (d) The inter-dot tunnel coupling is extracted by sweeping along the detuning  $\epsilon$ , recording the average charge occupancy and measuring the broadening of the transition.

where  $t_0 = 2 \mu s$  is the half-period of the square pulse. It produces an exponential rise and decay of the average sensor signal with time constants  $t_{1,1}$  and  $t_{1,2}$  whenever the gate voltage is changed. The prefactors and offsets of the exponential rise and decay are derived from the requirement that the curve is continuous. The same procedure is used to extract the tunnelling times to lead Y, with the only difference that now the square pulse has to force transitions between the charge configurations  $(1,0) \leftrightarrow (2,0)$  (or  $(1,1) \leftrightarrow (2,1)$ ).

Typical target values for  $t_{l,1(2)}$  range from 25 ns to 50 ns. Importantly, since all initialization methods addressed here require only tunneling to one lead (see Sec. III E–III F), the barrier to the other lead can be made less transparent to reduce relaxation (Fig. 3(c)).

#### D. Inter-dot tunnel coupling

The tunnel coupling  $t_c$  between the two dots is mostly controlled by the N and T virtual gates, and determines the strength of the exchange interaction between the two electrons, and therefore the energy splitting  $J(\epsilon)$  between the singlet S and the triplet  $T_0$  in the (1,1) configuration. In order to characterize the tunnel coupling, we measure the broadening of the inter-dot transition between the (2,0)-(1,1) charge configuration by sweeping the detuning  $\epsilon$  orthogonally across the (2,0)-(1,1)-transition (see Fig. 2(a)), and recording the average charge state<sup>46</sup> as shown in Fig. 3(d). We typically measure each detuning step for  $1 \,\mu s$  and average over 4000 scans for a total measurement time of 0.4 s. For simplicity, we extract the broadening of the transition by fitting Eq. (2) to the data, rather than using the physically correct model of an avoided crossing, as we find the difference between the two approaches to be marginal. The value extracted for the effective temperature w now represents the inter-dot tunnel coupling  $t_c$ . Good values for  $t_c$  for the operation of the qubit range from 18  $\mu eV$  to 24  $\mu eV$ , using an estimated lever arm of 9.8 V/eV. Smaller values of the tunnel coupling would lead to Zener-tunnelling when sweeping through the (2,0)-(1,1) transition, and should be therefore avoided.

This characterization method is limited by temperature broadening, which, in our set-up prevents tunnel couplings below 9  $\mu$ eV to be resolved. A similar approach to ours is used in Ref.<sup>34and35</sup> for an automated tuning of the interdot tunnel coupling, whereas an alternative approach for determining the inter-dot tunnel coupling based on time-resolved charge sensing is described in Ref.<sup>48</sup>. Furthermore, the tunnel coupling can also by extracted by photon assisted tunneling spectroscopy.<sup>49</sup> Compared to the presented method, both alternatives are time-consuming and thus less attractive for our purposes.

#### E. Locating the measurement point

The operation of a qubit relies on the ability to reliably initialize the qubit in a well known state and to accurately measure the qubit's final state.<sup>50</sup> Historically, the standard approach for initializing a spinqubit in a singlet state is based on the transition cycle  $T(1,1)\rightarrow(1,0)\rightarrow S(2,0)$ , which requires electron exchange with both reservoirs<sup>44</sup> (here and in the following the notation S(n,m) and T(n,m) indicates the singlet and the triplet state in the (n,m) charge configuration, respectively). Here, we present a modified version that only relies on tunnel coupling to one lead. Compared to the old approach, this procedure requires less tuning and enables simpler future device layouts. It also allows for an enhanced charge–detection readout–scheme<sup>51</sup> to counteract the visibility loss at high magnetic field gradients.<sup>52</sup>

We first need to locate the region of metastable (1,1)

triplets within the (2,0) ground state charge configuration, i.e. the area around point M in Fig. 4(a). The latter represents a high-resolution charge stability diagram, indicating the thresholds of the relevant transitions. To identify the region of metastable (1,1) triplets, we repeatedly apply the pulse scheme M-R<sub>1</sub>-R<sub>2</sub>-M, while sweeping through the RFX-RFY plane by adding a DC-offset to the RF-gates, waiting wait for 200 ns at points  $R_1$  and  $R_2$ .<sup>44,53</sup> Data acquired during the pulse sequence is discarded, and we read out the state of the system only at the final point M. If, during a scan, point M falls deep into one of the charge-stability regions, we then simply observe the same response as in a charge scan without pulses applied (see Fig. 2). However, if the pulse sequence M-R<sub>1</sub>-R<sub>2</sub>-M drives the system through three stability regions as indicated in Fig. 4(a), the measured signal will then have a value between the one corresponding to the (2,0) charge state and that of the (1,1) state. The reason is that when we step from  $R_1$  to  $R_2$ , we initialize at random either a singlet S(2,0) or a triplet T(2,0) state. If the system is in the T(2,0) state, then it tunnels into the (1,1) configuration when we step back to point M. Vice versa, if the system in  $R_2$  is in the S(2,0) state, it remains in this state. In this way, we map out the so-called measurement triangle (or trapezoid, if the singlet-triplet splitting is smaller than the inter-dot charge coupling, as in Fig. 4), i.e. the region of the RFX-RFY plane where Pauli spin blockade allows for spin-charge conversion.

To determine the position of the singlet reload point, we extend the pulse scheme to M-R<sub>1</sub>-R<sub>2</sub>-M-S-M (see Fig. 4(c)), by including an additional 100 ns pause at point S. When point S stays energetically between the (1,1)-T(2,0) and (1,1)-S(2,0) transitions (see Fig. 2), then electron exchange with the Y-reservoir will lead to the initialization of a (2,0) singlet state. If this is the case, measuring the state of the system back at point M will give a value of  $V_{\rm rf}$  corresponding to the (2,0) charge state, instead of the intermediate value observed with the M-R<sub>1</sub>-R<sub>2</sub>-M pulse scheme. Scanning the position of the pulse cycle over the RFX-RXY plane (the relative position of the points M, R<sub>1,2</sub> and S is kept fixed) maps out the area known as "mouse bite", visible in see Fig. 4(d).

Once the "mouse bite" has been identified, we also know a suitable position of point S for fast initialization of the qubit in the S(2,0) state. To further optimize the position of S, we repeat the pulse sequence M-R<sub>1</sub>-R<sub>2</sub>-M-S-M, but now sweeping the position of point S perpendicularly to the (2,0)-(1,0) transition-line, while keeping all other points of the sequence fixed. In particular, point M has to lay within the "mouse bite". As before, we measure the state of the system only in the final point M. The response signal  $V_{\rm rf}$  shows a plateau as a function of the position of point S, at the signal-level of the (2,0)charge ground state, see Fig. 5(a). The two ridges where the signal increase represent the onset of the transitions  $(1,0) \rightarrow S(2,0)$  and  $(1,0) \rightarrow T(2,0)$ , respectively. The optimal position of point S for the operation of the qubit lays symmetrically between these two points.



Figure 4. (a) High-resolution charge stability diagram around the (2,0)-(1,1)-transition used to define the two-electron spin qubit. Important points used to initialize the qubit in different states and for measurement, are marked by dots, and further explained in the main text. Transitions are labeled in white. (b) Diagram showing the energy relaxation cascade used to initialize the (1,1) ground state  $|\uparrow\uparrow\rangle$  at point T<sub>+</sub> (see Sec. III F).  $S_L \otimes \downarrow (\uparrow)$  denotes the state of a singlet state in the left quantum dot and an down (up)-state in the right dot. Arrows indicate relaxation via electron exchange with the (2,1)charge configuration. (c) Charge stability diagram measured using the pulse sequence  $M-R_1-R_2-M$  (see Sec. III E). The measurement trapezoid appears as an area where the readout signal  $V_{\rm rf}$  is between the values corresponding to the S(2,0) and to the (1,1) configurations (turquoise-yellowish area). The blurred boundaries of the readout trapezoid reflect a failure of the random load pulse sequence rather than instabilities. (d) Adding a wait time at point S after the pulse sequence from (c) maps out the "mouse-bite" (yellow area within the measurement trapezoid), i.e. the region of singlet reload within the measurement triangle (see Sec. IIIE).

In a last characterization scan, we fix point S at the optimal position and repeat the pulse sequence M-R<sub>1</sub>-R<sub>2</sub>-M-S-M now varying the waiting time t at point S. Again, we measure the state of the system only in the final point M. The longer we wait in point S, the higher the probability to initialize a singlet S(2,0), and therefore the lower the value of V<sub>rf</sub> measured at point M, see Fig.5(b). We fit these date with a simple exponential decay

$$V_{\rm rf}(t) = V_{\rm rf,0} + Ae^{-\frac{t}{t_{\rm load}}},\tag{4}$$

where  $V_{\rm rf,0}$ , A and  $t_{\rm load}$  are fit parameters. For a well– tuned dot, the singlet reload–time  $t_{\rm load}$  typically lies in the range of 10 to 50 ns. This characterization scan is complementary to the one presented in Sec. III C. It exploits the full time resolution of 1 ns of the AWG, as it is not limited by the bandwidth of the readout tank circuit.

Having identified an optimal singlet–reload point and characterised the singlet reload–time  $t_{\text{load}}$ , in the rest of this paper whenever write "initializing the qubit in the singlet S(2,0) state" we mean the following procedure: i) go to the optimal point S, ii) wait in this position for ~  $5 t_{\text{load}}$ , iii) move to measurement point M. Note that this initialization procedure requires only electron exchange with one lead, which means that only one tunnel barrier has to be tuned to find an optimal operation regime. Moreover, the initialization time is simply given by the tunnel coupling to this lead, and can be as fast as a few tens of nanoseconds.

## F. Locating the triplet $T_+$ reload point

A fundamental technique for the operation of qubits based on GaAs is dynamical nuclear polarization (DNP). This technique is used for stabilizing the surrounding bath of nuclear spin, and relies on the ability to initialize the (1,1) ground state  $T_+$ .<sup>28</sup> Originally, this was done exploiting both the (2,1)-(1,1) and the (1,0)-(1,1) transitions, i.e. allowing electron exchange with both  $leads^{38}$ . Here we report a different approach, again based on tunneling only to one lead. The trick is to exploit the relaxation cascade shown in Fig. 4(b), which characterizes the region of the stability diagram close to the (1,1)-(2,1)transition. In the presence of an external magnetic field  $B_{\rm ext}$ , the triplet  $T_+$  represents the ground state of the (1,1) charge configuration, and transitions from the (2,1)ground-state to the excited states of the (1,1) configuration are not energetically allowed close to the (1,1)-(2,1)boundary. Hence, if we initialize the qubit in the S(2,0)state, and then pulse to a point  $T_+$  close to the (1,1)-(2,1) transition, (see Fig. 2(a)) the qubit either ends up directly into the T<sub>+</sub> state, or it will eventually reach this state at the end of the relaxation cascade sketched in Fig. 2(b). Importantly, for this to happen we need to ensure that the exchange interaction satisfy the requirement  $B_{\text{ext}} > J(\epsilon) > \Delta B_z$ , which is necessary for having sufficient mixing between the  $|\uparrow\downarrow\rangle$ ,  $|\downarrow\uparrow\rangle$  states and the full relaxation to the  $T_+$  ground state. Here  $\Delta B_z$  is the difference of magnetic field in the two dots.

To find the optimal  $T_+$  reload point in the charge stability diagram, we perform the following sweep. We initialize the qubit in the S(2,0) state and then pulse to point  $T_+$  without crossing the upper triple point to avoid measurement artefacts. The distance between  $T_+$  and the upper triple-point has to be chosen such as to fulfil the energy requirement  $B_{\text{ext}} > J(\epsilon) > \Delta B_z$ . After a waiting time of 100 ns to allow energy relaxation, we switch back to the measurement point M and measure



Figure 5. (a) To determine the optimal position of the singlet reload point, we shift the position of point S in the pulse sequence M-R<sub>1</sub>-R<sub>2</sub>-M-S-M along the direction  $\delta_{\rm S}$  (see text and Fig. 4(a)). The optimal position for S, is in the middle of the plateau of low  $V_{\rm rf}$  values. (b) The singlet reload-time is extracted by applying the pulse sequence M-R<sub>1</sub>-R<sub>2</sub>-M-S-M, and measuring the triplet return-probability as a function of the waiting time t at point S. (c,d) The positions of the T<sub>+</sub> reload-point and of the ST<sub>+</sub> transition can be determined using the pulse sequences described in Sec. III F and III G respectively, and result in peaks in the measured  $V_{\rm rf}$  signals as function of the displacemente  $\delta_{\rm T}$  and of the detuning  $\epsilon$ .

the state of the system. We repeat this procedure while sweeping the position of point  $T_+$  by  $\delta_T$ , perpendicularly to the direction of the  $(1,1)\rightarrow(2,1)$  transition (Fig. 4(a)). The optimal position of point  $T_+$  appears as a maximum of  $V_{rf}$  as function of  $\delta_T$  (see Fig. 5(c)), indicating that indeed a triplet was initialized while waiting at point  $T_+$ . To extract the exact position of the reload point, we use a phenomenological model motivated by Eq. (2) and given by

$$V_{\rm rf}(\delta_{\rm T}) = V_{\rm rf,0} + \frac{1}{2} A_1 \left( 1 + \tanh\left(\frac{\delta_{\rm T} - \delta_{\rm tl,1}}{w}\right) \right) - \frac{1}{2} A_2 \left( 1 + \tanh\left(\frac{\delta_{\rm T} - \delta_{\rm tl,2}}{w}\right) \right)$$
(5)

to fit the data. The position of the  $T_+$  point is then given by  $(\delta_{tl,1} + \delta_{tl,2})/2$ .

## G. Locating the $ST_+$ -transition

In addition to the location of the  $T_+$ -reload point, it is also necessary to know the location of the S- $T_+$ -anticrossing to perform DNP. To find the latter, we follow Ref.  $^{43}$  and initialize the qubit in the singlet state, then change the detuning  $\epsilon$ , wait 100 ns at a given detuning, and then return to the measurement point and read out the final qubit state. When the detuning is at the  $S-T_+$ anti-crossing, the hyperfine and the spin-orbit interaction can turn the initialized S-state into an T<sub>+</sub>-state, giving rise to a maximum in the measured  $V_{\rm rf}$  as a function of  $\epsilon$ . Because the location of the ST<sub>+</sub> transition strongly depends on the local magnetic field, any unintentional polarization, for example, due to hyperfine mediated spin flips at the  $ST_{+}$  transition, shifts the precise position of the anti-crossing. To avoid this problem, we include pauses of a few milliseconds at the end of each  $\epsilon$ -sweep, to allow any unintentional polarization to relax. If needed, we average over a few different sweeps, and fit our data with a Gaussian model

$$V_{\rm rf}(\epsilon) = V_{\rm rf,0} + \delta V_{\rm rf}\epsilon + Ae^{-\frac{(\epsilon - \epsilon_{\rm stp})^2}{2w^2}} \tag{6}$$

to extract the position  $\epsilon_{\text{stp}}$  of the ST<sub>+</sub> transition ( $V_{\text{rf},0}$ ,  $\delta V_{\text{rf}}$ , A,  $\epsilon_{\text{stp}}$  and w are fit parameters).

Not only is this position crucial for the pulsed DNP scheme but, in combination with the  $T_+$  reload point, it is also used as an anchor point in the charge stability diagram. Adjusting the dot using the X and Y virtual gates to obtain the same values for the  $ST_+$  and  $T_+$  scan after a small charge–switching event usually restores all quantum dot parameters, and results in the same  $J(\epsilon)$ relation. Furthermore, the position of the  $ST_+$  crossing is used to automatically determine switching events<sup>20,54,55</sup> that shift the whole transition by several mV.

#### H. Tuning workflow

To summarize, once the (2,0)-(1,1) charge transition has been identified, the typical fine-tuning workflow of a  $ST_0$ -qubit starts by defining the virtual gates. The next step is to bring the tunnel couplings to the leads in the right regime. Then, the singlet reload point S and the measurement triangle are determined. The energy splitting  $J(\epsilon)$  of the qubit is subsequently tuned by adjusting the inter-dot tunnel coupling. A working scan of the S-T<sub>+</sub> transition as described in Sec. III G is a good indicator of a suitable inter-dot tunnel coupling for the operation of the qubit. Usually, tuning the tunnel couplings is an iterative procedure, as adjusting the T and N virtual gates used to tune the inter-dot coupling also affects the coupling to the leads a little. Finally, the position of the  $T_+$  reload point is determined. During the whole tuning procedure we periodically check the exact position of the (2,0)-(1,1) transition by recording a charge stability diagram. The triple points, which act as anchor points, are extracted automatically by a fit that includes a model of the charge transition, as described in Sec. III A. The lower triple point is used as a reference for the measurement point, and either an offset on the RF-gates or on the virtual gates X and Y is used to center the transition accordingly. The fit stability of all scans requires a signal-to-noise ratio of the order of 5 (measured as the ratio of a transition step size to the rmsfluctuation away from the transition in a charge stability diagram). To ensure a high sensitivity, we periodically check the sensing dot operating point (see Sec. A 1) by performing line scans through the charge stability diagram in Fig. 6) and adjust the sensor dot gate voltages accordingly. Manual retuning to restore the quantum dot parameters once the charge sensor becomes insensitive or a charge rearrangement occurs takes in general a few iterations of performing the various characterization scans and adjusting the gate voltages, and can be typically performed in a couple of minutes.

#### IV. CONCLUSION

This paper provides a detailed description of tuning and characterization routines that we use to realize a  $ST_0$  qubit in a GaAs double quantum dot. We describe efficient methods to determine the tunnel couplings between the dots and to the leads, and methods to locate the various points in the charge stability diagram that are needed for the operation of the qubit itself or for pulsed feedback DNP.

While all relevant quantitative double dot parameters are already obtained automatically, the decision of how to adjust the gate voltages is currently man-made by the operator, based on experience. A crucial next step is to also automate this step. One complication is that the effect of the T and N gates on the inter-dot tunnel coupling changes substantially in different regions of gate voltage space, or when charge rearrangements in the vicinity of the dot occur, including even sign changes. This behavior renders tuning algorithms based exclusively on precalibrated gradient information ineffective, but could be addressed with more sophisticated, adaptive approaches. For example, the use of a Kalman filter 56 to continuously update the response tensor based on the recent tuning history appears promising. Hence, we are confident that the procedures described here will be a very useful basis for reaching that goal. Such advances will be indispensable as soon as the number of qubits increases substantially. It will likely also be necessary and possible to detect data sets affected by charge rearrangements, e.g. by plausibility checks on the fit parameters and residuals.

While all measurements presented in this paper were performed on a GaAs double quantum dot operated as  $ST_0$  qubit, the only procedures that are GaAs-specific are those needed to set up DNP operating points. All other tuning methods are equally adaptable to Si-based devices.

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#### Appendix A: Coarse tuning of the quantum dots

In this appendix we cover the first step of tuning the device to either the (2,0)-(1,1) or the (1,1)-(0,2)-chargetransition. Additionally, we describe the tuning of the adjacent quantum dot used for charge sensing of the qubit dots. These methods have hardly changed compared to standard quantum transport measurements<sup>38</sup> and will need further refinement<sup>33</sup> for automation. They are included here for completeness. For the initial coarse tuning of the double dot, instead of using RF-reflectometry, we directly measure the conductance through the double dot and through the sensing dot (see Fig. 1). To do so, we apply a voltage bias of 100  $\mu$ V across the devices. The resulting currents are converted to voltages (named  $V_{\rm SD}$ and  $V_{\rm D}$  for the sensing dot and double dot, respectively) using a home-built IV-converter and measured with a lock-in amplifier.

#### 1. Tuning of the sensing dot

The first step in the tuning procedure is to set up charge detection through the sensing dot. This requires finding a set of voltages applied to the sensing dot gates SB1, SB2 and SP (gate names are defined in Fig. 1) such that the conductance through the dot is maximally sensitive to the local electrostatic potential. To do so, we measure  $V_{\rm rf}$  while performing a two-dimensional scan with the sensing dot gates SB2 vs. SB1&SP. Since  $V_{\rm rf}$ depends on the conductance through the dot, Coulomb oscillations appear in the measured signal when the applied voltages are sufficiently negative to make the source and drain barriers opaque. Fig. 6 shows a region in gate voltage space that shows the typical pattern of a single quantum dot.<sup>58</sup> In this particular sample, SP and SB1 were shorted and thus had to be kept on the same potential. Usually, SP can be used to fine-tune the dot and to shift it closer to the double quantum dot. To obtain the best charge sensitivity, the voltages applied to SB2



Figure 6. Characteristic charge stability–diagram of the sensing dot, measured with RF-reflectometry. Depending monotonically on the conductance through the dot,  $V_{\rm rf}$  shows Coulomb oscillations once the source and drain barriers are sufficiently opaque. Tuning the sensor to a sensitive position (see circle) allows for charge sensing of the nearby double quantum dot.

vs. SB1&SP have to be tuned to values where the slope of the Coulomb peak is steepest.

# 2. Locating the (2,0)-(1,1) or (0,2)-(1,1) charge transition

The second step is to determine the depletion and pinch-off voltages of the different gates that define the qubit double quantum dot. To do that, we directly measure the conductance through the double dot by applying a 100  $\mu$ V bias voltage,  $V_D$ , as shown in Fig. 1, and measuring the resulting current. Measuring the conductance as a function of the voltage applied pairwise to the gates N12 and T12, S1 and B1, S2 and B2 (see Fig. 1 for gate nomenclature), allows us to determine the depletion voltages.

The gate voltages are then set close to their depletion voltages and the device is tuned close to complete pinchoff. Next, we perform a two-dimensional scan over a couple of tens of mV with the gates B1 and B2. Usually we anticipate to first form a large single quantum dot and then separate it into two dots by applying more negative voltages on the T12 and N12 gates. If the tunnel barriers between the dot and leads X and Y are almost pinched off and have similar transmission probabilities, Coulomb blockade peaks should appear, showing the characteristic honeycomb pattern of a lateral double quantum dot.<sup>58</sup>



Figure 7. (a) Honeycomb pattern of the double dot resolved using the sensing dot. Background oscillations are caused by an imperfect compensation of the response of the sensor to gates B1 and B2. (b) Direct transport measurement through the double dot. Coulomb peaks are visible only for not too negative voltages. (c) Same as in (a), but this time using the side gates S1 and S2 instead of B1 and B2. This typically reduces the background oscillation in the transconductance of the sensing dot. The different intensities of the lines delineating the honey-comb pattern reflect the transparency of the tunnel barriers to the external leads

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Observing this honeycomb pattern in the direct current through the double dot close to pinch-off can be challenging because the current goes to zero and Coulomb peaks are hardly detectable, see Fig. 7(b). To study this regime, we then use the sensing dot. Due to the capacitive coupling between the double dot and the sensing dot, a change in the occupation of the double dot results in an abrupt change in the current through the sensing dot and therefore into a sharp signature in the transconductance  $dV_{SD}/dB_1$ , Fig. 7(a). When performing this type of scan, the voltage SB2 is adjusted to compensate the unintentional influence of the stepping gate B2 on the potential of the sensing dot. Similar scans can also be performed by using the side gates S1 and S2 instead of B1 and B2. This typically reduces the background oscillation in the transconductance of the sensing dot (see Fig. 7c), as the gates S1 and S2 have a weaker influence on the sensing dot than B1 and B2. Going towards more negative voltages eventually locates either the (2,0)-(1,1)or the (0,2)-(1,1) charge transition. Once a suitable transition has been found, we adjust S1 and S2 such that a recorded high-resolution charge stability diagram via RF-reflectometry using the RF-gates, RFX and RFY, is centered around the transition of interest (see Fig. 2 (a)).

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