A model for single electron decays from a strongly isolated quantum dot

J. Martorell
Dept. d'Estructura i Constituents de la Materia, Facultat Física, University of Barcelona, Barcelona 08028, Spain

D. W. L. Sprung and P. A. Machado
Department of Physics and Astronomy, McMaster University
Hamilton, Ontario L8S 4M1 Canada

C. G. Smith
Semiconductor Physics Group, Cavendish Laboratory, Madingley Road
Cambridge CB3 0HE, UK

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Recent measurements of electron escape from a non-equilibrium charged quantum dot are interpreted within a 2D separable model. The confining potential is derived from 3D self-consistent Poisson-Thomas-Fermi calculations. It is found that the sequence of decay lifetimes provides a sensitive test of the confining potential and its dependence on electron occupation.

73.23.Hk, 73.20.Dx, 73.40.Gk, 85.30.Vw, 71.10.-w

I. INTRODUCTION

In a recent experiment [1], a strongly isolated quantum dot was charged with excess electrons, and their sequential escapes were recorded over a one hour time period. This was repeated 150 times to obtain a statistical distribution of decay times. The dot is formed in an electron gas located at a depth of 70 nm in a GaAs–AlGaAs heterostructure. Its shape is defined by electrostatic confinement using a set of gates, as sketched in the insert to Fig. 1. The gate voltages were ramped up quickly, so that the dot retained a sizeable number of excess electrons when it was well isolated from the surrounding electron gas. The observations correspond to sequential tunnelling of (seven) electrons from the dot to the surroundings. The lifetimes extracted from the escape times distribution [1] are shown in Fig. 1. A striking quasi-linear dependence of the logarithm of the lifetime on electron number is apparent.

Sequential decays have been known and studied for over a century in the context of nuclear physics. The combined instances of alpha and beta decays from the heaviest elements are responsible for most natural radioactivity. The description of alpha decay in terms of tunnelling of alpha particles through a confining potential dates back to the 1920's (Gamow [2], Condon and Gurney [3]). Although the basic nature of the decay as a barrier penetration is well understood, accurate predictions for radioactive lifetimes are difficult because the process by which the escaping alpha-particle is preformed within the nucleus requires an understanding of four-body correlations. As a result, it is impossible to deduce accurate information on the barrier shape. Nevertheless, alpha decays have provided useful information on nuclear radii and the range and gross features of the nuclear interaction [4].

FIG. 1. Experimental lifetimes (in seconds) extracted from the decay sequences, as reported ref. [1]. The insert shows the gate arrangement which defines the dot.

It has become commonplace to say that a quantum dot is an artificial atom, but in fact the self-consistent potential confining electrons in a large dot has more in common with the mean field potential in a heavy nucleus: flat in the interior, with abrupt walls. An artificial nucleus is a more apt description, as will become clear in this paper. Indeed, the detection of sequential decays from an isolated quantum dot is a more favourable situation for study of the decay process, as the question of preforming the electron does not arise. Hence, we can more confidently test our knowledge of the confining barriers for electrons, as well as the profile, and dependence on occupation number, of the dot potential. We will analyze these aspects in this work, and show that these measurements of the lifetimes of “radioactive quantum dots” introduce new constraints on our ability to model...
their structure.

The present experiment has another significant advantage over nuclear decays: instead of counting incoherent decays from a large sample of identical nuclei, here a single dot is involved, and the correlation between consecutive events can be analyzed. In addition, it should be possible to design the shape, density and excitation energy of the dot within rather broad margins, so that future experiments on mesoscopic systems will be much more flexible than those in nuclear systems, where only those nuclei existing in nature, or created in sufficient numbers, can be studied. Thus, the study of electron decays from a quantum dot has the potential to reveal new features of the tunnelling process. This is a topic of currently renewed interest: see for example van Dijk and Nogami [5]. The type of simple model developed in this paper can be of great utility in such future studies.

In this work we will describe the decay process using analytic models which incorporate characteristics of the confinement potential extracted from realistic numerical simulations. As the dot contains about 300 electrons, Poisson-Thomas-Fermi calculations should be adequate to describe the electron density and the confining potential of the dot. With these in hand we have developed accurate analytic approximations for the confining potential that allow us to construct an envelope approximation wavefunction for the electrons in the dot, and to compute the electron lifetimes from a fully quantal expression for the transmission amplitude across the barrier.

Previous works which model a quantum dot have been concerned with the wave functions of confined states in the dot, the electron density distribution and the shape of the confining potential. For such purposes, only the inside of the barrier matters. It is when one looks at the escape of electrons from the dot that the barrier height, its width and shape become important; these are the new features explored in this paper. In section II we describe the development of our model, while in Section III we discuss the results for the sequence of lifetimes and compare them with experiment. Some details are relegated to two appendices.

II. MODELLING OF ISOLATED DOT DECAYS

A. Framework

The Poisson-Thomas-Fermi modelling is described in more detail elsewhere [6], so here we list only the main steps:

i) first, Poisson-Schrödinger (PS) and Poisson-Thomas-Fermi (PTF) simulations as described in [7] are performed for the ungated heterostructure. Our inputs for the PS simulation are the thickness and composition of each layer in the heterostructure, and the dopant concentration in the donor layer. From these we predict the density of the 2DEG. The only adjustable parameter is the donor ionization energy which is set to be $\epsilon_\Phi_x = 0.12 \, eV$, in order to reproduce the measured 2DEG density, $n_e = 2.74 \times 10^{11} \, cm^{-2}$. For the simpler Poisson-Thomas-Fermi scheme we employ a common relative permittivity $\varepsilon_r = 12.2$ for all layers of the heterostructure, which, combined with the parameters already used for the PS simulation, also reproduces the experimental $n_e$.

After this “fitting” the model has no other free parameters.

ii) For the gated structure we use the gate layout and voltages of the experiment. To solve the Poisson equation for the gated heterostructure one has to impose as a boundary condition the value of the electrostatic potential on the exposed surface of the heterostructure, and on the gates. We assume Fermi level pinning and choose the energy of the surface states as the zero of the energy scale. In this convention, the conduction band edge is set at $\epsilon V_g = 0.67 \, eV$ on the exposed surface. Under each gate the conduction band is set at $\epsilon V_{m,i} + \epsilon V_g$, where $V_g$ is the gate voltage and the metal semiconductor contact potential, $\epsilon V_{m,i}$, is taken as $0.81 \, eV$ [9]. The electrostatic potential due to the gates is then computed using semi-analytic expressions based on the work of Davies et al. [10] and [11]. Added to this are: a) the Coulomb potential (direct term) between the electrons, and a mirror term which imposes the boundary conditions at the surface, and b) the contribution from the fully ionized donor layer and its mirror term (see Sect. IIA of [8] for details of a similar example.) We neglect exchange and correlation effects, which are small.

iii) The connection between the confining potential defined by the conduction band edge and the electron density is completed by using the Thomas-Fermi approximation at zero temperature:

$$\rho_e(\mathbf{r}) = \frac{1}{3\pi^2} \left( \frac{2m^*}{\hbar^2} (E_F - \epsilon V(\mathbf{r})) \right)^{3/2}$$

The PTF iteration is performed starting from the ungated heterostructure densities as trial values.

B. Equilibrium dot

As a first step, we examine the dot in its final state after all the excess electrons have escaped. This corresponds to a PTF simulation with the same Fermi level, $E_F,\dot{\rho}_e = 0$, for the electrons in the dot and in the 2DEG outside the barriers. The gate voltages are taken from ref. [1] as $V_{PL} = -0.40 \, V$, $V_{C1} = V_{C2} = -0.44 \, V$ and $V_H = -0.7 \, V$. The predicted PTF 3D electron distribution $\rho_e(x,y,z)$ is more conveniently visualized in terms of a projected 2D density:

$$n_e(x,y) = \int_{z_3}^{z_1} \rho_e(x,y,z) \, dz,$$
where \( z_j \) is the junction plane. The \( n_c(x, y) \) distribution, shown in Fig. 2, has an approximately rectangular boundary, and its maximum value is close to the 2DEG density of the ungated heterostructure. In this calculation the dot contains 286 electrons.

![FIG. 2. The two dimensional PTF density, \( n_c(x, y) \), for a dot in equilibrium with the surrounding 2 DEG.](image)

C. Dot with excess electrons

To study these configurations we set the Fermi level inside the dot, \( E_{F, dot} \), higher than its value outside the barriers, \( E_{F, 2DEG} = 0 \). We can do so because the dot is well pinched off from the surrounding electron gas. We ran PTF simulations with equally spaced values for \( E_{F, dot} \) running from 0 to 17.5 meV in steps of 2.5 meV. The occupation \( Q \) of the dot increases linearly with \( E_{F, dot} \) at the rate 2.75 electrons per meV, giving occupations \( 286 \leq Q \leq 334 \).

The simulations also produce the confining potential for the electrons in the dot, \( eV(x, y, z) \). To reduce this to a two-dimensional function, \( U(x, y) \), we take a weighted average over the density profile in the \( z \) direction:

\[
U_{PTF}(x, y) = \frac{\int_{z_1}^{\infty} eV(x, y, z) \, P(z) \, dz}{\int_{z_1}^{\infty} P(z) \, dz}
\]

(3)

where

\[
P(z) = \int_{\Omega} \rho_c(x, y, z) \, dx \, dy.
\]

(4)

Here the domain of integration \( \Omega \) is a rectangle in the \( xy \) plane which extends a short distance into the surrounding electron gas, from \((x_i, y_i) = (-510 \text{ nm}, -255 \text{ nm})\) to \((x_f, y_f) = (510 \text{ nm}, 255 \text{ nm})\). This includes an area outside the dot where the 2DEG is still depleted by the gates. Although the computed \( V(x, y, z) \) is not separable, previous experience with Poisson-Schrödinger simulations of wires \[8\], \[12\] and circular dots, has shown us that the factorization ansatz leads to very good approximations when the \( z \) degree of freedom is integrated out as in eq.

3. This prescription to construct the 2D potential avoids the type of \textit{ad hoc} assumptions often made.

In Fig. 3 we show the \( U_{PTF}(x, y) \) corresponding to the equilibrium dot of Fig. 2. As expected from the gate layout shown in the inset to Fig. 1, it has two very high barriers running parallel to the \( x \) axis, one centered at \( y = 0 \) and the other that begins with a steep rise at \( y \approx 400 \text{ nm} \) (and shows clearly the mark of the three-fingered gate layout labelled \( C_1, C_2 \), and \( PLunger \) in Fig. 1). Tunnelling across these barriers is negligible. In addition there is a symmetric pair of barriers running parallel to the \( y \) axis, with maxima at \( x \approx \pm 238 \text{ nm} \) through which the electrons \textit{do} tunnel. In the interior, the potential is practically constant. Although these \( x \)-barriers have somewhat increasing height with increasing \( y \), the rectangular shape of the potential suggests using a separable approximation in cartesian coordinates:

\[
U_{PTF}(x, y) \approx U_z(x, y) = U(x) + W(y)
\]

(5)

![FIG. 3. Two dimensional confining potential, \( U_{PTF}(x, y) \) for the dot of Fig. 1.](image)

![FIG. 4. Continuous lines: sections at \( y = 200 \text{ nm} \) of the \( U_{PTF}(x, y) \) corresponding to \( E_{F, dot} = 0.0, 0.005, 0.010 \) and 0.015 eV. Dashed lines: analytic parametrization for \( U(x) \) as described in the text (the latter shown only for \( x > 0 \) for clarity.)](image)
We will interpret the experimental decay data using this separability ansatz. For the $W(y)$ barriers, which are basically impenetrable, we use two simple models described below. As a guide to a realistic choice for the $x$-dependent term we examine in Fig. 4 the profiles of $U_{PTF}(x,y)$ at a fixed value of $y = 200 \, \text{nm}$ in the middle of the dot. The profiles shown cover a range of occupations of up to forty excess electrons. In this range, the potential at the dot center increases linearly with $Q$, according to

$$U_b = 0.347Q - 118.4 \, \text{meV}.$$  \hspace{1cm} (6)

At large distances outside the dot, $U_{\infty} = -18.8 \, \text{meV}$ is constant. Similarly, the location of the barrier maximum and its height can be parametrized as:

$$x_b = \frac{238 - Q - 286}{16} \, \text{nm}$$

$$U_b = 0.117Q - 13.4 \, \text{meV}.$$  \hspace{1cm} (7)

Note that $dU_b/dQ \approx 1/3dU_0/dQ$ reflects the decrease of the screened Coulomb repulsion away from the center of the dot. Furthermore, we have found that the $x$-dependence can be very well reproduced (see Fig. 4) using the following analytic model:

$$U(x) = U_b + U_{MF}(x), \quad x > 0,$$

$$= U(0), \quad x < 0,$$

where

$$U_{MF} \equiv U_c \frac{\sin^2 \left( \frac{x-x_b}{y_b} \right)}{\cosh^2 \left( \frac{x-x_b}{y_b} - \mu \right)}.$$  \hspace{1cm} (8)

This potential form has the great advantage that the transmission coefficient for $U_{MF}$ is known analytically [13]. $U_{MF}$ is an asymmetric barrier which takes one value for $x << x_b$ and another value for $x >> x_b$:

$$U_{MF}(x_b) = 0$$

$$U_{MF}(\infty) \equiv \lim_{x \to \infty} U_{MF}(x) = U_c e^{2\mu}$$

$$U_{MF}(\infty) \equiv \lim_{x \to -\infty} U_{MF}(x) = U_c e^{-2\mu}.$$  \hspace{1cm} (9)

The parameters $U_b, U_c, \mu, x_b, w_b$ allow one to fit the barrier height, the potential floors inside and outside the dot, the barrier spacing and the barrier width. Since the barriers are spread quite far apart, in practice $x_b >> w_b$, so $U_{MF}(x = 0) \approx U_{MF}(\infty)$. In this case,

$$U_b \equiv U(0) \approx U_b + U_c e^{-2\mu}$$

$$U_{\infty} \equiv \lim_{x \to \infty} U(x) = U_b + U_c e^{2\mu}.$$  \hspace{1cm} (10)

Then we can solve for

$$\mu = \frac{1}{4} \ln \left( \frac{U_b - U_{\infty}}{U_b - U_b} \right) \quad \text{and}$$

$$U_c = -(U_b - U_{\infty}) e^{2\mu}.$$  \hspace{1cm} (11)

To determine the parameters appearing in eq. 8, we take the values of the PTF potential at the origin, $U_b$, well beyond the barrier, $U_{\infty}$, and the value $U_{x_1}$ at the barrier maximum $x = x_b$, and then plot $U(x)$ to find the best $w_b$, which turned out to be 48 nm. This gives a convenient analytic form for the confining potential, motivated by PTF, whose transmission coefficient is:

$$T = \frac{2\sinh(\pi k_+ \sinh(\pi k_-))}{\cosh(\pi(k_+ + k_-)) + \cosh(\pi\beta)},$$  \hspace{1cm} (12)

where

$$k_+ = \sqrt{\frac{2m^*}{\hbar^2}(E - U_0)}$$

$$\beta = \sqrt{\frac{2m^*}{\hbar^2}(2U_b - 2U_c - U_0 - U_{\infty})} w_b^2 - 1.$$  \hspace{1cm} (13)

Barrier shape $W(y)$: In Fig. 5 we examine a section of $U_{PTF}(x = 0, y)$ through the center of the dot. We use two approximate models, the simplest one being an infinite square well, of width $w_y \approx 350 \, \text{nm}$. The slightly fancier one is a truncated harmonic oscillator:

$$W_{tho}(y) = \begin{cases} 0 & (\text{flat bottom}) \\ -0.13 + \frac{1}{2} k_y (y - y_b)^2 & (\text{walls}) \end{cases}.$$  \hspace{1cm} (14)

with $y_b = 238 \, \text{nm}$ and $k_y = 7.35 \cdot 10^{-4} \, \text{nm}^{-2}$. As can be seen in Fig. 5 this parametrization (plus the constant term $U_0$) reproduces the main features of the $x = 0$ sections of the PTF potentials.

By combining eqs. 6 to 14 we determine a separable analytic potential model for the dot containing a desired number $Q$ of electrons. This removes the necessity of repeatedly solving the PTF equations for the self-consistent field, while studying the decay process.

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**FIG. 5.** Continuous lines: sections at $x = 0 \, \text{nm}$ of the $U_{PTF}(x,y)$ for the same $E_{F,\text{dot}}$ as in Fig. 3. Dashed lines: analytic parametrization of $W_{\text{tho}}(y) + U_0$ as described in the text.
D. Quasibound states of the dot

We construct the electron wave functions inside the dot in the envelope function approximation, using our parametrized potential, \( U_s(x,y) \). The single electron energies are

\[
E_{n_x,n_y} = E_{n_x} + E_{n_y}
\]

and the electron wave functions factorize

\[
\Psi_{n_x,n_y}(x,y) = \phi_{n_x}(x)\psi_{n_y}(y) .
\]

The factors satisfy 1D Schrödinger equations:

\[
-\frac{\hbar^2}{2m^*}\phi''_{n_x}(x) + U(x)\phi_{n_x}(x) = E_{n_x}\phi_{n_x}(x)
\]

\[
-\frac{\hbar^2}{2m^*}\psi''_{n_y}(y) + W(y)\psi_{n_y}(y) = E_{n_y}\psi_{n_y}(y)
\]

The second equation is for a confined wavefunction, easily solved by standard numerical methods. We label the solutions by the number of loops, \( n_y \), of the eigenfunction. For example, taking \( W(y) \) to have hard walls, the energy is

\[
E_{n_y,sw} = \frac{\hbar^2}{2m^*} \left( \frac{n_y \pi}{w_y} \right)^2 .
\]

For the truncated harmonic oscillator shape there is no similar analytic expression, but the dependence on \( n_y \) is similar.

The \( x \)-dependent equation describes 1D electrons confined in the dot by the “leaky barriers”. Weaker quasibound state solutions were computed using methods described in [14]. However, for levels corresponding to the long tunnelling lifetimes observed in the experiment, the energies and eigenfunctions can be computed well enough by the simpler prescription of setting the electron wavefunction to zero at the points \( \pm x_0 \) inside the barriers. Furthermore, if only the eigenvalues and lifetimes are needed, we have checked that the WKB quantization condition is adequate:

\[
\int_{x_1}^{x_2} \sqrt{\frac{2m^*}{\hbar^2}} \left( E(n_x) - U(x) \right) dx = \left( n_x - \frac{1}{2} \right) \pi
\]

In the Appendix we describe the determination of the lifetimes, \( \tau_{n_x} \). From here on the energies presented are obtained in the WKB approximation. The differences from the more accurate predictions using the true quasibound state energies can scarcely be seen on the scale of the graphs. For barrier penetrability we use eq. 12.

We “construct” the desired dot configuration with excess electrons by generating a \( U_s(x,y) \) for the chosen value of \( Q \), and filling the levels as follows: i) First we list the \( \{E_{n_x}, \tau_{n_x}\} \), in order of increasing \( n_x \) (and therefore of increasing energy and decreasing lifetime.) This list is truncated at an \( n_x = n_{x,\text{max}} \) whose lifetime is less than 0.01 sec. b) Next we form a list of 2D levels \( (n_x, n_y) \) by choosing those for which

\[
E_{n_x} + E_{n_y} \leq E_{n_x,\text{max}} + E_{n_y=1} .
\]

The levels in this list are occupied in order of increasing energy and according to Fermi statistics, see eqs. A5 A6. We choose the dot Fermi level so that the number of electrons is the desired \( Q \). It is supposed that, for the long lifetimes observed in the experiment, the electrons have time to lose energy by phonon collisions and occupy the quasibound states of lowest energy. Then, as described in Appendix A, we determine the lifetime for one electron to escape from the dot. This involves a weighted average of the level lifetimes, according to the occupancy of each level at the experimental temperature, \( T' = 100 \text{ mK} \).

To produce a sequence of decays for comparison to experiment we proceed as follows: i) we start with a dot containing a number of electrons, \( Q_0 \), chosen large enough so that the lifetime for one electron to escape is smaller than those observed in experiment. ii) We redetermine the barrier and dot configuration for \( Q = Q_0 - 1 \) electrons, as described in the above paragraph and determine again the corresponding lifetime for escape of one electron. This process is repeated to generate a sequence of decays that covers and extends beyond the range of lifetimes measured in experiment. From that list we choose as the first observed electron decay that corresponding to the \( Q \) whose lifetime is the first to be larger than \( t_0 = 25 \text{ seconds} \).

![FIG. 6. Calculated lifetimes (in seconds) when \( W(y) \) is either the truncated harmonic oscillator: stars, or a square well with \( w_y = 380 \text{ nm} \): + signs. The dotted line is the prediction of the two level model, eq. 22.](image)

III. RESULTS AND DISCUSSION

In Fig. 6 we show results from our model, using parameters chosen as described above, for a range of lifetimes extending over three orders of magnitude. The stars correspond to the truncated harmonic oscillator choice for
with a value \( w_2 = 380 \text{ nm} \) chosen to optimize the agreement with the other prescription in the range of experimental lifetimes, from 10 to 1000 seconds.) One sees that the trends are very similar. For \( Q \) in the neighbourhood of 304, the predicted decay lifetimes fall in the experimental range.

As already mentioned in Section II, our PTF simulations predict \( Q = 286 \) for the dot in equilibrium with the surrounding electron gas. This is also what we find with this separable model, as the curve of lifetimes shown in figure 6 extrapolates smoothly up to \( Q = 287 \), for which we predict a lifetime of \( \log_{10} \tau = 5.2 \), or 44 hours. After that, the Fermi level of the electrons inside the dot falls below that of the surrounding 2DEG and further decays are blocked. It should take almost two days for the dot to reach equilibrium with its surroundings.

Before attempting a more detailed comparison with the experimental data it is useful to examine the main features in our predicted sequences. First we focus on the linear behaviour for values \( Q < 300 \). (We have found similar behaviour in other ranges of \( Q \) when we use slightly different sets of parameters.) Such linear dependence occurs when our model produces a sequence of decays dominated by those from a single 1D electron level; i.e. corresponding to a fixed value of \( n_x \). To understand why, suppose that at zero temperature and for \( Q \) electrons, the occupied level with shortest lifetime is \((n_x, n_y)\), and that \((n'_x, n'_y)\) are occupied levels with higher energy and longer lifetime (this requires that at least \( n'_x < n_x, s \) for longer lifetime and \( n'_y > n_y \) for higher total energy.) When one forms the \( Q - 1 \) electron configuration according to the rules explained above, one of the \((n'_x, n'_y)\) levels will be empty, whereas the level \((n_x, s, n_y)\) will again be filled. In more physical terms: all the electrons with energy above that of the level with shortest lifetime will lose energy by phonon collisions and fall into the leaky level, from which they finally escape. Since the lifetime does not depend on \( n_y \), all the electrons with energy above that of the state \((n_x, s, n_y) = 1\) will escape through the same leaky 1D level, \( n_x, s \), which remains the favoured decay channel as long as it is occupied. Therefore the total probability for one electron to escape from the occupied states with quantum number \( n_x, s \), is the probability for a single 1D electron with energy \( E_{n_x, s} \), multiplied by the number of electrons in occupied states with the same quantum number \( n_x, s \): \( q_{n_x, s} \):

\[
\tau(Q) = \frac{\tau_{n_x, s}(Q)}{q_{n_x, s}(Q)} ,
\]

and when the occupation \( q_{n_x, s} \) of the leaky level is constant, the linear variation of \( \log_{10}(\tau) \) reflects that of the lifetime of the leaky level. This is where the 2D nature of the quantum dot asserts its presence, even though the decay appears to proceed only in one dimension.

In Fig. 7 we show the occupations of the two levels with the shortest lifetimes. One sees that when \( Q < 300 \) the occupation of the \( n_x = 13 \) level stays practically constant and \( n_x = 14 \) level remains empty. For higher values of \( Q \) both levels contribute significantly to the escape lifetime. In this situation:

\[
\tau(Q) = \frac{1}{q_{n_x}(Q) + q_{n_x}(Q)} .
\]

This is shown as the dotted curve in Fig. 6, and it accounts very well for the trend of the lifetimes predicted by the separable model.

![Fig. 7. Total occupation of levels with \( n_x = 13 \) (black squares) and \( n_x = 14 \) (open squares). The lines are drawn to guide the eye. The truncated harmonic oscillator model was used for \( W(y) \).](image)

Our separable model favours the appearance of the degeneracy in lifetime of states with the same \( n_x, s \). A non-separable model would lift that degeneracy and then the lifetime sequences should show a behaviour intermediate between the two situations discussed above. In particular, the sudden change of slope at \( Q = 302 \) in Fig. 6 would presumably spread over a wider range of values of \( Q \). Not surprisingly, the predicted lifetimes for the observed decays depend sensitively on details of the barrier shape. Those shown in Fig. 8 correspond to the square well choice for \( W(y) \), and our standard set of parameters. In addition we show how the lifetimes vary when the barrier width is changed by amounts ranging from +4% to -3% (from left to right). As can be seen, the exact value of each decay lifetime depends quite strongly on the barrier width, as expected for a tunnelling process. But the number of slow decays is much more stable: four or five in most of the cases shown, and in several cases their lifetimes are quite compatible with the experimental points. In particular it is remarkable that a 2% increase in the standard barrier width produces a sequence (third line from left) in excellent agreement with experiment (disconnected points shifted to extreme left).

There is a clear distinction between the lifetime trends
of the thicker and thinner barrier widths. In the latter one sees very clearly the transition between escape from the \( n_x = 13 \) and the \( n_x = 14 \) levels at \( Q = 302 \). For the thicker barrier widths, escape is dominated by the \( n_x = 14 \) levels that become progressively more occupied above \( Q = 302 \).

Finally we show in Fig. 9 predictions for the fast decays: their number and location in a graph such as that of figure 1 depends very sensitively on the time \( t_0 \) beyond which the experiment measures lifetimes. That is, as the dot is isolated there must be a burst of very short lived escapes, but after some seconds one reaches the stage where separate events can be recorded and the lifetimes deduced. The two dashed horizontal lines in Fig. 9 correspond to values of \( \log_{10}(25 \text{ s}) \) and \( \log_{10}(35 \text{ s}) \). As can be seen, when we explore the same range of barrier widths as in Fig. 8, the number of fast decays above that \( t_0 \) varies from 3 to 4. The overall trend seems to be consistent with experiment, in particular if the value of \( t_0 \) is increased towards the more pessimistic estimate of 35 s.

IV. SUMMARY AND CONCLUSIONS

Electron escape from a strongly isolated dot with excess electrons has been studied in the framework of the self-consistent Poisson-Schrödinger and Poisson-Thomas-Fermi approximations. Based on these calculations a rectangular separable potential model has been devised which incorporates the main features of the self-consistent field. Rearrangement effects are taken into account by recalculating the confining potential \( U_s(x, y) \) after each electron escape.

The use of a separable potential introduces certain correlations in the energy spectrum of the single electron orbitals. A more realistic confining potential would have a more rounded shape, which would remove the separability, and modify those correlations. In the same vein, the tunnelling in our simplified model is 1D, whereas the actual process is 2D.

We find it quite remarkable that despite all these simplifications the predictions turn out to be so satisfactory. The model therefore may be reliable for extrapolating to longer times. For instance we find that the isolated dot would hold one excess electron for as long as 44 hours. On such a time scale, one could use well isolated dots containing a few long lived electrons, to study their entangled states. This would open an interesting new approach to the implementation of quantum computation in semiconductor devices.

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We summarize here the expressions relating the lifetimes to the probability of transmission across the barrier. We follow the standard treatment and definitions for alpha decay in nuclear physics, as can be found for example in ref. [4].

Our potential $V_s(x,y)$ is separable, and the electron can escape only across the barriers in the $x$-direction. Therefore, we have adapted the expressions derived in [4] to the 1D situation.

The lifetime $\tau = 1/\lambda$ is the inverse of the “decay constant”, defined as the number of “decays” per second per parent “dot”. For one dot the electron wavefunction is normalized to unity over the volume inside the barriers, and $\lambda$ for a given level is just the outgoing flux at large distance.

When the decay probability is small, one can treat the electron as confined in the dot. Classically, its trajectory will oscillate between the right, $x_r$, and left, $x_l$ turning points, with a period

$$P = 2 \int_{x_l}^{x_r} \frac{dx}{v(x)},$$

where $v(x)$ is the classical electron velocity at energy $E_x$:

$$v(x) = \sqrt{\frac{2}{m^*}(E_x - U(x))}.$$  \tag{A2}

The flux $\lambda$ is then given by the frequency of hits against the barriers, $2/P$, times the transmission probability $T$ across a barrier, and therefore:

$$\tau = \frac{1}{\lambda} = \frac{1}{T} \int_{x_l}^{x_r} \frac{dx}{v}.$$  \tag{A3}

This expression is very convenient because the transmission coefficient eq. 12 for our parametrized potential, $U(x)$, is known analytically [13]. For more general barrier profiles and the long lifetimes of interest, one can use the WKB approach and its corresponding connection formulæ across the barrier (see e.g. Appendix D of [4]):

$$T_{WKB} \approx e^{\frac{2\omega}{\hbar}}$$

$$\omega = \int_{x_l}^{x_r} \kappa dx = \int_{x_l}^{x_r} \sqrt{\frac{2m^*}{\hbar^2}(U(x) - E_x)} \, dx.$$  \tag{A4}

If the WKB wave function is used inside the well to determine the period $P$, the same decay half-life is obtained as in eq. A3 above.

Since the dot is located inside a crystal at temperature $T'$, via phonon coupling the electrons in the dot should also be at the same temperature. The level occupations $f(E)$ are determined by Fermi statistics:

$$f(E) = [1 + e^{\frac{E - E_F}{k_B T'}}]^{-1},$$  \tag{A5}

where these are now 2D energies. The Fermi level is obtained from

$$Q = \sum_{i=1, n_x, n_y} 2f(E_i),$$  \tag{A6}

where the 2 accounts for spin degeneracy. For the ensemble of electrons in the dot, the flux $\lambda$ will now be the sum of fluxes for each occupied single particle level, weighted by the level occupancy:

$$\lambda = \sum_{i=1, n_x, n_y} 2f(E_i) \lambda_i$$  \tag{A7}

and the corresponding half-life is still $\tau = 1/\lambda$. In particular this argument applies in the $T' = 0$ limit, as we implicitly assumed in Section II to explain the sequence of lifetimes.

### APPENDIX B: LIFETIME DEPENDENCE ON $Q$

For a level of given $n_x$, the lifetime depends on $Q$ because the barrier characteristics change, and so does the level energy, $E_{n_x}$. The latter varies mainly because $U_0$ depends on $Q$, and this affects the transmission probability $T$. To good approximation

$$\frac{dE_{n_x}}{dQ} \approx \frac{dU_0}{dQ}.$$  \tag{B1}

Neglecting the dependence of the level lifetime on the period $P$, we can write

$$d \ln \tau \approx -d \ln T \frac{dE}{dQ}.$$  \tag{B2}

Taking the transmission probability $T$ from the WKB expression leads to

$$\frac{d\ln T}{dQ} = 2 \frac{d}{dQ} \int_{\text{barrier}} \sqrt{\frac{2m^*}{\hbar^2}(U(x) - E)} \, dx$$

$$= \int_{\text{barrier}} \sqrt{\frac{2m^*}{\hbar^2}(U(x) - E)} \left( \frac{dU(x)}{dQ} - \frac{dE}{dQ} \right) dx.$$  \tag{B3}

Noting $B1$, the second contribution to the integral depends linearly on the placement of the potential floor. However, the variation of the barrier shape $(dU(x)/dQ)$ cannot be neglected. Indeed, eq. 8 gives approximately

$$\frac{dU(x)}{dQ} = \frac{dU_0}{dQ} \left( \frac{1}{3} + 2 \left[ \frac{1}{3} e^{-\frac{2}{3}} \cosh^2 \left( \frac{3}{2} \frac{E - E_F}{\mu} \right) \right] \right)$$

$$\text{when } x < x_b,$$

$$\frac{dU(x)}{dQ} = \frac{dU_0}{dQ} \left( -\frac{1}{3} + 2 \left[ \frac{1}{3} e^{\frac{2}{3}} \cosh^2 \left( -\frac{3}{2} \frac{E - E_F}{\mu} \right) \right] \right)$$

$$\text{when } x > x_b.$$  \tag{B4}
Using B4, the contribution from $dU/dQ$ to the integral of eq. B3 is obtained with an accuracy better than 2%.

For the standard choice of parameters, and $Q$ in the range 300 to 310 the computed values of $d \log_{10} \tau /dQ$ turn out to be $\approx -0.14$ for the levels of interest. In Fig. 10 we plot the evolution of the level lifetimes with $Q$, compared to the expression ($Q_0 = 303$)

$$\log_{10} \tau_{n_x}(Q) = \log_{10} \tau_{n_x}(Q_0) - 0.14(Q - Q_0) \quad (B5)$$

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2. G. Gamow, Z. Phys. 51 (1928) 204.