Numerical testing of the Fowler–Nordheim equation for the electronic field emission from a flat metal and proposition for an improved equation

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The author uses a transfer-matrix technique to simulate field electronic emission from a flat metal. He compares, in particular, the results provided by this numerical scheme with those predicted by the standard Fowler–Nordheim equation. He considers for this study electric fields between 1 and 10 V/nm as well as work functions between 1.5 and 5 eV. The results demonstrate that the Fowler–Nordheim theory and the transfer-matrix calculations are globally in good agreement. With the Fermi energy of 10 eV considered in this work, the results provided by the standard Fowler–Nordheim equation are, however, systematically larger than the quantum-mechanical result, especially for low values of the work function and for high electric fields. This is essentially due to the fact the standard Fowler–Nordheim theory relies on the simple Jeffreys–Wentzel–Kramers–Brillouin approximation for evaluating the electronic transmission through the surface barrier of the emitter. A correction factor is thus established that enables the temperature-dependent version of the standard Fowler–Nordheim equation to match the exact quantum-mechanical result. © 2010 American Vacuum Society. [DOI: 10.1116/1.3455495]

I. INTRODUCTION

Field electron emission is a process by which electrons are emitted from a material because of the application of external fields. The process by which this emission occurs, in the cold-emission regime in which the thermal excitation of electrons to energies that are above the apex of the surface barrier can be neglected, turns out to be the quantum-mechanical tunneling of electrons through the surface barrier of the material. The work function of the material essentially determines the energies at which this tunneling takes place, while the role of the external field is essentially to reduce both the height and the width of the surface barrier. Reducing the work function or increasing the external field thus lead to an increased probability of tunneling and therefore to an increased emission of current. 1

The first successful model for the electronic emission achieved from a flat metal through a triangular barrier was proposed by Fowler and Nordheim in 1928. 2 This work was subsequently extended in order to also account for the image interaction between the emitted electrons and the metal. 3–5 This extension lead to the “standard Fowler–Nordheim (FN) equation” J = ar²²F² exp[−bυφ²/F] for the current density J achieved from a flat metal that is subjected to an external field F. In this expression, a = 1.541 434 × 10⁻⁶ A eV⁻² and b = 6.830 890 eV⁻²/² V nm⁻¹, where υ and t are the tabulated functions that account for the image interaction and φ is the work function of the emitter. 7,8 To account for a nonzero temperature T of the emitter, J must be multiplied by (πkₜT/d)²/sin(πkₜT/d), where d = hₑF/(2√2mₑφ) is Boltzmann’s constant, e is the elementary positive charge, m is the mass of the electron, and $h = h/2\pi$ with h the constant of Planck. 3,4 The resulting expression, $J_{\text{FN}} = [(πkₜT/d)²/sin(πkₜT/d)] × a t²F² exp[−bυφ²/F]$, is also referred to as the “Murphy–Good (MG) expression.”

The derivation of the standard Fowler–Nordheim equation relies on the use of the simple Jeffreys–Wentzel–Kramers–Brillouin (JWKB) approximation for evaluating the electronic transmission through the surface barrier of the emitter. As argued by Forbes, 13 this approximation does not provide the exact result for this transmission so that the standard Fowler–Nordheim equation should actually be completed by a prefactor in order to account for the discrepancy between the transmission provided by the simple JWKB approximation and a more exact quantum-mechanical approach. This conclusion was confirmed by numerical simulations in which we compared the transmission obtained using either the simple JWKB approximation or a transfer-matrix (TM) technique for the case of one-dimensional barriers that are relevant to field-emission problems. 14

Since the standard FN theory is so widely used by the field-emission community, it is important to deal with any issue that may question its validity. It is the objective of this work to confront the results provided by the FN theory with those provided by a more exact quantum-mechanical approach. In Sec. II, we present the TM methodology that is used for these numerical simulations. 16,17 In Sec. III, we then compare the results provided by the TM methodology and the FN theory. The results show that the results provided by these two approaches are globally in good agreement. For the Fermi energy of 10 eV considered in this work, the results provided by the standard FN equation are, however, systematically smaller than the exact solution by a factor of the order of 1.1–2.1 for the conditions considered. These
results hence confirm Forbes’s opinion that a prefactor should be included in the standard Fowler–Nordheim equation in order to account for these deviations. A correction factor is hence established that enables the Murphy–Good expression to match the exact quantum-mechanical result. A polynomial adjustment of this correction factor is finally provided.

II. METHODOLOGY

For the quantum-mechanical simulation of field emission, we consider a model that consists of three regions: (i) region I (z ≤ 0), which stands for the metal that provides the electrons, (ii) region II (0 ≤ z ≤ D), which describes the surface barrier through which the electrons have to tunnel in order to be emitted, and (iii) region III (z ≥ D), which represents the region in which the electrons are transmitted. This is the general framework used in our previous work and the theory presented in this section is actually valid for a three-dimensional barrier. For the applications considered in this article, we will, however, consider a flat surface so that the potential energy only changes along z. We assume that a potential difference V of 30 V is established across region II. The length D of region II is related to the field strength F considered by the formula D = V/F. We then define \( V_i = eFz - \phi - E_F \) as the potential energy in region I, \( V(z) = eFz - \phi \) as the potential energy in region II, and \( V_\text{III} = 0 \) as the potential energy in region III. In these expressions, \( e \) refers to the absolute value of the electronic charge, \( \phi \) is the work function of the metal, \( E_F \) is the Fermi energy, and \( \epsilon_0 \) is the electric constant. The potential energy \( V(z) \) is prevented from going to \(-\infty\) as \( z \to 0 \) by using the reference energy \( V_i \) in region I as cutting value.3,7,14,18 For typical metals, we have \( E_F = 10 \) eV and \( \phi = 4.5 \) eV.

We work in cylindrical coordinates and assume that the electrons of this model are confined in a cylinder with radius \( R \). The boundary states in regions I and III are then given by19

\[
\Psi_{m,j}^{\text{III}}(r,t) = \frac{RJ_m(k_{m,j} r) \exp(\text{i} m \phi)}{\sqrt{2 \int_0^R dp [J_m(k_{m,j} p)]^2}} \times e^{\pm \sqrt{2m/h^2}(E - V_\text{III})} \frac{k_{m,j}^2}{2} e^{-\text{i} E t / h},
\]

(1)

where \( E \) refers to the total energy and the \( \pm \) sign refers to the propagation direction relative to the z-axis. The \( J_m \) refer to the Bessel functions and the \( k_{m,j} \) refer to the radial component of the wave vector. The \( k_{m,j} \) are defined by either \( J_m(k_{m,j} R) = 0 \) or \( J''_m(k_{m,j} R) = 0 \), where \( J''_m \) refers to the derivative of the Bessel functions. For \( R \) sufficiently large, it does not matter which of these two conditions is actually considered. We get, however, a faster convergence as \( R \to \infty \) by taking the average of the results obtained using the two conditions separately. For the simulations presented in this work, we used a value of \( R = 14 \) nm. The \( m \) values were allowed to go as high as necessary (typically up to \( m = 100–160 \) depending on the field and work function considered).

One can then use Schrödinger’s equation to propagate these boundary states across region II. This leads to a set of scattering solutions of the form

\[
\Psi_{m,j}^{\text{III}}(r,t) = \Psi_{m,j}^{\text{III},+} + \sum_{m',j'} S_{m',j',m,j}^{+} \Psi_{m',j'}^{\text{III},+}
\]

which correspond to single incident states \( \Psi_{m,j}^{\text{III},+} \) in region I. The coefficients \( S_{m',j',m,j}^{+} \) correspond to the amplitudes of, respectively, the transmitted and reflected states. The way these solutions are established for a one-dimensional barrier is presented in Ref. 14. Technical details for the consideration of three-dimensional aspects and for the control of numerical instabilities are given in Refs. 20–23.

The current density provided by the metal in region I is then obtained by integrating the contributions of every scattering solution. Referring to previous work for technical details,19,20 this current density \( J \) is finally given by

\[
J = \frac{1}{\pi R^2} \frac{2e}{h} \int_0^\infty \sum_{m,j} f(E) \left| S_{m,j}^{\text{III},+} \right|^2 dE,
\]

(3)

where \( f(E) = 1/\{1 + \exp[(E - \mu)/k_BT]\} \) is the Fermi factor of the metal in region I, \( \mu = \mu_1 + E_F \) is the chemical potential, \( k_B \) is the constant of Boltzmann, and \( T = 300 \) K is the room temperature assumed in this work. \( v_{1,m,j} \) and \( v_{\text{III},m,j} \) refer finally to the group velocities in respectively Region I and III. The summations in Eq. (3) must only include propagative states.

III. APPLICATION

We consider first that the Fermi energy \( E_F \) is 10 eV and that the work function \( \phi \) is 4.5 eV. These are indeed the typical values for a metal. We also assume a room temperature \( T = 300 \) K. One can then compute the current density one would obtain from a perfect metal characterized by these parameters when the field strength \( F \) takes values between 1 and 10 V/nm. For this first situation, we actually consider that the potential energy in the intermediate region II of our model is either (i) a triangular (T) barrier \( V(z) = eFz - \phi - E_F \) or (ii) a Schottky–Nordheim (SN) barrier \( V(z) = eFz - \phi - (1/16 \epsilon_0 E_F)(e^2/z) \). The current density obtained in the first case is given by the original model of Fowler–Nordheim,2 while the current density obtained in the second case corresponds to the standard Fowler–Nordheim equation in which the image interaction is taken into account.4,6 When using either the elementary or the standard Fowler–Nordheim equation, we actually include the factor \((\pi k_B T/d) \sin(\pi k_B T/d) \) that accounts for the temperature \( T = 300 \) K of the emitter (this temperature-dependent version of the standard Fowler–Nordheim equation is also referred to as the “Murphy–Good expression”). We compare in both cases these analytical solutions with our transfer-matrix cal-

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These results are presented in Fig. 1. The transfer-matrix results turn out to be in excellent agreement with the analytical solutions provided by the FN theory (for both the triangular and the Schottky–Nordheim barrier). At this point, this essentially validates the results provided by the transfer-matrix technique.

The materials considered for current applications are characterized by work functions that can be smaller than the value of 4.5 eV considered so far. We thus represented in Fig. 2 the current density obtained from a flat metal when the work functions $\phi$ is 1.5, 2.5, 3.5, and 4.5 eV. We compare the results provided by the Murphy–Good expression and the transfer-matrix technique. The representation is restricted to fields $F$ that keep below the critical value $F_{\text{crit}} = (4\pi\epsilon_0\phi^2/e^2)$ for which the apex of the barrier corresponds to the Fermi level of the metal. This corresponds indeed to the limit of validity of the standard Fowler–Nordheim theory. Realistic devices would anyway not stand the currents that correspond to fields higher than that critical value. The MG and TM results are globally in good agreement, but a closer examination reveals that the Murphy–Good expression actually overestimates the quantum-mechanical result by a factor of the order of 1.1–2.1 for the conditions considered (this will be better illustrated in Fig. 4). This discrepancy is more significant for lower values of the work function and for higher values of the field strength $F$. These conditions are actually those that reduce both the height and the width of the surface barrier and therefore lead to higher emissions of current.

In Fig. 3, we provide the Fowler–Nordheim representation of the $J$–$F$ data presented in Fig. 2 [this representation consists in representing the $\log(J/F^2)$ data as a function of $1/F$]. This representation reveals as previously that the standard Fowler–Nordheim theory overestimates the quantum-mechanical result. It also shows that the Fowler–Nordheim representation of the $J$–$F$ data that corresponds to a flat metal is not a straight line as often expected. The deviations from a mere linear dependence of $\log(J/F^2)$ on $1/F$ are actually more important on the left side of this representation (strong fields $F$). They are also more significant for smaller values of the work function $\phi$. Deviations of this type are actually inherent to the standard Fowler–Nordheim theory because the functions $t$ and $v$ depend on the field strength $F$.

This was made explicitly clear in 1953, when Burgess et al. showed how values of the slope correction factor $s$ varied with field and work function. Space-charge effects are in-
voked to justify deviations that are higher than that expected from the standard FN theory. Our simulations reveal, however, that deviations from a straight line are actually inherent to the basic phenomenon (to an extend that exceeds that expected from the standard FN theory).

The deviations between the results provided by the standard Fowler–Nordheim theory and those provided by a more exact quantum-mechanical approach are essentially due to the fact the standard FN theory relies on the simple JWKB\(^9–12\) approximation for evaluating the electronic transmission through the surface barrier of the emitter. As shown in previous work,\(^14\) the simple JWKB approximation overestimates this transmission by a typical factor between 1.0 and 3.5 for the conditions considered in this work (the highest deviations correspond to small work functions \(\phi\) and to high electric fields \(F\)). One can account for these deviations by including an effective correction factor \(P_{\text{eff}}\) in the electronic transmission, thus yielding the Landau and Lifschitz\(^{25}\) expression \(T = P_{\text{eff}} \exp[-G]\) for this transmission \([G = (2\sqrt{2m\hbar})/z_w^2 |V(z) - E|^{1/2}dz\] with \(z_1\) and \(z_2\) the classical turning points of the potential barrier \(V(z)\) at the energy \(E\).\(^{13,14}\) The current densities \(J\) depend, however, on a whole range of energies (typically energies that are close to the Fermi level of the metal) and an averaging of these deviations actually occurs. The current densities provided by the standard Fowler–Nordheim theory are therefore overestimated by a factor that takes values between 1.1 and 2.1 for the conditions considered in this article (instead of a factor between 1.0 and 3.5 for the transmission coefficient).

It is actually remarkable that the standard Fowler–Nordheim theory provides results that keep so close to the exact solution despite the fact \(J\) ranges over orders of magnitude when \(\phi\) and \(F\) are changed. For the interpretation of \(J–F\) data, it is, however, essential to rely on an equation that is as exact as possible. It is therefore useful to provide the correction factor \(\lambda_{\text{MG}}\) to use with the Murphy–Good expression

\[
J_{\text{MG}} = \left[ (\pi k_b T/d) \sin(\pi k_b T/d) \right] \times \alpha^{-2} \phi^{-1} F^2 \exp \left[ -b \phi^{3/2} / F \right]
\]

in order to match the exact quantum-mechanical result (the adapted equation is therefore \(J = \lambda_{\text{MG}} \times (\pi k_b T/d) \times \alpha^{-2} \phi^{-1} F^2 \exp \left[ -b \phi^{3/2} / F \right]\)). This is done in Fig. 4, where we represented the ratio \(\lambda_{\text{MG}} = J_{\text{TM}}/J_{\text{MG}}\) between the current densities \(J_{\text{TM}}\) obtained using the TM technique and the values \(J_{\text{MG}}\) provided by the Murphy–Good expression. The representation corresponds to work functions \(\phi\) between 1.5 and 5 eV and to fields \(F\) between 1 and 10 V/nm (with, however, \(F < (4\pi e_0 \phi^2 / e^2)\) to keep in the range of parameters for which the standard FN theory holds). The data represented in Fig. 4 can actually be represented by the polynomial adjustment \(\lambda_{\text{MG}} = J_{\text{TM}}/J_{\text{MG}} = \sum_{i=0}^{6} \sum_{j=0}^{6} a_{ij} X^i Y^j\), where \(X = F - 5\) with \(F\) the field strength in V/nm and \(Y = \phi - 3.5\) with \(\phi\) the work function in eV. The coefficients \(a_{ij}\) are provided in Table I. This adjustment provides a mean absolute error of \(2.2 \times 10^{-4}\) on the data represented in Fig. 4, with a maximal absolute error of \(5.9 \times 10^{-3}\). This adjustment is represented with the original data in Fig. 4.

IV. CONCLUSION

A transfer-matrix technique was used for the quantum-mechanical simulation of electronic field emission from a flat metal. This study essentially aimed at testing the accuracy of the standard Fowler–Nordheim theory and at determining the correction factor to include in the Murphy–Good expression in order to match the exact quantum-mechanical result. The results show that, for typical values of the field strength \(F\) and of the work function \(\phi\), and for an assumed Fermi energy of 10 eV, the standard Fowler–Nordheim theory actually

Table I. Coefficients \(a_{ij}\) of the polynomial adjustment \(\lambda_{\text{MG}} = J_{\text{TM}}/J_{\text{MG}} = \sum_{i=0}^{6} \sum_{j=0}^{6} a_{ij} X^i Y^j\) for the prefactor \(\lambda_{\text{MG}}\) to use with the MG expression in order to match the current densities \(J_{\text{TM}}\) provided by the transfer-matrix technique. In this expression, \(X = F - 5\) with \(F\) the field strength in V/nm and \(Y = \phi - 3.5\) with \(\phi\) the work function in eV. This expression is restricted to 1 V/nm \(\leq F \leq 10\) V/nm, 1.5 eV \(\leq \phi \leq 5\) eV, and \(F < (4\pi e_0 \phi^2 / e^2)\).

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overestimates the current densities by a factor between 1.1 and 2.1. These deviations appear essentially for low work functions $\phi$ and for high electric fields $F$. They are responsible for the Fowler–Nordheim representation of the $J$–$F$ data achieved from a flat metal to deviate from an expected straight line (to an extent that exceeds that expected from the standard FN theory and without the implication of space charge effects). These deviations between the results provided by the standard Fowler–Nordheim theory and the quantum-mechanical result are essentially due to the fact that the standard FN theory relies on the simple JWKB approximation for evaluating the electronic transmission through the surface barrier of the metal. This approximation typically overestimates this transmission by a factor between 1.0 and 3.5 for the conditions considered in this work. One can, however, account for these deviations by including an effective correction factor $P_{\text{eff}}$ in the electronic transmission as well as a corresponding correction factor $\lambda^{\text{MG}}$ in the Murphy–Good expression. This work thus confirms the conclusions formulated by Forbes on the need to account for these prefactors in theories of field emission. We provided, in particular, an accurate determination of the correction factor $\lambda^{\text{MG}}$ to include in the Murphy–Good expression. A polynomial adjustment that enables $\lambda^{\text{MG}}$ to be calculated for practical but otherwise arbitrary values of $F$ and $\phi$ was given. This work hence clarifies the issue raised by the limits of the simple JWKB approximation. The results of this article should improve the accuracy of $J$–$F$ data analysis that is based on the standard FN theory.

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