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TUNNELLING FROM A MANY PARTICLE POINT OF VIEW

by

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December, 1962

This research was supported by the United States Air Force through the Air Force Office of Scientific Research of the Air Research and Development Command under Contract No. AF-AFOSR 62-46.
ABSTRACT

The Hamiltonian for a system of interacting electrons in the presence of a barrier is transformed into one which can be separated into three parts. Two of the parts describe the electrons on the right and left sides of the barrier, while the third is a transition term which allows tunnelling through the barrier. It is shown that it is not reasonable to make the right and left hand Hamiltonians commute, but the effect of this failure is easily accounted for in perturbation theory. In addition, corrections to the transition operator arising from inter-electronic interaction are written down and discussed briefly. They may be both quantitatively and qualitatively important.
Experiments in which electrons are observed to tunnel through insulating layers have played an increasingly important role in recent times. The tunnelling process has, of course, been well understood since the discovery of quantum mechanics. Bardeen has very recently pointed out, however, that in the case of a many-electron system, the tunnelling process is very conveniently regarded as in time-dependent perturbation theory. In other words, it is seen as a transition process from one set of nearly stationary states to another.

Let us keep in mind, to be definite, a system of interacting electrons enclosed in cubical boxes separated by a high, narrow barrier. It will be a convenient simplification in the later development if we suppose that the boxes and the barrier are symmetrically disposed, as in Fig. 1.

![Figure 1](image_url)

**Fig 1.**

Barrier Potential V(x) vs. x.
The most convenient formulation of the problem will involve showing that the complete Hamiltonian of the many electron system, including the insulating barrier, can be split into the form,

\[ H = H_R + H_L + T \]  \hspace{1cm} (1)

Here \( H_R \) is the Hamiltonian of the electrons to the "right" of the barrier, \( H_L \) the Hamiltonian for the left-hand electrons, and \( T \) is the operator which makes possible the transition of electrons from left to right. It is perhaps clear intuitively from the work of Bardeen that such a formulation of the problem exists, and indeed, several papers have appeared in which the Hamiltonian (1) was taken for granted. (3)

It is the purpose of this note to show in what sense (1) is actually correct. Perhaps this is not an entirely empty exercise since an instructive difficulty has to be overcome. Furthermore, it is possible to discuss in detail the effect of interactions on the above formulations, which is always difficult to do in an intuitive fashion.
II.

The dilemma which arises is as follows: In defining the "right-hand" and "left-hand" Hamiltonians, one naturally needs to define right and left-hand states. Now, first of all, these wavefunctions have to be in the domain of the original Hamiltonian operator, $H$. In the concrete model we have adopted this simply means that they must vanish at the sides of the boxes $|x| = a$, as well as being twice differentiable. Secondly, we would like the left-hand states to be complete on the left-hand side. This requirement insists that should an electron flow from a wire into the box to the left of $-b$, its wavefunction could be expressed in terms of left-hand states only. Thirdly, one should like, if possible, that all left-hand states be orthogonal to all right-hand states, in order that $H_R$ should commute with $H_L$. Finally, the states have to form a complete set when taken all together.

If one could produce such a set of (single-electron) wave functions $\Phi_k^r(x)$ and $\Phi_k^l(x)$, it would be a simple matter to expand the annihilation operator as

$$\psi(x) = \sum a_k \Phi_k^r(x) + b_k \Phi_k^l(x)$$

(2)

One would then replace the creation and annihilation operators in the second-quantized form of $H$,

$$H = \int \psi^\dagger(x) \left[ p^2/2m + V(x) \right] \psi(x) + \frac{i}{2} \int \psi^\dagger \psi^\dagger \psi \psi$$

(3)

by the expression (2). $H_R$ could be identified as that part of $H$ containing only "a" operators, $H_L$ would contain only "b" operators, and $T$ would be the remainder.

Unfortunately, no such states appear to exist. For, should all of the $\Phi_k^r(x)$ vanish identically for positive values of the coordinate, then either they do not form a complete set for negative $x$, or they are too singular at $x = 0$. And if they have support in the right-hand region, they cannot be orthogonal to the $\Phi_k^r(x)$ should these functions be complete.
We are in consequence forced to compromise the requirements on the wave functions. There are two approaches which can be taken. The first possibility is to maintain the requirement of mutual orthogonality, but to abandon the requirement of separate "right" and "left" completeness. Instead, the right and left states will be made "as complete as possible" in their respective regions.

If this is done, a mathematically convenient formalism results, since the operator $T$ can be treated systematically by ordinary perturbation theory. We shall show below a simple method for achieving this result. One must pay a price, however. In practice, the model which we are studying is not really of interest by itself. We also need a method of hooking up wires and batteries connecting the right-hand electrons with those on the left. Now wires and batteries are cumbersome to treat quantum mechanically, to say the least. It is essential to have a simple and convincing method of mathematically treating the classical part of the apparatus. These methods usually involve introducing electrons into the right-hand states and taking them out of the left-hand ones. However, the wires introduce electrons into the right-hand side. If the right-hand states are not complete, we have to take into account the possibility that electrons flowing into the right side partially go into the left-hand states. Methods may be advanced to get around this difficulty, but they seem rather contrived. The approach does contain the possibility of conveniently studying higher order effects, which is becoming fashionable.

Another approach is to abandon the orthogonality condition, requiring instead the states to be "as orthogonal as possible", and to keep the separate completeness of the right and left states. In that case, the commutator $[H_R, H_L]$ is an operator of the order of magnitude of $T$. 
III.

In the symmetrical model which we are considering, a particularly simple set of states springs to mind. The exact eigenstates of the single-particle Hamiltonian may be called \( \psi_k(x) \), \( \Omega_k(x) \). They have even and odd parities under the reflection of the \( x \)-coordinate, respectively. These states occur in pairs, the energy of the even (odd) state being \( E_k(\mp)T_{kk} \), to an excellent approximation, where \( T_{kk} \) happens to be the diagonal element of Bardeen's transition operator,

\[
T_{kk} = \left( a k \varepsilon_k / m V_0 \alpha \right) e^{\alpha \rho} (-2 b V_{2m}(V_0 - \varepsilon_k))
\]

and \( \varepsilon_k \) is the energy level calculated as if the barrier were very wide. With appropriate choices of phase for the \( \psi_k(x) \), \( \Omega_k(x) \), one naturally defines

\[
\sqrt{2} \phi_k^r(x) = \psi_k(x) \pm \Omega_k(x)
\]

The \( \phi_k^r(x) \) are large only in the right-hand region, but there is a long tail (proportional to \( T_{kk} \)) in the "wrong" region. Defining the \( a \) and \( b \) operators according to (2), the Hamiltonian becomes

\[
H = \sum_k \varepsilon_k a_k^+ a_k + W_R + \sum_k \varepsilon_k b_k^+ b_k + W_L
- \sum_k T_{kk} (a_k^+ b_k + b_k^+ a_k) + W_T
\]

We have not bothered to write out the contributions from the interaction term. This will be taken up below.

Neglecting the interaction terms then we can calculate the time derivative of the number of particles in right-hand states. We find

\[
\frac{d}{dt} N_R = (N_L^0 - N_R^0) T \sin 2\pi t
\]
We have assumed that \( T_{kk} \) is slowly varying with \( k \), and has average value \( T \). We have also supposed that \( N_R + N_L \) is constant, and that at \( t = 0 \), the numbers of particles present in the right (left) states is \( N_{o R(L)} \).

Formula (6) exhibits the oscillatory behavior well-known from elementary quantum mechanics. Such behavior is not to be expected in the context of real metals, where one would anticipate finding a constant transition rate for not-too-long times. If (6) does not show a constant transition rate at any time, it can only be a result of the two clever definition of the states \( \phi_{k}^{r(l)} \). We have put electrons into the long tail of the \( \phi_{k}^{r(l)} \) on the "wrong" side of the barrier, already accomplishing the greater part of that tunnelling which usually proceeds at a constant rate. If we look at the number of electrons in right-hand states we will not obtain a true picture of the tunnelling, at least not without further arguments. By the same token, one must handle gingerly any states which are not confined rather strictly to one side or the other.
The other alternative is to find a separation of the Hamiltonian, as in Eq. (1), based on states which are really confined to one side or the other. A natural set of states is given by the eigenstates of $H_1$, where

$$H_1 = p^2 + V(x)$$

and

$$V_1(x) = V(x), \quad x > -b$$

$$= V_0, \quad x < -b$$

This particular choice is convenient, but not mandatory. We call the eigenstates $\phi_k(x)$ and also define

$$\chi_k(x) = \phi_k(-x)$$

Each of the sets $\phi_k, \chi_k$ is complete in the entire space ($a > |x|$). However, to represent a function which has large amplitude on the left by $\phi_k$, it will be necessary to employ large $k$ values. Let us introduce $c_k$ and $d_k$ operators according to

$$\psi(x) = \sum_k c_k \phi_k(x) = \sum_k d_k \chi_k(x)$$

We next employ the representation

$$\psi(x) = \sum_m (\phi_m(x) + \chi_m(x)) \lambda_m \lambda_m$$

Then

$$\lambda_{km} = (\phi_k, \phi_m) - \sum_n (\phi_k \chi_n) \lambda_{kn} \lambda_{nm}$$

$$\lambda_{km} = (\phi_k, \chi_m) + \sum_n (\phi_k \chi_n) \mu_{kn} \mu_{nm}$$
We have next to compute \( (\phi_k, e_m) \). This matrix element is diagonal in all quantum numbers except the momentum component conjugate to \( x \). We shall suppress these other quantum numbers, as we have up to now.

The largest matrix elements between states of practically the same energy are found to be, after a tedious calculation,

\[
(\phi_k, e_m) = 2^{-\frac{1}{2}} \delta_{k_m} + (\phi_k, e_m - 2^{-\frac{1}{2}} \phi_m) = 2^{-\frac{1}{2}} \left[ \delta_{k_m} - \frac{T_{kk}}{(E_k - E_m)} + \frac{i}{2} \tilde{S}_{km} + \ldots \right]
\]

\[(\phi_k, e_m) = 2^{-\frac{1}{2}} \left[ \delta_{k_m} + \frac{T_{kk}}{(E_k - E_m)} - \frac{i}{2} \tilde{S}_{km} + \ldots \right]
\]

(15)

(16)

where we have introduced

\[
\tilde{S}_{km} = (\phi_k, x_m)
\]

The dots represent terms of higher order in \( T_{kk} \).

In addition, there are large matrix elements \( (\phi_k, e_m) \) when \( k \) is so large that the wave-length on the left is practically given by \( m \). In that region, however, we find \( (\phi_k, e_m) = 2^{-\frac{1}{2}} (\phi_k, x_m) \).

Thus we find to sufficient approximation, that

\[
\lambda_{kl} = 2^{-\frac{1}{2}} \left[ \delta_{kl} - \frac{T_{kl}}{(E_k - E_l)} - \frac{i}{2} \tilde{S}_{kl} \right]
\]

(18a)

\[
\mu_{kl} = 2^{-\frac{1}{2}} \left[ \delta_{kl} + \frac{T_{kl}}{(E_k - E_l)} + \frac{i}{2} \tilde{S}_{kl} \right]
\]

(18b)

and there are no large matrix elements to high energy states.
In the remainder of this section, we shall neglect all inter-electron interaction. Substitution of expressions (10) - (12) and (18) into the Hamiltonian yields at once

\[
H = \sum_{k,l,n} \left( \epsilon_k - T_{kk} \right) \lambda_{km} \left( c_m^+ d_m^+ \right) \lambda_{kn} \left( c_n^+ d_n^+ \right) \\
- \left( \epsilon_k + T_{kk} \right) \mu_{km} \left( c_m^+ d_m^+ \right) \mu_{kn} \left( c_n^+ d_n^+ \right)
\]

\[
= \sum_k \epsilon_k \left( c_k^+ c_k + d_k^+ d_k \right) \\
- \sum \frac{\tilde{T}_{\alpha \beta}}{\tilde{T}_{\alpha \beta}} \left( c_\alpha^+ d_\alpha + d_\alpha^+ c_\alpha \right) + \frac{\epsilon_k + \epsilon_\alpha}{2} \sum_{k,\alpha} \left[ c_k^+ d_\alpha + d_k^+ c_\alpha \right] \\
+ \sum_{k,m,n} \frac{\tilde{T}_{km} \tilde{T}_{mk}}{(\epsilon_k - \epsilon_m)(\epsilon_k - \epsilon_n)} \left( c_m^+ c_n + d_m^+ d_n \right)
\]

(19)

The last term, as well as others indicated by dots, may be dropped if we wish to keep only the terms of lowest order in the tunnelling exponential. The desired separation of the single-particle Hamiltonian has thus been achieved with

\[
H_R = \sum \epsilon_k c_k^+ c_k \quad \text{(20a)}
\]

\[
H_L = \sum \epsilon_k d_k^+ d_k \quad \text{(20b)}
\]
However,

\[
\left[ H_{R}, H_{L} \right] = \sum_{r,s} \varepsilon_r \varepsilon_s \xi_{rs} \left( c_r^+ d_s - d_r^+ c_s \right)
\]

which is of the order of \( T_{rs} \). This means that the eigenstates of \( H_{R} + H_{L} \) are not just the products of eigenstates of \( H_{R}, H_{L} \) separately.

Let us, for example, calculate the current through the barrier, \( J \), where

\[
J_{op} = \frac{e}{2mi} \int dy \, dz \left[ \psi_r^+(x,y) \frac{2}{i} \left( \psi_r(x,y) - \left( \frac{1}{2i} \frac{\partial}{\partial x} \psi_l(x,y) \right) \psi_l(x,y) \right]_{x=0}
\]

It may readily be shown that

\[
J_{op} = -e \sum_{c, k} \left( c_{k}^+ d_{k} - d_{k}^+ c_{k} \right) T_{kk}
\]

by the methods of Bardeen. We propose to calculate

\[
J = \langle \Phi(t), J_{op}, \Phi(t) \rangle
\]

where \( \Phi(t) \) is the state reducing at \( t = 0 \) to the product state

\[
\Phi(0) = \Phi_R \Phi_L |0\rangle
\]

\( \Phi_R |0\rangle \) being an eigenstate of \( H_{R} \).

It is easy to verify that the failure of \( H \) to commute with \( H_{L} \) is just compensated by the term containing \( f_{KL} \) in (19). Indeed

\[
(H_{R} + H_{L}) \Phi_R \Phi_L |0\rangle = \Phi_L H_{R} \Phi_R |0\rangle + \Phi_R H_{L} \Phi_L |0\rangle
\]

\[
+ \sum_{k} \varepsilon_{k} \xi_{rk} \left( c_{k}^+ d_{k} + d_{k}^+ c_{k} \right) \Phi_R \Phi_L |0\rangle
\]

to the first order tunnelling. In consequence, we may drop that term, at the same time assuming \( \{ c_{r}^+, d_{r} \} = 0 \). This leads to the usual formula for the current proportional to \( |T_{KL}|^2 \).
V.

We turn next to the problem of interactions. We shall treat here for simplicity only short range interactions of simple type. (The range should be small even in comparison with the widths of the barrier.)

Perhaps the easiest way to write down the terms of the interaction Hamiltonian is to employ the representation

$$\psi(x) = \sum_n \tilde{\phi}_n c_n + \sum_n \tilde{\chi}_n d_n.$$  \hfill (27)

We have introduced the notation

$$\tilde{\phi}_n = \phi_n - \sum_{m} s_{mn} \chi_n.$$  \hfill (28)

Here $s_{mn}$ is given by (17) with the understanding that the matrix elements connecting states of widely different energies are to be dropped.

We may immediately write down the interaction terms

$$W = \left\{ \frac{1}{2} \sum W_{pqrs}^{(0)} c^+_p c^+_q c^+_r c^+_s + \frac{i}{2} \sum W_{pqrs}^{(0)} d^+_p d^+_q d^+_r d^+_s \right\}
+ \left\{ \frac{1}{2} \sum W_{pqrs}^{(1)} [c^+_p d^+_q d^+_r d^+_s + d^+_p c^+_q c^+_r c^+_s + \text{hermitian conj.}] \right\}
+ \left\{ \frac{1}{2} \sum W_{pqrs}^{(2)} c^+_p c^+_q d^+_r d^+_s + \text{hermitian conj.} \right\}
+ \sum W_{pqrs}^{(3)} c^+_p d^+_q d^+_r c^+_s$$  \hfill (29)
Here, we have written
\[ W^{(0)}_{pqr} = \langle x-y \rangle_\phi \phi_p(x) \phi_q(y) \phi_r(y) \phi_r(x) \] (30a)
\[ W^{(1)}_{pqr} = \langle x-y \rangle_\phi \phi_p(x) \phi_q(y) \phi_q(y) \phi_r(x) - (p \leftrightarrow q) \] (30b)
\[ W^{(2)}_{pqr} = \langle x-y \rangle_\phi \phi_p(x) \phi_q(y) \phi_r(y) \phi_r(x) \] (30c)
\[ W^{(3)}_{pqr} = \langle x-y \rangle_\phi \phi_p(x) \phi_r(y) \phi_q(y) - \phi_r(x) \phi_r(y) \phi_r(x) \] (30d)

Because of the distinction between \( \phi \) and \( \phi \), we see that the usual interaction terms, \( W^{(0)} \), are slightly modified. However, this is of no interest. On the other hand, \( W^{(1)} \) is a term leading to tunneling. It will be small of order \( T \) and could be quite important numerically for strongly interacting systems. The remaining terms are of second order in \( T \) and should, for the sake of consistency, be dropped. \( W^{(2)} \) is of some interest, however, in that it shows that a direct tunneling of two particles is possible, even apart from the higher order contributions of \( T \) and \( W^{(1)} \).

We still have to take into account the fact that the large interaction terms, \( W^{(0)} \), on the first line (28) do not commute with one another. It is not difficult to show that this can be accounted for, at least in first order, by a change in \( W^{(1)} \). The result is that if we replace the second line of (29) by
\[ \frac{1}{2} \sum_{pqrst} \left[ W^{(1)}_{pqrst} (c_p^+ d_q^+ c_r^c c_s^c) + d_p^+ c_q^+ + c_p^+ c_q^+ c_r^c c_s^c \right] \] + \[ W^{(1)}_{pqrst} (d_p^+ d_q^+ c_r^c c_s^c + c_p^c c_q^+ c_r^c c_s^c) \] (31)

then we may regard the \( c \)'s and \( d \)'s as commuting operators. Expression (31) must be thought of as operating to the right. In (31), \( W^{(1)}_{pqrst} \) is the same as expression (30b) with the tilde removed from the \( \tilde{\chi} \) functions. \( W^{(1)}_{pqrst} \) is the same as (30b) with both the \( \tilde{\chi} \phi \) functions replaced by \( \chi \phi \) functions respectively.
In summary, we have succeeded in the aim of splitting the Hamiltonian as in Eq. (1). The parts $H_R$ and $H_L$ do not quite commute, but we have shown, at least in first order, that this can be completely accounted for by a change in the transition operator $T$. The transition term is the same as written down by Bardeen for free particles. We have explicitly found the correction to this in the presence of interactions. This correction might well be quantitatively significant.

The author would like to thank Professors Falk, Ferrell, and Glick for stimulating conversations.
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