Transmission coefficient of interacting few-body system in one dimensional space

Peng Guo a,b, *, Vladimir Gasarian a

a Department of Physics, California State University, Bakersfield, CA 93311, USA
b Kavli Institute for Theoretical Physics, University of California, Santa Barbara, CA 93106, USA

1. Introduction

Few particles interaction in one dimensional space (1D) has attracted a lot of attentions in various fields of physics both experimentally [1–6] and theoretically [7–14]. The increased interest in 1D system is largely motivated by recent advance on experimental techniques. These new experimental techniques not only make studies of low-dimensional interacting particles experimentally possible, but also provide opportunities to challenge our understanding of few-body systems. For instance, a peculiar bound state of atom pairs in an optical lattice with repulsive interaction between atoms has been observed recently [1]. Similar optical lattice technique is also used to create ultracold strongly interacting atoms [6]. Single electron transistor and small quantum dots may be other exciting examples as experimental realizable few-body systems [4]. In such systems, the quantized energy levels and the Coulomb interaction are comparable, so that only a small number of electrons can be confined to the region with the size of orders of the Fermi wavelength. It is hence commonly assumed that the electrons interact with each other only when they are on the same quantum dot with contact interactions. Therefore, a set of nearly isolated small quantum dots may be considered as an ideal system for the studies of few-body system, such as transport properties [15].

Quantum transport effect may play the crucial role in nanoscale semiconductor devices. The transport properties of non-interacting electrons are usually described by Landauer-Büttiker formula [16, 17], which establishes the relations between the conductance $G$ of non-interacting electrons in a quasi-1D wire and transmission coefficient: $G = \frac{e^2}{h} \sum_{n} T_n$, where $T_n$ stand for the transmission probability coefficient for the incoming state in $n$-th channel. However, there is a firm belief that the behavior of electrons in 1D wire may be strongly affected by few-body interactions, and even a small perturbation can change the scattering pattern of particles significantly, see, e.g. Refs. [11–14]. For interacting systems, electron-electron interaction may be incorporated within Landauer-Büttiker formula by including self-energy of electrons in a perturbation theory approach [18,19]. This approach is more or less Hartree-Fock type approximation by assuming that electron-electron interactions are elastic on average, so that single electron maintains the same momentum and all the inelastic effects are included in self-energy of electron as virtual processes. Clearly, the approach may work well for many-body systems at larger scale, however, for few-body system, there are the cases that inelastic process may become essential [11–14]. In this letter, we aim to present a general formalism for describing transport properties of interacting few-body system. The formalism is based on the physical transition $S$-matrix and unitarity relation of few-body system, hence, it is suitable not only for weakly interacting few-body system, such as electron-electron coulomb interaction, but also for strongly coupled systems that cannot be easily handled by perturbation theory.
We also would like to point out that there have been other approaches which have been developed in the past to incorporating electron-electron, or electron-atom interactions in transport properties, such as, approach based on Master equations, [20,21].

2. S-matrix and wave packet prescription

In order to describe transport properties of a quantum conductor, one of the key elements is to introduce properly defined transmission or reflection coefficients to reflect the probabilities for electrons tunneling through potential barriers. Normally, for the single electron in 1D, the task may be easily accomplished by study the scattering solutions of Schrödinger equation. The asymptotical wave function of single electron in 1D has quite a simple form with linear superposition of two components: forward- and backward-going plane waves. Although, it has been a well-known fact that the plane wave is not normalizable and doesn’t represent a physically realizable state. In the case of 1D single electron scattering, it is still possible to interpret the coefficients of two plane waves as physical probability amplitudes. For example, assuming a particle incident from left that is described by a forward-going plane wave: \( e^{ipx} (p > 0) \), asymptotic wave function in forward and backward directions are given respectively by,

\[
\Psi(x, p) \xrightarrow{x \to -\infty} [1 + it(p, p)] e^{ipx},
\]

\[
\xrightarrow{x \to +\infty} e^{ipx} + it(-p)e^{-ipx},
\]

where \( t(k, p) \) represents the scattering amplitude of particle. The transmission and reflection probabilities are thus associated to the coefficients of transmitted wave in forward direction and reflected wave in backward direction: \( T = |1 + it(p, p)|^2 \) and \( R = |it(-p)|^2 \) respectively. Probability conservation yields the relation: \( T + R = 1 \). Unfortunately, for multiple particles, even in one spatial dimension, the physical interpretation of coefficients in front of asymptotic wave functions become problematic. The reason is that the momenta among particles after scattering may be redistributed, and the coefficients that describe the momenta distribution is not normalizable due to plane wave description of asymptotic states. Using two-electron scattering off atoms as an example, with an incident plane wave of two electrons: \( e^{ipx} \), where \( p = (p_1, p_2) \) and \( \mathbf{x} = (x_1, x_2) \) are 2D vectors that represent electron’s momenta and positions respectively, the asymptotic form of two electrons wave function in forward direction is

\[
\Psi(x, p) \xrightarrow{\theta_k \to \theta_p} [2\pi \delta(\theta_k - \theta_p) + 2it(p\hat{x}, p)] \frac{e^{i(px + \mathbf{x}\cdot\hat{p})}}{\sqrt{2\pi px}},
\]

where \( \theta_k = \tan^{-1} \frac{x_1}{\xi_1}, \theta_p = \tan^{-1} \frac{x_2}{\xi_2}, \mathbf{x} = \frac{\xi}{\xi}, \xi = \sqrt{x_1^2 + x_2^2}, \) and \( p = \sqrt{p_1^2 + p_2^2} \). \( t(p\hat{x}, p) \) again represents the scattering amplitude of two electrons. The expression in Eq. (2) resembles the one electron case in forward direction, however, the divergent \( \delta(\theta_k - \theta_p) \) term in coefficient of outgoing spherical wave prevents direct physical interpretation of coefficient as probability amplitude. The divergence of \( \delta(\theta_k - \theta_p) \) term is due to the fact that the angular component of plane wave which describes the momenta distribution among two electrons is not normalizable. Plane wave description violates Heisenberg uncertainty principle and is not suitable to be used to represent physical states whose positions and momenta are both well determined.

On the other hand, it has been known that in single electron scattering case given in Eq. (1), the coefficients of transmitted and reflected waves can be identified as reduced S-matrix elements after removing energy conservation \( \delta \)-function constraint,

\[
S(k, p) = \delta_{k, p} + it(k, p),
\]

where \( \hat{k} = \frac{k}{|k|}, \hat{p} = \frac{p}{|p|} \) and \( |k| = |p| \). So that \( T = |S(p, p)|^2 \) and \( R = |S(-p, p)|^2 \), and probability conservation relation, \( T + R = 1 \), is equivalent to unitarity relation of S-matrix:

\[
\sum_{k=\pm p} |S(k, p)|^2 = 1.
\]

This connection is due to the fact that the transport behaviors are determined by scattering properties of electrons in a conductor. Similarly, in case of two electrons, the coefficient of outgoing spherical waves, see Eq. (2), can also be identified as reduced S-matrix element,

\[
S(k, p) = 2\pi \delta(\theta_k - \theta_p) + 2it(k, p),
\]

which satisfies unitarity relation,

\[
\int \frac{d\theta_k}{2\pi} S^*(k, p)S(k, p') = 2\pi \delta(\theta_k - \theta_p').
\]

As already mentioned previously, in multiple particles cases, because of normalization issue of plane wave description of asymptotic states, the unitarity relation of S-matrix in Eq. (6) clearly cannot be interpreted directly as probability conservation relation. In fact, according to quantum scattering theory [22], the unitarity relation of S-matrix satisfies the probability conservation condition, only when S-matrix element indeed represents the physical probability amplitude for the occurrence of transition. The S-matrix in Eq. (3) clearly doesn’t represent the physical probability amplitude. As will be made clear later on, when the physical probability amplitude condition is met, the transmission and reflection coefficients then may be introduced based on the physical probability interpretation of unitarity relation.

Above mentioned difficulty can be remedied by introducing wave packet prescription of physical asymptotic states. The wave packet prescription is only required for the incident asymptotic state, since the outgoing states will be averaged out for physical transition process. Moreover, because only reduced S-matrix elements after removing energy conservation constraint are used for the transport properties, it is sufficient to introduce the wave packet of multiple particles that only describes the momenta distribution among particles corresponding to a fixed total energy. Using again two electrons case as an example, the physical wave function with an incoming wave packet of two electrons may be defined by

\[
\Phi(x, p_0) = \int d\theta_p \frac{d\theta_p}{2\pi} \Psi(x, p)\Theta(\theta_p, \theta_{p_0}),
\]

where the function \( \Theta(\theta_p, \theta_{p_0}) \) describes the angular distribution of incoming wave packet peaked around a fixed angle \( \theta_{p_0} \), and is normalized according to relation,

\[
\int \frac{d\theta_p}{2\pi} |\Theta(\theta_p, \theta_{p_0})|^2 = 1.
\]

Hence, the physical S-matrix element is related to plane wave based S-matrix element by,

\[
S(\theta_k, \theta_{p_0}) = \int \frac{d\theta_p}{2\pi} S(\mathbf{k}, p)\Theta(\theta_p, \theta_{p_0}).
\]

The physical S-matrix element in Eq. (9) and the unitarity relation in Eq. (6) together yields a well-defined probability conservation relation.

\[
\int \frac{d\theta_k}{2\pi} |S(\theta_k, \theta_{p_0})|^2 = 1.
\]
Therefore, \(|S(\theta_0, \theta_\rho)|^2\) now can be used to represent the probability of transition from initial momenta configurations peak at \(\theta_0 \in [0, \frac{\pi}{2}]\) into final configuration \(\theta_k\). The transmission coefficient may be introduced by considering all configurations scattered into forward direction,

\[
\mathcal{T} = \int_{\mathbb{R}} \frac{d\theta_0}{2\pi} |S(\theta_0, \theta_\rho)|^2,
\]

and based on the probability conservation, the reflection coefficient thus is simply given by \(R = 1 - \mathcal{T}\).

The arguments and wave packet prescription can be generalized to multiple electrons as well. The plane wave based unitarity relation for \(D\) electrons has the similar form as in Eq. (6),

\[
\oint \frac{d\mathbf{k}}{(2\pi)^{D-1}} |S(\mathbf{k}, \mathbf{\rho}_0)|^2 = 1,
\]

where physical \(S\)-matrix element of \(D\)-electrons is given by

\[
S(\mathbf{k}, \mathbf{\rho}_0) = \int \frac{d\mathbf{k}}{(2\pi)^{D-1}} S(\mathbf{k}, \mathbf{p}) \Theta(\mathbf{\hat{p}}, \mathbf{\hat{p}}_0).
\]

3. Two electrons interaction in a crystal

In this section, as a specific example of our formalism, we use a simple solvable model to demonstrate some interesting features and transport properties of two interacting electrons in crystal. The dynamics of our model is given by Hamiltonian,

\[
\hat{H} = \hat{T} + V(X_1) + V(X_2) + U(X),
\]

where \(\hat{T} = -\frac{1}{2m} \sum_{i=1,2} \frac{\partial^2}{\partial x_i^2}\) is kinetic energy of two electrons. Two types of interactions are considered in this model: (1) \(V(X_1)\) describes the pair-wise interaction between \(i\)-th electron and atoms in crystal, and (2) \(U(X)\) represent the three-body interaction involving both electrons and atoms in crystal. \(U\)-type three-body potential may be approximated by contact interactions,

\[
U(X) = \sum_{a=0}^{N-1} U_0 \delta(x_1 - a_0) \delta(x_2 - a_0),
\]

where \(a_0\) are the locations of atoms and \(N\) denotes the total number of atoms, hence \(U\)-type potential only contribute when both electrons meet at same atom. The spin effect has been neglected in this model, and solutions for spin triplet and singlet may be achieved by symmetrization of wave function. The scattering solution of Schrödinger equation for two electrons system may be obtained by considering Lippmann-Schwinger equation, see e.g. [23].

The \(S\)-matrix may be introduced by studying the asymptotic wave function of two electrons system. In our case, asymptotic wave function is given by combination of both plane waves and spherical wave [23],

\[
\Psi(X, p) = \begin{cases} 
  e^{\rho_1(x_1) + \rho_2(x_2)} \rho_1(x_1, p_1) e^{\rho_2(x_2)} & \text{for } p \leq 0, \\
  e^{\rho_1(x_1) + \rho_2(x_2)} \rho_1(x_1, p_1) e^{\rho_2(x_2)} & \text{for } p > 0,
\end{cases}
\]

where \(\rho_1 = \frac{\partial}{\partial p}, \text{ and } \rho_2(p_1, p_2)\) denotes for the scattering amplitude of single electron. The three-body scattering amplitude, \(T_U\), is given by

\[
T_U(X, p) = \frac{-m}{2} \sum_{\alpha, \beta=0}^{N-1} \phi^\alpha(a_\alpha, k) [D^{-1}]_{\alpha, \beta} \phi(a_\beta, p),
\]

where \(a_\alpha = (a_\alpha, a_\alpha)\), and \(D\) matrix is defined as

\[
D_{\alpha, \beta} = 1 - \frac{\partial}{\partial a_\alpha} - G(a_\alpha, a_\beta).
\]

The wave function \(\phi(X, p)\) and Green’s function \(G(X, X')\) satisfy equations,

\[
\left[ -\hat{T} - V(X_1) - V(X_2) \right] \phi(X, p) = \phi(X, p),
\]

and

\[
\left[ -\hat{T} - V(X_1) - V(X_2) \right] G(X, X') = \delta(X - X'),
\]

respectively.

The plane waves in Eq. (17) are the result of pair-wise \(V\)-potentials between electrons and atoms in crystal. The coefficient of spherical wave, \(T_U\), stands for the “true” three-body \(U\)-type interaction when both electrons and atoms are involved in interaction. After removing energy conservation constraint, the reduced plane wave basis \(S\)-matrix are given by

\[
S(\mathbf{k}, \mathbf{\rho}_0) = \sum_{\delta=0, \ldots, \theta_k} 2\pi \delta(\theta_k - \delta) \delta(\mathbf{k}) + 2i T_U(\mathbf{k}, \mathbf{\rho}_0),
\]

where \(\theta_k = \tan^{-1} \frac{k_1}{k_1}, \theta_\rho = \tan^{-1} \frac{\rho_1}{\rho_1}\), and

\[
s_\delta(\mathbf{k}, \mathbf{\rho}_0) = \left[ \delta_0, \beta_1 + i \delta(k_1, p_1) \right] \left[ \delta_1, \beta_2 + i \delta(k_2, p_2) \right].
\]

stands for the reduced \(S\)-matrix with only the presence of \(V\)-potentials, where \(k_i = \frac{k_1}{p_1}\) and \(\beta_1 = \frac{\rho_1}{p_1}\). The reduced plane wave basis \(S\)-matrix satisfies unitarity relation given in Eq. (6). A simple choice of angular distribution function \(\Theta(\theta_\rho, \theta_{\rho_0})\) may be a gaussian function,

\[
\Theta(\theta_\rho, \theta_{\rho_0}) = \frac{(2\pi)^3}{\sqrt{\pi}} e^{-\frac{\theta_\rho^2}{4\sqrt{\pi}}},
\]

where \(\tau\) is a small parameter to control the width of peak of wave packet. Hence the physical \(S\)-matrix that describes the physical transition probability of two electrons is given by

\[
S(\theta_k, \theta_{\rho_0}) = \sum_{\delta=0, \ldots, \theta_k} s_\delta(\mathbf{k}, \mathbf{\rho}_0) \Theta(\theta_\rho, \theta_{\rho_0}) + 2i \oint \frac{d\theta_\rho}{2\pi} T_U(\mathbf{k}, \mathbf{\rho}_0) \Theta(\theta_\rho, \theta_{\rho_0}),
\]
and the transmission coefficient defined by Eq. (11) represents the probability of finding both two electrons scattered into forward direction.

3.1. U = 0 limit

In the case of U = 0, the $T_U(k, p) \rightarrow 0$, the unitarity relation of physical S-matrix in Eq. (10) is thus reduced to a simple form,

$$\sum_{k_i = \pm p_i} \left| s_V(k, p) \right|^2 = 1, \quad (26)$$

this is exactly what we expected for non-interacting two electrons. Assuming $(p_1 > 0, p_2 > 0)$, the transmission and reflection coefficients may be introduced by

$$T = \left| s_V(p_1, p_2; p_1, p_2) \right|^2.$$

$$R = \left| s_V(-p_1, p_2; p_1, p_2) \right|^2 + \left| s_V(p_1, -p_2; p_1, p_2) \right|^2 + \left| s_V(-p_1, -p_2; p_1, p_2) \right|^2. \quad (27)$$

Hence, $T$ does indeed describe the probability of finding both electrons in forward direction after scattering. The complication in reflection coefficient of two electrons are due to the fact that two electrons wave function now has four independent plane waves: $e^{i(k_p x_1 + k_p x_2)}$, in addition to both electrons in forward direction, which create three other scenarios: (i) $s_Y(-p_1, p_2; p_1, p_2)$ describe particle-2 moves forward and particle-1 is scattered backward; (ii) similarly, $s_Y(p_1, -p_2; p_1, p_2)$ is related to particle-1 moves forward and particle-1 is scattered backward; (iii) and $s_Y(-p_1, -p_2; p_1, p_2)$ is associate with both particles are scattered backward.

3.2. V = 0 limit

At another extreme limit, as $V \rightarrow 0$, $s_Y(k, p) \rightarrow \delta_{k_1 p_1} \delta_{k_2 p_2}$, $\phi(x, p) \rightarrow e^{ipx}$, and

$$G(x, x') = -\frac{m}{2} i H_0^{(1)}(p | x - x'|). \quad (28)$$

The physical S-matrix now has a form,

$$S(\theta, \theta_0) = \Theta(\theta, \theta_0) + 2 \int \frac{d\theta_0}{2\pi} T_U(k, p) \Theta(\theta_0, \theta_0), \quad (29)$$

where

$$T_U(k, p) = -\frac{m}{2} \sum_{\alpha, \beta = 0}^{N-1} e^{-ik\alpha} \left[ D^{-1} \right]_{\alpha, \beta} e^{ip\alpha}, \quad (30)$$

and

$$D_{\alpha, \beta} = \frac{1}{U_0} \delta_{\alpha, \beta} + \frac{m}{2} i H_0^{(1)}(p | \alpha - \beta |). \quad (31)$$

The diagonal matrix elements $D_{\alpha, \alpha}$ present another difficulty due to the ultraviolet divergence of Hankel function at origin,

$$H_0^{(1)}(pr) \rightarrow \frac{2i}{r} \left( \gamma_e + \ln \frac{p}{\Lambda} \right), \quad \Lambda = \frac{2}{r}. \quad (32)$$

where $\Lambda$ is served as ultraviolet regulator. Ultimately, the physical result should not depend on the choice or regulator and it will be set to $\Lambda \rightarrow \infty$. The ultraviolet divergence may be dealt with standard renormalization procedure [24,25]. The ultraviolet divergence in Hankel function at origin may be absorbed by bare coupling strength $U_0$, a scale dependent running renormalized coupling strength is hence introduced by

$$\frac{1}{U_R(\mu)} = \frac{1}{U_0} - \frac{m}{\pi} (\gamma_e + \ln \frac{\mu}{\Lambda}), \quad (33)$$

where $\mu$ stands for the renormalization scale, and $U_R(\mu)$ is the physical coupling strength measured at scale $\mu$. The diagonal matrix element $D$ is now given by

$$D_{\alpha, \alpha} = \frac{m}{2} i + \frac{1}{U_R(\mu)} - \frac{m}{\pi} \ln \frac{p}{\mu_B}. \quad (34)$$

The physical observable, $D$, shouldn’t depend on the renormalization scale $\mu$,

$$\frac{d}{d\mu} D_{\alpha, \alpha} = 0. \quad (35)$$

Hence it yields a equation for running coupling strength,

$$\frac{dU_R(\mu)}{d\ln \mu} = \frac{mU_R^2(\mu)}{\pi}, \quad (36)$$

and the solution of running coupling strength is given by

$$\frac{1}{mU_R(\mu)} = \frac{1}{mU_R^0} - \frac{1}{\pi} \ln \frac{\mu}{\mu_B}. \quad (37)$$

where the initial condition of physical observable $U_R^0 = U_R(\mu_B)$ is coupling strength measured at scale $\mu_B$. The scale dependence in $U_R(\mu)$ and $\frac{dU_R}{d\ln \mu}$ in Eq. (34) cancel out, so ultimately, physical observable, $D$, indeed doesn’t depend on the choice of renormalization scale $\mu$.

$$D_{\alpha, \alpha} = \frac{m}{2} i + \frac{1}{U_R^0} - \frac{m}{\pi} \ln \frac{p}{\mu_B}. \quad (38)$$

For the weak coupling $(U_R^0 \ll m^{-1})$, the $D$ matrix may be approximated by only diagonal elements: $D_{\alpha, \alpha} \sim \delta_{\alpha, \beta} \frac{1}{U_R^0}$, hence

$$T_U(k, p) \rightarrow -\frac{1}{2} \left( \frac{1}{mU_0^0} - \frac{1}{\pi} \ln \frac{p}{\mu_B} \right) e^{i\frac{\mu \Omega}{\sqrt{2}}} \sin \frac{\mu \Omega}{\sqrt{2}}. \quad (39)$$

where $\Omega \equiv \cos(\theta_p - \frac{\pi}{4}) - \cos(\theta_k - \frac{\pi}{4})$. We have also assumed that all atoms are separated with even distance: $\epsilon_0 = \frac{1}{2} \alpha, \alpha = 0, \ldots, N - 1$, where $L$ stands for the length of crystal. The transmission coefficient, in case of a shape peaked wave packet $(r \rightarrow 0)$, is now given by

$$T \rightarrow 1 - \frac{2\tau}{(2\pi)^2} \frac{1}{4} + \frac{1}{\pi mU_0^0} \frac{1}{\tau} \frac{2\tau}{\sin^2 \frac{\mu \Omega}{\sqrt{2}}} d\theta_k. \quad (40)$$

One of interesting feature in two interacting electrons case is that due to the U-type three-body interaction, the integrand expression in Eq. (40), $\frac{\sin \frac{\mu \Omega}{\sqrt{2}}}{\sin \frac{\mu \Omega}{\sqrt{2}}}$, shows the interference pattern and resembles to intensity distribution from an ideal grating with N slits in optics or the resistance of one-dimensional chains in Kronig-Penny-like models (see, e.g. [26]). In contrast, in the case of the single electron interacting with N numbers of contact interactions, even at weak coupling limit, the phase factors in forward scattering amplitude all cancel out. The transmission coefficient for single electron is independent of phase factors: $T = 1 - N^2 \frac{mU_0^0}{p^2}$, and shows no interference pattern.
4. Discussion and summary

In order to see the resemblance of multiple channels Landauer-Büttiker formula and multiple particles S-matrix formalism, let’s consider the case of single electron traveling in a quasi-1D wave guide along z-direction. The potential barrier is placed at center of wave guide, and the motion of electron in transverse direction is confined in a narrow tube. Hence, the energy spectra in transverse direction is discretized, the wave function is given by the product of a plane wave in z-direction, $e^{ipz}$, and bound state wave function in transverse direction, $\Phi_n(x, y)$, where $n$ refers to the $n$-th energy state, $\epsilon_n$, in transverse direction, and $p_n = \sqrt{2m(E - \epsilon_n)}$. Assuming initial incident electron is in $n$-th eigenstate, thus, the scattered wave function of electron in forward direction is given by

$$\Psi_n(x, E) \rightarrow \sum_{n'} S_{n,n'} \Phi_{n'}(x, y) e^{i\theta_n x},$$

(41)

where $S_{n,n'}$ is scattering S-matrix element between $n$-th and $n'$-th channels, and satisfies unitarity relation: $\sum_{n'} |S_{n,n'}|^2 = 1$. Since transverse wave function, $\Phi_n(x, y)$, is also well normalized according to

$$\int dxdy \Phi_n^*(x, y) \Phi_n(x, y) = \delta_{n,n'},$$

(42)

the coefficient of plane wave in z-direction, $S_{n,n'} \Phi_{n'}(x, y)$, may still be used to describe probability of physical transition process. Hence, the transmission coefficient in initial channel-$n$ may be defined as net result of coefficient square,

$$\mathcal{T}_n = \int dxdy \sum_{n'} S_{n,n'} \Phi_{n'}(x, y)|^2 = \sum_{n'} |S_{n,n'}|^2.$$  

(43)

In the case of two electrons, the situation is somehow similar, the two electrons wave function in forward direction is now described by outgoing spherical waves, $e^{i(p_x x + \frac{p_y y}{2})}$, propagating in radial direction, and the angular dependent physical S-matrix element,

$$\Phi(x, p_0) \rightarrow \Phi_{\theta_0} S(\theta_0) = e^{i(\frac{p_x x - \frac{p_y y}{2})}{\sqrt{2m p_x}}},$$

(44)

If each possible configuration of allowed momenta distribution among particles is labeled as a single channel, in multiple particles case, there are infinite channels. The scattering of multiple particles may be treated as a continuously distributed multiple-channel problem. Physical S-matrix element square, $|S(\theta_0, \theta_{p_0})|^2$, hence describe the transition probability between channel-$\theta_0$ and channel-$\theta_{p_0}$. The transmission coefficient in initial channel-$\theta_0$ is thus given by net result of all forward transitions,

$$\mathcal{T}_{\theta_0} = \int_0^\pi d\theta_0 \sum_{\theta_{p_0}} |S(\theta_0, \theta_{p_0})|^2.$$  

(45)

In summary, the transport properties of few-electron system is normally complicated by some new features due to multiple particles interaction effect, such as interference and diffraction. The proper approach of introducing transmission and reflection coefficient of few-electron system is discussed in present work based on the probability interpretation of physical unitarity relation of scattering S-matrix. The normalization paradox of unitarity relation is remedied by the wave packet description of incident physical states.

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