Continuum discretization methods in a composite-particle scattering off a nucleus: the benchmark calculations

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(Dated: June 13, 2008)

The direct comparison of two different continuum discretization methods towards the solution of a composite particle scattering off a nucleus is presented. The first approach – the Continuum-Discretized Coupled Channel (CDCC) method – is based on the differential equation formalism, while the second one – the Wave-Packet Continuum Discretization method – uses the integral equation formulation for the composite-particle scattering problem. As benchmark calculations we have chosen the deuteron on a $^{58}\text{Ni}$ target scattering (as a realistic illustrative example) at three different incident energies: high, middle and low. Clear non-vanishing effects of closed inelastic channels at small and intermediate energies are established. The elastic cross sections found in both approaches are very close to each other for all three considered energies.

PACS numbers: 25.10.+s, 25.45.De, 03.65.Nk, 21.45.+v

I. INTRODUCTION

Historically the main progress in studying the elastic and inelastic (breakup) composite projectile scattering off a heavy target was done within the Continuum-Discretized Coupled-Channel (CDCC) method [1-17] in the framework of the Schroedinger coupled-channel scheme. In the CDCC approach the total three-body scattering wave function (in initial versions of the method the projectile was assumed to be a two-fragment nucleus) is expanded in a complete basis of the fragments relative motion of the projectile and the unknown threedimensional elastic and breakup channel wave-functions of the projectile c.m. motion are obtained by solving a set coupled-channel equations. This approach has been successfully applied in the field of nuclear reactions to find the scattering amplitudes of many particular processes successfully applied in the field of nuclear reactions to find coupled-channel equations. As benchmark calculations we have chosen the deuteron on a $^{58}\text{Ni}$ target scattering (as a realistic illustrative example) at three different incident energies: high, middle and low. Clear non-vanishing effects of closed inelastic channels at small and intermediate energies are established. The elastic cross sections found in both approaches are very close to each other for all three considered energies.

Therefore above the three-body breakup threshold the total wave function $\Psi(\mathbf{r}, \mathbf{R})$ vanishes at $r \to \infty$ (similarly to the bound-state function) while the dependence of the total scattering wave function upon the projectile center of mass coordinate has the pure outgoing wave asymptotic behavior. Such an asymmetry for the motions along the two coordinates at energies above the breakup threshold looks not fully justified because in this energy region the proper asymptotic behavior over
all spatial variables might be important.

However, in case of the scattering of a weakly bound projectile (e.g., deuteron or $^7\text{Li}$) at relatively high incident energies ($E \gtrsim 40$ MeV) the motion along the projectile c.m. coordinate $R$ proceeds much more faster than the motion along the relative coordinate $r$ (if the projectile c.m. energy is much larger than its binding energy). In this case the breakup of the projectile takes place mostly outside the nuclear field of the target. In other words, the projectile is first excited into its internal-continuum states and then (already outside the reaction region) the excited projectile breaks up into its fragments. Within this picture, the breakup of the projectile in the target field can be treated as a particular case of inelastic scattering into the projectile continuum states. Then a proper averaging over energy bins for the fragments motion in the discretized continuum gives just a discrete representation for an exact spectral density function of this subsystem. As a result, this discretized density function enters all the elastic and breakup amplitudes. Thus, the applicability condition for the treatment of the above processes within the CDCC method should be as follows: $E_{\text{total}} \gg E_{\text{binding}}$. However, when the projectile kinetic energy $E_{\text{total}}$ is comparable or even a bit larger than its inner binding energy the other asymptotic channels should be taken into consideration and thus, the coupled-channel Schroedinger equation formalism should be replaced by the respective Faddeev approach.

Thus, within its scope of validity, the CDCC approach can be considered as a rather useful and practical working method to handle with the realistic problems of composite projectile scattering off a target. However it would be very important to test carefully the accuracy and predictive power of the CDCC results in some fully independent way based on a different approach. The present paper is devoted just to a careful comparison between the CDCC and one alternative method for some typical problem in the field.

The alternative approach which we have chosen for our study is a new integral equation approach based on a total continuum discretization [19, 20, 21, 22] which has been developed by two of the present authors some time ago. The approach is based on the Wave-Packet Continuum Discretization (WPCD) technique which allows to convert the three-body integral equations into the respective matrix equation with the matrix elements easily calculable in an analytical way. In the WPCD method the motion along both radial coordinates $r$ and $R$ is considered symmetrically, i.e. the respective averaged wave functions (stationary wave packets) vanishing at infinity along both coordinates are used. Such a fully discretized integral approach can be used because in the expression for the elastic scattering amplitude the total three-body resolvent operator $G(E)$ is bracketed from the left and from the right by fast decaying factors over all the spatial coordinates:

$$A_{\text{el}}(E) = \langle \phi_0, \psi_{C}^{(-)}(E-\epsilon_0) | \hat{V} + \hat{V} G(E) \hat{V} | \phi_0, \psi_{C}^{(+)}(E-\epsilon_0) \rangle,$$

where $| \phi_0, \psi_{C}^{(+)}(E-\epsilon_0) \rangle$ is the initial (final) projectile three-body wave function, so that the "external" interaction, $\hat{V}$, decaying along the c.m. coordinate $R$ and the projectile bound-state wave functions $\phi_0(r)$ decaying along the internal coordinate $r$ cuts effectively the asymptotic parts of the total resolvent $G(E)$. Thus, instead of the exact three-body resolvent operator $G(E)$, its respective wave-packet finite-dimensional approximation $G(E)$ can be favorably used [20, 22]. Therefore, in this integral wave-packet approach we do not need any asymptotic parts of scattering wave functions. All that is required is just the inner parts of the wave functions but with the proper normalization which can be well approximated via the stationary wave packets in the WPCD approach. It should be stressed that the effective cutoff of peripheric three-body resolvent parts does not depend on the total energy. Hence, if to neglect the stripping channels (similarly to the conventional CDCC method) and to treat the elastic and breakup channels only, the wave-packet method looks fully justified at low and high energies as well.

Such a ‘totally discretized’ integral technique has several evident advantages as compared to other formulations of the same problem. In addition to a convenient matrix scheme for a calculation of both the elastic and breakup amplitudes, it also allows to construct in an explicit form an optical (energy-dependent and non-local) effective potential for the projectile-target interaction which accounts for the intermediate inelastic processes properly [20, 22]. Another distinctive feature of the integral approach is a proper and straightforward inclusion of closed channels which are treated in this approach on the same footing as the open channels (see especially the section III).

So, to make the comparison between two approaches maximally broad we considered the deuteron elastic scattering by $^{58}\text{Ni}$ at three different incident energies: high, intermediate and low, where closed channel contributions are expected to be quite different. This reaction has been also used to compare the CDCC and Faddeev approaches in a recent work [23], showing very good agreement between both methods. It will be vividly demonstrated that the contribution of the closed channels, which are commonly neglected in CDCC calculations [1, 2, 3, 4, 5, 6, 7, 8, 9, 12], raises essentially when the projectile energy gets low. In principle, closed channels can be incorporated within the CDCC scheme, however, in this case the solution of the coupled equations becomes numerically unstable. In this situation, it is more convenient to solve the problem using the R-matrix method [24]. This is an equivalent way of solving the coupled equations, but with the advantage of being more stable numerically as compared to the traditional radial stepping methods. However, this reformulation demands more computational resources than the conventional coupled-channel scheme [11]. Thus, we made a comprehensive comparison for both open and closed channel contributions between the R-matrix CDCC and
WPCD approaches at different energies.

The paper is organized as follows. In Section II a brief description of the CDCC and WPCD approaches is presented. Section III contains numerical results obtained with these two approaches and the discussion is placed in Section IV. Finally, the summary of the paper is given in Section V.

II. BRIEF DESCRIPTION OF THE CDCC AND WPCD APPROACHES

The Hamiltonian describing the scattering of the two-fragment composite particle \{bc\} off the target nucleus \(A\) is taken in the form:

\[
H = h_{bc}(r) + h_C(R) + V_{b-A}(E_{lab}/2) + V_{c-A}(E_{lab}/2) + \Delta V_C,
\]

where \(h_{bc}\) is the subHamiltonian of the internal \(b-c\) motion (acting on the fragment relative coordinate \(r\)), \(h_C\) — the subHamiltonian of the projectile center of mass asymptotic motion including long-range point-like Coulomb interaction (acting along the c.m. coordinate \(R\)), \(V_{b-A}\) and \(V_{c-A}\) are energy dependent optical potentials for the fragment–target system taken at half the incident particle energy \(E_{lab}\) (for the nearly equal mass fragments). Finally, \(\Delta V_C\) is the additional short-range Coulomb interaction (acting along the c.m. coordinate) which is caused by the finite charge radius of the target nucleus (due to charge distribution in the target nucleus). To obtain the elastic scattering and breakup amplitudes for the above scattering problem with the three-body Hamiltonian \(H\) one has to solve either the three-body Schroedinger equation with proper asymptotic boundary conditions or the system of the respective three-body Lippmann–Schwinger equations.

A. Continuum discretized coupled channel method (a differential equation formulation).

In the CDCC approach the continuous spectrum of the \(h_{bc}\) subHamiltonian is discretized by dividing the continuous momentum distribution into a finite number of non-overlapping bins \([k_{i-1}, k_i)\) with corresponding averaged (within each bin) continuum wave functions \(\{\phi_i\}\) and respective averaged ‘channel eigenenergy’ values \(\epsilon^*_i = \langle \phi_i | h_{bc} | \phi_i \rangle\). To simplify the notation we omit here the angular variables but they are assumed to be included. Then, the total three-body wave function of the Hamiltonian \(H\) is expanded over the set of averaged wave functions describing the projectile internal spectrum (including the projectile bound state \(|\phi_0\rangle\) with the binding energy \(\epsilon_0^*\)):

\[
|\Psi(E)\rangle = \sum_{i=0}^{N} |\phi_i, \chi_i\rangle,
\]

where \(\{R|\chi_i\rangle \equiv \chi_i(R)\) are the channel wave functions which define the elastic \((i = 0)\) and breakup \((i \neq 0)\) amplitudes. The bin wave functions \(\{\phi_i\}\) are typically constructed by averaging the true continuum states, \(\phi(k)\) within the bin interval:

\[
|\phi_i\rangle = \frac{1}{\sqrt{\Delta_i}} \int_{k_{i-1}}^{k_i} f(k)|\phi(k)\rangle dk, \quad i = 1, \ldots, N
\]

where \(\Delta_i\) is a normalization constant and \(f(k)\) is a weight function. For non-resonant continuum the common choice is \(f(k) = 1\), in which case \(\Delta_i = k_i - k_{i-1}\).

Applying the expansion to the initial Schroedinger equation for the Hamiltonian \(H\) one gets a system of coupled differential equations for the unknown functions \(\chi_i(R)\):

\[
[h_C(R) + \Delta V_C(R) + V_{ii}(R) - (E - \epsilon_i)]\chi_i(R) = \sum_{i', j=1}^{N} V_{ii'}\chi_{i'}(R),
\]

where \(V_{ii'}(R) \equiv \langle \phi_i | V_{b-A} + V_{c-A} | \phi_{i'} \rangle\) are coupling potentials. This system is solved with the following boundary conditions:

\[
\chi_i(R) \sim I(K_i, R)\delta_{i0} - \sqrt{\frac{K_i}{K_0}} S_{i0} O(K_i, R),
\]

where \(I\) and \(O\) are the Coulomb incoming and outgoing waves, \(K_i\) is the c.m. wave number which is related to the ‘channel eigenenergy’ \(\epsilon^*_i\) by energy conservation, i.e. \(E = \epsilon^*_i + \frac{k^2}{2m}\) (here \(M\) is deuteron-target reduced mass) and \(S_{i0}\) are \(S\)-matrix elements for the elastic \((i = 0)\) and inelastic \((i \neq 0)\) scattering.

B. Wave-packet continuum discretization method

In the WPCD approach one applies an analogous discretization of the continuous spectrum of the \(h_{bc}\) subHamiltonian but making use of energy bins \([\epsilon_{i-1}, \epsilon_i)\). In a similar way, one defines a set of stationary wave packets \((WP)\) as integrals of the exact scattering wave

\[1\] The integral is taken over the relative r coordinate only.
\[2\] For closed channels \((\epsilon_i > E)\) the boundary condition should be defined with an exponentially decaying and rising functions. In the standard CDCC approach the inclusion of closed channels causes numerical instabilities due to the unavoidable appearance of exponentially rising asymptotic parts, so that the closed channels are usually not taken into account in these calculations. To avoid this problem the combination of the CDCC and R-matrix approaches can be used.
\[3\] It is possible to make just the same energy and momentum bins in the WPCD and CDCC approaches using the interrelation \(\epsilon_i = \frac{\hbar^2 k_i^2}{2m}\), where \(m\) is the reduced mass of the two projectile constituents.
functions $|\phi(E)\rangle$ of the $h_{bc}$ sub-Hamiltonian over the respective energy bins:

$$|Z_i\rangle = \frac{1}{\sqrt{D_i}} \int_{\epsilon_{i-1}}^{\epsilon_i} |\phi(E)\rangle dE, \quad i = 1, \ldots, N$$

where $D_i = \epsilon_i - \epsilon_{i-1}$ are the bin widths and the corresponding 'channel eigenenergies' are just the bin midpoints: $\epsilon_i^* = \frac{1}{2}(\epsilon_i + \epsilon_{i-1})$. The set of these WP states should be supplemented with the bound state $|Z_0\rangle \equiv |\phi_0\rangle$. The main difference in this discretization procedure with respect to the conventional CDCC one is that we use here a full set of WP states which includes both open and closed channels.

Besides that we build a partition of the continuous energy spectrum of the Coulomb sub-Hamiltonian $h_C$ acting along the projectile c.m. coordinate $R$ and employ the corresponding set of Coulomb stationary wave packets (CPs) constructed by averaging the regular Coulomb continuum wave functions $|\psi^C(E)\rangle$ over energy bins $[\mathcal{E}_{j-1}, \mathcal{E}_j]$: $|X_j^C\rangle = \frac{1}{\sqrt{\Delta_j}} \int_{\mathcal{E}_{j-1}}^{\mathcal{E}_j} |\psi^C(E)\rangle dE,$

where $\Delta_j \equiv \mathcal{E}_j - \mathcal{E}_{j-1}$ are the bin widths. Finally, we construct the three-body wave-packet (TWP) basis set for the elastic channel asymptotic Hamiltonian,

$$H_{bc} = h_{bc} \oplus h_C,$$

as the direct product of the states $|Z_i\rangle$ and $|X_j^C\rangle$ (including all the required partial wave couplings):

$$|S_{ij}\rangle \equiv |Z_i, X_j\rangle, \quad i = 0, \ldots, N; \quad j = 1, \ldots, M.$$  

Such a three-body packet basis (which is also of the $L_2$ type) is very convenient for the subsequent investigations. One of its main advantages is that the matrix of the channel resolvent $G_{bc} \equiv [E + i0 - H_{bc}]^{-1}$ is diagonal in this basis. Moreover, the corresponding matrix elements of the three-body resolvent operator have an explicit analytical form and they depend on the spectrum discretization parameters (i.e. $\epsilon_i$ and $\mathcal{E}_j$ values) only.

Due to the fact that the interactions between the projectile constituents and the target are given by complex optical-model potentials the contribution of the intermediate rearrangement (viz. stripping) channels to the elastic scattering or breakup should be of minor importance. Therefore, in order to treat the projectile elastic scattering in the leading order, one can take into account only those intermediate breakup states whose asymptotic behavior are defined by the three-body channel Hamiltonian $H_{bc}$. So, the total three-body scattering wave function $|\Psi(E)\rangle$ in this case can be found from a single Lippmann–Schwinger (LS) equation:

$$|\Psi(E)\rangle = |\Psi_d(E)\rangle + G_{bc}(E)V_{bc}|\Psi(E)\rangle,$$

where $V_{bc} \equiv V_{0-A} + V_{c-A} + \Delta V_C$ is the "external" short-range interaction and $|\Psi_d(E)\rangle \equiv |Z_0, \psi^C(E - \epsilon_0^*)\rangle$ is the outgoing wave function for the projectile in its ground state. The long-range Coulomb part for the point-like Coulomb interaction between the projectile and target is included into the $G_{bc}$ operator. Next, we make the projection of Eq. (10) onto the wave-packet subspace and expand the total wave function in the three-body wave-packet basis for the Hamiltonian $H_{bc}$:

$$|\Psi(E)\rangle \approx |\hat{\Psi}(E)\rangle = \sum_{ij} C_{ij}|S_{ij}\rangle.$$  

Then Eq. (10) can be rewritten in a finite-dimensional form

$$|\hat{\Psi}(E)\rangle = |S_{0j0}\rangle + G_{bc}(E)\hat{V}|\hat{\Psi}(E)\rangle,$$

where $|S_{0j0}\rangle \equiv Z_0, X_j^C$ is the packet state corresponding to the outgoing wave function $|\Psi_d(E)\rangle$ (the index $j_0$ is defined by the energy rule: $E \in [\mathcal{E}_{j_0-1} - \epsilon_0^* \mathcal{E}_{j_0} + \epsilon_0^*)$. The interaction operator $\hat{V}$ with a hat is the wave packet projection of the external interaction $V_{bc}$

$$\hat{V} \equiv \sum_{ij, i'j'} |S_{ij}\rangle \langle S_{ij}|V_{ij, i'j'},$$

whose matrix elements in the TWP basis can be interpreted as generalized coupling potentials:

$$V_{ij, i'j'} \equiv \langle S_{ij}|V_{0-A} + V_{c-A} + \Delta V_C|S_{i'j'}\rangle$$

and may be calculated either analytically or numerically. Thus, in this approach solving the three-body LS equation is reduced to the solution of the simple matrix equation

It should be stressed that the matrix equation in our case is strongly distinguished from that which can be deduced from a direct matrix reduction of the three-body LS equation using some quadrature mesh-points because all the matrix elements in our case have been averaged over the energy bins corresponding to both the active coordinates.
the breakup amplitude can be readily found as a non-diagonal (off-shell) matrix element of the same matrix 
\[ V^{-1} - G_{bs} \]^{-1} in the WP basis [20, 22]. The above matrix is essentially a finite-dimensional analog of the exact transition operator matrix.

Here it is appropriate to summarize the main distinctive features which distinguish the wave-packet approach from the conventional CDCC method:

(i) The wave-packet basis for the projectile bound and continuum internal states is chosen to go up to high excitation energies, so that not only open-channel but also closed-channel contributions are fully included into the WP three-body calculations.

(ii) The 'internal' and 'external' motions of the projectile are taken into consideration in the WP technique on 'equal footing', i.e. using the bases of wave packets averaged over the respective energy bins for the motions along both independent radial coordinates. So that the total three-body resolvent in the transition matrix elements is cut effectively at large distances by the interaction potentials and also by the bound-state functions of the projectile in the initial and final states.

(iii) The above mentioned cutoff of the total three-body resolvent is intimately related to the integral formulation of the WP approach, in which a proper asymptotic behavior of coupling potentials and also an explicit matching procedure between inner and asymptotic wave functions are not required.

(iv) The kernel of the respective matrix equation in the WP approach includes rather smoothed (on energy) matrix elements due to the integration over energy bins for the motions along all active coordinates. This important feature of WP matrix kernels can readily be extended to the solution of the general Faddeev equations using the WP-technique [20].

(v) The integral formulation of the three- and few-body scattering problem makes it possible to reformulate easily the initial problem in terms of the Feshbach projection operator technique [22]. This reformulation allows to reduce the elastic scattering of a composite projectile to a simple potential scattering by a non-local and energy-dependent potential and moreover to construct this potential in an explicit analytical form.

So, it would be very instructive to make a direct comparison between the WPCD and the conventional CDCC results for some standard test problem.

III. COMPARISON BETWEEN THE CDCC AND WPCD: THE BENCHMARK CALCULATIONS AT DIFFERENT INCIDENT ENERGIES

As a general numerical test of the two methods we have chosen the popular case of the elastic scattering of the deuteron off $^{58}$Ni target at three different incident energies, namely, $E_{lab} = 80, 21.6$ and 12 MeV. These detailed calculations allow not only to make more wide comparison between two approaches but also to deduce from this comparison the role of the closed channels at different energies.

For the sake of simplicity, the n-p interaction is assumed here to have the Gaussian form [4]:

\[ V(r) = -V_0 e^{-\beta r^2}, \quad V_0 = 66.99 \text{ MeV}, \quad \beta = 0.415 \text{ fm}^{-2}. \] (16)

Also, keeping in mind the comparison purposes between the two alternative methods, we restrict ourselves here to s-waves for the n-p relative motion 7.

The nucleon-nucleus interactions are represented by local optical-model potentials of Woods-Saxon type. In particular, we employ here the potentials from the Koning–Delaroche global fits [23] evaluated at half of the deuteron incident energy. The mass values $m_n = 1.0087$ amu (for the neutron), $m_p = 1.0078$ amu (for the proton) and $m_{58}$Ni = 57.9353 amu (for the target) are adopted here.

A. Choice for parameter values of the momentum distribution in the CDCC method

Following the standard procedure, in the CDCC calculations the continuum was truncated by defining a maximum excitation $E_{max}$, and divided into $N$ energy bins, evenly spaced in the linear momentum. For each bin, a representative wave function is constructed according to Eq. 3 with $f(k) = 1$. The number of bins $N$ was determined in order to achieve convergence of the elastic angular distribution. For the two lower energies, $E_{lab} = 12$ MeV and $E_{lab} = 21.6$ MeV, the calculations include both open and closed channels. Thus, the continuum was represented by a set of $N_{open}$ bins up the maximum available c.m. energy, and by $N_{closed}$ bins from this energy to $E_{max}$, and hence $N = N_{open} + N_{closed}$.

B. Choice for parameter values of the wave-packet bases for the WPCD calculations

To simplify the construction of the n-p relative motion WPs $\{|Z_i\rangle\}$ and to get simultaneously the bound state

\[ \text{7} \] Higher n-p partial waves would be certainly important to compare the theoretical results with the respective experimental data.
wave function $|Z_0\rangle$ we apply here a single diagonalization procedure for the $\hbar \omega_0$ subHamiltonian on the simple Gaussian basis $|\psi_n\rangle_{\alpha_n=1}$, i.e. we assume that the n-p wave packets are expanded as follows:

$$|Z_i\rangle \approx \sum_{n=1}^{N_G} A_n |\psi_n\rangle, \quad \psi_n(r) = B_n e^{-\alpha_n r^2},$$

(17)

where $B_n$ are normalization factors. The scale parameters $\alpha_n$ are defined on the generalized Tchebyshev grid

$$\alpha_n = \alpha_0 \left[ \tan \left( \frac{(2n-1)}{i} \right) \right]^t, \quad n = 1, \ldots, N_G, \quad (18)$$

where $\alpha_0$ is the common scale parameter and $t$ defines the distribution sparseness for the Gaussian 'frequencies' $\alpha_n$. As $t$ become larger the mesh points $\alpha_n$ are simultaneously extended to lower and higher values, so that more longer-range and shorter-range Gaussian components appear in the basis. The energy distribution $\epsilon_i^t$ (and the corresponding momenta $k_i = \sqrt{2m\epsilon_i^t / \hbar^2}$) obtained with such a basis occurred to be highly non-homogeneous (i.e. non-equidistant), but it covers practically all the energy spectrum and provides well converged results for the scattering amplitudes [19]. Figure 1 shows the energy and corresponding momentum distributions for the following parameter values chosen for the grid: $N_G = 20$, $t = 2$, $\alpha_0 = 0.1 \text{ fm}^{-2}$. In practical calculations we found that it is quite sufficient to include only the first $N$ basis functions with maximum bin energy $\epsilon_N$ less than some chosen maximal energy value $E_{\text{max}}$ of the included closed channels. For example, it is clear from Fig. 1 that a basis containing the first $N = 13$ functions is enough for $E_{\text{max}} = 100$ MeV.

As for the WP basis to describe the n-p pair c.m. motion we employ (for all required partial waves $L$) a set of the exact Coulomb packets of the dimension $M$ constructed also on the Tchebyshev grid with $t = 1$, which corresponds to the energy distribution:

$$E_j = E_0 \tan \left( \frac{(2j-1)}{4 M} \right), \quad j = 1, \ldots, M, \quad (19)$$

where $E_j$ are the end points of the energy bins ($E_0 = 0$) and $E_0$ is the incident energy of the projectile in the c.m. frame.

C. Results of calculations

In this section we will present the particular results for the differential elastic cross sections calculated at different incident energies.

1. $E_{\text{lab}} = 80$ MeV

At this relatively high energy, we found that the breakup effect contribution is rather small, producing just a minor correction to the folding model result (i.e. the calculation omitting continuum channels). Also it was shown previously [2] that only about half of the open channels (i.e. $E_{\text{max}} \approx 40$ MeV) play a significant role at this relatively high energy. So one can conclude that the effect of closed channels can be ignored here. In Fig. 2 it is shown the comparison between the elastic differential cross sections obtained within the CDCC, the WPCD and the folding model. For these calculations the WP bases with dimensions $N = 13$ (for n-p relative motion)$^8$ and $M = 500$ (for deuteron c.m. motion) are used. The number of partial waves required for the deuteron-target relative motion is $L_{\text{max}} = 45$. For the CDCC calculations, the continuum was truncated at $E_{\text{max}} = 70$ MeV and divided into $N = 20$ bins, uniformly distributed in the linear momentum $k$. The set of coupled equations were solved using the Numerov method, and matched to their asymptotic solution at 20 fm. These calculations were performed with the code FRESCO [11].

As it follows from Fig. 2 the results of both discretization methods are in very good agreement for all angles. It should be emphasized that despite the highly non-homogeneous bin distributions for the internal $n-p$ subsystem as well as for the deuteron-target subHamiltonian lead to well converged results of the WPCD calculations. So, it is not necessary to use in practical calculations just equidistant momentum bin distributions as it is usually done [4, 7, 8]. Moreover, the application of some special energy distribution allows to decrease significantly the number of the packet basis functions required for convergence. Thus, to summarize, despite of the quite different...

$^8$ The Gaussian basis [17] with $N_G = 20$, $t = 2$, $\alpha_0 = 0.1 \text{ fm}^{-1}$ has been used for this calculation.
The elastic cross sections calculated with the WPCD approach are in very good agreement for all partial waves. In this figure we have included also the calculation corresponding to the folding model results are presented in Fig. 7. It is evident that the results for the differential cross sections found with the WPCD and R-matrix CDCC methods with similar $E_{\text{max}}$ values are almost identical. So, it can be concluded that the details of the discretization distributions in both methods do not play an important role in the convergence and the most significant factor here is just the $E_{\text{max}}$ value.

The converged elastic cross sections (calculated with the highest value, $E_{\text{max}} = 139$ MeV) are presented in Fig. 3. The curves corresponding to both discretization methods are hardly distinguishable. In this figure we have used the three-body WP sets of dimensions $N = 15$ (constructed from the Gaussian basis [17] with $N_G = 30$, $t = 3$, $\alpha_0 = 0.1$ fm$^{-1}$) and $M = 150$ are sufficient, and the maximum deuteron c.m. angular momentum value is only $L_{\text{max}} = 15$. For the CDCC calculations we used $N_{\text{open}} = 15$ and $N_{\text{closed}} = 10$ (for $E_{\text{max}} \sim 80$ MeV). Again, the R-matrix technique was used to solve the coupled equations, with a matching radius of $R_m = 30$ fm.

The elastic cross sections calculated with the WPCD and the R-matrix CDCC approaches for different $E_{\text{max}}$ values are shown in Figs. 6a, b and c. The comparison between the converged (with respect to $E_{\text{max}}$ values) cross sections calculated with both methods along with the folding model results are presented in Fig. 7.

It is evident that the influence of the CC is the largest at backward angles (Fig. 3a). For the other values of $E_{\text{lab}} = 21.6$ MeV. The effect of closed channels

At this intermediate energy, the influence of closed channels (CC) on the elastic scattering amplitude becomes visible. Our calculations have shown that the main part of this effect is seen at backward angles. Though the CC effect has been commonly neglected in the conventional CDCC calculations, it is worth to study carefully the convergence of the elastic cross section when the number of closed channels and the corresponding maximum energy $E_{\text{max}}$ are increased.

To prepare this calculations in the WPCD scheme we have used the three-body WP sets of dimensions $N = 15$ (constructed with the Gaussian basis [17] with $N_G = 30$, $t = 3$, $\alpha_0 = 0.1$ fm$^{-1}$) and $M = 150$. The number of partial waves required at this energy is $L_{\text{max}} = 25$. To incorporate the closed channels into the CDCC framework a special combination of the coupled-channel scheme and the R-matrix approach has been used [24]. In contrast to the Numerov method, where the radial functions $\chi_i(R)$ are obtained by direct integration of the coupled differential equations [4], in the R-matrix method a basis set of energy eigenstates is first obtained by solving the diagonal parts of the coupled equations, with the basis functions all having fixed logarithmic derivatives at a given distance $R_m$. Next, the unknown functions $\chi_i(R)$ are expanded in this basis. Although this procedure is computationally more demanding, it leads to more stable results when closed channels are taken into account. In order to achieve convergence at this incident energy, we required $N_{\text{open}} \sim 20$ and $N_{\text{closed}} \sim 15$ bins, for a maximum excitation energy $E_{\text{max}} \sim 140$ MeV. The R-matrix radius was fixed to $R_m = 30$ fm.

In Fig. 3 the comparison between the R-matrix CDCC and the WPCD elastic cross sections calculated at almost equal $E_{\text{max}}$ values is presented. For $E_{\text{max}} = 20$ MeV, there are only open-channels and the conventional Numerov procedure has been employed to solve the CDCC equations (see Fig. 3a). For the other values of $E_{\text{max}}$ (Fig. 3b and c), the R-matrix method has been employed, while the WPCD technique has not been changed. It is evident that the results for the differential cross sections found with the WPCD and R-matrix CDCC methods with similar $E_{\text{max}}$ values are almost identical. So, it can be concluded that the details of the discretization distributions in both methods do not play an important role in the convergence and the most significant factor here is just the $E_{\text{max}}$ value.

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We finally present in this section the calculations at the relatively low energy of 12 MeV. In this case the dimensions of the WP sets $N = 15$ (constructed from the Gaussian basis [17] with $N_G = 30$, $t = 3$, $\alpha_0 = 0.1$ fm$^{-1}$) and $M = 150$ are sufficient, and the maximum deuteron c.m. angular momentum value is only $L_{\text{max}} = 15$. For the CDCC calculations we used $N_{\text{open}} = 15$ and $N_{\text{closed}} = 10$ (for $E_{\text{max}} \sim 80$ MeV). Again, the R-matrix technique was used to solve the coupled equations, with a matching radius of $R_m = 30$ fm.

The elastic cross sections calculated with the WPCD and the R-matrix CDCC approaches for different $E_{\text{max}}$ values are shown in Figs. 6a, b and c. The comparison between the converged (with respect to $E_{\text{max}}$ values) cross sections calculated with both methods along with the folding model results are presented in Fig. 7.

It is evident that the influence of the CC is the largest here and this effect is seen not only at backward angles but also at intermediate angles ($\theta_{\text{c.m.}} \sim 90^\circ - 100^\circ$). The small discrepancies between the WPCD and R-matrix
CDCC results observed at most backward angles might be ascribed to the different n-p continuum discretization parameters, but we cannot rule out that they can be related to numerical inaccuracies of either of the two methods. In any case, this effect, which has not been seen in the two previous cases, begins to play some role at rather low incident energies.

**IV. DISCUSSION**

The close agreement between the WPCD and R-matrix CDCC approaches found in the present work can be interpreted as a good indicator that the basic assumptions (or approximations) adopted in the WPCD method, in particular the employment of orthonormalized stationary wave packets instead of exact non-normalizable three-body scattering wave functions are well justified and correct. This fact implies some further consequences in the whole many-body scattering theory. Actually, the usage of the total continuum discretization on the basis of normalizable (of $L_2$ type) stationary wave packets is equivalent to the formulation of the many-body scattering problem in a finite multi-dimensional box and leads a treatment of the original scattering problem analogous to the much more simple bound-state problem. In turn, the last conclusion allows to treat the many-body scattering problem in similar way as the many-body bound states, e.g. by expanding the $L_2$-type stationary wave packets (constructed from the exact unknown scattering
wave functions) on some convenient $L_2$ basis widely used in bound state problems (for example, the many-body harmonic oscillator basis). The first results which two of the present authors \cite{20} have obtained with the application of this technique for the solution of the Faddeev equations above the three-body breakup threshold seems to support this expectation.

Another nice feature of the WPCD approach is the possibility to reformulate in a straightforward way the whole problem in terms of the Feshbach projection technique \cite{22} and to include all the breakup channels into the respective Feshbach polarization non-local potential. After this, one can readily use the full applicability of the WPCD approach to the resulted non-local interactions (in contrast to the conventional coupled-channel method) and to reduce in a standard way the complicated many-body scattering problem to the simple matrix equations \cite{26}.

On the other hand, the very close agreement between the conventional CDCC approach and the fully alternative wave-packet integral method should remove any doubts reported in literature \cite{16,18} concerning the reliability and convergence properties of the Schroedinger coupled-channel scheme in the framework of the three-body scattering problem.

V. CONCLUSION

In this work, we have compared two essentially different approaches based on the general idea of continuum discretization for solving three-body scattering problems. The CDCC method was proposed to solve the coupled-channel problem in the framework of the Schroedinger equation formalism. This approach uses a discretization of the projectile internal continuum only, along with an explicit matching between inner and asymptotic scattering wave functions to extract the multichannel $S$-matrix.

On the other hand, the WPCD approach is based on the total three-body continuum discretization which allows to use the integral equation formalism of the scattering theory. The special wave-packet technique allows to construct finite-dimensional analogs for all the scattering operators and to use only inner parts of wave functions in order to find the observables.

The direct comparison of both methods leads to the following conclusions:

(i) The two methods of continuum discretization discussed in this work produce very close results for the composite particle elastic scattering at rather high and also at low energies.

(ii) Different types of discretization of the projectile internal subHamiltonian continuum used in both methods do not lead to visible discrepancies at high and intermediate energies. Small differences in the elastic cross sections arise only at low incident energies. An employment of a highly non-homogeneous

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FIG. 6: Elastic $d+^{58}\text{Ni}$ cross section at $E_{\text{lab}} = 12$ MeV calculated within the CDCC (dotted line) and WPCD (dashed line) approaches for different maximum values of the excitation energy, $E_{\text{max}}$: a) $E_{\text{max}} = 10.2$ MeV, b) $E_{\text{max}} = 24.5$ MeV, c) $E_{\text{max}} = 36$ MeV. The full curves correspond to the converged CDCC results.

FIG. 7: The same as in Fig. 4 but for $E_{\text{lab}} = 12$ MeV.
dicretization distribution in the WPCD approach still leads to well-converged results.

(iii) The effect of closed channels is not seen at high incident energies. This effect begins to play a visible role at intermediate energies (at the most backward angles) and can not be ignored as it was assumed in some previous CDCC calculations. The convergence of the calculated cross section when increasing the number of included CCs depends only on the maximum excitation energy and is not sensitive to the details of the continuum dicretization.

(iv) The effect of CC on the elastic scattering becomes more significant when the incident energy of the projectile decreases.

(v) The very close agreement between the results obtained with these two different approaches at various energies, from low to high, seems to indicate that the three-body wave-packet technique can be considered as a rather reliable and numerically accurate method for calculations of composite projectile scattering by nuclei. Simultaneously, one can confirm again that the conventional (or R-matrix) CDCC method leads to quite reliable and well-converged results for the elastic and inelastic scattering of composite projectiles.

Acknowledgements The Russian authors (V.I.K. and O.A.R.) are thankful to Dr. V.N. Pomerantsev for numerous fruitful discussions. They appreciate very much partial financial supports from the RFBR grant 07-02-00609, the joint RFBR–DFG grant 08-02-91959 and the President grant MK-202.2008.2. A.M.M. acknowledges a research grant from the Junta de Andalucía and financial support by the DGICYT under project FPA2006-13807-c02-01.

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