Reaction Mechanisms of Pair Transfer

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50 Years of Nuclear BCS
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Chapter 1

Reaction mechanisms of pair transfer

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1. Introduction

Much of the evidence for nucleonic pairing in nuclei comes from energy expectation values, but important further information comes from the *transfer* of pairs of nucleons to or from another nucleus of known structure. Indeed, much of what we have learned about nuclear properties have been derived from experiments involving the collision, or reaction, between two nuclei. In this regard, a more fundamental understanding of nuclear reactions has been, and will continue to be (especially in the FRIB era), crucial to the nuclear physics community. This chapter focuses on the theory, calculation and model results for the reactions mechanisms of pair transfer.

Here we consider the reaction mechanisms for pair transfer between two nuclei, for reactions that we can describe as \( A(B+2, B)A+2 \). Here, the two nucleons may be two neutrons, two protons, or a proton and a neutron, and are transferred from core \( B \) to core \( A \). The nucleons may transfer either in one *simultaneous* step, or one after the other *sequentially*. If a distinguishable proton and a neutron are transferred, then both proton-then-neutron and neutron-then-proton routes need to be considered. Furthermore, these sequential and simultaneous routes contribute amplitudes that all add *coherently*. This feature enables us to probe the nature of coherent two-nucleon superpositions in nuclei. Conversely, these superpositions, coupling orders and phase conventions have all to be defined consistently in a
good calculation.

Subsequent sections will therefore consider the definition of two-nucleon overlap functions, their coordinate transformations, the definition of transfer matrix elements along with zero-range approximations and non-orthogonality corrections. Finally, some results are shown to illustrate the coherence effects in the reaction mechanisms of pair transfers.

In the last 50 years, a significant number of papers have been presented in which absolute differential cross sections have been calculated, and compared with experimental results.\textsuperscript{1–19} Traditionally, for example in,\textsuperscript{9} the theory predictions have fallen well below the experimental data. This ratio has been called the ‘unhappiness factor’.\textsuperscript{20,21} Most previous calculations modeled the transfer of a dineutron as a single cluster. Only from Charlton\textsuperscript{8} were sequential transfer contributions considered. We see now that modern calculations are in considerably better agreement with experiment.

2. Bound states and vertex functions

The general theory of nucleon pair bound states defines the overlap function \( \phi_J^I(\mathbf{r}, \rho) = \langle \Phi_A(I)|\Phi_{A+2}(J) \rangle \) in terms of the Jacobi coordinates \( \mathbf{r} \) between the two nucleons, and \( \rho \) between their center of mass (cm) and the core \( A \). The core spin is \( I \) and the spin of the \( A+2 \) composite state is \( J \). When intrinsic spins \( s_1, s_2 \) are also included in a particular coupling order such as \(|\{L, (\ell, (s_1 s_2)S)j,J12,I,J\rangle \rangle \langle \mu I \nu I | JM \rangle \langle L \Lambda jm_{12}|J12M12|\ell \mu S \Sigma jm_{12}\rangle \langle s_1 \sigma_1 s_2 \sigma_2 S \Sigma \rangle \rangle \) (1)

for some radial wave function \( u_{12}(r, \rho) \) that can be given either as a cluster product of single-particle wave functions \( u_{12}(r, \rho) = \Phi_L(r)\phi_\ell(\rho) \), input directly as a two-dimensional distribution e.g. from a Faddeev bound-state calculation, or calculated from the correlated sum of products of single-particle states with independent coordinates. These two-nucleon wave functions will in general be the eigenstates of a three-body bound state Schrödinger equation

\[ [T_\mathbf{r} + T_\rho + V_{1A} + V_{2A} + V_{12} - \epsilon] \phi_J^I(\mathbf{r}, \rho) = 0, \]

where the \( V_{iA} \) are the potentials between nucleon \( i \) and the core, and \( V_{12} \) is the pairing interaction between the two nucleons.
Such two-particle states that come from shell-model calculations or from Sturmian-basis calculations are usually described by means of the \( |r_1, r_2\rangle \) coordinates, and then transformed internally into the centre-of-mass coordinates \( |r, \rho\rangle \) of Eq. (1) using \( r_i = x_i r + y_i \rho \). For equal mass particles, \( x_1 = x_2 = 1 \), and \( y_1 = -y_2 = \frac{1}{2} \). This describes a pair state as

\[
\varphi_{12}(r_1, r_2) = \sum_i c_i \langle \ell_1(i), s_1 i | j_1(i), (\ell_2(i), s_2) j_2(i); J_{12} \rangle.
\]

The coefficients \( c_i \) for correlated basis states \( i \) and the single-particle wave functions \( \varphi_{\ell sj}(r) \) contain all the physics information about the bound state needed to do a transfer calculation. Shell model codes can produce the coefficients \( c_i \) needed here in terms of previously calculated eigenstates of the \( A \) and the \( A+2 \) systems.

The vertex functions of these bounds states are defined to be these bound state wave functions \( \phi_{\ell sj}(r, \rho) \) multiplied by the potentials which have zero effects after the transfer step is performed and all exit channel nuclei have completely separated. These potentials are therefore the sum \( V \) of the binding potentials \( V_{sp} = V_{\ell sj}(r_i) \) to give \( V = V_{sp}^{1A} + V_{sp}^{2B} \). (These are the individual potentials that should appear in the bound-state equation \( [T_r + V_{\ell sj}(r) - \varepsilon] \varphi_{\ell sj}(r) = 0 \).) The vertex function does not include the nucleon-nucleon pair interaction \( V_{12}(r_1 - r_2) \), since this potential produces binding effects in both the initial and final bound states. We denote by \( V\phi_{\ell sj}(r, \rho) \) the vertex function after transformation into Jacobi coordinates by the same method used to transform the wave function itself.

### 3. Post and prior coupling forms

We now consider the Hamiltonian \( \mathcal{H} \) for the whole system of \( A+B+2 \) nucleons and described by system wave function \( \Psi \). Let the various \( A+2 \) and \( B+2 \) bound states be denoted by \( \Phi_i \) for various \( i \). Then we may expand \( \Psi \) in terms of the \( \Phi_i \) with some coefficients \( \psi_i(R_i) \) depending on the two-body separation vectors \( R_i \), as \( \Psi = \sum_i \psi_i(R_i) \Phi_i \).

The model Schrödinger’s equation \( [\mathcal{H} - E]\Psi = 0 \) when projected separately onto the different basis states \( \Phi_j \), yields the set of equations

\[
[E_j - H_j] \psi_j(R_j) = \sum_{i \neq j} \langle \Phi_j | [\mathcal{H} - E] | \Phi_i \rangle \psi_i(R_i).
\]

which couple together the unknown wave functions \( \psi_i(R_i) \). The channel Hamiltonians are defined by \( H_j - E_j = \langle \Phi_j | [\mathcal{H} - E] | \Phi_j \rangle \) such that the \( E_j \) are the asymptotic kinetic energies in channel \( j \).
The matrix element \( \langle \Phi_j | \mathcal{H} - E | \Phi_i \rangle \) has two different forms, depending on whether we expand

\[
\mathcal{H} - E = H_j - E_j + V_j \quad \text{(the ‘post’ form)}
\]

\[
= H_i - E_i + V_i \quad \text{(the ‘prior’ form)}.
\]

The name (post or prior) is determined by whether it is the initial or final channel whose Hamiltonian is used. The above Eq. (4), as set up, has \( j \) as the initial channel and \( i \) as the final channel for the indicated coupling. Thus

\[
\langle \Phi_j | \mathcal{H} - E | \Phi_i \rangle = V_{ji}^{\text{post}} + [H_j - E_j]K_{ji} \quad \text{(post)}
\]

or

\[
= V_{ji}^{\text{prior}} + K_{ji}[H_i - E_i] \quad \text{(prior)}
\]

(5)

where

\[
V_{ji}^{\text{post}} \equiv \langle \Phi_j | V_j | \Phi_i \rangle, \quad V_{ji}^{\text{prior}} \equiv \langle \Phi_j | V_i | \Phi_i \rangle, \quad K_{ji} \equiv \langle \Phi_j | \Phi_i \rangle.
\]

(6)

The overlap function \( K_{ji} = \langle \Phi_j | \Phi_i \rangle \) in Eq. (5) arises from the non-orthogonality between the basis states \( \Phi_i \) and \( \Phi_j \) if these are in different mass partitions. The \( K_{ji} \) are non-local operators that go to zero asymptotically. (Within the same partition, the \( \Phi_i \) are inelastic states, and form an orthogonal set.)

The first-order DWBA matrix element use entrance \( \psi_i \) and exit \( \psi_j \) channel wave functions satisfying \( [H_i - E_i] \psi_i = 0 \) and \( [H_j - E_j] \psi_j = 0 \) respectively. Its matrix element is

\[
T_{ji}^{(1)} = \langle \psi_j^{(-)} | \Phi_j | \mathcal{H} - E | \Phi_i \psi_i^{(+)} \rangle
\]

(7)

The prior form of this is

\[
T_{ji}^{(\text{prior})} = \langle \psi_j^{(-)} | \Phi_j | H_i - E_i + V_i | \Phi_i \psi_i^{(+)} \rangle
\]

\[
= \langle \psi_j^{(-)} | \Phi_j | V_i | \Phi_i \psi_i^{(+)} \rangle + \langle \psi_j^{(-)} | \Phi_j | H_i - E_i | \psi_i^{(+)} \rangle
\]

\[
= \langle \psi_j^{(-)} | \Phi_j | V_i | \Phi_i \psi_i^{(+)} \rangle + 0
\]

\[
= \langle \psi_j^{(-)} | V_{ji}^{\text{prior}} | \psi_i^{(+)} \rangle.
\]

(8)

Similarly, the equivalent post form is

\[
T_{ji}^{(\text{post})} = \langle \psi_j^{(-)} | \Phi_j | H_j - E_j + V_j | \Phi_i \psi_i^{(+)} \rangle
\]

\[
= \langle \psi_j^{(-)} | \Phi_j | V_j | \Phi_i \psi_i^{(+)} \rangle + \langle \psi_j^{(-)} | [H_j - E_j] \Phi_j | \Phi_i \psi_i^{(+)} \rangle
\]

\[
= \langle \psi_j^{(-)} | \Phi_j | V_j | \Phi_i \psi_i^{(+)} \rangle + 0
\]

\[
= \langle \psi_j^{(-)} | V_{ji}^{\text{post}} | \psi_i^{(+)} \rangle.
\]

(9)
Thus the non-orthogonality term disappears in first-order DWBA. Post
and prior first-order DWBA matrix elements can be made to exactly agree
numerically, if sufficient care is taken to ensure convergence of the non-local
form factors.

Let a second-order DWBA matrix element use entrance channel \( i \), exit
channel \( k \), and some intermediate channel \( j \), as \( i \to j \to k \). The propagation
in the intermediate channel may be described in terms of the Greens
function \( G_j \), or equivalently within an iterated coupled-channels set. The
two-step DWBA matrix element is

\[
T_{ki}^{(2)} = \langle \psi_k^- | \langle \Phi_k | H - E | \Phi_j \rangle G_j \langle \Phi_j | H - E | \Phi_i \rangle | \psi_i^+ \rangle.
\]  

(10)

Now there are four matrix elements that may be calculated, according to
the first and the second Hamiltonian form chosen: post-post, post-prior,
prior-post, and prior-prior. The terms prior and post for each step are used
to refer to the initial or final channels of that step, not the overall incoming
or outgoing channels. In ‘prior-post’, the prior refers to the first step, and
the post refers to the second step.

The post-post form of this, for example, is

\[
T_{ki}^{(post,prior)} = \langle \psi_k^- | \langle \Phi_k | H_k - E_k + V_k | \Phi_j \rangle G_j \langle \Phi_j | H_j - E_j + V_j | \Phi_i \rangle | \psi_i^+ \rangle.
\]

(11)

Here the \( [H_k - E_k] \) can operate on the final \( \psi_k \) to give zero, but little can
simplify the \( [H_j - E_j] \) since \( [H_j - E_j] G_j \neq 0 \) always. Thus

\[
T_{ki}^{(post,prior)} = \langle \psi_k^- | V_{kj}^{post} G_j V_{ki}^{post} | \psi_i^+ \rangle + \langle \psi_k^- | V_{kj}^{post} [H_j - E_j] K_{ji} | \psi_i^+ \rangle.
\]

(12)

This second term is called a ‘non-orthogonality term’ since it involves the
bound-state non-orthogonality overlaps \( K_{ji} = \langle \Phi_j | \Phi_i \rangle \), which is significant
when \( R_i \) and \( R_j \) are both within the range of the bound states.

Similar analyses for post-prior and prior-prior two-step DWBA expression
also have non-orthogonality terms in the final form. The prior-post
form, however, is

\[
T_{ki}^{(prior,post)} = \langle \psi_k^- | \langle \Phi_k | H_k - E_k + V_k | \Phi_j \rangle G_j \langle \Phi_j | H_i - E_i + V_i | \Phi_i \rangle | \psi_i^+ \rangle.
\]

(13)

Here the \( [H_i - E_i] \) can also operate on the initial \( \psi_i \), to give zero, as well
as \( [H_k - E_k] \) on the \( \psi_k \), so we have the simplest form

\[
T_{ki}^{(prior,post)} = \langle \psi_k^- | V_{kj}^{post} G_j V_{ji}^{prior} | \psi_i^+ \rangle.
\]

(14)

The non-orthogonality terms can thus be made to disappear in second-order
DWBA if the first and second steps use the prior and post interactions.
respectively. When they are included as necessary, the results should be the same whatever post or prior forms are used.

In third and higher-order transfer calculations, some non-orthogonality terms will always be present, but most pair transfer mechanisms can be well modeled as two-step processes.

4. Two-nucleon transfer interaction

We now consider the specific transfer matrix element $V_{ji}^{\text{prior}} = \langle \Phi_j | V_i | \Phi_i \rangle$. Given an expression for this prior form, we may calculate the post interaction easily as $V_{ji}^{\text{post}} = (V_{ij}^{\text{prior}})^\dagger$. Take $\Phi_j$ to refer to the bound states of nucleus $A+2$ outside core $A$, and $\Phi_i$ analogously for nucleus $B$.

The transfer interaction has therefore the non-local matrix element

$$V_{ji}(R_j, R_i) = \langle \phi^{J_A}_I(r_A, \rho_A) | V_{1B}^{\text{sp}} + V_{2B}^{\text{sp}} + U_{AB} - U_i | \phi^{J_B}_I(r_B, \rho_B) \rangle. \quad (15)$$

As is usual in transfer operators, there are three kinds of potentials appearing here. First there are the binding potentials $V_{1B}^{\text{sp}}(r_{1B}) + V_{2B}^{\text{sp}}(r_{2B})$. Since these binding potentials always appear while multiplied by their bound state wave functions, we need only store and use the vertex functions defined in section 2. Second there is the ‘core-core’ potential $U_{AB}(R_{AB})$ between the core nuclei $A$ and $B$. Finally is subtracted an optical potential. In this prior form we subtract the optical potential in the initial channel, $U_i(R_i)$. The difference $U_{AB} - U_i$ of the two optical potentials is called the remnant term, and is sometimes taken to be small.

The integrals in Eq. (15) include integrating over the two-nucleon separation $r$ as well as over their cm distance $\rho_A$ from the core $A$. The $r$ coordinate appears in both the initial and final states, and so is not labeled by $A$ or $B$. This has the important consequence that neither the distance nor the angle of the $r$ coordinate is changed in the transfer. Neither, therefore, is their relative angular momentum $\ell$, and, for similar reasons, nor their spin couplings $S$ and total angular momentum $j$. The two neutron transfer can hence be viewed as the transfer of a ‘structured particle’ $(\ell, (s_1s_2)S)j$, and then becomes similar to single-particle transfers of above. This means that when we also integrate over the coordinates $\rho_A$ and $\rho_B$, we can use the standard procedures already developed for one-particle transfer interactions.
5. Coordinate transformations

The transfer mechanism requires the pair wave function to be expressed in the form of Eq. (1), so independent-particle forms of Eq. (3) have to be transformed in their coordinates as

\[ \varphi_{12}(\mathbf{r}_1, \mathbf{r}_2) = \sum_i c_i \sum_{LTSJ} |L, (\ell, (s_1 s_2)S)j; J_{12}T \rangle \phi_{L(TS)12}^{j1 j2}(r, \rho) \] (16)

A particular basis state \( i \) in the \((r, \rho)\) coordinates is

\[ \phi_{L(TS)12}^{j1 j2}(r, \rho) = \langle L, (\ell, (s_1 s_2)S)j; J_{12} \parallel (\ell_1(i), s_1)j_1(i), (\ell_2(i), s_2)j_2(i); J_{12} \rangle \]

\[ \times \langle [Y_{L1}(\tilde{r})Y_{L2}(\tilde{\rho})]_\lambda \parallel [\varphi_{i1 j_1}(\mathbf{r}_1)\varphi_{i2 j_2}(\mathbf{r}_2)]_{J_{12}T} \rangle \] (17)

where (suppressing \( i \) indices for clarity)

\[ \langle L, (\ell, (s_1 s_2)S)j; J_{12} \parallel (\ell_1, s_1)j_1, (\ell_2, s_2)j_2; J_{12} \rangle = \sum_\lambda \hat{\lambda} \hat{J}_{12} \]

\[ \times \left[ \binom{\ell_1}{r_1} \binom{\ell_2}{s_2} \binom{\lambda}{S} \binom{\lambda}{L} \right] \frac{1}{2^{[1+(\delta_{\ell_1, s_1} - \delta_{\ell_1, j_1})]}} \hat{J} \hat{\lambda} W(L \ell J_{12} S; \lambda j)(-1)^{\ell - L - \lambda} \] (18)

The radial overlap integral can be derived by means of harmonic-oscillator expansions,23 with the Bayman-Kallio expansion24 or using the Moshinsky solid-harmonic expansion.25 This last method gives

\[ K_{LL T1T2}^\lambda(r, \rho) \] (20)

\[ = \langle [Y_{L1}(\tilde{r})Y_{L2}(\tilde{\rho})]_\lambda \parallel [\varphi_{T1}(\mathbf{r}_1)\varphi_{T2}(\mathbf{r}_2)]_\lambda \rangle \]

\[ = \sum_{n_1 n_2} \left( \begin{array}{c} 2\ell_1+1 \\ 2n_1 \end{array} \right) \left( \begin{array}{c} 2\ell_2+1 \\ 2n_2 \end{array} \right) (x_1 r_1)^{\ell_1-n_1} (y_1 \rho)^{n_1} (x_2 r_2)^{\ell_2-n_2} (y_2 \rho)^{n_2} \]

\[ \times \sum_Q q^{Q}_{L1 T2}(r, \rho) (2Q+1) \hat{\ell}_1 \hat{\ell}_2 \hat{\ell}_1 - n_1 \hat{\ell}_2 - n_2 \hat{\ell} \]

\[ \times \sum_{A_1 A_2} \left( \begin{array}{c} \ell_1-n_1 \end{array} \begin{array}{c} n_2 A_1 \\ 0 \end{array} \right) \left( \begin{array}{c} \ell_2-n_2 \end{array} \begin{array}{c} n_1 A_2 \\ 0 \end{array} \right) \left( \begin{array}{cc} \Lambda_1 & L Q \\ 0 & 0 \end{array} \right) \left( \begin{array}{cc} \Lambda_2 & \ell Q \\ 0 & 0 \end{array} \right) \]

\[ \times (-1)^{\ell_1+\ell_2+L+\Lambda_1 A_2 (2\Lambda_1 + 1)(2\Lambda_2 + 1)} W(\Lambda_1 A_1 \ell_1; Q \lambda) \]

\[ \times \left( \begin{array}{c} \ell_1-n_1 \\ n_1 \end{array} \begin{array}{c} \ell_2-n_2 A_2 \\ 0 \end{array} \right) \left( \begin{array}{c} \Lambda_1 \ell_1 \ell_2 \lambda \end{array} \right) \] (21)

where \( \binom{a}{b} \) is a binomial coefficient. The kernel function \( q^{Q}_{i1 T2}(r, \rho) \) is the Legendre expansion of the product of the two radial wave functions in terms...
of \( \varphi \), the cosine of the angle between \( \mathbf{r} \) and \( \rho \):

\[
q_{r_1, r_2}^Q(r, \rho) = \frac{1}{2} \int_{-1}^{+1} \frac{\varphi_{s_1 s_1 j_1}(r_1)}{r_1} \frac{\varphi_{s_2 s_2 j_2}(r_2)}{r_2} P_Q(u)du
\]

6. Zero-range and other approximations

The coupling potentials \( V_{ji}(\mathbf{R}_j, \mathbf{R}_i) \) of Eq. (15) are non-local, in the sense that in general the initial and final radii, \( \mathbf{R}_j \) and \( \mathbf{R}_i \), will be different. They will not only have different magnitudes, but also different directions. In the early days of transfer modeling, the calculations became much more practical if a zero-range approximation could be found, in which the coupling was restricted to \( \mathbf{R}_j = \alpha \mathbf{R}_i \) for some constant \( \alpha \) (which need not be unity).

When the projectile is a light ion such as \(^3\text{H} \) or \(^4\text{He} \) for nucleus \( A + 2 \), then the binding potential sum \( V_{1B}^{sp} + V_{2B}^{sp} \) will have short range. We may therefore consider approximating

\[
[V_{1B}^{sp} + V_{2B}^{sp}] \phi_{B}^{jB}(\mathbf{r}, \rho_B) \sim D_0 \delta(\rho_B) \phi_{nn}^B(\mathbf{r})
\]

for some nucleon-nucleon wave function \( \phi_{nn}(\mathbf{r}) \) that we are free to choose. This a zero-range approximation. Note that it is only \( \rho_B \) which needs to have zero range, not \( \mathbf{r} \). The constant \( D_0 \) is called the zero-range constant.

If, furthermore, we can neglect the remnant term \( U_{AB} - U_i \), then the transfer coupling of Eq. (15) can be simplified as

\[
V_{ji}(\mathbf{R}_j, \mathbf{R}_i) = \langle \phi_{ji}^{ja}(\mathbf{r}, \rho_A)|V_{1B}^{sp} + V_{2B}^{sp}|\phi_{ji}^{jB}(\mathbf{r}, \rho_B) \rangle
\]

\[
= \langle \phi_{ji}^{ja}(\mathbf{r}, \rho_A)|D_0 \delta(\rho_B) \phi_{nn}^B(\mathbf{r}) \rangle
\]

\[
= D_0 \langle \phi_{nn}^B(\mathbf{r})|\phi_{ji}^{ja}(\mathbf{r}, \rho_A) \rangle \delta(\rho_B)
\]

\[
= D_0 \langle \phi_{nn}^B(\mathbf{r})|\phi_{ji}^{ja}(\mathbf{r}, \rho_A) \rangle \delta(\beta \left( \mathbf{R}_j - \frac{A}{A+2} \mathbf{R}_i \right))
\]

since

\[
\mathbf{R}_j = \frac{A}{A+2} \mathbf{R}_i = \rho_B/\beta \quad \text{for} \quad \beta = \frac{2(A+B+2)}{(A+2)(B+2)}
\]

That is, we arrive at a ‘form factor’ \( \langle \phi_{nn}^B(\mathbf{r})|\phi_{ji}^{ja}(\mathbf{r}, -\mathbf{R}_j) \rangle \) that is local in \( \mathbf{R}_j = \frac{A}{A+2} \mathbf{R}_i = -\rho_A \) because of the delta function \( \delta(\rho_B) \). To find the form factor, we need to determine the average nucleon-nucleon relative wave function \( \phi_{nn}^B(\mathbf{r}) \) in the light ion, and project the heavy-nucleus two-body wave function \( \phi_{ji}^{ja}(\mathbf{r}, \rho_A) \) onto this relative motion. This gives a function only of the distance \( \rho_A = \mathbf{R}_j \) and the angles. The kinematics for this zero-range approximation are identical to those for the one-body transfer of a
Table 1. Two-neutron overlap function for $(^{122}\text{Sn}|^{124}\text{Sn})$

| State      | Value       |
|------------|-------------|
| $1g_{7/2}$ | 0.62944     |
| $2d_{5/2}$ | 0.59927     |
| $2d_{3/2}$ | 0.71913     |
| $3s_{1/2}$ | 0.51892     |
| $1h_{11/2}$ | $-1.24399$ |

mass-2 cluster from core $B$ to core $A$. A local-energy approximation may be used to improve the treatment of the finite range of the vertex function, just as for one-body transfers.

This makes clear the conclusion stated at the end of section 4, namely that transfer reactions only probe in the unknown nucleus those components of $nn$ relative motion that already exist in the known nucleus. Since the known light nuclei $^3\text{H}$, $^3\text{He}$ and $^4\text{He}$ have predominantly $s$-wave relative motion between the two transferred nucleons, our transfer reactions will only probe pairing states of $s$-wave relative motion in the target. The magnitude of the transfer cross section will be proportional to the form factor overlap $\langle \phi^B_{nn}(r)|\phi^A_J(r,\rho_A) \rangle$.

Zero-range approximations can be also used for some of the sequential steps involving these light nuclei, but not all of them if we are using ‘prior-post’ couplings to avoid non-orthogonality corrections. For stripping reactions such as $(t,p)$, the first prior $(t,d)$ step has no good zero-range approximation, and for pickup reactions such as $(p,t)$, the second post $(d,t)$ step must be treated in full finite range for the same reason.

7. Results

In this short paper we will focus on the reaction mechanisms for the pair transfer $^{124}\text{Sn}(p,t)^{122}\text{Sn}$ at 25 MeV, using the overlap function shown in Table 1 we find by overlapping the shell-model wave functions for the ground states of $^{122}\text{Sn}$ and $^{124}\text{Sn}$. The theory of these overlaps was explained by Cohen and Kurath for $p$-shell nuclei. We compare with the experimental data of Guazzoni et al.

We use the triton potential of Li, the deuteron potential of Daehnick, and the proton potential of Chapel Hill. All the two-neutron wave functions are constructed within the half-separation-energy approximation. For a triton wave function we use the pure $s^2$ configuration found by the product of eigenstates at the half-separation energy (4.24 MeV) in a Woods-
Saxon potential with $V = 77.83$ MeV, $R = 0.95$ fm, and $a = 0.65$ fm. The Sn wave functions shown in Table 1 are found at the half-separation energy (7.219 MeV) in a WS potential with $r = 1.17$ fm, and $a = 0.75$ fm that has the fixed spin-orbit component $V_{so} = 6.2$ MeV, $r = 1.01$ fm, and $a = 0.75$ fm.

The complete cross section prediction is shown in Fig. 1. We see the excellent agreement between theory and experiment as already published. Now we see that, with good shell-model overlaps and proper finite-range and sequential contributions, the unhappiness factors are much closer to unity.

To see the importance of the non-orthogonality terms, and hence of choosing ‘prior-post’ couplings if non-orthogonality terms are to be avoided, Fig. 2 plots the different sequential cross sections for all possible combinations of post and prior for the two steps. The prior-post solid curve is the dot-dashed curve in Fig. 1. The other curves are all different from this one, and cannot be simply added to the simultaneous amplitude to get the correct result. This also implies that no complete calculation with only zero-range couplings is possible.
Finally, it is instructive to look at the interference effects between the various simultaneous and sequential contributions. To display these coherence effects, I choose to plot the scattering amplitude at zero degrees for the non-spin-flip amplitude $m_p = m_t = 1/2$ (the only non-zero amplitude at this angle). Fig. 3 plots all the simultaneous and sequential contributions from the different components listed in Table 1, along with their coherent sums. We see that all the contributions to the simultaneous transfer are constructively coherent, as are all the contributions to the total sequential amplitude. This constructive coherence follows from the signs of the amplitudes in Table 1, and reflects the significant pairing enhancement in $^{124}$Sn. The total sequential and simultaneous amplitudes are not completely coherent with each other, however. This reflects the important of the deuteron channel with its own specific optical potential, and is also shown by their slightly different shapes of angular distributions in Fig. 1.

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Fig. 3. Simultaneous (short dash), sequential (dot-dash) and simultaneous+sequential (solid line) amplitudes at zero degrees for the reaction $^{124}\text{Sn}(p,t)^{122}\text{Sn}$ at 25 MeV. The short lines show the individual contributions from the wave function components of Table 1, and the longer lines with symbols are their coherent sums.

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