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TRANSFORMATION OF A NUCLEON-NUCLEON POTENTIAL OPERATOR INTO ITS SU(3) TENSOR FORM USING GPUS

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Abstract. Starting from the matrix elements of a nucleon-nucleon potential operator provided in a basis of spherical harmonic oscillator functions, we present an algorithm for expressing a given potential operator in terms of irreducible tensors of the SU(3) and SU(2) groups. Further, we introduce a GPU-based implementation of the latter and investigate its performance compared with a CPU-based version of the same. We find that the CUDA implementation delivers speedups of 2.27x – 5.93x.

1. Introduction. A major challenge in computational nuclear physics is to solve the Schrödinger equation for a system of nucleons interacting via realistic nuclear interactions that are consistent with the underlying theory of quantum chromodynamics. Configuration-interaction approaches convert the Schrödinger equation into a matrix eigenvalue problem. A Lanczos algorithm is then applied to find a few of the lowest-lying eigenvalues and eigenstates of the nuclear Hamiltonian matrix, whose elements are expressed in terms of a large many-nucleon basis that span a physically relevant subspace of the Hilbert space, the so-called model space.
The symmetry-adapted no-core shell model [2, 3] (SA-NCSM) is a configuration-interaction approach that employs group-theoretical methods based on the SU(3) and SU_S(2) symmetry groups (the subscript S denotes spin, see ahead) to reorganize the model space and nuclear interactions with respect to exact and partial symmetries of nuclei. In particular, the symmetry-adapted basis states of the SA-NCSM are classified according to their transformation properties with respect to the following group reduction chain:

\[
\text{SU}(3) \supset \text{SO}(3) \times \supset \text{SU}(2) \supset U(1) \supset \text{SU}_S(2)
\]

Note that the bottom line in (1) displays quantum numbers associated with a given symmetry group. Namely, a generic irreducible representation (irrep) of the SU(3) group is labeled by a pair of non-negative integers \((\lambda, \mu)\). The total orbital angular momentum \(L\), associated with the SO(3) subgroup of SU(3), is a good quantum number in this scheme. In addition, by coupling the orbital angular momentum \(L\) to the complementary spin degree of freedom \(S\), associated with the SU_S(2) group, one can achieve a classification scheme with good total angular momentum \(J = L + S\). The label \(M_J\) is the projection of \(J\) along the \(z\)-axis. The multiplicity label \(\kappa\) reflects the fact that multiple occurrences of \(L\) are possible within the \((\lambda, \mu)\) irrep of SU(3).

For a given \((\lambda, \mu)\) and \(L\) (all non-negative integers), the multiplicity index \(\kappa\) runs from 1 to \(\text{kmax}(\lambda, \mu, L)\) given by

\[
\text{kmax}(\lambda, \mu, L) = \max\left(0, \left\lfloor \frac{\lambda + \mu + 2 - L}{2} \right\rfloor \right) - \max\left(0, \left\lfloor \frac{\lambda + 1 - L}{2} \right\rfloor \right) - \max\left(0, \left\lfloor \frac{\mu + 1 - L}{2} \right\rfloor \right)
\]

with \(\lfloor \ldots \rfloor\) denoting the largest integer.

The essential input for the SA-NCSM calculation, which is unique to this theoretical framework, is a nucleon-nucleon potential operator \(\hat{V}\) expanded in terms of SU(3) and SU_S(2) irreducible tensors. Finding this expansion is a computationally intensive task. The main goal of this article is to introduce a GPU-based implementation of this task.

2. Expansion formula. The essential input for the SA-NCSM approach is a nucleon-nucleon potential operator \(\hat{V}\) expressed in terms of SU(3) and SU_S(2) irreducible tensors that are formed from the harmonic oscillator (HO) annihilation and creation operators,

\[
\hat{V} = \sum_{n_a n_b} \sum_{n_c n_d} \sum_{\lambda_f \mu_f} \sum_{\lambda_i \mu_i} \sum_{\lambda_0 \mu_0} \sum_{S_f S_i S_0 \kappa_0 \rho_0} C \left( n_a, n_b, (\lambda_f \mu_f), n_c, n_d, (\lambda_i \mu_i), (\lambda_0 \mu_0), S_f, S_i, S_0, \kappa_0, \rho_0 \right)
\]

with \(C\) denoting the coefficient.
Here $a_{\frac{1}{2}}^{(n,0)}$ represents an irreducible tensor labeled by $(n,0)$ and $\frac{1}{2}$ tensor characters with respect to the SU(3) and SU$_3$(2) groups. Components of this tensor are fermion creation operators acting in the HO shell $n$. Similarly, $a_{\frac{1}{2}}^{(0,n)}$ denotes an irreducible tensor whose components are related to fermion annihilation operators in HO shell $n$ [5]. The symbol $\times$ signifies SU(3) and SU(2) tensorial coupling. The square brackets $[\ldots]$ denote SU(3) and SU(2) irreducible tensor operators [5]. The irreducible tensor operators are labeled by $(\lambda_f \mu_f)$, $(\lambda_i \mu_i)$, and $(\lambda_0 \mu_0)$ quantum numbers given by the computationally inexpensive SU(3) coupling rules [6], which determine the decomposition of the Kronecker product of two SU(3) irreps in a direct sum of SU(3) irreps,

$$
(n_a 0) \times (n_b 0) = \sum_{\oplus} (\lambda_f \mu_f) \tag{4}
$$

$$
(0 n_d) \times (0 n_c) = \sum_{\oplus} (\lambda_i \mu_i) \tag{5}
$$

$$
(\lambda_f \mu_f) \times (\lambda_i \mu_i) = \sum_{\oplus} (\lambda_0 \mu_0). \tag{6}
$$

The symbol $\rho_0$ in (3) denotes a multiplicity label needed to distinguish between multiple occurrences of a $(\lambda_0 \mu_0)$ irrep in the Kronecker product $(\lambda_f \mu_f) \times (\lambda_i \mu_i)$. Irreps $(\lambda_i \mu_i)$ and $(\lambda_f \mu_f)$ in the Kronecker products (4) and (5) are always unique and no labels $\rho$ are needed. The irreducible tensors in the expansion (3) are also labeled by the spin quantum numbers $S_f, S_i, S_0$. For a two-body operator, they can take the following values $(S_f, S_i, S_0) \in \{(0, 0, 0) (0, 1, 1) (1, 1, 0) (1, 1, 1) (1, 1, 2)\}$.

Each term in the expansion (3) contains an irreducible tensor component multiplied by its strength (a real number) that can be computed as

$$
C (n_a, n_b, (\lambda_f \mu_f), n_c, n_d, (\lambda_i \mu_i), (\lambda_0 \mu_0), S_f, S_i, S_0, \kappa_0, \rho_0)
$$

$$
= (-1)^{S_f+S_i} \sqrt{(2S_i+1)(2S_f+1)(2S_0+1)}
$$

$$
\sum_{l_a, j_a; l_b, j_b; J} \langle n_a l_a j_a, n_b l_b j_b; J|\hat{V}|n_c l_c j_c, n_d l_d j_d; J\rangle
$$

$$
(-1)^{n_a+n_c-j_a-j_c} \sqrt{2J+1} \sqrt{(2j_a+1)(2j_b+1)} \sqrt{(2j_c+1)(2j_d+1)}
$$

$$
\sum_{L_i, l_f} \sqrt{(2L_i+1)(2L_f+1)} \left\{ \begin{array}{c}
L_f \\
l_f
\end{array} \right\} \left\{ \begin{array}{c}
l_f \\
L_i
\end{array} \right\} \left\{ \begin{array}{c}
l_c \\
L_i
\end{array} \right\} \left\{ \begin{array}{c}
l_c \\
L_f
\end{array} \right\}
$$

$$
\sum_{\kappa_0, \kappa_f} ((\lambda_f \mu_f) \kappa_f L_f) \langle (0 n_d) l_d; (0 n_c) l_c || (\lambda_i \mu_i) \kappa_i L_i
$$

$$
(\lambda_f \mu_f) \kappa_f L_f; (\lambda_i \mu_i) \kappa_i L_i || (\lambda_0 \mu_0) \kappa_0 S_0 \rangle_{\rho_0}. \tag{7}
$$
It is important to note that the input data for this calculation are provided in form of the matrix elements (real numbers) of a nuclear potential operator \( \hat{V} \) in two-nucleon \( J \)-coupled HO basis, \((n_d l_d j_d, n_b l_b j_b; J || n_c l_c j_c, n_d l_d j_d; J)\). The Kronecker delta \( \delta_{l_d l_b} \) and \( \delta_{l_c l_d} \) are equal to one for proton-proton and neutron-neutron matrix elements, and vanish for proton-neutron matrix elements. The factors in brackets are the Wigner 9j and 6j coefficients (real numbers). In the CPU version of the code, we adopt the \texttt{WIGXJF} library [4] for their accurate evaluation. The double-barred factors, denoted by symbol \((-; -; -)_-\), are SU(3) \( \supset \) SO(3) Wigner coefficients (real numbers) that are computed by optimized numerical subroutines [1]. Computing coupling coefficients on-the-fly is computationally expensive. To improve the algorithm performance, we store coefficients in fast lookup data structures (map from STL library).

3. **Algorithm.** As indicated by the summation order in relation (3), we iterate over combinations of HO shell numbers, \( n_a, n_b, n_c, n_d \) (line 2 in Algorithm 1). For each combination, we apply SU(3) coupling rules to find direct sum decompositions \( \sum \otimes (\lambda_f \mu_f) \) and \( \sum \otimes (\lambda_i \mu_i) \) (lines 3 and 4). Next, we iterate over SU(3) irreps in each decomposition (line 5 and line 7) and obtain the required SU(3) Wigner coefficients \( W^f_{(n_a, n_b, \lambda_f, \mu_f)} \) and \( W^i_{(n_a, n_c, \lambda_i, \mu_i)} \) (lines 6 and 8)

\[
\langle (n_0) 0_a; (n_0) 0_b || (\lambda_f \mu_f) \kappa_f L_f \rangle
\]
\[
\langle (0 n_d) l_d; (0 n_c) l_c || (\lambda_i \mu_i) \kappa_i L_i \rangle.
\]

Consequently, we apply SU(3) coupling rules for the Kronecker product \((\lambda_f \mu_f) \times (\lambda_i \mu_i)\) to obtain a direct sum decomposition \( \sum \otimes (\lambda_0 \mu_0) \) (line 9). For each irrep \((\lambda_0 \mu_0)\), we calculate the multiplicity \( \rho^\text{max}_0 \) (line 11) and obtain SU(3) Wigner coefficients \( W^0_{(\lambda_f, \mu_f, \lambda_i, \mu_i, \lambda_0, \mu_0)} \) (line 12)

\[
\langle (\lambda_f \mu_f) \kappa_f L_f; (\lambda_i \mu_i) \kappa_i L_i || (\lambda_0 \mu_0) \kappa_0 L_0 \rangle \rho_0.
\]

**Algorithm 1:** Determination of irreducible tensors

```plaintext
1   transformInteractionToSU3Tensors(inputData)
2   for each \((n_a, n_b, n_c, n_d)\) \in \text{inputData} do
3       \((n_0) 0_a \times (n_0) 0_b \rightarrow \sum \otimes (\lambda_f \mu_f)\)
4           \((0 n_d) l_d \times (0 n_c) l_c \rightarrow \sum \otimes (\lambda_i \mu_i)\)
5       for each \((\lambda_f \mu_f) \in \sum \otimes (\lambda_f \mu_f)\) do
6           obtain \(\langle (n_0) 0_a; (n_0) 0_b || (\lambda_f \mu_f) \kappa_f L_f \rangle \equiv W^f_{(n_a, n_b, \lambda_f, \mu_f)}\)
7           for each \((\lambda_i \mu_i) \in \sum \otimes (\lambda_i \mu_i)\) do
8           obtain \(\langle (0 n_d) l_d; (0 n_c) l_c || (\lambda_i \mu_i) \kappa_i L_i \rangle \equiv W^i_{(n_a, n_c, \lambda_i, \mu_i)}\)
9           \((\lambda_f \mu_f) \times (\lambda_i \mu_i) \rightarrow \sum \otimes (\lambda_0 \mu_0)\)
10          for each \((\lambda_0 \mu_0) \in \sum \otimes (\lambda_0 \mu_0)\) do
11             compute \(\rho_0^\text{max}\) for SU(3) coupling \((\lambda_f \mu_f) \times (\lambda_i \mu_i) \rightarrow (\lambda_0 \mu_0)\)
12             obtain \(\langle (\lambda_f \mu_f) \kappa_f L_f; (\lambda_i \mu_i) \kappa_i L_i || (\lambda_0 \mu_0) \kappa_0 L_0 \rangle \rho_0 \equiv W^0_{(\lambda_f, \mu_f, \lambda_i, \mu_i, \lambda_0, \mu_0)}\)
13             C(n_a, n_b, \lambda_f, \mu_f, n_c, n_d, \lambda_i, \mu_i, \lambda_0, \mu_0, \ldots) \leftarrow tensorStrengthCoeffs(n_a, n_b, n_c, n_d, inputData, W^f_{(\lambda_f)}, W^i_{(\lambda_i)}, W^0_{(\lambda_0), \rho_0^\text{max}})
14           end
15       end
16   end
17 end
```
The SU(3) Wigner coefficients are utilized in the last step (line 13), where tensor strengths
\[ C(n_a, n_b, (\lambda_f \mu_f), n_c, n_d, (\lambda_i \mu_i), (\lambda_0 \mu_0), S_f, S_i, S_0, \kappa_0, \rho_0) \]
are computed using relation (7) for all allowed spin combinations, \( S_f, S_i, S_0 \), as well as multiplicities \( \kappa_0 \) and \( \rho_0 \). For details see the Algorithm 2. Note that the first sum in (7) corresponds to the iteration over all two-body matrix elements with the given combination \( n_a, n_b, n_c, n_d \) of HO shell numbers (see line 3 in the Algorithm 2).

Algorithm 2: Computation of the tensor strength coefficients.

```plaintext
1 tensorStrengthCoeffs(n_a, n_b, n_c, n_d, inputData, W_f, W_i, W_0, \rho_0^{max})
2 results ← 0
3 for each V^{n_a,n_b,n_c,n_d}_{l_a,l_b,l_j,l_d} ∈ inputData do
4    for each (S_f, S_i, S_0) do
5        \( \kappa_0^{max} ← \max(\lambda_0, \mu_0, S_0) \)
6        for each \( L_f = |l_a - l_b| \ldots l_a + l_b \) do
7            \( C_f = \begin{pmatrix} l_a & l_b & L_f \\ 1 & 1 & S_f \\ j_a & j_b & J \end{pmatrix} \)
8            \( \kappa_f^{max} ← \max(\lambda_f, \mu_f, L_f) \)
9            for each \( L_i = |l_c - l_d| \ldots l_c + l_d \) do
10               \( C_i = \begin{pmatrix} l_c & l_d & L_i \\ 1 & 1 & S_i \\ j_c & j_d & J \end{pmatrix} \)
11               \( \kappa_i^{max} ← \max(\lambda_i, \mu_i, L_i) \)
12               for each \( \kappa_0 = 1, \ldots, \kappa_0^{max} \) do
13                  for each \( \rho_0 = 1, \ldots, \rho_0^{max} \) do
14                     results(S_f, S_i, S_0, \kappa_0, \rho_0) += results(S_f, S_i, S_0, \kappa_0, \rho_0) + V^{n_a,n_b,n_c,n_d}_{l_a,l_b,l_j,l_d} C_f C_i \)
15                     \((-1)^{S_f+S_i+S_0+L_i} \sqrt{(2S_i+1)(2S_f+1)(2S_0+1)(2L_i+1)(2L_f+1)} \)
16                     \( \sum_{\kappa_j=1}^{\kappa_j^{max}} \sum_{\lambda_j=1}^{\kappa_j} W_f(n_a,n_b,\lambda_f,\mu_f)(l_a,l_b,\kappa_f,L_f) W_i^{\dagger}(n_d,n_c,\lambda_i,\mu_i)(l_d,l_c,\kappa_i,L_i) \)
17                     \( W_0(\lambda_f,\mu_f,\lambda_i,\mu_i,\lambda_0,\rho_0) \)
18                  end
19              end
20          end
end
```

4. Implementation on GPUs. As there is currently no GPU-based code for calculating the required SU(3) Wigner coefficients, these and all additional input required by Algorithm 2 are determined using a CPU-based architecture.

Our first attempt to perform the Algorithm 2 on GPU was only parallelization of the loop over matrix elements (line 3 in Algorithm 2). In this approach, each CUDA thread stores its own private instance for resulting data and the parallel reduction across all threads was performed at the end of the calculation.
Unfortunately, such an approach does not provide enough parallelism to sustain a reasonable occupancy of the GPU. To overcome this we process all irreducible tensors at the same time. For this purpose, we first generate for given HO shell numbers \( n_a, n_b, n_c, n_d \) a complete set of all possible SU(3) quantum numbers of irreducible tensors, denoted as \( \{(\lambda_f \mu_f) (\lambda_i \mu_i) (\lambda_0 \mu_0)\} \), we also compute the required SU(3) coupling coefficients and store both in a buffer that is optimized for efficient transfer of data from CPU to GPU. In the next subsection we describe it in details.

4.1. \textbf{Irreducible tensors triplets buffers.} We preallocate arrays long enough on both CPU and GPU. During the computations on the CPU we treat the array on CPU like stacks while keeping pointers to particular data records. The data layout is depicted on Figure 1.

| Record Idx. | Wigner coefficients | Irreps |
|-------------|---------------------|--------|
| 1           | \( W^f(n_a,n_b,\lambda_f^i,\mu_f^i) \) | \( (\lambda_f^i,\mu_f^i) \) |
| 2           | \( W^f(n_c,n_d,\lambda_c^j,\mu_c^j) \) | \( (\lambda_c^j,\mu_c^j) \) |
| 3           | \( W^0(\lambda_f^i,\mu_f^i,\lambda_1^1,\mu_1^1,\lambda_0^0,\mu_0^0) \) | \( (\lambda_0^0,\mu_0^0) \) |
| 4           | \( W^0(\lambda_1^1,\mu_1^1,\lambda_1^2,\mu_1^2,\lambda_0^0,\mu_0^0) \) | \( (\lambda_0^0,\mu_0^0) \) |

... 

| \( i_1 \) | \( W^0(\lambda_f^i,\mu_f^i,\lambda_2^2,\mu_2^2,\lambda_0^0,\mu_0^0) \) | \( (\lambda_0^0,\mu_0^0) \) |
| \( i_1+1 \) | \( W^f(n_c,n_d,\lambda_c^j,\mu_c^j) \) | \( (\lambda_c^j,\mu_c^j) \) |
| \( i_1+2 \) | \( W^0(\lambda_f^i,\mu_f^i,\lambda_1^1,\mu_1^1,\lambda_0^0,\mu_0^0) \) | \( (\lambda_0^0,\mu_0^0) \) |
| \( i_1+3 \) | \( W^0(\lambda_1^1,\mu_1^1,\lambda_1^2,\mu_1^2,\lambda_0^0,\mu_0^0) \) | \( (\lambda_0^0,\mu_0^0) \) |

... 

| \( i_2 \) | \( W^0(\lambda_f^i,\mu_f^i,\lambda_2^2,\mu_2^2,\lambda_0^0,\mu_0^0) \) | \( (\lambda_0^0,\mu_0^0) \) |
| ... | ... | ... |
| \( i_3 \) | \( W^f(n_c,n_d,\lambda_c^j,\mu_c^j) \) | \( (\lambda_c^j,\mu_c^j) \) |
| \( i_3+1 \) | \( W^0(\lambda_f^i,\mu_f^i,\lambda_1^1,\mu_1^1,\lambda_0^0,\mu_0^0) \) | \( (\lambda_0^0,\mu_0^0) \) |
| \( i_3+2 \) | \( W^0(\lambda_1^1,\mu_1^1,\lambda_1^2,\mu_1^2,\lambda_0^0,\mu_0^0) \) | \( (\lambda_0^0,\mu_0^0) \) |

... 

| \( i_4 \) | \( W^0(\lambda_f^i,\mu_f^i,\lambda_2^2,\mu_2^2,\lambda_0^0,\mu_0^0) \) | \( (\lambda_0^0,\mu_0^0) \) |
| \( i_4+1 \) | \( W^f(n_c,n_d,\lambda_c^j,\mu_c^j) \) | \( (\lambda_c^j,\mu_c^j) \) |
| \( i_4+2 \) | \( W^0(\lambda_f^i,\mu_f^i,\lambda_1^1,\mu_1^1,\lambda_0^0,\mu_0^0) \) | \( (\lambda_0^0,\mu_0^0) \) |
| \( i_4+3 \) | \( W^0(\lambda_1^1,\mu_1^1,\lambda_1^2,\mu_1^2,\lambda_0^0,\mu_0^0) \) | \( (\lambda_0^0,\mu_0^0) \) |

... 

| \( i_{\text{max}} \) | \( W^0(\lambda_f^i,\mu_f^i,\lambda_2^2,\mu_2^2,\lambda_0^0,\mu_0^0) \) | \( (\lambda_0^0,\mu_0^0) \) |

\textbf{Figure 1.} Data layout of buffer used for transfer of the Wigner coupling coefficients and quantum numbers of irreducible tensors computed on CPU to GPU.

Superscripts \( N_f, N_i \) and \( N_0 \) denote number of irreps \( \sum_{a} (\lambda_f \mu_f) \), \( \sum_{a} (\lambda_i \mu_i) \) and \( \sum_{a} (\lambda_0 \mu_0) \) respectively. Each record has different size which is stored within the
Algorithm 3: Buffered determination of irreducible tensors

1. `transformInteractionToSU3TensorsWithBuffer(inputData)`
2. for each \((n_a, n_b, n_c, n_d) \in \text{inputData}\) do
3. allocate buffer \(B_W\) for Wigner coefficients and quantum numbers
4. allocate buffer \(B_{ref}\) for tuples of references into \(B_W\)
5. `recordIdx := 1`
6. \((n_a 0) \times (n_b 0) \rightarrow \sum_{\lambda f \mu f} (\lambda f \mu f)\)
7. \((0 n_d) \times (0 n_c) \rightarrow \sum_{\lambda i \mu i} (\lambda i \mu i)\)
8. for each \((\lambda f \mu f) \in \sum_{\lambda f} (\lambda f \mu f)\) do
9. obtain \((n_a 0)l_a; (n_b 0)l_b; (\lambda f \mu f) \kappa_f L_f \equiv W^f_{(n_a, n_b, \lambda f, \mu f)}\)
10. \(B_W \leftarrow \left( (\lambda f, \mu f), W^f_{(n_a, n_b, \lambda f, \mu f)} \right)\)
11. \(W_{idx}^f := \text{recordIdx}\)
12. `recordIdx + +`
13. for each \((\lambda f \mu f) \times (\lambda i \mu i) \rightarrow \sum_{\lambda f \mu f} (\lambda \mu 0)\) do
14. obtain \((0 n_a)l_a; (0 n_c)l_c; (\lambda i \mu i) \kappa_i L_i \equiv W^i_{(n_a, n_c, \lambda i, \mu i)}\)
15. \(B_W \leftarrow \left( (\lambda i, \mu i), W^i_{(n_a, n_c, \lambda i, \mu i)} \right)\)
16. \(W_{idx}^i := \text{recordIdx}\)
17. `recordIdx + +`
18. for each \((\lambda f \mu f) \times (\lambda i \mu i) \rightarrow \sum_{\lambda f \mu f} (\lambda \mu 0)\) do
19. compute \(p^\text{max}_0\) for SU(3) coupling \((\lambda f \mu f) \times (\lambda i \mu i) \rightarrow (\lambda \mu 0)\)
20. obtain \((\lambda f \mu f) \kappa_f L_f; (\lambda i \mu i) \kappa_i L_i; (\lambda \mu 0) \kappa_0 L_0 \equiv W^0_{(\lambda f, \lambda i, \mu f, \lambda i, \mu i, \lambda \mu 0)}\)
21. \(B_W \leftarrow \left( (\lambda \mu 0), W^0_{(n_a, n_b, \lambda \mu 0)} \right)\)
22. \(W_{idx}^0 := \text{recordIdx}\)
23. `recordIdx + +`
24. \(B_{ref} \leftarrow (W^f_{idx}, W^i_{idx}, W^0_{idx}, p^\text{max}_0)\)
25. end
26. end
27. end
28. transfer \(B_W\) and \(B_{ref}\) on GPU
29. `tensorStrengthCoeffsWithBuffer(n_a, n_b, n_c, n_d, inputData, B_W, B_{ref})`
30. end

Note that on the line 25 we create triplet of references (and \(p^\text{max}_0\)) to records holding appropriate combination

\[
\left( (\lambda f, \mu f), W^f_{(n_a, n_b, \lambda f, \mu f)}; (\lambda i, \mu i), W^i_{(n_a, n_b, \lambda i, \mu i)}; (\lambda \mu 0), W^0_{(n_a, n_b, \lambda \mu 0)}; p^\text{max}_0 \right)
\]
which we insert into a buffer $B_{ref}$. Each such triplet represents one irreducible tensor as depicted on Figure 2 (here we omit $\rho_0^{\text{max}}$). This buffer is transferred on GPU as well. The size of results $\text{results}(S_f, S_i, S_0, \kappa_0, \rho_0)$ computed on the line 14

| Record Idx. | Irreducible tensors references triplets |
|-------------|-----------------------------------------|
| 1           | (1, 2, 3)                               |
| 2           | (1, 2, 4)                               |
| $j_1$       | (1, $i_1$)                              |
| $j_1 + 1$   | (1, $i_1 + 1$, $i_1 + 2$)               |
| $j_1 + 2$   | (1, $i_1 + 1$, $i_1 + 3$)               |
| $j_2$       | (1, $i_1 + 1$, $i_2$)                   |
| $j_3$       | (1, $i_3$, $i_3 + 1$)                   |
| $j_3 + 1$   | (1, $i_3$, $i_3 + 2$)                   |
| $j_4$       | (1, $i_3$, $i_4$)                       |
| $j_4 + 1$   | ($i_4 + 1$, $i_4 + 2$, $i_4 + 3$)       |
| $j_{\text{max}}$ | ($i_4 + 1$, $i_4 + 2$, $i_{\text{max}}$) |

**Figure 2.** Data layout of buffer holding records with triplets of references to records in buffer $B_W$. Each such triplet thus represents one irreducible tensor.

in Algorithm 2 can change for different $(\lambda_0 \mu_0)$ since it depends on $\kappa_0^{\text{max}}$ and $\rho_0^{\text{max}}$. Therefore the results are managed in a similar way as $B_W$. We use preallocated array $B_R$ and for each tensor given by $\{(\lambda_f \mu_f)(\lambda_i \mu_i)(\lambda_0 \mu_0)\}$ we store related $\kappa_0$, $\rho_0$ a reference to $B_R$. Different CUDA blocks processing the same irreducible tensor have their own copies which are reduced on CPU at the end.

4.2. **Tensor strengths computation.** The computation of the tensor strengths is then performed on GPU using Algorithm 4. In this algorithm, each CUDA thread (of one dimensional CUDA blocks) takes one matrix element (of given combination $n_a, n_b, n_c, n_d$ of HO shell numbers, line 12) and one irreducible tensor (lines 13, 15, 19 and 23) and it performs summation over all combinations of quantum numbers $\{(\lambda_f \mu_f)(\lambda_i \mu_i)(\lambda_0 \mu_0)\}$ (lines 25–29). These combinations together with related Wigner coefficients are fetched from the buffer $B_W$ (lines 15, 19 and 23) via $B_{ref}$ (line 13). This leads to a much better GPU utilization as the number of CUDA threads needed is equal to the size of the $\{(\lambda_f \mu_f)(\lambda_i \mu_i)(\lambda_0 \mu_0)\}$ set multiplied by the number of two-body matrix elements for given values $n_a, n_b, n_c, n_d$ of HO shells. Finally, on the line 34, we perform parallel reduction inside CUDA blocks to sum-up contributions of all matrix elements mapped to given CUDA block. Intermediate results from particular CUDA blocks are at the end transferred on the host system to be summed-up over all matrix elements on CPU.
Algorithm 4: GPU computation of the tensor strength coefficients.

1. tensorStrengthCoeffsWithBuffer\( (n_a, n_b, n_c, n_d, inputData, BW, B_{ref}) \)
2. results \( \leftarrow 0 \), matrixElements = \([inputData(n_a, n_b, n_c, n_d)]\)
3. tensors = \([B_{ref}]\), matrixElementsBlocks = \([\text{matrixElements}/\text{blockSize}]\)
4. blocks = \(\text{matrixElementsBlock} \times \text{tensors}\)
5. create blocksCount blocks of CUDA threads
6. for each CUDA thread \( t \) do
7. tensorIdx = blockIdx/matrixElementsBlocks
8. matrixElementsBlockIdx = blockIdx \% matrixElementsBlocks
9. matrixElementIdx = matrixElementsBlockIdx \times \text{blockDim} + \text{threadIdx}
10. \( W_{idos}^f, W_{idr}^i, W_{idc}^0, \rho_{0\text{max}}^\text{ref} \) \( \leftarrow B_{ref}[\text{tensorIdx}] \)
11. for each \((S_f, S_i, S_0)\) do
12. \( \lambda_0, \mu_0 \) \( \leftarrow BW[W_{idos}^f] \)
13. \( \kappa_0^\text{max} \leftarrow \text{kmax}(\lambda_0, \mu_0, S_0) \)
14. for each \( L_f = |l_a - l_b| \ldots l_a + l_b \) do
15. \( C_f = \{ l_a, l_b, l_f \} \)
16. \( \lambda_j, \mu_j \leftarrow BW[W_{idr}^i] \)
17. \( \kappa_j^\text{max} \leftarrow \text{kmax}(\lambda_j, \mu_j, L_f) \)
18. for each \( L_i = |l_c - l_d| \ldots l_c + l_d \) do
19. \( C_i = \{ l_c, l_d, l_i \} \)
20. \( \lambda_j, \mu_j \leftarrow BW[W_{idc}^0] \)
21. \( \kappa_j^\text{max} \leftarrow \text{kmax}(\lambda_j, \mu_j, L_i) \)
22. for each \( \kappa_0 = 1, \ldots, \kappa_0^\text{max} \) do
23. for each \( \rho_0 = 1, \ldots, \rho_0^\text{max} \) do
24. \[ (-1)^{S_f+S_i+S_0+L_i} \sqrt{(2S_i+1)(2S_f+1)(2S_0+1)(2L_i+1)(2L_f+1)} \]
25. \[ \sum_{n_a=1}^{\kappa_0^\text{max}} \sum_{n_j=1}^{\kappa_j^\text{max}} W_{idos}^f(n_a, n_b, \lambda_f, \mu_f, L_f) W_{idr}^i(n_b, n_c, \lambda_i, \mu_i, L_i) W_{idc}^0(\lambda_j, \mu_j, \lambda_0, \mu_0, \kappa_0, \kappa_j, L_i, \kappa_i, S_0, \rho_0, \rho_0, \lambda_0, \mu_0) \]
26. end
27. end
28. end
29. for each \( \kappa_0 = 1, \ldots, \kappa_0^\text{max} \) and \( \rho_0 = 1, \ldots, \rho_0^\text{max} \) do
30. results\((S_f, S_i, S_0, \kappa_0, \rho_0, \text{matrixElementsBlockIdx}) = \)
31. reduce\((\text{results}(S_f, S_i, S_0, \kappa_0, \rho_0, t))\)
32. end
33. end
34. end

4.3. Multiple GPU parallelization. To support these computations on distributed memory parallel systems including multiple GPU systems, we implemented a parallel algorithm based on the MPI library. The loop over all \( n_a, n_b, n_c, n_d \) values of HO shells is distributed among collaborating MPI processes. As the runtime is
a strongly varying function of HO shell numbers, we use a single master process for distribution of the workload among collaborating processes. If multiple MPI processes run on a single computing node, they share its GPU resources, i.e. more MPI processes may access one GPU. We did not develop hybrid OpenMP/MPI algorithm since the amount of data shared between threads processing different HO shells would not be significant.

5. Performance results. To evaluate the performance, we conducted experiments using two different systems. The first one is the Blue Waters system, which is equipped with 4228 Cray XK7 nodes that consist of one 8-core AMD Opteron 6276 Interlagos Processor and one NVIDIA K20 GPU with 2688 CUDA cores, 6 GB of memory and 250 GB/s memory bandwidth. The second system is made up of a single node containing two 16-core IBM Power9 CPUs that are connected by high-speed interface NVLink to a single NVIDIA V100 GPU. This GPU has 5120 CUDA cores, 32 GB of memory and 900 GB/s memory bandwidth.

We measured the performance for different values of $N_{\text{max}}$ parameter that defines the size of two-nucleon Hilbert space in which the interaction matrix elements are provided. This means that for a given $N_{\text{max}}$ cutoff parameter, we are considering only two-nucleon matrix elements $\langle n_a l_a j_a , n_b l_b j_b ; J \| \hat{V} \| n_c l_c j_c , n_d l_d j_d ; J \rangle$ such that $0 \leq (n_a + n_b) \leq N_{\text{max}}$ and $0 \leq (n_c + n_d) \leq N_{\text{max}}$. In particular, $N_{\text{max}} = 8$ corresponds to 2,300,760 input two-nucleon matrix elements, $N_{\text{max}} = 10$ corresponds to 8,143,452 input two-nucleon matrix elements, $N_{\text{max}} = 12$ corresponds to 24,578,196 input two-nucleon matrix elements, and $N_{\text{max}} = 14$ corresponds to 65,615,742 input two-nucleon matrix elements.

Table 1 shows the performance results obtained on the Blue Waters system. The computations were carried out using 1, 2, 4, 8, 16, and 64 nodes, each with 8 MPI processes. One of all MPI processes was the master process assigning HO shells to other processes. The fourth column in Table 1 provides efficiency evaluated with respect to the number of nodes (we do not take the master process into account). Our master-slave workload distribution attains a very good scalability in particular for larger values of $N_{\text{max}}$, that is, for larger sets of interaction matrix elements. The last column in Table 1 shows speedups achieved by the GPU accelerated implementation compared to the CPU version of the code. We see that our GPU implementation delivers speedups ranging from 2.44 to 5.93. The speedups tend to decrease for increasing values of $N_{\text{max}}$ parameter. This is caused by the fact that increasing the $N_{\text{max}}$ parameter entails a rapid increase in the number of SU(3) coupling coefficients required. As a result, a more significant portion of the runtime is spent computing on CPUs. (This further suggests that a fresh look into the possibility of developing a parallel SU(3) package could lead to further gains.)

Table 2 shows the performance obtained on the Power9 system with 32 MPI processes. These results corroborate the performance behavior observed on the Blue Waters system.

6. Conclusion. We presented the algorithm for computing the expansion of a nucleon-nucleon potential operator in terms of the SU(3)×SU_{\text{3g}}(2) irreducible tensors. We described its implementation that combines MPI and CUDA parallelization. Enabling GPU using CUDA leads to a performance improvement. In particular, we achieved speedups ranging from 2.44 to 5.93 on Blue Waters, and almost a factor of 3 speedups for the system consisting of the two 16-core Power9 CPUs connected to a single NVIDIA V100 GPU.
| $N_{\text{max}}$ | MPI proc. | CPU only | CPU+GPU |
|-----------------|-----------|----------|---------|
|                 | Time [s]  | Efficiency | Time [s] | Speed-up |
| 8               | 295.3    | –         | 75.3    | 3.92     |
| 15+1            | 137.6    | 1         | 36.8    | 3.73     |
| 31+1            | 66.5     | 1         | 17.5    | 3.79     |
| 63+1            | 39.4     | 0.83      | 9.23    | 4.26     |
| 127+1           | 32.8     | 0.56      | 6.35    | 5.17     |
| 255+1           | 31.0     | 0.52      | 5.22    | 5.94     |
| 10              | 2219     | –         | 648     | 3.42     |
| 15+1            | 1034     | 1         | 318     | 3.24     |
| 31+1            | 499      | 1         | 151     | 3.28     |
| 63+1            | 248      | 0.99      | 74      | 3.32     |
| 127+1           | 165      | 0.74      | 43      | 3.75     |
| 255+1           | 138      | 0.59      | 32      | 4.25     |
| 12              | 13083    | –         | 4493    | 2.91     |
| 15+1            | 6097     | 1         | 2116    | 2.88     |
| 31+1            | 2943     | 1         | 1054    | 2.79     |
| 63+1            | 1447     | 1         | 515     | 2.80     |
| 127+1           | 776      | 0.92      | 269     | 2.88     |
| 255+1           | 565      | 0.68      | 169     | 3.33     |
| 14              | 64865    | –         | 26104   | 2.48     |
| 15+1            | 30227    | 1         | 12204   | 2.47     |
| 31+1            | 14596    | 1         | 5944    | 2.45     |
| 63+1            | 7179     | 1         | 2932    | 2.44     |
| 127+1           | 3581     | 0.99      | 1461    | 2.45     |
| 255+1           | 2142     | 0.83      | 838     | 2.55     |

Table 1. Performance results obtained on the Blue Waters system. Notation +1 in the column with MPI processes denotes one master process which just assigns HO shells to other MPI processes. The master process is not taken into account for efficiency evaluation.

| $N_{\text{max}}$ | CPU only | CPU+GPU |
|-----------------|----------|---------|
|                 | Time [s]  | Time [s] | Speed-up |
| 8               | 41.65    | 15.78   | 2.63     |
| 10              | 274.14   | 97.99   | 2.79     |
| 12              | 1649.7   | 611.1   | 2.69     |
| 14              | 7761.9   | 3407.6  | 2.27     |

Table 2. Performance results obtained for 32 MPI processes running on two 16-core IBM Power9 CPUs and a single NVIDIA V100 GPU.

The source code is available on Gitlab: [https://gitlab.com/oberhuber.tomas/np-interactions](https://gitlab.com/oberhuber.tomas/np-interactions).

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