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Quantum Monte Carlo for large chemical systems: Implementing efficient strategies for petascale platforms and beyond

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Various strategies to implement efficiently QMC simulations for large chemical systems are presented. These include: i.) the introduction of an efficient algorithm to calculate the computationally expensive Slater matrices. This novel scheme is based on the use of the highly localized character of atomic Gaussian basis functions (not the molecular orbitals as usually done), ii.) the possibility of keeping the memory footprint minimal, iii.) the important enhancement of single-core performance when efficient optimization tools are employed, and iv.) the definition of a universal, dynamic, fault-tolerant, and load-balanced framework adapted to all kinds of computational platforms (massively parallel machines, clusters, or distributed grids). These strategies have been implemented in the QMC=Chem code developed at Toulouse and illustrated with numerical applications on small peptides of increasing sizes (158, 434, 1056 and 1731 electrons). Using 10k–80k computing cores of the Curie machine (GENCI-TGCC-CEA, France) QMC=Chem has been shown to be capable of running at the petascale level, thus demonstrating that for this machine a large part of the peak performance can be achieved. Implementation of large-scale QMC simulations for future exascale platforms with a comparable level of efficiency is expected to be feasible.

I. INTRODUCTION

Quantum Monte Carlo (QMC) is a generic name for a large class of stochastic approaches solving the Schrödinger equation by using random walks. In the last forty years they have been extensively used in several fields of physics including nuclear physics,[1] condensed-matter physics,[2] spin systems,[3] quantum liquids,[4] infrared spectroscopy,[5, 6] etc. In these domains QMC methods are usually considered as routine methods and even in most cases as state-of-the-art approaches. In sharp contrast, this is not yet the case for the electronic structure problem of quantum chemistry where QMC[7, 8] is still of confidential use when compared to the two well-established methods of the domain (Density Functional Theory (DFT) and post-Hartree-Fock methods). Without entering into the details of the forces and weaknesses of each approach, a major limiting aspect of QMC hindering its diffusion is the high computational cost of the simulations for realistic systems.

However — and this is the major concern of this work — a unique and fundamental property of QMC methods is their remarkable adaptation to High Performance Computing (HPC) and, particularly, to massively parallel computations. In short, the algorithms are simple and repetitive, central memory requirements may be kept limited whatever the system size, and I/O flows are negligible. As most Monte Carlo algorithms, the computational effort is almost exclusively concentrated on pure CPU (“number crunching method”) and the execution time is directly proportional to the number of Monte Carlo steps performed. In addition, and this is a central point for massive parallelism, calculations of averages can be decomposed at will: \( n \) Monte Carlo steps over a single processor being equivalent to \( n/p \) Monte Carlo steps over \( p \) processors with no communication between the processors (apart from the initial/final data transfers). Once the QMC algorithm is suitably implemented the maximum gain of parallelism (ideal scalability) should be expected.

A most important point is that mainstream high-level quantum chemistry methods do not enjoy such a remarkable property. Essentially based on iterative schemes defined with the framework of linear algebra they involve the manipulation and storage of extremely large matrices and their adaptation to extreme parallelism and low-memory-footprint implementation is intrinsically problematic.

Now, in view of the formidable development of computational platforms, particularly in terms of number of computing cores (presently up to a few hundreds of thousands and many more to come) the practical bottleneck associated with the high computational cost of QMC is expected to become much less critical and thus QMC may become in the coming years a method of practical use for treating chemical problems out of the reach of present-day approaches. Following this line of thought a number of QMC groups are presently working on implementing strategies allowing their QMC codes to run efficiently on very large-scale parallel computers.[9–11] Essentially, most strategies rely on massive parallelism and on some efficient treatment (“linear-scaling”-type algorithms) for dealing with the matrix computations and manipulations that represent the most CPU-expensive part of the algorithm.

Here, we present several strategies implemented in the QMC=Chem code developed in our group at the University of Toulouse.[12] A number of actual simulations realized on the Curie machine at the French GENCI-
TGCC-CEA computing center with almost ideal parallel efficiency in the range 10 000–80 000 cores and reaching the petascale level have been realized.

The contents of this paper is as follows. In The first section, a brief account of the QMC method employed is presented. Only those aspects essential to the understanding of the computational aspects discussed in this work are given. In second section, the problem of computing efficiently the Slater matrices at the heart of the QMC algorithm (computational hot spot) is addressed. A novel scheme taking advantage of the highly-localized character of the atomic Gaussian basis functions (not the molecular orbitals as usually done) is proposed. A crucial point is that the approach is valid for an arbitrary molecular shape (e.g. compact molecules), there is no need of considering extended or quasi-one-dimensional molecular systems as in linear-scaling approaches. The third section discusses the overall performance of the code and illustrates how much optimizing the single-core performance of the specific processor at hand can be advantageous. The fourth section is devoted to the way our massively parallel simulations are deployed on a general computational platform and, particularly, how fault-tolerance is implemented, a crucial property for any large-scale simulation. Finally, a summary of the various strategies proposed in this work is presented in the last section.

II. THE QUANTUM MONTE CARLO METHOD

In this work we shall consider a variant of the Fixed-Node Diffusion Monte Carlo (FN-DMC) approach, the standard quantum Monte Carlo method used in computational chemistry. Here, we shall insist only on the aspects needed for understanding the rest of the work. For a complete presentation of the FN-DMC method the reader is referred, e.g to [2],[7], or [8] and references therein.

A. Fixed-node diffusion Monte Carlo (FN-DMC)

Diffusion Monte Carlo. In a diffusion Monte Carlo scheme, a finite population of “configurations” or “walkers” moving in the $3N$-dimensional space ($N$, number of electrons) is introduced. A walker is described by a $3N$-dimensional vector $\mathbf{R} \equiv (r_1, r_2, ..., r_N)$ giving the positions of the $N$ electrons. At each Monte Carlo step each walker of the population is diffused and drifted according to

$$\mathbf{R}’ = \mathbf{R} + \tau \mathbf{b} (\mathbf{R}) + \sqrt{2} \eta$$

where $\tau$ is a small time-step, $\eta$ a Gaussian vector ($3N$ independent normally distributed components simulating a free Brownian diffusion), and $\mathbf{b} (\mathbf{R})$ the drift vector given by

$$\mathbf{b} (\mathbf{R}) = \frac{\nabla \psi_T (\mathbf{R})}{\psi_T (\mathbf{R})} \tau$$

where $\psi_T$, the trial wave function, is a known computable approximation of the exact wavefunction. At the end of this drift/diffusion step each walker is killed, kept unchanged, or duplicated a certain number of times proportionally to the branching weight $w$ given by

$$w = e^{-\frac{\tau}{2} [(E_L(\mathbf{R}’) - E_T) + (E_L(\mathbf{R}) - E_T)]]}$$

where $E_T$ is some reference energy and $E_L$ the local energy defined as

$$E_L(\mathbf{R}) \equiv \frac{H \psi_T (\mathbf{R})}{\psi_T (\mathbf{R})} \tau$$

The population is propagated and after some equilibrium time it enters a stationary regime where averages are evaluated. As an important example, the exact energy may be obtained as the average of the local energy.

The Fixed-Node approximation. Apart from the statistical and the short-time (finite time-step) errors which can be made arbitrary small, the only systematic error left in a DMC simulation is the so-called Fixed-Node (FN) error. This error results from the fact that the nodes of the trial wavefunction [defined as the $(3N-1)$-dimensional hyper-surface where $\Psi_T (\mathbf{R}) = 0]$ act as infinitely repulsive barriers for the walkers [divergence of the drift vector, Eq.(2)]. Each walker is thus trapped forever within the nodal pocket delimited by the nodes of $\Psi_T$ where it starts from. When the nodes of $\psi_T$ coincide with the exact nodes, the algorithm is exact. If not, a variational fixed-node error is introduced. However, with the standard trial wavefunctions used, this error is in general small,[13] a few percent of the correlation energy for total energies.

B. Parallelizing FN-DMC

Each Monte Carlo step is carried out independently for each walker of the population. The algorithm can thus be easily parallelized over an arbitrary number of processors by distributing the walkers among the processors, but doing this implies synchronizations of the CPUs since the branching step requires that all the walkers have first finished their drifted-diffusion step.

To avoid this aspect, we have chosen to let each CPU core manage its own population of walkers without any communication between the populations. On each computing unit a population of walkers is propagated and the various averages of interest are evaluated. At the end of the simulation, the averages obtained on each processor are collected and summed up to give the final answers. Regarding parallelism the situation is thus ideal since, apart from the negligible initial/final data transfers, there are no communications among processors.

The only practical problem left with FN-DMC is that the branching process causes fluctuations in the population size and thus may lead to load-balancing problem.
among processors. More precisely, nothing prevents the population size from decreasing or increasing indefinitely during the Monte Carlo iterations. To escape from this, a common solution consists in forcing the number of walkers not to deviate too much from some target value for the population size by introducing a population control step. It is usually realized by monitoring in time the value of the reference energy $E_T$ via a feedback mechanism, see e.g. [14]. The price to pay is the introduction of some transient load imbalances and inter-processor communications/synchronization to redistribute walkers among computing cores, inevitably degrading the parallel speed-up. This solution has been adapted by several groups and some tricks have been proposed to keep this problem under control.[9–11, 15]

Here, we propose to avoid this problem directly from the beginning by employing a variant of the FN-DMC working with a constant number of walkers. Several proposals can be found in the literature, e.g. [16, 17]. Here, we shall employ the method described in Ref.[17]. In this approach the branching step of standard DMC is replaced by a so-called reconfiguration step. Defining the normalized branching weights as follows

$$p_k = \frac{w_k}{\sum_{i=1}^{M} w_i}$$ (5)

the population of walkers is “reconfigured” by drawing at each step $M$ walkers among the $M$ walkers according to the probabilities $p_k$. At infinite population, this step reduces to the standard branching step where walkers are deleted or duplicated proportionally to the weight $w$. At finite $M$ the normalization factor $\sum_{i=1}^{M} w_i$ is no longer constant and a finite-population bias is introduced. To remove this error and recover the exact averages, a global weight given as the product of the total population weights for all preceding generations must be included into the averages. Note that this algorithm enables the possibility to use small walker populations on each core since there is no finite-population bias (typically, we use 10 to 100 walkers per core). For all details the reader is referred to [17].

C. Critical CPU part

At each Monte Carlo step the CPU effort is almost completely dominated by the evaluation of the wavefunction $\Psi_T$ and its first and second derivatives (computational hot spot). More precisely, for each walker the values of the trial wavefunction, $\Psi_T$, its first derivatives with respect to all 3N-coordinates [drift vector, Eq.(2)], and its Laplacian $\nabla^2 \Psi_T$ [kinetic part of the local energy, Eq.(4)] are to be calculated. It is essential that such calculations be as efficient as possible since in realistic applications their number may be very large (typically of the order of $10^9 - 10^{12}$).

A common form for the trial wavefunction is

$$\Psi_T(\mathbf{r}) = e^{J(\mathbf{R})} \sum_{K=(K_1,K_2)} c_K \text{Det}_{K_1}(\mathbf{r}_1, ..., \mathbf{r}_{N_1}) \text{Det}_{K_2}(\mathbf{r}_{N_1+1}, ..., \mathbf{r}_{N}).$$ (6)

where the electron coordinates of the $N_1$ (resp. $N_2$) electrons of spin $\uparrow$ (resp. $\downarrow$) have been distinguished, $N = N_1 + N_2$. In this formula $e^{J(\mathbf{R})}$ is the Jastrow factor describing explicitly the electron-electron interactions at different levels of approximations. A quite general form may be written as

$$J(\mathbf{R}) = \sum_\alpha U^{(e-n)}(r_{i\alpha}) + \sum_{i,j} U^{(e-c)}(r_{ij}) + \sum_{\alpha i,j} U^{(e-c-n)}(r_{ij}, r_{i\alpha}, r_{j\alpha}) + ...$$ (7)

where $r_{ij} = |\mathbf{r}_i - \mathbf{r}_j|$ is the inter-electronic distance and $r_{i\alpha} = |\mathbf{r}_i - \mathbf{Q}^{\alpha}|$, the distance between electron $i$ and nucleus $\alpha$ located at $\mathbf{Q}^{\alpha}$. Here $U$’s are simple functions and various expressions have been employed in the literature. The Jastrow factor being essentially local, short-ranged expressions can be employed and the calculation of this term is usually a small contribution to the total computational cost. As a consequence, we shall not discuss further the computational aspect of this term here.

The second part of the wavefunction describes the shell-structure in terms of single-electron molecular orbitals and is written as a linear combination of products of two Slater determinants, one for the $\uparrow$ electrons and the other for the $\downarrow$ electrons. Each Slater matrix is built from a set of molecular orbitals $\phi_i(\mathbf{r})$ usually obtained from a preliminary DFT or SCF calculations. The $N_{\text{orb}}$ molecular orbitals (MOs) are expressed as a sum over a finite set of $N_{\text{basis}}$ basis functions (atomic orbitals, AOs)

$$\phi_i(\mathbf{r}) = \sum_{j=1}^{N_{\text{basis}}} a_{ij} \chi_j(\mathbf{r})$$ (8)

where the basis functions $\chi_j(\mathbf{r})$ are usually expressed as a product of a polynomial and a linear combination of Gaussian functions. In the present work the following standard form is employed

$$\chi(\mathbf{r}) = (x - Q_x)^{n_x} (y - Q_y)^{n_y} (z - Q_z)^{n_z} g(\mathbf{r})$$ (9)
with
\[ g(\mathbf{r}) = \sum_k c_k e^{-\gamma_k (\mathbf{r} - \mathbf{Q})^2}. \] (10)

Here \( \mathbf{Q} = (Q_x, Q_y, Q_z) \) is the vector position of the nucleus-center of the basis function, \( \mathbf{n} = (n_x, n_y, n_z) \) a triplet of positive integers, \( g(\mathbf{r}) \) is the spherical Gaussian component of the AO, and \( \gamma_k \) its exponents. The determinants corresponding to spin \( \uparrow \)-electrons are expressed as
\[ \text{Det}_{K_\uparrow}(\mathbf{r}_1, ..., \mathbf{r}_{N_\uparrow}) = \text{Det} \begin{pmatrix} \phi_{i_1}(\mathbf{r}_1) & \cdots & \phi_{i_{N_\uparrow}}(\mathbf{r}_{N_\uparrow}) \\ \vdots & \ddots & \vdots \\ \phi_{i_{N_\uparrow}}(\mathbf{r}_1) & \cdots & \phi_{i_{N_\uparrow}}(\mathbf{r}_{N_\uparrow}) \end{pmatrix} \] (11)
where \( K_\uparrow \) is a compact notation for denoting the set of indices \( \{i_1, ..., i_{N_\uparrow}\} \) specifying the subset of the molecular orbitals used for this particular Slater matrix. A similar expression is written for spin \( \downarrow \)-electrons.

In contrast to the calculation of the Jastrow factor, the evaluation of the determinantal part of the wavefunction and its derivatives is critical. To perform such calculations we employ a standard approach\[7\] consisting in calculating the matrices of the first and second (diagonal) derivatives of each molecular orbital \( \phi_i \) with respect to the three space variables \( l = x, y, z \) evaluated for each electron position \( \mathbf{r}_j \), namely
\[ D_{i,j}^{(1)} = \frac{\partial \phi_i(\mathbf{r}_j)}{\partial x_l} \] (12)
\[ D_{i,j}^{(2)} = \frac{\partial^2 \phi_i(\mathbf{r}_j)}{\partial x_l^2} \] (13)
and then computing the inverse \( D^{-1} \) of the Slater matrices defined as \( D_{ij} = \phi_i(\mathbf{r}_j) \). The drift components and the Laplacian corresponding to the determinantal part of the trial wavefunction are thus evaluated as simple vector-products
\[ \frac{1}{\text{Det}(\mathbf{R})} \frac{\partial \text{Det}(\mathbf{R})}{\partial x_l^i} = \sum_{j=1,N} D_{i,j}^{(1)} D_{j}^{-1} \] (14)
\[ \frac{1}{\text{Det}(\mathbf{R})} \frac{\partial^2 \text{Det}(\mathbf{R})}{\partial x_l^i} = \sum_{j=1,N} D_{i,j}^{(2)} D_{j}^{-1} \] (15)

From a numerical point of view, the computational time \( T \) needed to evaluate such quantities as a function of the number of electrons \( N \) scales as \( \mathcal{O}(N^3) \)
\[ T = \alpha N^3 + \beta N^3. \] (16)

The first \( N^3 \)-term results from the fact that the \( N^2 \) matrix elements of the Slater matrices are to be computed, each element being expressed in terms of the \( N_{\text{basis}} \sim N \) basis functions needed to reproduce an arbitrary delocalized molecular orbital. The second \( N^2 \)-term is associated with the generic cubic scaling of any linear algebra method for inverting a general matrix.

III. EXPLOITING THE HIGHLY LOCALIZED CHARACTER OF ATOMIC BASIS FUNCTIONS

As seen in the previous section, one of the two computational hot spots of QMC is the calculation of the derivatives of the determinantal part of the trial wave function for each electronic configuration \( (\mathbf{r}_1, ..., \mathbf{r}_N) \) at each Monte Carlo step. To be more precise, the \( N \text{orb} \) molecular orbitals (MO) used in the determinantal expansion (6) are to be computed (here, their values will be denoted as \( C_1 \)) together with their first derivatives with respect to \( x \), \( y \), and \( z \) (denoted \( C_2, C_3, C_4 \)) and their Laplacians (denoted \( C_5 \)). Calculations are made in single precision using an efficient matrix product routine we describe now. The matrix products involve the matrix of the AO coefficients \( a_{ij} \), Eq.(8) (here denoted as \( A \)) the matrix of the atomic Gaussian basis functions evaluated at all electronic positions, \( \chi_i(\mathbf{r}) \) (denoted \( B_1 \)), their first derivatives (denoted \( B_2, B_3, B_4 \)) and Laplacians (denoted \( B_5 \)). The five matrix products are written under the convenient form
\[ C_i = AB_i \quad i = 1, 5 \] (17)
Note that matrix \( A \) remains constant during the simulation while matrices \( B \) and \( C \) depend on electronic configurations. The matrix sizes are as follows: \( N \text{orb} \times N \) for the \( C_1 \)'s, \( N \text{orb} \times N_{\text{basis}} \) for \( A \), and \( N_{\text{basis}} \times N \) for \( B \). In practical applications \( N \text{orb} \) is of the order of \( N \) while \( N_{\text{basis}} \) is greater than \( N \) by a factor 2 or 3 for standard calculations and much more when using high-quality larger basis sets. The expensive part is essentially dominated by the \( N_{\text{basis}} \) multiplications. The total computational effort is thus of order \( N \text{orb} \times N \times N_{\text{basis}}, \text{i.e.} \sim \mathcal{O}(N^3) \).

The standard approach proposed in the literature for reducing the \( N^3 \)-price is to resort to the so-called linear-scaling or \( \mathcal{O}(N) \)-techniques.\[18–23\] The basic idea consists in introducing spatially localized molecular orbitals instead of the standard delocalized (canonical) ones obtained from diagonalization of reference Hamiltonians (usually, Hartree-Fock or Kohn-Sham). Since localized orbitals take their value in a finite region of space — usually in the vicinity of a fragment of the molecule — the number of basis set functions \( N_{\text{basis}} \) needed to represent them with sufficient accuracy becomes essentially independent of the system size (not scaling with \( N \) as in the case of canonical ones). In addition to this, each electron contributes only to a small subset of the localized orbitals (those non-vanishing in the region where the electron is located). As a consequence, the number of non-vanishing matrix elements of the \( C_i \) matrices no longer scales as \( N \text{orb} \times N \sim N^2 \) but linearly with \( N \). Furthermore, each matrix element whose computation was proportional to the number of basis set used, \( N_{\text{basis}} \sim N \), is now calculated in a finite time independent of the system size. Putting together these two results, we are led to a linear dependence of the computation of the \( C_i \) matrices upon the number of electrons.
Here, we choose to follow a different path. Instead of localizing the canonical molecular orbitals we propose to take advantage of the localized character of the underlying atomic Gaussian basis set functions. The advantages are essentially three-fold:

1. the atomic basis set functions are naturally localized independently of the shape of the molecule. This is the most important point since the localization procedures are known to be effective for chemical systems having a molecular shape made of well-separated sub-units (for example, linear systems) but much less for general compact molecular systems that are ubiquitous in chemistry.

2. the degree of localization of the standard atomic Gaussian functions is much larger than that obtained for molecular orbitals after localization (see results below)

3. by using the product form, Eq.(17), the localized nature of the atomic Gaussian functions can be exploited very efficiently (see next section).

In practice, when the value of the spherical Gaussian part \( g(r) \) of an atomic orbital function \( \chi(r) \) is smaller than a given threshold \( \epsilon = 10^{-8} \), the value of the AO, its gradients and Laplacian are considered null. This property is used to consider the matrices \( B_1, \ldots, B_5 \) as sparse. However, in contrast with linear-scaling approaches, the MO matrix \( A \) is not considered here as sparse. We shall come back to this point later. To accelerate the calculations, an atomic radius is computed as the distance beyond which all the Gaussian components \( g(r) \) of the atomic orbitals \( \chi(r) \) centered on the nucleus are less than \( \epsilon \). If an electron is farther than the atomic radius, all the AO values, gradients and Laplacians centered on the nucleus are set to zero.

The practical implementation to perform the matrix products is as follows. For each electron, the list of indices (array indices in what follows) where \( g(r) > 0 \) is calculated. Then, the practical algorithm can be written as

\[
\begin{align*}
C_1 &= 0. \\
C_2 &= 0. \\
C_3 &= 0. \\
C_4 &= 0. \\
C_5 &= 0.
\end{align*}
\]

```plaintext
do i=1, Number of electrons
  do k=1, Number of non-zero AOs for electron i
    do j=1, Number of molecular orbitals
      C1(j,i) += A(j,indices(k,i))*B1(k,i)
      C2(j,i) += A(j,indices(k,i))*B2(k,i)
      C3(j,i) += A(j,indices(k,i))*B3(k,i)
      C4(j,i) += A(j,indices(k,i))*B4(k,i)
      C5(j,i) += A(j,indices(k,i))*B5(k,i)
  end do
end do
```

(where \( x += y \) denotes \( x = x + y \)).

This implementation allows to take account of the sparsity of the \( B \) matrices, while keeping the efficiency due to a possible vectorization of the inner loop. The load/store ratio is 6/5 (6 load-from-memory instructions, 5 store-to-memory instructions) in the inner loop: the elements of \( B_n \) are constant in the inner loop (in registers), and the same element of \( A \) is used at each line of the inner loop (loaded once per loop cycle). As store operations are more expensive than load operations, increasing the load/store ratio improves performance as will be shown in the next section. Using this algorithm, the scaling of the matrix products is expected to drop from \( \mathcal{O}(N^3) \) to a scaling roughly equal to \( \mathcal{O}(N^2) \) (in a regime where \( N \) is large enough, see discussion in the next section). Let us now illustrate such a property in the applications to follow.

The different systems used here as benchmarks are represented in Figure 1. The trial wavefunctions used for describing each system are standard Hartree-Fock wavefunctions (no Jastrow factor) with molecular orbitals expressed using various Gaussian basis sets. System 1 is a copper complex with four ligands having 158 electrons and described with a cc-pVDZ basis set. System 2 is a polypeptide taken from reference[24] (434 electrons and 6-31G* basis set). System 3 is identical to System 2 but using a larger basis set, namely the cc-pVTZ basis set. System 4 is the 1ZE7 molecule from the Protein Data Bank (1056 electrons, 6-31G*). System 5 is the 1AMB molecule from the Protein Data Bank (1731 electrons, 6-31G*).

Table IV shows the level of sparsity of the matrices \( A \) (\( A_{ij} \equiv a_{ij} \)) and \( B_i \) (\( B_{1ij} \equiv \chi_i(r_j) \)) for the five systems (matrices \( B_n \) with \( n > 1 \) behave as \( B_1 \) with respect to sparsity). As seen the number of basis set functions employed is proportional to the number of electrons with a factor ranging from about 2.2 to 6.8.

Regarding the matrix \( A \) of MO coefficients the results
are given both for standard canonical (delocalized) MOs and for localized orbitals. To get the latter ones, different localization schemes have been applied.  

However, they essentially lead to similar results. Here, the results presented are those obtained by using the Cholesky decomposition of the density matrix expressed in the AO basis set. As seen the level of sparsity of the matrix $A$ is low. Although it increases here with the system size it remains modest for the largest size (there are still about one third of non-zero elements). Of course, such a result strongly depends on the type of molecular system considered (compact or not compact) and on the diffuse character of the atomic basis set. Here, we have considered typical systems of biochemistry.

Next, the level of sparsity of the $B$ matrices is illustrated. The percentage of non-zero values of $\chi_i(r_j)$ has been obtained as an average over a Variational Monte Carlo run. In sharp contrast with MOs the atomic orbitals are much more localized, thus leading to a high level of sparsity. For the largest system, only 3.9% of the basis function values are non-negligible.

In the last column of the table the maximum number of non-zero elements obtained for all columns of the matrix (i.e., a value of electron position) during the entire Monte Carlo simulation is given. A first remark is that this number is roughly constant for all system sizes. A second remark is that the percentage of non-zero values is only slightly greater than the average, thus showing that the $B$ matrices can be considered sparse during the whole simulation, not only in average. As an important consequence, the loop over the number of non-zero AOIs for each electron in the practical algorithm presented above (loop over $k$ index) is expected to be roughly constant as a function of the size at each Monte Carlo step. This latter remark implies for this part an expected behavior of order $O(N^2)$ for large $N$. Let us now have a closer look at the actual performance of the code.

IV. OVERALL PERFORMANCE OF QMC$\equiv$CHEM

When discussing performance several aspects must be considered. A first one, which is traditionally discussed, is the formal scaling of the code as a function of the system size $N$ ($N$ ~ number of electrons). As already noted, due to the innermost calculation, products, and inversion of matrices, such a scaling is expected to be cubic, $O(N^3)$. However, there is a second important aspect, generally not discussed, which is related to the way the expensive innermost floating-point operations are implemented and on how far and how efficiently the potential performance of the processor at hand is exploited. In what follows we shall refer to this aspect as “single-core optimization”. It is important to emphasize that such an aspect is by no way minor and independent on the previous “mathematical” one.

To explicit this point, let us first recall that the computational time $T$ results essentially from two independent parts, the first one resulting from the computation of the matrix elements, $T_1 \sim \alpha N^3$ and the second one from the inversion of the Slater matrix, $T_2 \sim \beta N^3$. Now, let us imagine that we have been capable of devising a highly efficient linear-scaling algorithm for the first contribution such that $T_1 \sim \epsilon N \ll T_2$ within the whole range of system sizes $N$ considered. We would naturally conclude that the overall computational cost $T \sim T_2$ is cubic. In the opposite case where a very inefficient linear-scaling algorithm is employed for the first part, $T \sim T_1 \gg T_2$, we would conclude to a linear-scaling type behavior. Of course, mathematically speaking such a way of reasoning is not correct since scaling laws are only meaningful in the asymptotic regime where $N$ goes to infinity. However, in practice only a finite range of sizes is considered (here, between 2 and about 2000 active electrons) and it is important to be very cautious with the notion of scaling laws. A more correct point of view consists in looking at the global performance of the code in terms of total CPU time for a given range of system sizes, a given compiler, and a given type of CPU core.

Finally, a last aspect concerns the memory footprint of the code whose minimization turns out to be very advantageous. Indeed, the current trend in supercomputer design is to increase the number of cores more rapidly than the available total memory. As the amount of memory per core will continue to decrease, it is very likely that programs will need to have a low memory footprint to take advantage of exascale computers. Another point is that when less memory is used less electrical power is needed to perform the calculation: data movement from the memory modules to the cores needs more electrical power than performing floating point operations. Although at present time the power consumption is not yet a concern to software developers, it is a key aspect in present design of the exascale machines to come.

In this section, the results discussed will be systematically presented by using two different generations of Intel Xeon processors. The first processor, referred to as Core2$^2$, is an Intel Xeon 5140, Core2 2.33 GHz, Dual core, 4 MiB shared L2 cache. The second one, referred to as Sandy Bridge, is an Intel Xeon E3-1240 at 3.30 GHz, Quad core, 256 KiB L2 cache/core, 8 MiB shared L3 cache (3.4 GHz with turbo). Note also that the parallel scaling of QMC being close to ideal (see next section), single-core optimization is very interesting: the gain in execution time obtained on the single-core executable is directly transferred to the whole parallel simulation.

A. Improving the innermost expensive floating-point operations

For the Core2 architecture, the practical algorithm presented above may be further improved by first using the unroll and jam technique, which consists in unrolling the outer loop and merging multiple outer-loop iterations
in the inner loop:

```fortran
  do i=1, Number of electrons
    do k=1, Number of non-zero AOs for electron i, 2
      do j=1, Number of molecular orbitals
        C1(j,i) += A(j,indices(k,i))*B1(k,i) + &  
                    A(j,indices(k+1,i))*B1(k+1,i)
        C2(j,i) += A(j,indices(k,i))*B2(k,i) + &  
                    A(j,indices(k+1,i))*B2(k+1,i)
      end do
    end do
  end do
```

To avoid register spilling, the inner loop is split in two loops: one loop computing $C_1, C_2, C_3$ and a second loop computing $C_4, C_5$. The load/store ratio is improved from 6/5 to 5/3 and 4/2.

For the Sandy Bridge architecture, the external body is unrolled four times instead of two, and the most internal loop is split in three loops: one loop computing $C_1, C_2$, a second loop computing $C_3, C_4$, and a third loop computing $C_5$. The load/store ratio is improved from 6/5 to 6/2 and 5/1.

Then, all arrays were 256-bit aligned using compiler directives and the first dimensions of all arrays were set to a multiple of 8 elements (if necessary, padded with zeros at the end of each column) in order to force a 256-bit alignment of every column of the matrices. These modifications allowed the compiler to use only vector instructions to perform the matrix products, both with the Streaming SIMD Extension (SSE) or the Advanced Vector Extension (AVX) instruction sets. The x86_64 version of the MAQAO framework[29] indicates that, as the compiler unrolled twice the third loop ($C_5$), these three loops perform 16 floating point operations per cycle, which is the peak performance on this architecture.

Finally, to improve the cache hit probability, blocking was used on the first dimension of $B_n$ (loop over $k$). In each block, the electrons (columns of $B$) are sorted by ascending first element of the indices array in the block. This increases the probability that columns of $A$ will be in the cache for the computation of the values associated with the next electron.

The results obtained using the Intel Fortran Compiler XE 2011 are presented in table I for both the Core2 and the Sandy Bridge architectures. The single-core double-precision Linpack benchmark is also mentioned for comparison. The results show that the full performance of the matrix products is already reached for the smallest number of basis functions used. For example, for systems 3 and 4 corresponding to the same molecule with a different number of basis functions, the multiplicative part is still dominant for the larger basis set ($\beta$-strand with cc-pVTZ) while it is no longer true for the smaller basis set ($\beta$-strand with 6-31G*). In figure 2 a plot of the total computational time for the Sandy Bridge core as a function of the number of electrons is presented. A standard fit of the curve with a polynomial form $N^\gamma$ leads to a $\gamma$-value of about 2.5. However, as discussed above such a power is not really meaningful. From the data of table II it is easy to extract the pure contribution related to the inversion and a factor very close to 3 is obtained, thus illustrating that for this linear algebra part we are in the asymptotic regime. For the multiplicative part, the pure $N^2$ behavior is not yet recovered and we are in an intermediate regime. Putting together these two situations

![FIG. 2: Single-core scaling with system size.](image-url)
TABLE I: Single core performance (GFlops/s) of the matrix products (single precision), inversion (double precision) and overall performance of QMC=Chem (mixed single/double precision). The percentage of the peak performance is given in parentheses. Core2: Intel Xeon 5140, Core2 2.33GHz, Dual core, 4MiB shared L2 cache. Sandy Bridge: Intel Xeon E3-1240, Sandy Bridge 3.30GHz, Quad core, 256KiB L2 cache/core, 8MiB shared L3 cache (3.4GHz with turbo).

<table>
<thead>
<tr>
<th>RAM (MiB)</th>
<th>Smallest system</th>
<th>β-Strand</th>
<th>β-Strand TZ</th>
<th>1ZE7</th>
<th>1AMB</th>
</tr>
</thead>
<tbody>
<tr>
<td>Core2</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>QMC step(s)</td>
<td>0.0062</td>
<td>0.0391</td>
<td>0.0524</td>
<td>0.2723</td>
<td>0.9703</td>
</tr>
<tr>
<td>Inversion</td>
<td>15%</td>
<td>31%</td>
<td>21%</td>
<td>47%</td>
<td>58%</td>
</tr>
<tr>
<td>Products</td>
<td>25%</td>
<td>23%</td>
<td>35%</td>
<td>21%</td>
<td>18%</td>
</tr>
<tr>
<td>Sandy Bridge</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>QMC step(s)</td>
<td>0.0026</td>
<td>0.0119</td>
<td>0.0187</td>
<td>0.0860</td>
<td>0.3042</td>
</tr>
<tr>
<td>Inversion</td>
<td>12%</td>
<td>26%</td>
<td>17%</td>
<td>42%</td>
<td>52%</td>
</tr>
<tr>
<td>Products</td>
<td>24%</td>
<td>22%</td>
<td>32%</td>
<td>21%</td>
<td>20%</td>
</tr>
</tbody>
</table>

TABLE II: Single-core memory consumption and elapsed time for one VMC step. Values in % represent the percentage of the total CPU time. Core2: Intel Xeon 5140, Core2 2.33GHz, Dual core, 4MiB shared L2 cache. Sandy Bridge: Intel Xeon E3-1240, Sandy Bridge 3.30GHz, Quad core, 256KiB L2 cache/core, 8MiB shared L3 cache (3.4GHz with turbo).

leads to some intermediate scaling around 2.5.

2. Sparsity

In our practical algorithm, for the matrix products we have chosen to consider the $B$ matrices as sparse as opposed to the $A$ matrix which is considered dense. The reason for that is that considering the matrix $A$ sparse would not allow us to write a stride-one inner loop. In single precision, SSE instructions executed on Intel processors can perform up to 8 instructions per CPU cycle (one 4-element vector ADD instruction and one 4-element vector MUL instruction in parallel). Using the latest AVX instruction set available on the Sandy Bridge architecture, the width of the SIMD vector registers have been doubled and the CPU can now perform up to 16 floating point operations per cycle. A necessary condition for enabling vectorization is a stride-one access to the data. This implies that using a sparse representation of $A$ would disable vectorization, and reduce the maximum number of floating operations per cycle by a factor of 4 using SSE (respectively 8 using AVX). If matrix $A$ has more than 25% (respectively 12.5%) non-zero elements, using a sparse representation is clearly not the best choice. This last result is a nice illustration of the idea that the efficiency of the formal mathematical algorithm depends on the core architecture.

Now, let us consider the inversion step which is the dominant CPU-part for the big enough systems (here, for about a thousand electrons and more). In Table I the performance in GFlops/s of the inversion step is presented for both processors. For comparisons the theoretical single-core peak and single-core Linpack performance are given. The first column gives the overall performance of the code while the second column is specific to the inversion part. As seen the performance of both parts increases with the number of electrons. For largest systems the performance represents more than 50% of the peak performance of each processor. For the largest system the whole code has a performance of about 54% of the peak performance for the Core2, and about 61% for the Sandy Bridge. The performance is still better for the inversion part: 60% for the Core2 and 67% for the Sandy Bridge.

4. Determinant calculation compared to spline interpolation

Most authors use three-dimensional spline representations of the molecular orbitals in order to compute in constant time the values, first derivatives and Laplacians of one electron in one molecular orbital, independently of the size of the atomic basis set. This approach seems efficient at first sight, but the major draw-
back is that the memory required for a single processor can become rapidly prohibitive since each molecular orbital has to be pre-computed on a three-dimensional grid. To overcome the large-memory problem, these authors use shared memory approaches on the computing nodes, which implies coupling between the different CPU cores. In this paragraph, we compare the wall time needed for spline interpolation or computation of the values, first derivatives and Laplacians of the wave function at all electron positions.

Version 0.9.2 of the Einspline package[31] was used as a reference to compute the interpolated values, gradients and Laplacians of 128 MOs represented on $23 \times 21 \times 29$ single precision arrays. The “multiple uniform splines” set of routines were used. To evaluate the value, gradient and Laplacian of one molecular orbital at one electron coordinate, an average of 1200 CPU cycles was measured using LIKWID[32] on the Core2 processor versus 850 CPU cycles on the Sandy Bridge processor. Even if the interpolation is done using a very small amount of data and of floating point operations, it is bound by the memory latency. Indeed, the needed data is very unlikely to be in the CPU cache and this explains why the number of cycles per matrix element is quite large. As our code uses a very small amount of memory, and as the computationally intensive routines are very well vectorized by the compiler, the computation of the matrix elements is bound by the floating point throughput of the processor.

The number of cycles needed to build the $C_1 \ldots C_5$ matrices is the number of cycles needed for one matrix element scaled by the number of matrix elements $N^2_2 + N^2_2$. Table III shows the number of CPU cycles needed to build the full $C_1 \ldots C_5$ matrices for a new set of electron positions using spline interpolation or using computation. The computation includes the computation of the values, gradients and Laplacians of the atomic orbitals (matrices $B_1 \ldots B_5$) followed by the matrix products.

Using a rather small basis set (6-31G*), the computation of the matrices in the 158-electron system is only 10% slower than the interpolation on the Core2 architecture. Using a larger basis set (cc-pVTZ), the computation is only 57% slower.

As the frequency is higher in our Sandy Bridge processor than in our Core2 processor, we would have expected the number of cycles of one memory latency to increase, and therefore we would have expected the Einspline package to be less efficient on that specific processor. One can remark that the memory latencies have been dramatically improved from the Core2 to the Sandy Bridge architectures and the number of cycles for the interpolation decreases.

The full computation of the matrix elements benefits from the improvement in the memory accesses, but also from the enlargement of the vector registers from 128 bits to 256 bits. This higher vectorization considerably reduces the number of cycles needed to perform the calculation such that in the worst case (the largest basis set), the full computation of the matrix elements takes as much time as the interpolation. In all other cases, the computation is faster than the spline interpolation. Finally, let us mention that as the memory controller is directly attached to the CPU, on multi-socket computing nodes the memory latencies are higher when accessing a memory module attached to another CPU (NUMA architecture).

V. PARALLELISM: IMPLEMENTING A UNIVERSAL, DYNAMIC, AND FAULT-TOLERANT SCHEME

Our objective was to design a program that could take maximum advantage of heterogeneous clusters, grid environments, the petaflops platforms available now and those to come soon (exascale).

To achieve the best possible parallel speed-up on any hardware, all the parallel tasks have to be completely decoupled. Feldman et al have shown that a naive implementation of parallelism does not scale well on commodity hardware.[33] Such bad scalings are also expected to be observed on very large scale simulations. Therefore, we chose an implementation where each CPU core realizes a QMC run with its own population of walkers independently of all the other CPU cores. The run is divided in blocks over which the averages of the quantities of interest are computed. The only mandatory communications are the one-to-all communication of the input data and the all-to-one communications of the results, each result being the Monte Carlo average computed with a single-core executable. If a single-core executable is able to start as soon as the input data is available and stop at any time sending an average over all the computed Monte Carlo steps, the best possible parallel speed-up on the machine can always be obtained. This aspect is detailed in this section.

A. Fault-tolerance

Fault-tolerance is a critical aspect since the mean time before failure (MTBF) increases with the number of hardware components: using $N$ identical computing nodes for a single run multiplies by $N$ the probability of failure of the run. If one computing node is expected to fail once a year, a run using 365 computing nodes is not expected to last more than a day. As our goal is the use both of massive resources and commodity clusters found in laboratories, hardware failure is at the center of our software design.

The traditional choice for the implementation of parallelism is the use of the Message Passing Interface (MPI).[34] Efficient libraries are proposed on every parallel machine, and it is probably the best choice in most situations. However, all the complex features of MPI are not needed for our QMC program, and it does not really fit our needs: in the usual MPI implementations, the
TABLE III: Number of million CPU cycles needed for the computation of the values, gradients and Laplacians of the molecular orbitals using the Einspline package and using our implementation for the Core2 and the Sandy Bridge micro-architectures. The ratio QMC=Chem / Einspline is also given.

<table>
<thead>
<tr>
<th>System</th>
<th>Core2 QMC=Chem Einspline ratio</th>
<th>Sandy Bridge QMC=Chem Einspline ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Smallest system</td>
<td>16.7 15.0 1.11</td>
<td>9.2 10.6 0.87</td>
</tr>
<tr>
<td>β-Strand TZ</td>
<td>177.3 113.0 1.57</td>
<td>81.7 80.1 1.02</td>
</tr>
<tr>
<td>12ZE7</td>
<td>783.5 669.1 1.17</td>
<td>352.0 473.9 0.74</td>
</tr>
<tr>
<td>1AMB</td>
<td>2603.0 1797.8 1.45</td>
<td>1183.9 1273.5 0.93</td>
</tr>
</tbody>
</table>

FIG. 3: Overview of the QMC=Chem architecture.

whole run is killed when one parallel task is known not be able to reach the MPI_Finalize statement. This situation occurs when a parallel task is killed, often due to a system failure (I/O error, frozen computing node, hardware failure, etc). For deterministic calculations where the result of every parallel task is required, this mechanism prevents from unexpected deadlocks by immediately stopping a calculation that will never end. In our implementation, as the result of the calculation of a block is a Gaussian distributed random variable, removing the result of a block from the simulation is not a problem since doing that does not introduce any bias in the final result. Therefore, if one computing node fails, the rest of the simulation should survive.

We wrote a simple Python TCP client/server application to handle parallelism. To artificially improve the bandwidth, all network transfers are compressed using the Zlib library,[35] and the results are transferred asynchronously in large packets containing a collection of small messages. Similarly, the storage of the results is executed using a non-blocking mechanism. The computationally intensive parts were written using the IRPF90 code generator,[36] to produce efficient Fortran code that is also easy to maintain. The architecture of the whole program is displayed in figure 3.

B. Program interface

Our choice concerning the interaction of the user with the program was not to use the usual “input file and output file” structure. Instead, we chose to use a database containing all the input data and control parameters of the simulation, and also the results computed by different runs. A few simple scripts allow the interaction of the user with the database. This choice has several advantages:

- The input and output data are tightly linked together. It is always possible to find to which input corresponds output data.
- If an output file is needed, it can be generated on demand using different levels of verbosity.
- Graphical and web interfaces can be trivially connected to the program.
- Simple scripts can be written by the users to manipulate the computed data in a way suiting their needs.

Instead of storing the running average as the output of a run, we store all the independent block-averages in the database, and the running averages are post-processed on demand by database queries. There are multiple benefits from this choice:

- Checkpoint/restart is always available.
- It is possible to compute correlations, combine different random variables, etc, even when the QMC run is finished.
- Combining results computed on different clusters consists in simply merging the two databases, which allows automatically the use of the program on computing grids.[37]
- Multiple independent jobs running on the same cluster can read/write in the same database to communicate via the file system. This allows to gather more and more resources as they become available on a cluster, or to run a massive number of tasks in a best effort mode.[38]

C. Error-checking

We define the critical data of a simulation as the input data that characterizes uniquely a given simulation. For instance, the molecular coordinates, the molecular
orbitals, the Jastrow factor parameters are critical data since they are fixed parameters of the wave function during a QMC run. In contrast, the number of walkers of a simulation is not critical data for a VMC run since the results of two VMC simulations with a different number of walkers can be combined together. A 32-bit cyclic redundancy code (CRC-32 key) is associated with the critical data to characterize a simulation. This key will be used to guarantee that the results obtained in one simulation will never be mixed with the results coming from another simulation and corrupt the database. It will also be used to check that the input data has been well transferred on every computing node.

D. Program execution

When the program starts its execution, the manager process runs on the master node and spawns two other processes: a data server and a main worker process.

At any time, new clients can connect to the data server to add dynamically more computational resources to a running calculation, and some running clients can be terminated without stopping the whole calculation. The manager periodically queries the database and computes the running averages using all the blocks stored in the database. It controls the running/stopping state of the workers by checking if the stopping condition is reached (based for example on the wall-clock time, on the error bar of the average energy, a Unix signal, etc).

When running on super-computers, the main worker process spawns one single instance of a forwarder on each computing node given by the batch scheduler system using an MPI launcher. As soon as the forwarders are started the MPI launcher terminates, and each forwarder connects to the data server to retrieve the needed input data. The forwarder then starts multiple workers on the node with different initial walker positions.

Each worker is an instance of the single-core Fortran executable, connected to the forwarder by Unix pipes. Its behavior is the following:

```c
while ( .True. ) {
    compute_a_block_of_data();
    send_the_results_to_the_forwarder();
}
```

Unix signals SIGTERM and SIGUSR2 are trapped to trigger the `send_the_results_to_the_forwarder` procedure followed by the termination of the process. Using this mechanism, any single-core executable can be stopped immediately without losing a single Monte Carlo step. This aspect is essential to obtain the best possible speed-up on massively parallel machines. Indeed, using the matrix product presented in the previous section makes the CPU time of a block non-constant. Without this mechanism the run would finish when the last CPU finishes, and the parallel efficiency would be reduced when using a very large number of CPU cores.

While the workers are computing the next block, the forwarder sends the current results to the data-server using a path going through other forwarders. The forwarders are organized in a binary tree as displayed in figure 4: every node of the tree can send data to all its ancestors, in order to deal with possible failures of computing nodes. This tree-organization reduces the number of connections to the data server, and also enlarges the size of the messages by combining in a single message the results of many forwarders.

At the end of each block, the last walker positions are sent from the worker to the forwarder. The forwarder keeps a fixed-sized list of \(N_{\text{kept}}\) walkers enforcing the distribution of local energies: when a forwarder receives a set of \(N\) walkers, it appends the list of new walkers to its \(N_{\text{kept}}\) list, and sorts the \(N + N_{\text{kept}}\) list by increasing local energies. A random number \(\eta\) is drawn to keep all list entries at indices \(\frac{\eta + i(N + N_{\text{kept}})}{N_{\text{kept}}}, i = \{1, \ldots, N_{\text{kept}}\}\) After a random timeout, if the forwarder is idle, it sends its list of walkers to its parent in the binary tree which repeats the list merging process. Finally, the data server receives a list of walkers, merges it with its own list and writes it to disk when idle. This mechanism ensures that the walkers saved to disk will represent homogeneously the whole run and avoids sending all the walkers to the data server. These walkers will be used as new starting points for the next QMC run.

Using such a design the program is robust to system failures. Any computing node can fail with a minimal impact on the simulation:

- If a worker process fails, only the block being computed by this worker is lost. It does not affect the forwarder to which it is linked.
- If a forwarder fails, then only one computing node is lost thanks to the redundancy introduced in the binary tree of forwarders.
- The program execution survives short network disruption (a fixed timeout parameter). The data will arrive to the data server when the network becomes operational again.
- The disks can crash on the computing nodes: the temporary directory used on the computing nodes is a RAM-disks (/dev/shm).
The shared file system can fail as the single-core static executable, the python scripts and input files are broadcast to the RAM-disks of the compute nodes with the MPI launcher when the run starts.

Redundancy can be introduced on the data server by running multiple jobs using the same database. Upon a failure of a data server, only the forwarders connected to it will be lost.

In the case of a general power failure, all the calculations can be restarted without losing what has already been stored in the database. Finally, we have left the possibility of using different executables connected to the same forwarder. This will allow a combined use of pure CPU executables with hybrid CPU/GPU and CPU/MIC executables, to use efficiently all the available hardware. The extension to hybrid architectures will be the object of a future work.

E. Parallel speed-up

The benchmarks presented in this section were performed on the Curie machine (GENCI-TGCC-CEA, France). Each computing node is a dual socket Intel Xeon E5-2680: 2×(8 cores, 20 MiB shared L3-cache, 2.7 GHz) with 64 GiB of RAM. The benchmark is a DMC calculation of the β-strand system with the cc-PVTZ basis set (Table IV) using 100 walkers per core performing 300 steps in each block. Note that these blocks are very short compared to realistic simulations where the typical number of steps would be larger than 1000 to avoid the correlation between the block averages.

1. Intra-node

The CPU consumption of the forwarder is negligible (typically 1% of the CPU time spent in the single-core executables). The speed-up with respect to the number of sockets is ideal. Indeed the single-core binaries do not communicate between each other, and as the memory consumption per core is very low, each socket never uses memory modules attached to another socket. When multiple cores on the same socket are used we observed a slow-down for each core due to the sharing of the L3-cache and memory modules. Running simultaneously 16 instances of the single-core binaries on our benchmark machine yields an increase of 10.7% of the wall-clock time compared to running only one instance. For a 16-core run, we obtained a 14.4x speed-up (the Turbo feature of the processors was de-activated for this benchmark).

2. Inter-node

In this section the wall-clock time is measured from the very beginning to the very end of the program execution using the standard GNU time tool. Hence, the wall-clock time includes the initialization and finalization steps. The initialization step includes:

- Input file consistency checking
- Creating a gzipped tar file containing the input files (wave function parameters, simulation parameters, a pool of initial walkers), the Python scripts and static single-core executable needed for the program execution on the slave nodes
- MPI Initialization
- Broadcasting the gzipped tar file via MPI to all the slave nodes
- Extracting the tar file to the RAM-disk of the slave nodes
- Starting the forwarders
- Starting the single-core instances

Note that as no synchronization is needed between the nodes, the computation starts as soon as possible on each node.

The finalization step occurs as follows. When the data server receives a termination signal, it sends a termination signal to all the forwarders that are leaves in the tree of forwarders. When a forwarder receives such a signal, it sends a SIGTERM signal to all the single-core binary instances of the computing node which terminate after sending to the forwarder the averages computed over the truncated block. Then, the forwarder sends this data to its parent in the binary tree with a termination signal, and sends a message to the data server to inform it that it is terminated. This termination step walks recursively through the tree. When all forwarders are done, the data server exits. Note that if a failure happened on a node during the run, the data server never receives the message corresponding to a termination of the corresponding forwarder. Therefore when the data server receives the termination signal coming from the forwarders tree, if the data server is still running after a given timeout it exits.

We prepared a 10-minutes run for this section to compute the parallel speed-up curve as a function of the number of 16-core nodes given in Figure 5. The data corresponding to this curve are given in Table V. The reference for the speed-up is the one-node run. The speed-up for N nodes is computed as:

\[
\frac{t_{\text{CPU}}(N)/t_{\text{Wall}}(N)}{t_{\text{CPU}}(1)/t_{\text{Wall}}(1)}
\]

The initialization time was 9 seconds for the single node run, and 22 seconds for the 1000 nodes run. The finalization time was 13 seconds for the single node run, and 100 seconds for the 1000 nodes run.

Apart from the initialization and finalization steps (which obviously do not depend on the total execution
TABLE IV: System sizes, percentage of non-zero molecular orbital coefficients and average percentage of non-zero Atomic Orbital values. \(a\) Zero MO coefficients are those below \(10^{-5}\). These are given for localized orbitals, and for canonical orbitals in parentheses. \(b\) Zero AO matrix elements are those for which the radial component of the basis function has a value below \(10^{-8}\) for given electron positions.

| N \(N_{\text{basis}}\) | Smallest system \(\beta\)-Strand \(\beta\)-Strand TZ 1ZE7 1AMB |
|-------------------------|------------------------------------------------|------------------------------------------------|------------------------------------------------|------------------------------------------------|
| 158                     | 434                                          | 434                                          | 1056                                        | 1731 |
| \% of non-zero\(a\)    | 81.3%                                        | 48.4%                                        | 73.4%                                       | 49.4% | 37.1% |
| MO coefficients \(a_{ij}\) \((A_{ij} \neq 0)\) | (99.4%)                                      | (76.0%)                                      | (81.9%)                                     | (72.0%) | (66.1%) |
| Average \% of non-zero basis functions \(\chi_i(r_j)\) \((B_{1ij} \neq 0)\) | 36.2%                                        | 14.8%                                        | 8.2%                                        | 5.7% | 3.9% |
| Average number of non-zero elements per column of \(B_{1ij}\) | 146                                          | 142                                          | 241                                         | 135 | 152 |

TABLE V: Data relative to the scaling curve (Figure 5). CPU time is the cumulated CPU time spent only in the Fortran executables, Wall time is the measured wall-clock time, including initialization and finalization steps (serial). The 10-minutes run were measured, and the longer runs are estimated from the 10-minutes run data. Two checks were measured for the 60-minutes runs with 1 and 100 nodes (in italics).

<table>
<thead>
<tr>
<th>Number of 16-core Nodes</th>
<th>10 minutes</th>
<th>60 minutes</th>
<th>180 minutes</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CPU (s)</td>
<td>Wall (s)</td>
<td>Speed-up</td>
</tr>
<tr>
<td>1</td>
<td>9627</td>
<td>625</td>
<td>1.0</td>
</tr>
<tr>
<td>10</td>
<td>95721</td>
<td>627</td>
<td>9.9</td>
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<tr>
<td>25</td>
<td>239628</td>
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</tr>
<tr>
<td>50</td>
<td>477295</td>
<td>631</td>
<td>49.1</td>
</tr>
<tr>
<td>100</td>
<td>952388</td>
<td>636</td>
<td>97.2</td>
</tr>
<tr>
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<td>637</td>
<td>190.5</td>
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<tr>
<td>400</td>
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<td>648</td>
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</tr>
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<td>500</td>
<td>4479967</td>
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<td>453.7</td>
</tr>
<tr>
<td>1000</td>
<td>8233981</td>
<td>713</td>
<td>749.7</td>
</tr>
</tbody>
</table>

FIG. 5: Parallel speed-up of QMC-Chem with respect to 16-core compute nodes (Reference is one 16-core node).

Finally, a production run was made using 76,800 cores of Curie (4,800 nodes) on the \(\beta\)-strand molecule with a cc-pVTZ basis set via 12 runs of 40 nodes, and a sustained performance of 960 TFlops/s was measured. All the details and scientific results of this application will be presented elsewhere.[39]
VI. SUMMARY

Let us summarize the main results of this work. First, to enhance the computational efficiency of the expensive innermost floating-point operations (calculation and multiplication of matrices) we propose to take advantage of the highly localized character of the atomic Gaussian basis functions, in contrast with the standard approaches using localized molecular orbitals. The advantages of relying on atomic localization have been illustrated on a series of molecules of increasing sizes (number of electrons ranging from 158 to 1731). In this work, it is emphasized that the notion of scaling of the computational cost as a function of the system size has to be considered with caution. Here, although the algorithm proposed is formally quadratic it displays a small enough prefactor to become very efficient in the range of number of electrons considered. Furthermore, our implementation of the linear-algebra computational part has allowed to enlighten a fundamental issue rarely discussed, namely the importance of taking into consideration the close links between algorithmic structure and CPU core architecture. Using efficient techniques and optimization tools for enhancing single-core performance, this point has been illustrated in various situations. Remark that this aspect is particularly important: as the parallel speed-up is very good, the gain in execution time obtained for the single-core executable will also be effective in the total parallel simulation.

In our implementation we have chosen to minimize the memory footprint. This choice is justified first by the fact that today the amount of memory per CPU core tends to decrease and second by the fact that small memory footprints allow in general a more efficient usage of caches. In this spirit, we propose not to use 3D-spline representation of the molecular orbitals as usually done. We have shown that this can be realized without increasing the CPU cost. For our largest system with 1731 electrons, only 313 MiB of memory per core was required. As a consequence the key limiting factor of our code is only the available CPU time and neither the memory nor disk space requirements, nor the network performance. Let us re-emphasize that this feature is well aligned with the current trends in computer architecture for large HPC systems.

Finally, let us conclude by the fact that there is no fundamental reason why the implementation of such a QMC simulation environment which has been validated at petaflops level could not be extended to exascale.

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13. A word of caution is necessary here. Although the fixed-node error on total energies is indeed usually very small compared with typical errors of standard computational chemistry methods, this error can still be large enough to have a non-negligible impact on small energy differences of interest in chemistry (binding energies, energy barriers, electronic affinities, etc.). Accordingly, to have to our disposal nodal hypersurfaces of sufficient quality for a general molecular system remains an important issue of QMC approaches.
35. Http://zlib.net.
38. When cluster resources are unused, a QMC job starts. When another user requests the resources, the QMC job is killed.