BioEM: GPU-accelerated computing of Bayesian inference of electron microscopy images

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Abstract

In cryo-electron microscopy (EM), molecular structures are determined from large numbers of projection images of individual particles. To harness the full power of this single-molecule information, we use the Bayesian inference of EM (BioEM) formalism. By ranking structural models using posterior probabilities calculated for individual images, BioEM in principle addresses the challenge of working with highly dynamic or heterogeneous systems not easily handled in traditional EM reconstruction. However, the calculation of these posteriors for large numbers of particles and models is computationally demanding. Here we present highly parallelized, GPU-accelerated computer software that performs this task efficiently. Our flexible formulation employs CUDA, OpenMP, and MPI parallelization combined with both CPU and GPU computing. The resulting BioEM software scales nearly ideally both on pure CPU and on CPU+GPU architectures, thus enabling Bayesian analysis of tens of thousands of images in a reasonable time. The general mathematical framework and robust algorithms are not limited to cryo-electron microscopy but can be generalized for electron tomography and other imaging experiments.

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PROGRAM SUMMARY

Program Title: BioEM.
Journal Reference:
Catalogue identifier: GNU GPL v3.
Programming language: C++, CUDA.
Operating system: Linux.
RAM: Problem dependent, 100 MB – 4 GB.
Supplementary material: see online supplementary material.
External routines/libraries: Boost 1.55, FFTW 3.3.3, MPI.
Subprograms used: Situs 2.7.2[1] routine for reading .mrc images.
Running time: Problem dependent, ~ 0.1 ms per parameter set.
Distribution format: GIT repository or zip archive.
Nature of problem: Analysis of electron microscopy images.
Solution method: GPU-accelerated Bayesian inference with numerical grid sampling.

1. Introduction

Cryo-electron microscopy has revolutionized structural biology [2, 3] providing structures of chemical-motors, such as ATP synthase, ion-channels, and transporters, at atomic-level resolution [4–10]. Due to its near-native conditions and single-molecule character, cryo-EM is a powerful technique with great potential. In fact, the structures of many biomolecules that are difficult to characterize using X-ray crystallography or nuclear magnetic resonance have now been resolved with cryo-EM.

These advances are feasible because of new technologies with time-resolved direct electron detection cameras [11], the development of novel image processing methods [12–15], and the use of accelerated computing capacities of multi-core processors and hardware accelerators such as GPUs (Graphics Processing Units) [16–19]. Direct electron detection cameras record an image rapidly as an ensemble of time-dependent frames (or movies), with an unprecedentedly low electron dose, that capture the evolution of the individual particles in time [20]. This time-resolution makes it possible to characterize...
the effects of beam-induced motion and radiation damage. Novel image processing algorithms filter the contributions of each individual frame according to a precalculated energy filter or an in-situ B-factor assignment, and discard (or low-weight) the damaged frames. However, not only single time-frames are discarded, but also many complete particle images. Successful 3D classification algorithms group the images into classes that generate different 3D maps, and the images that determine the map with the highest resolution correspond, in most cases, to just a small fraction of the total.

Despite many improvements, there are still several scenarios in which cryo-EM algorithms face challenges to acquire high-resolution information. Most methods rely on the hypothesis that all molecular orientations are sampled equally. This is not the case for some hydrophobic or large systems (of size comparable to the ice thickness), which acquire preferred orientations due to the carbon grid. For disordered or heterogeneous complexes (having multiple binding partners and affinities), it is difficult to obtain sufficiently many particles to cover all orientations for each conformation, and computationally challenging to assign a single orientation to one of the multiple configurations. A major difficulty lies in generating the initial 3D models that the algorithms use as initial alignment references. Thus, despite the enormous advances, the effort to extend the reach of cryo-EM is ongoing.

In this work, we provide a computational tool to harness the power of cryo-EM as a true single-molecule technique. Our algorithm performs a Bayesian inference of electron microscopy (BioEM), accelerated by GPUs, to calculate the posterior probability of a set of structural models given a set of experimental images. In contrast to standard reconstruction algorithms, we perform a forward calculation, building a realistic image from a model and compare it to an unmodified particle image using a likelihood function. The BioEM posterior probability calculated in this way allows us to accurately rank and discriminate sets of structural models.

The paper is organized as follows: we first describe the mathematical framework to create a calculated image from a model, and the Bayesian technique to obtain the posterior probability for a set of particle images. Then, we introduce the BioEM algorithm and describe its main routines as well as the parallelization scheme. We demonstrate the performance and scalability of the BioEM program on a high-performance computing (HPC) cluster using two benchmark sets of particle images. Lastly, we discuss the
limitations of the method and point out future perspectives.

2. Mathematical formulation

The core of the BioEM method \cite{35} is the calculation of the posterior probability of an ensemble of structural models, \( m \in M \), given a set of experimental images, \( \omega \in \Omega \). The posterior is defined in the usual manner, as the product of a prior probability for the various parameters (e.g., orientations), and a likelihood function \( L \) defined as the probability of observing the measured image given the model and parameters, \( P(\text{model}|\text{data}) \propto \int d\theta p(\text{model}, \text{parameters } \theta)L(\text{data}|\text{model}, \text{parameters } \theta) \), with parameters integrated out. The only requirement on the models is that an EM projection image can be calculated for them. Models can thus be represented in a variety of ways, from atomic coordinates to 3D electron density maps to simple geometric shapes. The key idea is to perform forward calculations of 2D EM projection-image intensities \( I^{\text{cal}} \) for given models and then compare them to the observed image intensities \( I^{\text{obs}}_\omega \). The calculation of the posterior takes into account the relevant factors in the experiment (denoted by \( \theta \)), such as the molecule orientation, interference effects, uncertainties in the particle center, normalization and offset in the intensities and noise.

In Fig. 1 we show a schematic representation of how \( I^{\text{cal}} \) is constructed. We start from the 3D electron density \( \rho(x, y, z|m, \varphi) \) of model \( m \) in a particular orientation \( \varphi \), as given by three Euler angles or four quaternions. Under the weak-phase approximation \cite{36}, a 2D-projection density is determined by integration along \( z \),

\[
I^{(0)}(j, k|m, \varphi) = \int_{-\infty}^{\infty} dz \int_{y_k-\Delta y/2}^{y_k+\Delta y/2} dy \int_{x_j-\Delta x/2}^{x_j+\Delta x/2} dx \rho(x, y, z|m, \varphi) ,
\]

where \((x_j, y_k)\) are the 2D positions corresponding to pixel \( j, k \) of width \( \Delta x \) and \( \Delta y \). The discrete Fourier transform of an image \( I \) is defined as

\[
\hat{I}(s) \equiv \hat{I}(l, n) \equiv \mathcal{F}(I) = \sum_{j=1}^{N_x} \sum_{k=1}^{N_y} I(j, k) e^{2\pi i (lj+nk)} .
\]

Here and in the following, we interchangeably index Fourier space with reciprocal space vectors \( s = 2\pi(l/\Delta x N_x, n/\Delta y N_y) \) or the corresponding integer indices \( l \) and \( n \), whichever is more convenient for the operation at hand.
Analogously, in real space we use either physical positions \( r = (x, y) = (j\Delta x, k\Delta y) \) or the corresponding indices \( j, k \). \( N_x \) and \( N_y \) are the numbers of pixels in \( x \) and \( y \) directions, respectively.

The inverse Fourier transform is

\[
I(j, k) \equiv \mathcal{F}^{-1}(\hat{I}) = \frac{1}{N_{\text{pix}}} \sum_{l=1}^{N_x} \sum_{n=1}^{N_y} \hat{I}(l, n) e^{-2\pi i(lj+nk)},
\]

(3)

where \( N_{\text{pix}} = N_x N_y \) is the total number of pixels.

We account for interference and inelastic scattering effects in EM imaging through the contrast transfer function (CTF) and the envelop function (Env), respectively, which are both assumed to be radially dependent for simplicity. The product of CTF and Env is the Fourier-space equivalent of the real-space point spread function (PSF),

\[
\text{CTF}(s)\text{Env}(s) = \mathcal{F}(\text{PSF}),
\]

(4)

where \( s = |s| \). As functional forms, we assume \( \text{CTF}(s|a, A) = -A \cos(as^2/2) - \sqrt{1 - A^2} \sin(as^2/2) \) and \( \text{Env}(s|b) = e^{-bs^2/2} \) with coefficients \( a, A, \) and \( b \). [37]

In Fourier space, the interference effects on the ideal image are then accounted by

\[
\hat{I}^{(1)}(s|m, \varphi, a, A, b) = \hat{I}^{(0)}(s|m, \varphi) \text{CTF}(s|a, A)\text{Env}(s|b),
\]

(5)

which corresponds to a convolution with the PSF in real space,

\[
I^{(1)}(r|m, \varphi, a, A, b) = \sum_{r'} I^{(0)}(r'|m, \varphi) \text{PSF}(|r - r'||a, A, b).
\]

(6)

Implicit in this procedure is the assumption that the pixel size is small compared to significant variations in the PSF.

The calculated image also accounts for uncertainties in the particle position, and variations in the imaging conditions

\[
I^{\text{cal}}(r|m, \varphi, a, A, b, d, N, \mu) = NI^{(1)}(r + d|m, \varphi, a, A, b) - \mu,
\]

(7)

where \( d = (d_x, d_y) \) is a translation vector shifting the image and thus the particle center pixel by pixel, \( N \) scales the intensity, and \( \mu \) is an intensity offset. The likelihood function \( L(\omega|m, \theta) \) of model \( m \) with image parameters
\[ \theta = (\varphi, a, A, b, d, N, \mu, \lambda), \]

establishes a measure of similarity between \( I_{\text{cal}} \)
and \( I_{\text{obs}} \),

\[
L(\omega|m, \theta) = \frac{\exp \left( -\sum_{j,k} \left[ I_{\text{obs}}(j, k) - I_{\text{cal}}(j, k|m, \varphi, a, A, b, d, N, \mu) \right]^2 / 2\lambda^2 \right)}{(2\pi\lambda^2)^{N_{\text{pix}}/2}}.
\]

Here we assumed for simplicity uncorrelated Gaussian noise in each pixel with zero mean and standard deviation \( \lambda \).

Figure 1: Steps in building a realistic image starting from a 3D model: rotation \( \varphi \), projection, point spread function convolution \((a, A, b)\), center displacement \(d\), and integrated-out parameters of normalization \(N\), offset \(\mu\) and standard deviation of the noise \(\lambda\). The likelihood function establishes the similarity between the calculated image and the observed experimental image.

The likelihood function, \( L(\omega|m, \theta) \), measures the agreement between individual observed and calculated images for fixed parameters \( \theta \). In contrast to maximum likelihood techniques \cite{38} that determine a single optimal parameter set, we perform a Bayesian analysis covering a wide range of possible parameter sets. This range, and the weight of individual sets, is defined by the prior probability \( p(\theta) \) (see the Supplementary Information for details about the priors implemented in the code), which is combined with the prior of the model, \( p_M(m) \). The Bayesian posterior probability of a model, \( m \), given an image, \( \omega \), is then a weighted integral over the product of prior and
likelihood,
\[ P_{m\omega} \propto \int L(\omega|m, \theta)p_M(m)p(\theta)d\theta . \] (9)

The posterior probability of an ensemble containing multiple models \((m \in M)\) with relative weights \(w_m\) (where \(\sum m w_m = 1\)), given a set of images \((\omega \in \Omega)\), is then
\[ P(M|\Omega) \propto \prod_{\omega=1}^{\Omega} \sum_{m=1}^{M} w_m P_{m\omega} , \] (10)
i.e., the product over independent images of weighted sum over ensembles.

3. Algorithm and Optimization

To evaluate the posterior probabilities \(P_{m\omega}\) in Eq. 10 for every model \(m \in M\) and image \(\omega \in \Omega\), we have to compute the integral in Eq. 9 over the parameters \(\varphi, a, A, b, d, N, \mu, \) and \(\lambda\). As shown in ref. [35], we can integrate noise \(\lambda\), normalization \(N\), and offset \(\mu\) analytically. The resulting analytical expression of the posterior probability \(P_{m\omega}(\varphi, a, A, b, d)\) as a function of the remaining parameters is shown in Eq. S8 of the Supplementary Information. Average and mean squared averages of the intensities of the observed and calculated images (Eqs. S3 - S6) can be precomputed. Additionally, Eq. S8 involves the estimation of the cross-correlation between the calculated and observed images, \(C_{co}\) from Eq. S7. The remaining integrals in Eq. 9 over orientations parametrized by \(\varphi\), the PSF with parameters \((a, A, b)\), and center displacements \(d\) are evaluated numerically. Importantly, for each parameter combination (with the possible exception of \(d\); see below), the cross-correlation between the \(I_{cad}\) and \(I_{obs}\) has to be calculated in a computationally demanding step.

The data dependency in the construction of the calculated image is sketched in Fig. 1. The model must be first rotated, then projected, then convoluted, and then displaced. The loop over the experimental images is, in principle, independent but it is best to nest this loop inside the loop over the model rotations and PSF convolutions. Then, the same rotation, projection, and convolution do not have to be computed repeatedly for every observed image.

Listing[] presents a pseudo-code of the BioEM algorithm. Here, the subroutine compute_probability computes the posterior probability from Eq. S8, and norm_images[] is an array with precomputed \(C_o\) and \(C_{oo}\) (from Eqs. S3
and S5) for all observed images.

```
for (r = 0;r < num_rot;r++) {
    m_rot = do_rotation(r, model);
    m_proj = do_projection(m_rot);
    for (c = 0;c < num_convolutions;c++) {
        m_conv = do_psf_convolution(m_proj, c);
        for (d = 0;d < num_displacements;d++) {
            m_disp = do_2d_displacement(m_conv, d);
            norm_model = compute_norm(m_disp);
            for (w = 0;w < num_images;w++) {
                cross_corr = compute_cross_corr(m_disp, image[w]);
                image_probability[w] += compute_probability
                    (norm_images[w], norm_model, cross_corr);
            }
        }
    }
}
```

Listing 1: Pseudocode of the numerical integration.

In the following, we describe the optimizations applied during the development, including some technical details. The measurements presented in this section have been taken on our development system, which hosts a typical, workstation-class Intel Core i7-980 3.33 GHz 6-core CPU and an NVIDIA GTX Titan GPU. This is complemented in Section 4 by a systematic study of the parallel efficiency and the GPU acceleration, performed on a HPC cluster with server-class CPUs and GPUs.

3.1. Model Rotation

The BioEM algorithm offers two representations of model orientations in 3D space: with Euler angles or with quaternions. Whereas the Euler angles are most commonly used in EM image processing softwares, we have found it more suitable to use quaternions. The advantage is that with the quaternions uniform sampling of the group $SO(3)$ of rotations in 3D space is more readily implemented [39].

3.2. Imaging Effects

After the rotation and projection, we construct the Fourier transform of the ideal image, $\mathcal{F}(I(0))$, from Eq. 1. Interference and imaging effects are taken into account by multiplication with CTF and Env in Fourier space according to Eq. 5, or by convolution with the PSF in real space (which can be advantageous, e.g., if the PSF is zero everywhere except near the origin). In the code, the PSFs to be sampled over, and their Fourier transforms, are
precalculated in the form of radially symmetric 2D images, and stored in memory.

3.3. Center Displacement

To evaluate the integral over the image translation vectors \( \mathbf{d} \) we have implemented two alternative approaches suitable for small and large sets of \( \mathbf{d} \), respectively. In a real space formulation, the cross-correlation is simply calculated for different displacements \( \mathbf{d} \). In Fourier space, we use the fact that the cross-correlation of the calculated and the observed images shifted by \( \mathbf{d} \) (denoted by \( C_{co}(\mathbf{d}) \)), is given by

\[
C_{co}(\mathbf{d}) = \mathcal{F}^{-1}[\mathcal{F}(I^{obs}) \cdot \mathcal{F}(I^{(1)})](\mathbf{d}),
\]

where \( \mathcal{F}^{-1} \) is the inverse Fourier transformation and the dot indicates the complex product of Fourier components at equal \( \mathbf{s} \) vectors, i.e.,

\[
C_{co}(d_x, d_y) = \frac{1}{N_{pix}} \sum_{l=1}^{N_x} \sum_{n=1}^{N_y} \hat{I}^{obs}(l, n) \overline{\hat{I}^{(1)}(l, n)} e^{-2\pi i (ld_x + nd_y)},
\]

where the overline indicates the complex conjugate. In this way, we can take advantage of the fact that the convolution with the PSF carried out in Fourier space produces \( \hat{I}^{(1)} \), and that \( \hat{I}^{obs} \) can be precalculated. Hence, the only computationally intense part is the Fourier backtransformation \( \mathcal{F}^{-1} \) in Eq. \( \text{11} \). If we compare the two algorithm variants, the real-space one requires the computation of as many cross-correlations as there are displacements. The Fourier variant requires one Fourier backtransformation, irrespective of the number of displacements. Naturally, the first version should have advantage with few displacements, while variant two will be superior for many displacements. In fact, Fig. \( \text{2} \) shows that the Fourier version is superior in most cases, both on the CPU and the GPU.

The Fourier variant differs from the real-space version in that it wraps around the borders of the experimental image instead of just cropping it. However, comparing the results for certain larger datasets, we can conclude that this does not pose a problem, because the particle is generally located in the center of the image while the region around it (the borders) consists mostly of noise. Moreover, using the Fourier variant, we have the advantage that we do not have to convert the calculated image back to real space for computing the cross-calculation, which saves some of the Fourier transforms.
We still need the constants $C_0$ and $C_{oo}$, but these can be easily computed in Fourier space using Plancherel’s theorem, e.g., $C_{oo} \propto \sum |F(I^{obs})|^2$.

We perform the Fourier-transformation using the Fast Fourier Transform (FFT) libraries \texttt{fftw} on the CPU and \texttt{cuFFT} on the GPU. Since all images (calculated and observed) are real, we use the fast real-Hermitian FFT variant offered by these libraries. The computing time of the FFT algorithms depends on the image size (see Supplementary Fig. S1). Box-sizes that are powers of 2 per dimension or follow standard EM/FFT suggestions \cite{41} have the best performance.

3.4. Numerical Precision

We computed the numerical integral in Eq. 9 using single and double precision with and without the the Kahan summation algorithm \cite{42}. We found that all options lead to the same numerical result, mainly, because the likelihood function is sharply peaked around its maxima and few parameter sets contribute to the summation. Thus, we set as default the fastest setup which uses single precision without the Kahan summation. All benchmarks shown in this paper are based on this single-precision setting.
3.5. Blocking

A common technique to optimize memory access patterns of nested loops (for instance for matrix operations) is blocking, which improves the cache usage. Alternatively, if the memory access pattern does not (or almost not) depend on one of the loops, one can place that loop as the innermost loop. We apply this optimization for the real-space variant. The loop over the displacements shifts the memory access only by one entry per iteration, so we use this loop as the innermost loop (i.e., exchange the loops over $w$ and $d$ in Listing 1). Since, for the Fourier variant, the FFTs are computationally dominant, we do not need the blocking optimization in this case.

3.6. Vectorization

For the real-space version, we have written the code to support compiler auto vectorization. By checking the disassembly of the object file we verified that the compiler vectorized the code exactly in the way we intend. Since optimized FFT libraries use vectorization, we do not have to take action in this respect for the Fourier variant.

3.7. Parallelization

In order to speed up processing of the very compute-intense BioEM task, we parallelize the processing on top of the vectorization. We consider parallelization over the cores inside one compute node, parallelization over multiple compute nodes, and usage of parallel accelerator devices like GPUs to speed up processing.

3.7.1. CPU Usage

It is desirable to parallelize the inner loop over the images on a shared memory architecture. This way, we need to compute rotation, projection, and convolution only once, and we can then reuse the calculated image $I^{\text{cal}}$ for comparison with all observed images $I^{\text{obs}}$. We employ OpenMP to process the comparison of $I^{\text{cal}}$ to all images in parallel. The outermost loop over the rotations does not have any dependencies (except for the source data), hence we have chosen to parallelize it via the Message Passing Interface (MPI) to support utilization of many compute nodes in parallel. Fig. 3 shows a schematic representation of the parallelization approach. Each grid cell represents a single rotation to a single experimental image, inside which the integrals over the projection, PSF convolution, and center displacement are performed (zoomed square Fig. 3). The Fourier algorithm calculates the
center displacement and cross-correlation to the experimental image simultaneously. If there are only a few experimental images, then also the rotation, projection, and Fourier transformation of $I_{\text{cal}}$ take a non negligible time (up to 30% for the small dataset from the Results section). In order to speed this up, BioEM can use OpenMP to precalculate all Fourier transformed images $I_{\text{cal}}$ in parallel.

3.7.2. GPU Usage

In recent years, Graphics Processing Units (GPUs) have shown significant speedup in many applications, among them the FFT, which is heavily used by BioEM. In order to leverage this potential, we have adapted BioEM to run on GPUs. BioEM can use CUDA for the cross-correlation step, which essentially consists of an image multiplication in Fourier space and a Fourier back-transformation. We did not consider bringing more of the steps to the GPU because the other parts are not time critical and can be processed well by the CPU. We use a pipeline where the CPU can prepare the next rotations and convolutions while the GPU runs the comparison of the last $I_{\text{cal}}$ to all observed images. We also arrange the remote direct memory access (RDMA) transfer of the new data from the host to the GPU in the pipeline.
asynchronously via CUDA streams. In this way, BioEM keeps executing GPU kernels 100% of the time and there is no GPU idle time. In order to use both CPU and GPU to the full extent, BioEM splits the work between GPU and CPU and uses both processor types jointly for the comparison step. Fig. 4 shows the computational time as a function of the number of the observed images. Except for very small sets of images, the processing time depends linearly on the number of observed images. Here the transition to linear scaling occurs between 128 and 256 images and is, thus, below what one would encounter in typical applications.

Table 1 summarizes the evolution of the code’s performance over subsequent development phases, starting out from a first, serial prototype version executed on a single CPU core (Intel Core i7, 3 GHz) to a full, GPU-accelerated multicore node. We would like to emphasize that the huge overall speedup (3 or 4 orders of magnitude for different setups with 130 or 11000 images, respectively) is to a large degree due to algorithmic optimizations pointed out in Sections 3.3 to 3.6 and OpenMP parallelization of the CPU version. The latter defines the performance baseline for a fair comparison of the GPU version which delivers a speedup by a factor of 5 in this case (6-core CPU vs. Titan GPU).
3.8. Extra Features

In addition to calculating the posterior probability of a model given a set of experimental images, the code provides several extra features:

- **Synthetic map:** print the synthetic map corresponding to a specific parameter set, $\theta$.
- **Maximizing parameters:** report the grid value parameters that give a maximum of the posterior probability.
- **Posterior for orientations:** obtain the posterior probability as a function of the rotational (Euler or quaternion) angles.

Details on using each feature are provided in the user manual.

Table 1: Performance evolution of BioEM through subsequent optimization phases (from top to bottom, first column) as measured on a workstation with an Intel Nehalem Core i7-980 6-core CPU (3.33 GHz) and an NVIDIA GTX Titan GPU. To test the flexibility of the code, we used two different setups (second column): 11000 images (224 × 224 pixels) of the $F_{420}$-reducing hydrogenase system [23, 43], and 130 images (170 × 170 pixels) of the chaperonin GroEL [44], both with one PSF, one orientation and one center displacement. The third column states the absolute runtime and the forth and fifth column give the relative speedup compared to the previous, or initial code version, respectively.

| Version                          | Number of Images | Time (incremental) | Speedup (cumulative) |
|----------------------------------|------------------|--------------------|----------------------|
| First prototype version          | 130              | ≈ 2 s              | -                    |
|                                  | 11000            | ≈ 1200 s           | -                    |
| First BioEM C++ real-space version | 130             | 0.533 s            | 4.00                 |
|                                  | 11000            | 312.4 s            | 4.00                 |
| Vectorized C++ real-space version with optimizations from Section 3 | 130              | 0.083 s            | 6.39                 |
|                                  | 11000            | 42.62 s            | 7.33                 |
| Fourier-space version            | 130              | 0.120 s            | 0.69                 |
|                                  | 11000            | 8.256 s            | 5.18                 |
| Real-Hermitian FFT version       | 130              | 0.048 s            | 2.49                 |
|                                  | 11000            | 2.809 s            | 2.94                 |
| Parallelization with OpenMP (6 cores) | 130              | 0.011 s            | 4.46                 |
|                                  | 11000            | 0.581 s            | 4.84                 |
| GPU usage with CUDA (1 NVIDIA Titan GPU) | 130              | 0.00213 s         | 5.07                 |
|                                  | 11000            | 0.106 s            | 5.48                 |
| Combined CPU / GPU usage         | 130              | 0.00208 s          | 1.02                 |
|                                  | 11000            | 0.091 s            | 1.16                 |
3.9. Margin for Improvements

Using the CUDA profiler, we measured that more than 75% of the compute time is spent in the cuFFT library for both datasets. The fraction of CPU time spent in fftw is even larger. The majority of the remaining time is used for pixel-wise multiplication of the images in Fourier space, which is by definition memory-bandwidth limited, in particular on the GPU. It is a small inefficiency in this respect that the images are stored to memory after the multiplication and then read again for the FFT, but this can hardly be avoided due to the use of the FFT libraries. The FFT libraries themselves are already well optimized, thus the margin for an additional speedup is limited.

4. Performance

This section presents a performance evaluation of the BioEM software, focusing on parallel efficiency and GPU performance obtainable on a typical high-performance compute cluster that is employed for production runs with BioEM. The tests were performed on the high-performance system Hydra of the Max-Planck-Society, operated by the Max-Planck Computing and Data Facility in Garching, Germany. It consists of dual-socket nodes equipped with Intel Xeon E5-2680 v2 CPUs (20 physical cores per node with 2 hyperthreads per core) and interconnected with a high-performance network (FDR InfiniBand). A subset of the nodes is equipped with two NVidia K20X “Kepler” GPUs each. For the benchmarks on the Hydra system, Intel compiler suite XE 2014, CUDA 5.5 and FFTW 3.3.3 were used on top of the Linux operating system SLES11.

We selected a benchmark set with 100 (denoted by “small”) and 2000 (denoted by “large”) experimental particle images of the 1.2 MDa F_{420} reducing hydrogenase (Frh) system [23, 43]. Each particle image had 224 × 224 pixels with 1.32 Å of pixel size. An all-atom structure (∼82000 atoms) built from the 3D density map [23] was used as the reference model. The numerical integrals in Eq. 9 were performed over grids with 13500 orientations, 64 PSFs, and 400 center displacements. This parameter setup is consistent with a case without prior knowledge of the symmetry of the system or the orientations of the particle images. Thus, we did not take advantage of the 12-fold symmetry of the Frh complex to reduce the orientational search. However, in practical applications this can be easily implemented, by searching over a restricted set of Euler angles or corresponding quaternions.
Fig. 5 shows that we can achieve almost perfect linear scaling with the number of physical cores in a node for both datasets. BioEM allows us to parallelize over the cores inside a node in several ways: over the observed images via OpenMP, or over the orientations via MPI, or with a combination of MPI and OpenMP in a hybrid setup. The figure compares OpenMP to MPI scalability. There are two effects that limit pure OpenMP scaling. First, there are unavoidable non-uniform memory access (NUMA) effects because common global data (e.g., the calculated images) are stored only once and thus are scattered over NUMA domains. This becomes apparent in Fig. 5 which shows that both the large (blue, solid line) and the small setup (red, solid line) show very good scalability up to the maximum of 10 cores of a NUMA domain. Second, in particular for the small dataset, synchronization after the computation of the likelihood limits the performance. In contrast, these aspects do not affect the pure MPI setup with each process mapped to an individual core. The MPI configurations (dashed lines) exhibit nearly perfect scaling up to the maximum of 20 physical cores in the node because both datasets have sufficiently many orientations for MPI parallelization. Thus, we note as a side result that memory bandwidth is not a limiting factor in this context. Since BioEM is compute bound by the FFTs, Hyper-Threading yields a small but non-negligible improvement.

![Figure 5: BioEM speedup compared to a single thread as a function of the number of employed CPU cores using the OpenMP (solid lines), over the observed images, or MPI (dashed lines), over the orientations, parallelization for both the small and large datasets. The shaded region indicates the use of Hyper-Threading.](image)
An important advantage of the OpenMP parallelization is its smaller memory footprint. While in the OpenMP case the threads share a copy of the observed images, each process in the MPI case has its own copy. The large dataset, for example, requires 35 GB (1.75 GB per MPI-process), which can already be prohibitive on some of today’s HPC clusters and the memory requirement increases further with larger datasets. Moreover, with a plain MPI parallelization the total number of orientations poses a strict upper limit on the number of MPI tasks. This would ultimately limit the strong scalability of BioEM, in particular for smaller problems with only very few orientations. The steadily increasing number of cores per CPU and the stagnating (or even decreasing) per-core performances will further exacerbate these constraints in the future.

Fig. 6 and 7 provide an overview of the performance, defined as the inverse runtime, obtained by employing different parallelization and GPU-acceleration options implemented by BioEM for the small and large datasets respectively on multiple nodes of the Hydra cluster. Both figures show multiple curves for different execution configurations. The curves distinguish between CPU-only configurations, GPU-only configurations, and a combined configuration that uses both GPUs and all CPU cores of each node. In the GPU-only case, two curves for one and for two GPUs per node are shown. We present three curves in the CPU-only case that differ in the parallelization approach. We show curves with pure OpenMP parallelization inside the nodes (1 process, 40 threads), pure MPI parallelization (40 single-threaded processes per node), and a hybrid MPI-OpenMP configuration of two MPI processes with 20 OpenMP threads each per node. All configurations obtain a close-to-perfect linear scaling with the number of nodes due to the absence of both communication and load-imbalances in the implementation.

The hybrid MPI-OpenMP parallelization adds flexibility for the parallelization and helps to contain the memory footprint. It can reduce the number of MPI processes per node to only a few and multiple cores per MPI process can be used efficiently by the OpenMP parallelization. For instance, assigning two MPI processes per two-socket node (i.e. one per NUMA domain) avoids all NUMA limitations and reduces the memory footprint enormously (e.g., from 35 GB to around 2 GB per node for the large dataset). On top of that, this shifts the strong scaling limit from the number of orientations to the number of orientations multiplied by the number of cores per socket. Thus, the hybrid approach can speed up computations employing larger computational resources when the number of orientations limits the
Figure 6: BioEM wall-clock performance (inverse runtime) as a function of the number of compute nodes for the small dataset, with 100 observed images, employing different parallelization and GPU-acceleration options. Solid lines are CPU-only configurations, and squares with dashed lines are GPU-only configurations. The combined workload configuration (open circles and dashed lines) had 60% of the observed images on 2 GPU devices, and the remaining 40% on the CPU with 20 OpenMP threads per node.

MPI scaling. In that respect, the new Intel Xeon Phi many-core CPU (codename: “Knights Landing”) with about 240 threads and a scarce resource of 16 GB of high-bandwidth memory, presumably to be separated into four NUMA domains, might be a powerful and energy-efficient architecture for operating the BioEM software in the hybrid MPI-OpenMP setup.

The large dataset comprising more images allows the GPU a better exploitation of its parallel architecture. For the small benchmark task, one GPU is as fast as one of the two 10-core Ivy-Bridge processors. Processing the large dataset, the GPU runs twice as efficiently achieving the performance of a full Ivy-Bridge node with two processors. (Consider that the small set with only 100 images represents more or less a lower bound, so GPU-processing works for all real cases.) In both cases, two GPUs reach twice the performance of a single GPU.

The GPU achieves a smaller portion of its theoretical peak performance
compared to the processor, because reading the images from global memory saturates the memory bandwidth. Here, the processor can play the strength of its larger caches. We note that whereas CPUs can easily hold tens of thousands of images, GPUs are slightly limited in memory (~ 6 GB for a K20x). However, the limited memory size is no real restriction because processing the images takes place independently. The GPU can process subsets of the images step-by-step, and the host can combine the results later on. Since processing of such a subset of up to 6 GB of images takes on the order of minutes, the overhead for additional transfers and repeated projections, etc., of the model is negligible.

BioEM also allows us to split the workload among CPU and GPU, i.e., the observed images are split into two sets: one goes to the GPU, and the other is analyzed on the CPU using OpenMP. We find that this workload-sharing improves the performance significantly. The full capacity of the node
is utilized, profiting from the 2 GPU devices and all the cores in the node. The optimal splitting ratio depends on the specific problem and hardware. For these setups, the fastest setting was 60% GPU for the small, and 65% GPU for the large datasets. Due to synchronization issues, the combined configuration does not achieve the sum of the individual CPU and GPU performances. (For instance, the combined configuration on 32 nodes achieves 84% of the sum of the individual performances of the large dataset.)

Our performance assessment shows that the optimal execution setup depends on the problem. By trend, the MPI parallelization works better with many orientations, while OpenMP needs many observed images. A hybrid setup is often the best compromise and scales almost perfectly linear with the number of cores. For most cases, we recommend employing one MPI process per NUMA domain.

For instance, a complete analysis of 10000 images with 13500 orientations and 64 PSFs takes approximately 140 minutes on 16 Ivy-Bridge nodes. The same analysis can be performed within 55 minutes if the nodes are accelerated by two K20x GPUs. This demonstrates that our new software can efficiently handle the analysis of the large amounts of experimental particles used in electron microscopy. We estimate that per grid point (one PSF and one orientation), using the Fourier-algorithm, the code takes approximately 0.1 ms per image of ~50000 pixels, with the exact runtime depending on the specific setup: number of pixels, model size, parameter ranges, grid points, etc. In the Supplementary Information, we present some more estimates on the BioEM software runtime.

5. Discussion

The BioEM method provides an alternative approach to structurally characterize biomolecules using electron microscopy images. By calculating the posterior probability of a model with respect to each individual image, it avoids information loss in averaging or classification, and allows us to compare structural models according to their posterior probability. Bayesian analysis methods, such as Relion [14, 15], have been enormously successful in EM, contributing much to the resolution revolution [2]. However, the main use of Relion is in reconstructing 3D densities from projection images, and not to rank or compare existing structural models. It also differs from BioEM in the integration scheme and optimization algorithms (for a comparison see Supplementary Table S1). BioEM requires relatively few images to discrimi-
nate the correct model within a pool of plausible structures (e.g., <1500 particles for GroEL [35]), whereas, for full 3D reconstructions, Relion typically requires tens to hundreds of thousands of particles and costly computational resources to implement the multiple methods that select, classify, and polish the particles as well as refine the 3D maps. Beyond applications in studies of highly dynamic systems [35], we envision that BioEM can complement traditional 3D reconstruction techniques in the first steps of classification by assigning accurate orientations and single-particle PSF estimations, and in the last steps of refinement by validating the final 3D models. In addition, the BioEM method can be applied to problems where reconstruction techniques fail, e.g., when there are few particle images that acquire preferred orientations or when the system is flexible.

The mathematical framework can also be extended to analyze individual time-dependent frames from direct electron-detection cameras or electron tomography tilt-series (see the Supplementary Information). We foresee that the BioEM method can be generalized to other types of imaging experiments, such as atomic force or light microscopy after appropriate modifications of the forward calculation of $I_{\text{cal}}$.

A possible limitation of the method is that structural models are required. However, the models can be constructed using low-resolution data and hybrid modeling, e.g., by combining coarse-grained maps with components from homologous PDB domains and models from simulations. However, because the set of models is incomplete (the normalization in Eq. 9 is missing), BioEM cannot give an absolute estimation of the posterior probability but rather a relative value. Thus, model comparison is essential in the BioEM framework.

The BioEM software scales almost ideally with the number of CPU cores and has excellent performance on both CPU and GPU architectures. The code has been optimized for a fast, and accurate analysis of tens of thousands of images, as is required in electron microscopy, and is sufficiently flexible to adjust to diverse research necessities.

In order to cope with the growing heterogeneity of GPU-accelerated systems (in terms of number of CPUs and GPUs within a node and their relative performance) we plan in the future to add an autotuning feature to BioEM which dynamically chooses the optimal distribution of the workload between the CPUs and the GPUs of a node. The distribution can be continuously adjusted based on measurement of the current CPU and GPU image processing rate. Moreover, the autotuning could suggest a good setting for the number of MPI processes per node for the hybrid MPI-OpenMP mode.
The performance of BioEM is dominated to a great extent by the FFT libraries \cite{40} (fftw on the CPU and cuFFT on the GPU, respectively), which are well optimized, leaving little margin for performance improvements. Specifically, we run three consecutive steps for computing the cross-correlation and posterior probability: multiplication in Fourier space, fast Fourier backtransformation, and evaluation of the analytic formula. The multiplication of the images in Fourier space can saturate the memory bandwidth of the GPU. Since we use the cuFFT library which performs the FFT as a black box, the intermediate data must be stored before and after the FFT. However, we only need certain Fourier coefficients but cuFFT computes all of them. It could be possible to modify the cuFFT to extract only the relevant coefficients. However, we have not pursued this yet considering the challenges required to modify and maintain such code.

A different optimization is more promising. It is possible to parallelize over the projections via OpenMP instead of MPI using mutexes to control the probability updates of the observed images. Such an implementation should achieve the same performance as the MPI version but keep the small memory footprint of the OpenMP version. This would be optimal to analyze images where the molecular orientations are not distributed randomly but are correlated. For example, in electron tomography correlations arise in different tilt images of the same particle.

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References

References

\cite{1} W. Wriggers, Conventions and workflows for using situs, Acta Cryst. Sect. D Biol. Cryst. 68 (4) (2012) 344–351. doi:{10.1107/S0907444911049791}
[2] W. Kühlbrandt, The resolution revolution, Science 343 (6178) (2014) 1443–1444. doi:{10.1126/science.1251652}.

[3] X.-C. Bai, G. McMullan, S. H. W. Scheres, How cryo-EM is revolutionizing structural biology, Trends Biochem. Sci. 40 (1) (2015) 49–57. doi:{10.1016/j.tibs.2014.10.005}.

[4] M. Allegretti, N. Klusch, D. J. Mills, J. Vonck, W. Kuehlbrandt, K. M. Davies, Horizontal membrane-intrinsic alpha-helices in the stator a-subunit of an F-type ATP synthase, Nature 521 (7551) (2015) 237–240. doi:{10.1038/nature14185}.

[5] J. Wu, Z. Yan, Z. Li, C. Yan, S. Lu, M. Dong, N. Yan, Structure of the voltage-gated calcium channel cav1.1 complex, Science 350 (6267) (2015) aad2395. doi:10.1126/science.aad2395.

[6] M. Liao, E. Cao, D. Julius, Y. Cheng, Structure of the TRPV1 ion channel determined by electron cryo-microscopy, Nature 504 (7478) (2013) 107–112. doi:{10.1038/nature12822}.

[7] Z. Yan, X.-C. Bai, C. Yan, J. Wu, Z. Li, T. Xie, W. Peng, C.-c. Yin, X. Li, S. H. W. Scheres, Y. Shi, N. Yan, Structure of the rabbit ryanodine receptor RyR1 at near-atomic resolution, Nature 517 (7532) (2015) 50–55. doi:{10.1038/nature14063}.

[8] A. Brown, A. Amunts, X.-C. Bai, Y. Sugimoto, P. C. Edwards, G. Murshudov, S. H. W. Scheres, V. Ramakrishnan, Structure of the large ribosom al subunit from human mitochondria, Science 346 (6210) (2014) 718–722. doi:{10.1126/science.1258026}.

[9] P. Lu, X.-C. Bai, D. Ma, T. Xie, C. Yan, L. Sun, G. Yang, Y. Zhao, R. Zhou, S. H. W. Scheres, Y. Shi, Three-dimensional structure of human gamma-secretase, Nature 512 (7513) (2014) 166–170. doi:{10.1038/nature13567}.

[10] Y. He, C. Yan, J. Fang, C. Inouye, R. Tjian, I. Ivanov, E. Nogales, Near-atomic resolution visualization of human transcription promoter opening, Nature 533 (7603) (2016) 359–365. doi:{10.1038/nature17970}.
[11] A. R. Faruqi, R. Henderson, Electronic detectors for electron microscopy, Curr. Opin. Struct. Biol. 17 (5) (2007) 549–555. doi:{10.1016/j.sbi.2007.08.014}

[12] J. M. de la Rosa-Trevin, J. Oton, R. Marabini, A. Zaldivar, J. Vargas, J. M. Carazo, C. O. S. Sorzano, Xmipp 3.0: An improved software suite for image processing in electron microscopy, J. Struct. Biol. 184 (2) (2013) 321–328. doi:{10.1016/j.jsb.2013.09.015}

[13] G. Tang, L. Peng, P. R. Baldwin, D. S. Mann, W. Jiang, I. Rees, S. J. Ludtke, EMAN2: An extensible image processing suite for electron microscopy, J. Struct. Biol. 157 (1) (2007) 38–46. doi:{10.1016/j.jsb.2006.05.009}

[14] S. H. W. Scheres, RELION: Implementation of a Bayesian approach to cryo-EM structure determination, J. Struct. Biol. 180 (2012) 519–530. doi:{10.1016/j.jsb.2012.09.006}

[15] S. H. W. Scheres, A Bayesian view on cryo-em structure determination, J. Mol. Biol. 415 (2012) 406–418. doi:{10.1016/j.jmb.2011.11.010}

[16] X. Li, N. Grigorieff, Y. Cheng, GPU-enabled FREALIGN: Accelerating single particle 3D reconstruction and refinement in Fourier space on graphics processors, J. Struct. Biol. 172 (2010) 407–412. doi:{10.1016/j.jsb.2010.06.010}

[17] T. V. Hoang, X. Cavin, P. Schultz, D. W. Ritchie, gEMpicker: a highly parallel GPU-accelerated particle picking tool for cryo-electron microscopy, BMC Struct. Biol. 13 (2013) 25. doi:{10.1186/1472-6807-13-25}

[18] H. D. Tagare, A. Barthel, F. J. Sigworth, An adaptive expectation-maximization algorithm with GPU implementation for electron cryomicroscopy, J. Struct. Biol. 171 (3) (2010) 256–265. doi:{10.1016/j.jsb.2010.06.004}

[19] J. E. Stone, R. McGreevy, B. Isralevitz, K. Schulten, GPU-accelerated analysis and visualization of large structures solved by molecular dynamics flexible fitting, Farad. Discuss. 169 (2014) 265–283. doi:{10.1039/c4fd00005f}
[20] X. Li, S. Q. Zheng, K. Egami, D. A. Agard, Y. Cheng, Influence of electron dose rate on electron counting images recorded with the K2 camera, J. Struct. Biol. 184 (2013) 251–260. doi:{10.1016/j.jsb.2013.08.005}

[21] S. H. W. Scheres, Beam-induced motion correction for sub-megadalton cryo-EM particles, Elife 3 (2014) e03665. doi:{10.7554/eLife.03665}

[22] X. Li, P. Mooney, S. Zheng, C. R. Booth, M. B. Braunfeld, S. Gubbens, D. A. Agard, Y. Cheng, Electron counting and beam-induced motion correction enable near-atomic-resolution single-particle cryo-EM, Nat. Methods 10 (6) (2013) 584–590. doi:{10.1038/nmeth.2472}

[23] M. Allegretti, D. J. Mills, G. McMullan, W. Kuehlbrandt, J. Vonck, Atomic model of the F-420-reducing [NiFe] hydrogenase by electron cryo-microscopy using a direct electron detector, Elife 3 (2014) e01963. doi:{10.7554/eLife.01963}

[24] Z. Wang, C. F. Hryc, B. Bammes, P. V. Afonine, J. Jakana, D.-H. Chen, X. Liu, M. L. Baker, C. Kao, S. J. Ludtke, M. F. Schmid, P. D. Adams, W. Chiu, An atomic model of brome mosaic virus using direct electron detection and real-space optimization, Nature Commun. 5 (2014) 5808. doi:{10.1038/ncomms5808}

[25] X.-C. Bai, I. S. Fernandez, G. McMullan, S. H. W. Scheres, Ribosome structures to near-atomic resolution from thirty thousand cryo-EM particles, Elife 2 (2013) e00461. doi:{10.7554/eLife.00461}

[26] F. J. Sigworth, Principles of cryo-EM single-particle image processing, Microscopy 65 (1) (2016) 57–67. doi:{10.1093/jmicro/dfv370}

[27] P. Rosenthal, R. Henderson, Optimal determination of particle orientation, absolute hand, and contrast loss in single-particle electron cryo-microscopy, J. Mol. Biol. 333 (2003) 721–745. doi:{10.1016/j.jmb.2003.07.013}

[28] R. van Antwerpen, Preferred orientations of LDL in vitreous ice indicate a discoid shape of the lipoprotein particle, Arch. Biochem. Biophys. 432 (1) (2004) 122–127. doi:{10.1016/j.abb.2004.08.031}
[29] J. Zhang, M. L. Baker, G. F. Schroeder, N. R. Douglas, S. Reissmann, J. Jakana, M. Dougherty, C. J. Fu, M. Levitt, S. J. Ludtke, J. Frydman, W. Chiu, Mechanism of folding chamber closure in a group II chaperonin, Nature 463 (7279) (2010) 379–383. doi:{10.1038/nature08701}.

[30] R. M. B. Hoffman, T. M. A. Blumenschein, B. D. Sykes, An interplay between protein disorder and structure confers the Ca2+ regulation of striated muscle, J. Mol. Biol. 361 (4) (2006) 625–633. doi:{10.1016/j.jmb.2006.06.031}.

[31] J. Peschek, N. Braun, J. Rohrberg, K. C. Back, T. Kriehuber, A. Kas- tenmueller, S. Weinkauf, J. Buchner, Regulated structural transitions unleash the chaperone activity of alpha B-crystallin, Proc. Natl. Acad. Sci. USA 110 (40) (2013) E3780–E3789. doi:{10.1073/pnas.1308898110}.

[32] N. Elad, D. K. Clare, H. R. Saibil, E. V. Orlova, Detection and separation of heterogeneity in molecular complexes by statistical analysis of their two-dimensional projections, J. Struct. Biol. 162 (2008) 108–120. doi:{10.1016/j.jsb.2007.11.007}.

[33] D. Elmlund, H. Elmlund, High-resolution single-particle orientation refinement based on spectrally self-adapting common lines, J. Struct. Biol. 167 (2009) 83–94. doi:{10.1016/j.jsb.2009.04.009}.

[34] D. Elmlund, H. Elmlund, Cryogenic electron microscopy and single-particle analysis, Annu. Rev. Biochem. 84 (2015) 499–517. doi:{10.1146/annurev-biochem-060614-034226}.

[35] P. Cossio, G. Hummer, Bayesian analysis of individual electron microscopy images: Towards structures of dynamic and heterogeneous biomolecular assemblies, J. Struct. Biol. 184 (3) (2013) 427–437. doi:{10.1016/j.jsb.2013.10.006}.

[36] R. Wade, A brief look at imaging and contrast transfer, Ultramicroscopy 46 (1992) 145–156. doi:{10.1016/0304-3991(92)90011-8}.

[37] P. A. Penczek, Image restoration in cryo-electron microscopy, Methods Enzymol. 482 (2010) 35–72. doi:{10.1016/S0076-6879(10)82002-6}.
[38] F. J. Sigworth, P. C. Doerschuk, J.-M. Carazo, S. H. W. Scheres, An introduction to maximum-likelihood methods in cryo-EM, in: Methods Enzymol., Vol. 482, Elsevier, 2010, pp. 263–294. doi:{10.1016/S0076-6879(10)82011-7}

[39] A. Yershova, S. Jain, S. M. LaValle, J. C. Mitchell, Generating uniform incremental grids on SO(3) using the Hopf fibration, Int. J. Robot. Res. 29 (7) (2010) 801–812. doi:{10.1177/0278364909352700}

[40] M. Frigo, S. G. Johnson, The design and implementation of FFTW3, Proc. IEEE 93 (2) (2005) 216–231. doi:{10.1109/JPROC.2004.840301}

[41] G. Tang, L. Peng, P. R. Baldwin, D. S. Mann, W. Jiang, I. Rees, S. J. Ludtke, EMAN2: An extensible image processing suite for electron microscopy, J. Struct. Biol. 157 (1) (2007) 38–46. doi:{10.1016/j.jsb.2006.05.009}

[42] W. Kahan, Further remarks on reducing truncation errors, Commun ACM 8 (1) (1965) 40–48. doi:{10.1145/363707.363723}

[43] D. J. Mills, S. Vitt, M. Strauss, S. Shima, J. Vonck, De novo modeling of the F-420-reducing [NiFe]-hydrogenase from a methanogenic archaeon by cryo-electron microscopy, Elife 2 (2013) e00218. doi:{10.7554/eLife.00218}

[44] S. J. Ludtke, M. L. Baker, D.-H. Chen, J.-L. Song, D. T. Chuang, W. Chiu, De novo backbone trace of GroEL from single particle electron cryomicroscopy, Structure 16 (2008) 441–448. doi:{10.1016/j.str.2008.02.007}