Reference Manual: Functional Tensor-Train Chebyshev Method for Multidimensional Quantum Dynamics Simulations

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Introduction

We present tensor-train functional tensor-train Chebyshev (FTTC) codes for the simulation of exact molecular dynamics in high dimensionality. These codes employ the continuous analogue of the low-rank tensor train decomposition (a relative of matrix product states).

An example is given for simulation of a highly multidimensional model of hydrogen bonding in adenine-thymine DNA base pairs.

Dependencies

The C functional tensor-train Chebyshev (FTTC) code requires the following:

- C compiler
- CMake
- Compressed Continuous Computation (C3) library (which requires BLAS, LAPACK, and SWIG-python [optional])

The Python functional tensor-train Chebyshev (FTTC) code requires the following:

- Miniconda
- Compressed Continuous Computation (C3) library

The tensor-train Chebyshev (TTC) code requires the following:

- MATLAB (tested for R2018a and R2021a)
- H-Tucker Tensor Toolbox (path must be included in MATLAB code at runtime)
- TT-Toolbox (path must be included in MATLAB code at runtime)

Comparison of the tensor-train and function-train codes is facilitated with the following:

• Gnuplot

C Functional Tensor-Train Chebyshev (FTTC) Code Instructions

Code for C FTTC propagation is included in the folder FTTC. Details on installation of dependences and troubleshooting are included in the readme.txt file contained in the folder.

The program is compiled with the commands:

cmake .

make

The program is then run as follows:

./fttc

The program has been tested for macOS Mojave and Big Sur. The following steps can be employed for troubleshooting if compilation or runtime errors occur.

- If errors concerning missing c3axpy arguments in 2dpro.c appear when cmake is run and if epsilon is zero, add NULL as the missing argument in all instances.
- If the file Accelerate is not found by the compiler, but Accelerate.tbd exists in any folder, copy the Accelerate.tbd file into the same folder and rename it Accelerate.
- When updating the cmake file, remove outdated cmake output as follows:

rm - r./CMakeFilesCMakeCache.txt.

- If make yields an error that c3/array.h is missing, -1c3 cannot be found, etc., try the following:
 - Ensure that c3 is downloaded. Then update the version of cmake as follows:cmake_minimum_required(VERSION 3.0)
 - Change the final argument of target_link_libraries to the following: \$ {c3lib}.
 - Update the location of the C3 library lib files in CMakeLists.txt inside the if APPLE loop as follows:

```
set (CMAKE_LIBRARY_PATH ${CMAKE_LIBRARY_PATH} /usr/local/lib)
```

- Update the location of the C3 library lib files in CMakeLists.txt inside the if APPLE loop as follows:

```
find_library(c3lib c3 PATHS /usr/local/lib)
include_directories(/usr/local/include)
```

Propagation is controlled according to the following parameters:

- NCHEB Number of Chebyshev polynomials
- NE Number of grid points in each dimension
- NX Total number of grid points in the $x_1 x_2$ grid $(i.e., NE^2)$
- DIM Number of dimensions
- NSTEPS Number of propagation steps
- NDUMP Number of time steps after which the wavefunction is saved to file
- dis Initial displacement parameter
- dt Time step
- \bullet m Mass
- 1b Minimum value of the position-space grid
- ub Maximum value of the position-space grid

The functional tensor-train approximation can be modulated by changing:

- init rank Initial rank
- round_eps Rounding accuracy parameter

and the final arguments of the functions:

- function_train_round_maxrank_all Rounding maximum rank
- c3approx set adapt kickrank Cross approximation kick rank
- c3approx_set_cross_maxiter Cross approximation maximum iterations
- c3approx_set_cross_tol Cross approximation tolerance parameter
- c3approx_set_round_tol Cross approximation rounding tolerance parameter
- c3approx_set_adapt_maxrank_all Cross approximation maximum rank

The program outputs the files:

- timings Time required for each propagation step
- xi Autocorrelation function at each time step (real and imaginary parts)
- norm Norm at each time step
- wave.* Probability density at the *th multiple of printing frequency NDUMP

Example output is included in folder ExampleOutput. The output can be visualized with the scripts described in the final section of this reference manual.

The initial wavefunction can be modified in function GS and GSfix, and the potential can be modified in function Vpot (which requires an update of the maximum and minimum potential energy values Vmax and Vmin, respectively).

Python Functional Tensor-Train Chebyshev (FTTC) Code

Instructions

The Python FTTC propagation code is provided in the folder entitled FTTCPython. A readme.txt file is included with information on troubleshooting installation of the required C3 package. The program has been tested for macOS Big Sur.

Installation of the required c3py library is facilitated by creating a conda environment as follows:

```
conda create - n c3env python = 3.7 numpy scipy conda activate c3env
```

The c3py library is then set up with the following commands:

```
python setup.py
build python setup.py install
pip install matplotlib
```

Thereafter the environment is entered with:

```
conda activate c3env
```

To run the program, use the command:

```
python fttc.py
```

The propagation parameters follow those of the C program:

- NCHEB Number of Chebyshev polynomials
- NE Number of grid points in each dimension
- NX Total number of grid points in the $x_1 x_2$ grid $(i.e., NE^2)$
- DIM Number of dimensions
- NSTEPS Number of propagation steps
- NDUMP Number of time steps after which the wavefunction is saved to file
- dis Initial displacement parameter
- dt Time step
- \bullet m Mass
- xmin Minimum value of the position-space grid

• xmax – Maximum value of the position-space grid

The functional tensor-train approximation can be modulated by changing:

- init_rank Initial rank
- round_eps Rounding accuracy parameter
- rmax Maximum rank

and the final arguments of the functions:

- maxrank Cross approximation maximum rank
- kickrank Cross approximation kick rank
- maxiter Cross approximation maximum iterations
- cross_tol Cross approximation tolerance parameter
- round tol Cross approximation rounding tolerance parameter

Likewise, the program outputs the files:

- xi.npy Autocorrelation function at each time step (real and imaginary parts)
- norm.npy Norm at each time step
- wave * .npy Probability density at the *th multiple of printing frequency NDUMP

Example output is included in the FTTCPython folder in the subfolder ExampleOutput. Alongside the data files are scripts for visualization of the results. The scripts can be employed in the same fashion as the comparison scripts discussed in the final section of this reference manual.

The initial wavefunction can be changed with functions GS and GSfix (ex., to begin with a displaced initial state with position given by the parameter dis). The potential can also be changed by modifying function Vpot and its corresponding maximum potential energy value Vmax and minimum potential energy value Vmin.

Tensor Train Chebyshev (TTC) Code Instructions

Code for simulation of TTC dynamics is contained in the folder TTC. Tensor-train split operator Fourier transform (TT-SOFT) dynamics are also performed for comparison.

Dynamics

Tensor train Chebyshev propagation is performed by running tt_CHEBSOFT.m in MATLAB.

Propagation parameters are defined as follows:

- d Dimensionality
- ns Number of time steps
- dump Number of time steps after which the wavefunction is saved to file
- eps Precision of tensor train rounding
- ceps Precision of tensor train rounding for exponential
- rmax Maximum tensor train rank
- alpha Harmonic oscillator eigenstate parameter $\alpha = m\omega/\hbar$, which determines the width of the initial wavefunction
- x0 and x0array Position parameters for the initial wavefunction
- p0 Initial momentum
- \bullet nx = 2^q Number of grid divisions nx in position space as specified by the integer number of quantics q
- np Number of grid divisions in momentum space
- L Length of the simulation grid in each physical dimension

- dx Position grid spacing
- dp Momentum grid spacing
- tau Time step
- Npoly Number of Chebyshev polynomials
- m Mass in each physical dimension (currently only implemented for unit mass)
- x Position grid in each direction
- p Momentum grid in each direction
- verticalscale Vertical stretch parameter for DNA potential
- gamma Coupling parameter for DNA potential

The code visualizes the $x_1 - x_2$ slice of the potential energy surface for two-, three-, and four-dimensional potential energy surfaces. Figures comparing TTC and SOFT are printed to screen: the autocorrelation function (real and imaginary parts) at each time step; the probability density every dump time steps; and, after the run is complete, the norm at each time step. In addition, the following output files are created:

- resultofstep * *.mat Saved MATLAB variables at time step * * +1
- resultofstepfinal.mat Saved MATLAB variables after propagation

Example output images and files are given in the folder entitled ExampleOutput.

The code requires H-Tucker Tensor Toolbox and TT-Toolbox to be included in the path specified at the beginning of the tt_CHEBSOFT.m file. The potential energy surface can be changed by revising the definition of the tensor train PE_tt and the minimal and maximal energies Emin and Emax. An alternative initial wavefunction can be defined in the function in file psi0.m.

Conversion of Output Data Files

To facilitate the comparison of tensor train and function train data, after propagation is complete, the GnuplotConversion.m code is run in MATLAB given the following input data files held in the same folder:

- resultofstepfinal.mat Saved variables at final time
- resultofstep * *.mat Saved variables at time step **

Example output files are provided in folder ExampleOutput.

The code produces the following data files in Gnuplot format:

- norm.dat Norm at each time (column 1) for SOFT (column 2) and TT (column 3)
- autocorrelation.dat Autocorrelation at each time (column 1) for SOFT (real part in column 2 and imaginary part in column 3) and TTC (real part in column 4 and imaginary part in column 5)
- wave **.dat Absolute value squared of the density in directions x_0 (column 1) and x_1 (column 2) at chosen time step ii=**+1 for SOFT (column 3) and TTC (column 4), where ii=**+1 corresponds to the ii^{th} element in array numlist

The path at the beginning of the file must correspond to the location of the TT-Tensor Toolbox library. The conversion code can be implemented for visualization of other time steps through modification of the array numlist, which contains the number of each time step under consideration. The conversion code is currently implemented in fifty dimensions, as slices of the density are computed as follows:

```
 v1 = reshape \left( abs \left( full \left( G_{tt} \left( : ; : ; ... \right. \right. \right. \right. \right. \\  nx/2 , ... \\  nx/2 , ... \\  nx/2 , ... \\  nx/2 , ... \\  x/2 , nx/2 , ... \\  x/2 , nx/2 , ... \\  x/2 , nx/2 , ... \\  x/2 , nx/2 , ... \\  x/2 , nx/2 , nx/
```

where the 48 nx/2 entries refer to the index of the fixed position of the bath modes. The fixed position of the bath modes can be changed by replacing the nx/2 with the index of choice, and the calculation can be updated for B bath modes by replacing the 48 nx/2 entries with B nx/2 entries.

Visualization of these data files with Gnuplot is discussed in "FTTC/TTC/SOFT Comparison Instructions" at the end of this reference manual.

FTTC/TTC/SOFT Comparison Instructions

The results of the C FTTC and TTC simulations can be visualized with the Gnuplot scripts in folder Comparisons. For visualization, the TTC program data files wave. *.dat and autocorrelation.dat must be included in the folder TTC, and the C FTTC program data files wave. * and xi must be included in the folder FTTC.

The probability density comparison is created with the following command:

gnuplot wavefunctionsnapshots.gpt

which produces the files wavefunctionsnapshots.eps and wavefunctionsnapshots.pdf.

The autocorrelation function comparison is created with the following command:

gnuplot autocorrelationcompare.gpt

which produces the files autocorrelationcompare.eps and autocorrelationcompare.pdf.

Example output is given in folder ${\tt ExampleOutput}$ for verification.