TBTK - Tight Binding ToolKit

A c++ library for solving tight-binding models

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Model

Designed to solve any bilinear Hamiltonian

$$H=\sum_{ij}a_{ij}c_i^{\dagger}c_j.$$

Model: Indices and HoppingAmplitudes

Flexible indices allows for general geometries to be defined that are not necessarily indexable by a single type of index. Theses are generally reffered to as 'to' and 'from' indices, and are used to specify 'HoppingAmplitudes' a_{ij} .

$$H=\sum_{ij}a_{ij}c_i^{\dagger}c_j.$$

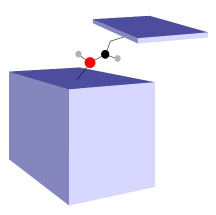
Index	Also known as	Example
i	to	$\{subsystem, x, y, spin\}$
j	from	{subsystem, x, y, z, orbital, spin}

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Model: Example

Consider for example a system consisting of three subsystems of different dimension.



Model: Example

A possible indexing scheme might look like this.

```
3D bulk: \{0, x, y, z, \text{ orbital}, \text{spin}\},
1D molecule: \{1, x, \text{ orbital}, \text{spin}\},
2D sheet: \{2, x, y, \text{spin}\}.
```

Model: Important functions

```
Setup model
    Model model;
Add hopping amplitude
    model.addHA(HoppingAmplitude(-mu, {x, y, s}, {x, y, s}));
Add hopping amplitude and Hermitian conjugate
    model.addHAAndHC(HoppingAmplitude(-t,
                                        \{x+1, y, s\},\
                                        \{x, y, s\}));
```

Construct model. Hilbert space is created.

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model.construct():

Model: Example

```
Model model;
for(int x = 0; x < SIZE_X; x++){
    for(int y = 0; y < SIZE_Y; y++){
        for(int s = 0; s < 2; s++){
            model.addHA(-mu, \{x, y, s\}, \{x, y, s\});
            if(x+1 < SIZE X)
                model.addHAAndHC(-t, \{x+1, y, s\}, \{x, y, s\});
model.construct();
```

Solvers

- DiagonalizationSolver (Eigenvalues and eigenvectors)
- ChebyshevSolver (Green's function)

DiagonalizatioSolver

The DiagonalizationSolver can be used to

- Calculate eigenvalues and eigenvectors.
- Self-consistently or non self-consistently.
- Calculate custom physical quantities by accessing the wave function directly using physical indices...
- ...or use a "property extractor" to extract common properties such as eigenvalues, DOS, spin-polarized LDOS, magnetization, etc.
- Have been tested for Hamiltonians with a basis size of up to 14000

dSolver.setSCCallabck(scCallback):

```
Create diagonalization solver
    DiagonalizationSolver dSolver;
Set model
    dSolver.setModel(&model);
Diagonalize Hamiltonian. Is done self-consistently if a
self-consistenct callback has been set
    dSolver.run();
Set self-consisteny callback, where scCallback is a user
defined callback function
```

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```
Set maximum number of self-consistency iterations
    dSolver.setMaxIterations(MAX_ITERATIONS);
Get eigenvalues
    dSolver.getEigenValues();
Get amplitude of eigenstate n at index (x, y, s)
    dSolver.getAmplitude(n, {x, y, s});
Create property extractor
    PropertyExtractor pe(&dSolver);
Get eigenvalues
    double *ev = pe.getEV();
```

```
Calculate DOS with a upper and lower energy cutoff of U_LIM
and L_LIM, respectively, and a resolution of RESOLUTION
number of points between L_LIM and U_LIM.
   double *dos = pe.calculateDOS(U_LIM, L_LIM, RESOLUTION);
Calculate electron density. Will be calculated for all x and
y, and spin-indices will be summed to create a total density.
   double *density = pe.calculateDensity(
                          {IDX_X, IDX_Y, IDX_SUM_ALL},
                          {SIZE_X, SIZE_Y, 2});
```

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Calculate spin-polarized LDOS. Will be calculated for all y along the line $x = SIZE_X/2$. Note that IDX_X is used for the y-index here. IDX_X, IDX_Y, IDX_Z, does for the property extractor refer to the first, second, and third loop index, and the y index is the first loop index when x is fixed. The upper and lower energy cutoff is set by U_LIM and L_LIM, respectively, and RESOLUTION number of points are used between the two limits.

```
double *sp_ldos = pe.calculateSP_LDOS(
                      {SIZE_X/2, IDX_X, IDX_SPIN},
                      {1, SIZE_Y, 2}
                      U LIM, L LIM, RESOLUTION):
```

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DiagonalizationSolver: Example

```
//Non self-consistently
DiagonalizationSolver dSolver;
dSolver.setModel(&model);
dSolver.run()
//Self-consistently (requires scCallback to be defined).
DiagonalizationSolver dSolver;
dSolver.setModel(&model);
dSolver.setSCCallback(scCallback):
dSolver.setMaxIterations(MAX_ITERATIONS);
dSolver.run():
```

Chebyshev solver

$$G_{ij}(E) = \frac{-2i}{\sqrt{1-E^2}} \sum_{n}^{\infty} b_{ij}^{(n)} e^{-in \operatorname{arccos}(E)}.$$

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- Calculates the Green's function.
- Can be done either on CPU or GPU
- Works for Hamiltonians with a basis size of at least 4 million

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```
Create Chebyshev solver
    ChebyshevSolver cSolver;
Set model
    cSolver.setModel(&model);
Calculate Chebyshev coefficeints for G_{ij}, where
i = (x, y, 0), j = (x+1, y, 1)
    complex<double> coefficients[NUM_COEFFICIENTS];
    cSolver.calculateCoefficeints({x, y, 0}, {x+1, y, 1},
                                   coefficients.
                                   NUM COEFFICIENTS):
    cSolver.calculateCoefficeintsGPU(\{x, y, 0\}, \{x+1, y, 1\},
                                      coefficients.
                                      NUM COEFFICIENTS):
```

```
Evaluate Green's function without using lookup table
    complex<double> greensFunction[ENERGY_RESOLUTION];
    cSolver.generateGreensFunction(greensFunction,
                                    coefficients,
                                   NUM COEFFICIENTS.
                                   ENERGY_RESOLUTION);
Setup lookup table
    cSolver.generateLookupTable(NUM_COEFFICIENTS,
                                ENERGY RESOLUTION):
Load lookup table to GPU
    cSolver.loadLookupTableGPU();
```

cSolver.destroyLookupTableGPU();