NQFT

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ONE

NQFT

1.1 nqft package

1.1.1 Submodules

1.1.2 nqft.functions module

This module contains global functions used in the expression and operation on Hubbard hamiltonian.

It is also used more globally to perform operations on data files containing spectral function.

 $\texttt{nqft.functions.add_column}(\mathit{file:}\ \mathit{str}, \mathit{column:}\ \mathit{array}, \mathit{idx:}\ \mathit{int}) \rightarrow \mathsf{None}$

Insert new data to given text file as a column.

Parameters

- **file** (str, default=None) File in which add column.
- **column** (*np.array*, *default=None*) Data column to add.
- idx (int, default=0) Index of the new column inside current file.

 $nqft.functions.delta(j: int, k: int) \rightarrow float$

Kronecker delta function.

Parameters

- **j** (int, default=None) First indice.
- **k** (int, default=None) Second indice.

Returns

_

Return type

float (0.0 or 1.0)

 $nqft.functions.find_nearest(array, value) \rightarrow int$

Finds index in given array of the closest value of parameter 'value'.

Parameters

- array (array-like, shape=(m, n), default=None) Array in which search for the value.
- value (int, float, default=None) Value to search for in array.

Returns

idx – Index at which user can find the closest value in array.

Return type

int

nqft.functions.flatten_fermi_arc($save_path='./nqft/Data/fermi_arc_data_1D/'$, type='text') \rightarrow None Saves Peter's Fermi arc numpy 2D arrays files as 1D arrays numpy files so they can be read in Rust/C.

Parameters

- **save_path** (*str*, *default="./nqft/Data/fermi_arc_data_1D/"*) Path where to save flatten arrays.
- **type** (*str*, *default="text"*) Format in which save flattened arrays (ex: type="npy" leads to .npy files)

Return type

None

 $nqft.functions.plot_hall(files=['./nqft/Data/hall.txt'], x='doping') \rightarrow Figure$

Plots hall coefficient as a function of interaction.

Parameters

- **files** (list[str], size=N, default=["./nqft/Data/hall.txt"]) Text files to plot.
- **x** (*str*, *default='doping'*) Type of x coordinate used to plot (setting up the legend label).

 $nqft.functions.read_fermi_arc(path='./nqft/Data/fermi_arc_data/') \rightarrow dict$

Reads Peter's data on spectral weight at Fermi level for a given number of sites.

Parameters

path (str, default="./nqft/Data/fermi_arc_data/") - Path to data directory.

Returns

arcs – A dict containing all spectral functions.

Return type

dict, size=6

 $nqft.functions.read_locals(shape: tuple[int], interaction: float) \rightarrow dict$

Reads local model's spectral functions for given shape and electronic density.

Parameters

- **shape** (tuple, size=2, default=None) Shape of the clusters.
- **filling** (*int*, *default=None*) Number of electrons in each cluster.

Returns

spectrums – Dictionnary containing all spectrums (values) associated with a specific density (keys).

Return type

dict

 $nqft.functions.scalar(m: Qobj, n=None) \rightarrow float$

Computes scalar product for Fock space vectors such as

 $scalar(m, n) = \langle m \mid n \rangle$.

Parameters

- m (qutip.Qobj, default=None) Bra on which perform scalar product.
- n (qutip.Qobj, default=None) Ket on which perform scalar product.

Returns

- Result of scalar product.

Return type

int, float

1.1.3 nqft.hall_effect module

Hall effect in cuprates with an incommensurate collinear spin-density wave.

Bases: object

Model instance to determine Hall coefficient and density from tight-binding hamiltonian.

hoppings

Hopping amplitude coefficients.

Type

tuple, size=3, default=None

broadening

Lorentzian broadening module.

Type

float, default=None

omega

Frequency at which we observe the fermi surface.

Type

float, default=None

mus

Chemical potential interval values and the interval between each element of an hypothetical array.

```
Type tuple, size=3, default=(-4, 4, 0.02)
```

resolution

Resolution of phase space (k_x, k_y) .

Type

int, default=4

use_peters

Determines if model's based on Peter R. spectral functions (fermi arcs)

Type

bool, default=False

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```
get_density() \rightarrow array
     Computes electron density.
          Returns
              density – Electron density.
          Return type
              np.array, size=M
get_hall_nb() \rightarrow array
     Computes Hall number.
          Returns
              n H – Hall number.
          Return type
              np.array, size=M
plot\_spectral\_weight(mu: float, key=None) \rightarrow Figure
     Ouputs the spectral weight as a 2D numpy array.
          Returns
              - - 2D graph of spectral weight.
          Return type
              plt.figure
sigma_ii(variable: str) \rightarrow array
     Computing longitudinal conductivity at zero temperature in the zero-frequency limit when interband tran-
     sitions can be neglected.
          Parameters
              variable (str, default=None) – Axis on which compute conductivity.
          Returns
              conductivity
          Return type
              float
sigma_ij() \rightarrow array
     Computing transversal conductivity at zero temperature in the zero-frequency limit when interband transi-
     tions can be neglected.
          Returns
              conductivity
          Return type
```

nqft.hall_effect. $dE(hop_amps: tuple, kx: ndarray, ky: ndarray, mu: array) <math>\rightarrow$ tuple Outputs model's energies and it's derivatives.

Parameters

float

- hop_amps (tuple, default=None) Hopping amplitudes coefficients.
- **kx** (np.ndarray, shape=(N, N), default=None) kx space as a 2D array.
- **ky** (np.ndarray, shape=(N, N), default=None) ky space as a 2D array.
- **mu** (*np.array*, *size=M*, *default=None*) Chemical potential values array.

```
Returns
```

E, **d**Es – Energies and it's derivatives in a tuple.

Return type

tuple, size=2

 $nqft.hall_effect.fit_lin(x: array, a: float, b: float) \rightarrow array$

Linear function.

nqft.hall_effect.**get_spectral_weight**(*omega: float, eta: float, E: ndarray*) → ndarray

Ouputs the spectral weight as a 3D numpy array.

Parameters

- omega (float, default=None) Frequency at which we observe the fermi surface.
- eta (float default=None) Lorentzian broadening module.
- **E** (np.ndarray. shape=(M, N, N), default=None) Eigenenergies of the system as a 3D numpy array.

Returns

A – Spectral weight.

Return type

np.ndarray, shape=(M, N, N)

nqft.hall_effect.timeit(func)

1.1.4 nqft.hamiltonian module

This module tests Qutip python library functions, objects and attributes in the context of Hubbard model using square fermion networks.

```
class nqft.hamiltonian.Network(sites_nb: int)
```

Bases: object

A class representing a fermionic many-body problem Network

sites_nb

Number of sites of the network

Type

int, default=None

vaccum

Vaccum state of 2D Fock space.

Type

Qobj, shape=(2, 1)

creation

Creation operator in second quantization formalism.

Type

Qobj, shape=(2, 2)

anihilation

Anihilation operator in second quantization formalism.

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I

Identity operator.

Type

Qobj, shape=(2, 2)

 $get_e(H: Qobj, states: list) \rightarrow float$

Outputs a matrix element (energy) from the hamiltonian using given kets on which perform projection.

Parameters

- H (Qobj, shape=(4^self.sites, 4^self.sites), default=None) Fermions network hamiltonian.
- states (array-like, shape=(2, 1), default=None) Vectors used to process scalar product on H.

(ex: states=[bra, ket] as integers to convert from binairy)

Returns

E – Representation of projected vectors on H (energy).

Return type

int

Examples

```
>>> N = Network(sites_nb=2)
>>> Hamiltonian = N.get_hamiltonian(model="Hubbard", U=2, t=1)
>>> N.get_e(H=Hamiltonian, states=[15, 15])
2.0
```

```
get_hamiltonian(model: str, U: int, **kwargs) \rightarrow Qobj
```

Outputs the hamiltonian of fermion network using specified many-body model.

Parameters

- **model** (*str*, *default=None*) Fermions network configuration.
- **U** (int, default=None) Module of interaction between fermions.
- **kwargs –

t: int, default=None

Probability amplitude for fermions to jump.

Returns

H – Tensor object representing hamitonian for given 'model'.

Return type

```
qutip.Qobj, shape=(4^SITES, 4^SITES)
```

Examples

```
>>> N = Network(sites_nb=4)
>>> N.get_hamiltonian(model="Hubbard", U=1, t=1)
Quantum object: dims = [[2, 2, 2, 2, 2, 2, 2, 2],
[2, 2, 2, 2, 2, 2, 2]], shape = (256, 256), type = oper,
isherm = True Qobj data =
[[0. 0. 0. ... 0. 0.
                          0.]
[0. 0. -1. ... 0.
                      0.
                          0.]
         0. ... 0.
[0.-1.
[ 0.
          0. ... 3. -1.
          0. ... -1.
[ 0.
                      3.
                          0.]
      0.
          0. ...
                  0.
```

```
get_state(state: int, type='ket') \rightarrow Qobj
```

Gives array-like representation of given state using Qutip 'tensor' operation.

Parameters

- **state** (*int*, *default=None*) Integer representing state number in Fock space.
- **type** (*str*, *default='ket'*) Type of vector outputed (bra, ket)

Returns

bra, **ket** – Full vector representation.

Return type

```
qutip.Qobj, shape=(4^SITES, 1)
```

Examples

```
>>> N = Network(sites_nb=1)
>>> self.get_state(state=2)
Quantum object: dims = [[2, 2], [1, 1]], shape = (4, 1), type = ket
Qobj data =
[[0.]
[0.]
[1.]
[0.]]
```

 $lanczos(H: Qobj, iterations: int, init_state=None) \rightarrow tuple$

Implementation of Lanczos algorithm for Network hamiltonian.

Parameters

- H (Qobj, shape=(4^self.sites, 4^self.sites), default=None) Fermions network hamiltonian.
- **iterations** (*int*, *default=None*) Number of iterations on which perform the algorithm.
- init_state (QObj, shape=(4^self.sites, 1) default=random) Initial quantum state to start the first iteration.

Returns

- Respectively the maximum eigenvalue and the associated eigenvector.

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```
Return type tuple, shape=(1, 2)
```

Examples

```
>>> N = Network(sites_nb=2)
>>> Hamitonian = N.get_hamiltonian(model="Hubbard", U=1, t=1)
>>> N.lanczos(H=Hamiltonian, iterations=10)
(2.561552779602264, Quantum object: dims = [[10], [1]],
shape = (10, 1), type = ket Qobj data =
[[0.43516215]
 [0.78820544]
 [0.43516215]
 Γ0.
            1
 Γ0.
            1
 [0.
            ]
 Γ0.
 [0.
            ]
 Γ0.
            1
 [0.
            ]])
```

1.1.5 nqft.qcm module

This module is dedicated to 'pyqcm' experimentation.

Parameters

Return type dict

```
class nqft.qcm.QcmModel(shape: tuple[int], filling: int, interaction: float, hoppings: tuple[float], broadening:
                               float, w: float, mu: float, resolution: int, tiling_shift: bool, show_spectrum=False,
                               overwrite=False)
      Bases: object
      Docs
      \texttt{get\_cluster\_averages}(operators: list[str]) \rightarrow dict
           Computes single cluster operator averages.
                Parameters
                    operators (list, default=None) – Operators to keep in output dictionnary.
                Returns
                    out_dict - Dictionnary containing operator names as keys and tuples as values representing
                    operator average and it's variance.
                Return type
                    dict
      get_lattice_averages(operators: list[str]) \rightarrow dict
           Computes lattice operator averages.
```

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operators (list, default=None) – Operators on which get average.

_ – Dictionnary containing operator names as keys and average as values.

```
plot_spectrums(peter\_key: str, type='contourf', save=False) \rightarrow Figure
```

Opens spectrums from (2x2, 3x4, 4x4) models and Peters spectrums array to compare the plot for given parameters.

Parameters

- peter_key (str, default=None) Determines which of Peter's array to compare. ('N24', 'N28', 'N30', 'N32', 'N36')
- **type** (*str*, *default='contourf'*) Spectral plot type (contourf, pcolormesh).
- **save** (*bool*, *default=False*) Saves of not the output plot.

Returns

Return type

matplotlib.pyplot.Figure object

```
nqft.qcm.build\_matrix(shape: tuple) \rightarrow list
```

Gives a coordinates matrix of a cluster having shape[0]*shape[1] sites.

Parameters

```
shape (tuple, shape=(2, 1), default=None) – Shape of sites network.
```

Returns

array - Nested lists of coordinates.

Return type

list, shape=(*shape)

Examples

```
>>> build_matrix(shape=(2, 2))
[[0, 0, 0], [1, 0, 0], [0, 1, 0], [1, 1, 0]]
```

```
nqft.qcm.get_hall\_coeff(spectral\_weight: ndarray, hoppings: tuple[float], x\_coord=None, file='./nqft/Data/hall.txt') <math>\rightarrow float
```

Computes Hall coefficient for given parameters and writes it to a file using specified x coordinate.

Parameters

- **spectral_weight** (*np.ndarray*, *shape=(N, M)*, *default=None)* Spectral function used to compute Hall coefficient.
- hoppings (tuple[float], size=3, default=None) Hopping amplitudes corresponding to spectral function.
- **x_coord** (*float/int*, *default=None*) X coordinate to use if writting Hall coefficient to a file. (Makes plotting easier and faster)
- **file** (*str*, *default="./nqft/Data/hall.txt"*) Path to file in which write Hall coefficient and given x coord.

Returns

n h – Hall coefficient as a float.

Return type

float

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1.1.6 Module contents

1.2 tests package

1.2.1 Submodules

1.2.2 tests.test_nqft module

```
tests.test_nqft.test_hamiltonian()
tests.test_nqft.test_version()
```

1.2.3 Module contents

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TWO

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