

pytheas Documentation

Release 1.1.1

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Introduction

Pytheas is a Python package for creating, running and postprocessing electrodynamic simulations. It is based on open source software Gmsh for creating geometries and mesh generation, and GetDP for solving the underlying partial differential equations with the finite element method.

It features built in models of:

- periodic media in 2D and 3D with computation of diffraction efficiencies
- scattering analysis in 2D and 3D
- Bloch mode analysis of metamaterials
- treatment of open geometries with perfectly matched layers
- tools to define arbitrary permittivity distributions
- quasi-normal mode analysis
- two scale convergence homogenization
- tools for topology optimization in 2D
- built-in refractive index database

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Installation

The easiest way to get started is to install via PyPi:

pip install pytheas-pip

User guide

3.1 pytheas.periodic2D: 2D metamaterials

The *pytheas.periodic2D* module implements the resolution of the scalar wave equation for TE and TM polarization for mono-periodic stuctures in 2D:

- subject to an incident plane wave (diffraction problem) with calculation of the diffraction efficiencies, absorption and energy balance.
- eigenvalues and eigenmodes (modal analysis)

3.1.1 Classes

Periodic2D([analysis, pola, A, lambda0,])	A class for a finite element model of a 2D mono-
	periodic medium.

pytheas.Periodic2D

```
class pytheas. Periodic 2D (analysis='direct', pola='TE', A=1, lambda0=1, lambda_mesh=1, theta_deg=0, d=0.8, h_sup=1, h_sub=1, h_layer1=0.1, h_layer2=0.1, h_des=1.0, h_pmltop=1.0, h_pmlbot=1.0, a_pml=1, b_pml=1, eps_sup=(1+0j), eps_layer1=(1+0j), eps_layer2=(1+0j), eps_des=(1+0j), eps_incl=(1+0j), mu_incl=(1+0j), mu_des=(1+0j))
```

A class for a finite element model of a 2D mono-periodic medium.

The model consist of a single unit cell with quasi-periodic boundary conditions in the *x* direction enclosed with perfectly matched layers (PMLs) in the *y* direction to truncate the semi infinite media. From top to bottom:

- PML top
- superstrate (incident medium)
- layer 2
- design layer: this is the layer containing the periodic pattern, can be continuous or discrete

- layer 1
- substrate
- PML bottom

Parameters

- **analysis** (str, default "direct") Analysis type: either "direct" (plane wave) or "modal" (spectral problem)
- **pola** (str, default "TE") Polarization case: either "TE" (E along z) or "TM" (H along z)
- A (float, default 1) Incident plane wave amplitude
- lambda0 (float, default 1) Incident plane wave wavelength in free space
- lambda_mesh (float, default 1) Wavelength to use for meshing
- **theta_deg** (*float*, *default* 0) Incident plane wave angle (in degrees). Light comes from the top (travels along -y if normal incidence, theta_deg=0 is set)
- d (float, default 0.8) Periodicity
- **h_sup** (*float*, *default 1*) Thickness superstrate
- h_sub (float, default 1) Thickness substrate
- h_layer1 (float, default 0.1) Thickness layer 1
- h_layer2 (float, default 0.1) Thickness layer 2
- **h_des** (*float*, *default 1*) Thickness layer design
- h_pmltop (float, default 1) Thickness pml top
- h_pmlbot (float, default 1) Thickness pml bot
- a_pml (float, default 1) PMLs complex y-stretching parameter, real part
- **b_pml** (*float*, *default 1*) PMLs complex y-stretching parameter, imaginary part
- eps_sup (complex, default (1 0 * 1j)) Permittivity superstrate
- eps_sub (complex, default (1 0 * 1j)) Permittivity substrate
- eps_layer1 (complex, default (1 0 * 1j)) Permittivity layer 1
- eps_layer2 (complex, default (1 0 * 1j)) Permittivity layer 2
- eps_des (complex, default (1 0 * 1j)) Permittivity layer design
- eps_incl (complex, default (1 0 * 1j)) Permittivity inclusion

cleanup()

Remove gmsh/getdp/python generated files from the temporary folder

compute_solution()

Compute the solution of the FEM problem using getdp

diffraction_efficiencies(cplx_effs=False, orders=False)

Postprocess diffraction efficiencies.

Parameters

- **cplx_effs** (*bool*) If *True*, return complex coefficients (amplitude reflection and transmission). If *False*, return real coefficients (power reflection and transmission)
- orders (bool) If True, computes the transmission and reflection for all the propagating diffraction orders. If False, returns the sum of all the propagating diffraction orders.

Returns A dictionary containing the diffraction efficiencies.

Return type dict

get_field_map(name)

Retrieve a field map.

Parameters name (str) – Choose between "u" (scattered field), "u_tot" (total field)

Returns A 2D complex array of shape (Nix, Niy)

Return type array

initialize()

Initialize the problem parameters.

make_inclusion(points, lcar='lc_incl', **kwargs)

Make a diffractive element geometry from points.

Parameters

- **points** (array of size (Npoints, 2)) The points defining the simply connected 2D geometry of the object.
- lcar (str (default "lc_incl")) Caracteristic length for the mesh.
- **kwargs (dict) Extra arguments.

make_mesh(other option=None)

Mesh the geometry using qmsh.

Parameters other_option (str) – Extra flag to pass to qmsh.

Returns The content of the .msh file.

Return type str

mk_tmp_dir()

Create a temporary directory

open_gmsh_gui(pos list=None)

Open gmsh GUI to visualize geometry and postprocessing results.

Parameters $pos_list(list)$ – A list of .pos files giving the views to load. By default it will render all the generated views.

postpro_absorption()

Compute the absorption coefficient

Returns Q - Absorption coefficient

Return type float

postpro_fields(filetype='txt', postop='postop_fields')

Compute the field maps and output to a file.

Parameters

- **filetype** (str, default "txt") Type of output files. Either "txt" (to be read by the method get field map in python) or "pos" to be read by qmsh/qetdp.
- postop (str, default "postop_fields") Name of the postoperation

postprocess(postop)

Run getdp postoperation.

Parameters postop (str) – Name of the postoperation to run.

rm_tmp_dir()

Remove the temporary directory

update_params()

Update the dictionary of parameters and the corresponding file

Examples using pytheas.Periodic2D

• Simulating diffraction by a 2D metamaterial

3.2 pytheas.scatt2D: 2D scattering

The *pytheas*. *scatt2D* module implements the resolution of the scalar wave equation for TE and TM polarization in 2D:

- subject to an incident plane wave or line source (diffraction problem)
- eigenvalues and eigenmodes (modal analysis)

3.2.1 Classes

Scatt2D()

A class for a finite element model of a 2D medium

pytheas.Scatt2D

class pytheas.Scatt2D

A class for a finite element model of a 2D medium

A = None

incident plane wave amplitude

Type flt

Ni_theta = None

number of theta points for computing the angular dependance of the modal coupling coefficients

Type int

$Nibox_x = None$

number of x interpolation points on the design box

Type int

```
number of y interpolation points on the design box
         Type int
Nin2f_x = None
    number of x interpolation points for near to far field calculations
Nin2f_y = None
    number of y interpolation points for near to far field calculations
         Type int
Nix = None
    number of x points for postprocessing field maps
         Type int
a_pml = None
    PMLs parameter, real part
         Type flt
analysis = None
     analysys type (either "direct" or "modal")
         Type str
b_pml = None
    PMLs parameter, imaginary part
         Type flt
beam_flag = None
    beam?
cleanup()
     Remove gmsh/getdp/python generated files from the temporary folder
compute_solution(res_list=None)
    Compute the solution of the FEM problem using getdp
dom_des = None
     design domain number (check .geo/.pro files)
eps_des = None
     permittivity scattering box
         Type flt
eps_host = None
     permittivity host
         Type flt
eps_incl = None
     permittivity inclusion
         Type flt
eps_sub = None
     permittivity substrate
```

 $Nibox_y = None$

```
Type flt
```

$h_pml = None$

thickness pml

Type flt

$hx_des = None$

x - thickness scattering box (design)

Type flt

hy_des = None

y - thickness scattering box

Type flt

initialize()

Initialize the problem parameters.

lambda0 = None

incident plane wave wavelength in free space

Type flt

lambda0search = None

wavelength around which to search eigenvalues

Type flt

lambda_mesh = None

wavelength to use for meshing

Type flt

ls_flag = None

line source position

make_inclusion(points, lcar='lc_incl', **kwargs)

Make a diffractive element geometry from points.

Parameters

- **points** (array of size (Npoints, 2)) The points defining the simply connected 2D geometry of the object.
- lcar (str (default "lc_incl")) Caracteristic length for the mesh.
- **kwargs (dict) Extra arguments.

make_mesh(other_option=None)

Mesh the geometry using gmsh.

Parameters other_option (str) – Extra flag to pass to gmsh.

Returns The content of the .msh file.

Return type str

mk_tmp_dir()

Create a temporary directory

nb_slice = None

number of y slices points for postprocessing diffraction efficiencies

Type int

neig = None

number of eigenvalues searched for in modal analysis

Type int

open_gmsh_gui(pos list=None)

Open gmsh GUI to visualize geometry and postprocessing results.

Parameters $pos_list(list)$ – A list of .pos files giving the views to load. By default it will render all the generated views.

pola = None

polarisation of the incident plane wave (either "TE" or "TM")

Type str

postpro_fields(filetype='txt', postop='postop_fields')

Compute the field maps and output to a file.

Parameters

- **filetype** (str, default "txt") Type of output files. Either "txt" (to be read by the method get_field_map in python) or "pos" to be read by gmsh/getdp.
- **postop** (str, default "postop_fields") Name of the postoperation

postprocess(postop)

Run getdp postoperation.

Parameters postop (str) – Name of the postoperation to run.

rm_tmp_dir()

Remove the temporary directory

scan_dist_ratio = None

such that scan_dist = min(h_sup, hsub)/scan_dist_ratio

Type flt

theta_deg = None

incident plane wave angle (in degrees). Light comes from the top (travels along -y if normal incidence, *theta_deg=0* is set)

Type flt

update_params()

Update the dictionary of parameters and the corresponding file

xnn = None

coords of point for PostProcessing

ypp = None

coords of point for PostProcessing

Examples using pytheas.Scatt2D

• Simulating diffraction by an object in 2D

3.3 pytheas.tools: tools and utilities

Input/output and utilities.

3.3.1 Submodules

femio	Tools for gmsh/getdp control and input/output.
utils	Shared utility functions used in pytheas.

pytheas.tools.femio

Tools for gmsh/getdp control and input/output.

Mesh the model using Gmsh

pytheas.tools.femio.postpro_commands(postop, path_pro, path_mesh, path_pos=None, verbose=0)

Generate a command list for postprocessing by GetDP (see main.pro file in ./base folder for default available postprocessings, or to add your own)

Parameters

- **postop** (str) The name of the postoperation to perform.
- path_pro (str) Path to the .pro file
- $path_mesh (str)$ Path to the .msh file
- path_pos (str , optional) Path to a file to be read by gmshread.
- verbose (int) verbosity level
- to None. (Defaults) -

Returns The list of strings to be oscommanded.

Return type list

pytheas.tools.utils

Shared utility functions used in pytheas.

```
pytheas.tools.utils.normalize(x)

Normalize an array between 0 and 1
```

Parameters x (array-like) – the quantity to be normalized

Returns x_norm – normalized array

Return type array-like

Examples

4.1 Material examples

Examples to show how to retrieve complex refractive index from a database, generating material patterns.

4.1.1 Importing refractive index from a database

Retrieve and plot the refractive index of a material in the refractive index.info data.

```
import numpy as np
from pytheas import refractiveindex as ri
import matplotlib.pyplot as plt
```

We can get the refractive index from tabulated data or a formula using the database in the pytheas.material module. We will import the measured data from the reference Johnson and Christy [JC1972]. We first specify the file yamlFile we want to import:

```
yamlFile = "main/Au/Johnson.yml"
```

We then get the wavelength bounds from the data (in microns) and create a wavelength range to interpolate:

```
bounds = ri.get_wl_range(yamlFile)
print(bounds[0], bounds[1])
lambdas = np.linspace(0.4, 0.8, 300)
```

Out:

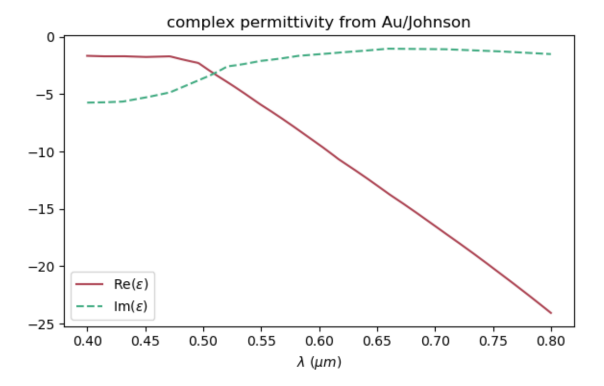
```
0.1879 1.937
```

Then get the refractive index data:

```
ncomplex = ri.get_complex_index(lambdas, yamlFile)
epsilon = ncomplex ** 2
```

And finally plot it:

```
fig, ax = plt.subplots(1, figsize=(6, 4))
ax.plot(lambdas, epsilon.real, "-", c="#ad4453", label=r"Re($\varepsilon$)")
ax.plot(lambdas, epsilon.imag, "--", c="#44ad84", label=r"Im($\varepsilon$)")
ax.set_xlabel(r"$\lambda$ ($\mu m$)")
ax.set_title("complex permittivity from " + yamlFile[5:][:-4])
ax.legend(loc=0)
plt.tight_layout()
```



Total running time of the script: (0 minutes 0.566 seconds)

Estimated memory usage: 13 MB

4.2 Periodic 2D examples

Examples to show how to simulate a mono periodic medium (metamaterial) with the finite element method and postprocessing the results (fields maps and diffraction efficiencies).

4.2.1 Simulating diffraction by a 2D metamaterial

Finite element simulation of the diffraction of a plane wave by a mono-periodic grating and calculation of diffraction efficiencies.

First we import the required modules and class

```
import numpy as np
import matplotlib.pyplot as plt

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```

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```
from pytheas import genmat
from pytheas import Periodic2D
```

Then we need to instanciate the class Periodic2D:

```
fem = Periodic2D()
```

The model consist of a single unit cell with quasi-periodic boundary conditions in the *x* direction enclosed with perfectly matched layers (PMLs) in the *y* direction to truncate the semi infinite media. From top to bottom:

- PML top
- superstrate (incident medium)
- layer 1
- design layer: this is the layer containing the periodic pattern, can be continuous or discrete
- layer 2
- substrate
- PML bottom

We define here the opto-geometric parameters:

```
mum = 1e-6 #: flt: the scale of the problem (here micrometers)
fem.d = 0.4 * mum #: flt: period
fem.h_sup = 1.0 * mum #: flt: "thickness" superstrate
fem.h_sub = 1.0 * mum #: flt: "thickness" substrate
fem.h_layer1 = 0.1 * mum #: flt: thickness layer 1
fem.h_layer2 = 0.1 * mum #: flt: thickness layer 2
fem.h_des = 0.4 * mum #: flt: thickness layer design
fem.h_pmltop = 1.0 * mum #: flt: thickness pml top
fem.h_pmlbot = 1.0 * mum #: flt: thickness pml bot
fem.a_pml = 1 #: flt: PMLs parameter, real part
fem.b_pml = 1 #: flt: PMLs parameter, imaginary part
fem.eps_sup = 1 #: flt: permittivity superstrate
fem.eps_sub = 3 #: flt: permittivity substrate
fem.eps_layer1 = 1 #: flt: permittivity layer 1
fem.eps_layer2 = 1 #: flt: permittivity layer 2
fem.eps_des = 1 #: flt: permittivity layer design
fem.lambda0 = 0.6 * mum #: flt: incident wavelength
fem.theta_deg = 0.0 #: flt: incident angle
fem.pola = "TE" #: str: polarization (TE or TM)
fem.lambda_mesh = 0.6 * mum #: flt: incident wavelength
#: mesh parameters, correspond to a mesh size of lambda_mesh/(n*parmesh),
#: where n is the refractive index of the medium
fem.parmesh_des = 15
fem.parmesh = 13
fem.parmesh_pml = fem.parmesh * 2 / 3
fem.type_des = "elements"
```

We then initialize the model (copying files, etc...) and mesh the unit cell using gmsh

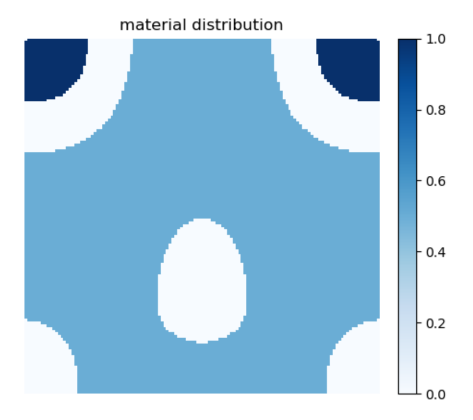
```
fem.getdp_verbose = 0
fem.gmsh_verbose = 0
(continues on next page)
```

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```
fem.initialize()
mesh = fem.make_mesh()
```

We use the genmat module to generate a material pattern

```
genmat.np.random.seed(100)
mat = genmat.MaterialDensity()  # instanciate
mat.n_x, mat.n_y, mat.n_z = 2 ** 7, 2 ** 7, 1  # sizes
mat.xsym = True  # symmetric with respect to x?
mat.p_seed = mat.mat_rand  # fix the pattern random seed
mat.nb_threshold = 3  # number of materials
mat._threshold_val = np.random.permutation(mat.threshold_val)
mat.pattern = mat.discrete_pattern
fig, ax = plt.subplots()
mat.plot_pattern(fig, ax)
```



We now assign the permittivity

```
fem.register_pattern(mat.pattern, mat._threshold_val)
fem.matprop_pattern = [1.4, 4 - 0.02 * 1j, 2] # refractive index values
```

Now we're ready to compute the solution:

```
fem.compute_solution()
```

Finally we compute the diffraction efficiencies, absorption and energy balance

```
effs_TE = fem.diffraction_efficiencies()
print("efficiencies TE", effs_TE)
```

Out:

```
efficiencies TE {'R': 0.42749531344001657, 'T': 0.45592852708133014, 'Q': 0.1177478267191832, 'B': 1. 
→00117166724053}
```

It is fairly easy to switch to TM polarization:

```
fem.pola = "TM"
fem.compute_solution()
effs_TM = fem.diffraction_efficiencies()
print("efficiencies TM", effs_TM)
```

Out:

Total running time of the script: (0 minutes 3.381 seconds)

Estimated memory usage: 14 MB

4.3 Scattering 2D examples

Examples to show how to simulate a the 2D scattering off an object subject to a plane wave or line source harmonic excitation.

4.3.1 Simulating diffraction by an object in 2D

Finite element simulation of the diffraction by an object illuminated by a plane wave or a line source. Calculation of scattering width and getting the field maps.

```
import numpy as np
import matplotlib.pyplot as plt
from pytheas import Scatt2D

plt.ion()
pi = np.pi
```

Then we need to instanciate the class Scatt2D:

```
fem = Scatt2D()
fem.rm_tmp_dir()
```

```
# We define first the opto-geometric parameters:
mum = 1 #: flt: the scale of the problem (here micrometers)
fem.lambda0 = 0.6 * mum #: flt: incident wavelength
fem.pola = "TE" #: str: polarization (TE or TM)
fem.theta_deg = 30.0 \# 0: coming from top (y>0)
fem.hx_des = 1.0 * mum #: flt: x thickness box
fem.hy_des = 1.0 * mum #: flt: y thickness box
fem.h_pml = fem.lambda0 #: flt: thickness pml
fem.space2pml_L, fem.space2pml_R = fem.lambda0 * 2, fem.lambda0 * 2
fem.space2pml_T, fem.space2pml_B = fem.lambda0 * 2, fem.lambda0 * 2
fem.eps_des = 1 #: flt: permittivity design box
fem.eps_host = 1.0
fem.eps_incl = 11.0 - 1e-2 * 1j
#: mesh parameters, correspond to a mesh size of lambda_mesh/(n*parmesh),
#: where n is the refractive index of the medium
fem.lambda_mesh = 0.6 * mum #: flt: incident wavelength
fem.parmesh_des = 10
fem.parmesh_incl = 10
fem.parmesh = 10
fem.parmesh_pml = fem.parmesh * 2 / 3
fem.Nix = 101
fem.Niy = 101
```

Here we define an ellipsoidal rod as the scatterer:

```
def ellipse(Rinclx, Rincly, rot_incl, x0, y0):
    c, s = np.cos(rot_incl), np.sin(rot_incl)
    Rot = np.array([[c, -s], [s, c]])
    nt = 360
    theta = np.linspace(-pi, pi, nt)
    x = Rinclx * np.sin(theta)
    y = Rincly * np.cos(theta)
    x, y = np.linalg.linalg.dot(Rot, np.array([x, y]))
    points = x + x0, y + y0
    return points

rod = ellipse(0.4 * mum, 0.2 * mum, 0, 0, 0)
fem.inclusion_flag = True
```

Initialize, build the scatterer, mesh and compute the solution:

```
fem.initialize()
fem.make_inclusion(rod)
fem.make_mesh()
fem.compute_solution()
```

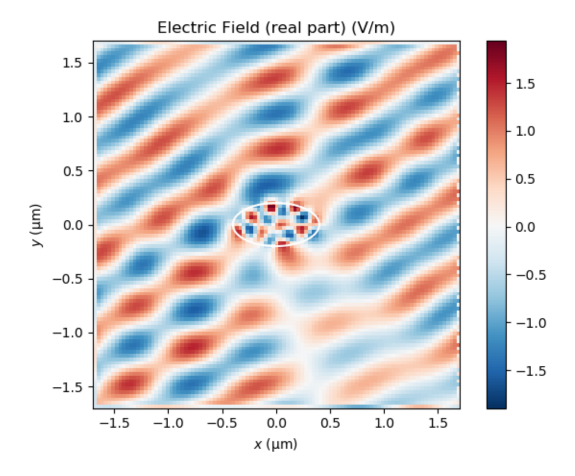
Get the electric field and plot it:

```
fem.postpro_fields()
u_tot = fem.get_field_map("u_tot.txt")
fig, ax = plt.subplots()
E = u_tot.real
```

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```
plt.imshow(E, cmap="RdBu_r", extent=(fem.domX_L, fem.domX_R, fem.domY_B, fem.domY_T))
plt.plot(rod[0], rod[1], "w")
plt.xlabel(r"$x$ ($\rm \mu$m)")
plt.ylabel(r"$y$ ($\rm \mu$m)")
plt.title(r"Electric Field (real part) (V/m)")
plt.colorbar()
plt.tight_layout()
```

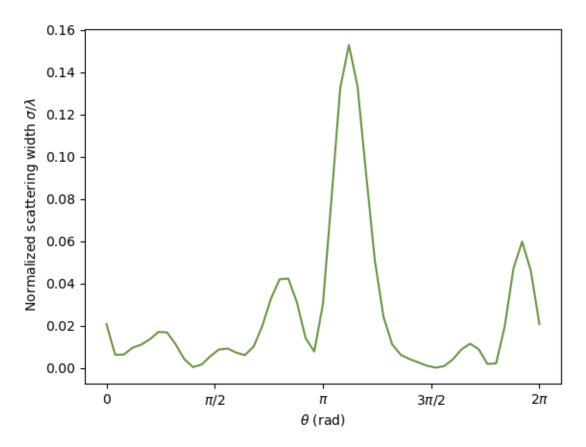


Do a near to far field transform and get the normalized scattering width:

```
ff = fem.postpro_fields_n2f()
theta = np.linspace(0, 2 * pi, 51)
scs = fem.normalized_scs(ff, theta)

fig, ax = plt.subplots()
plt.plot(theta / pi, scs, "-", c="#699545")
plt.xlabel(r"$\theta$ (rad)")
plt.ylabel(r" Normalized scattering width $\sigma/\lambda$")
ax.xaxis.set_ticks([0, 0.5, 1, 1.5, 2])
ax.xaxis.set_ticklabels(["0", "$\pi/2$", "$\pii\pi", "$3\pi/2$", "$2\pi$"])

scs_integ = np.trapz(scs, theta) / (2 * pi)
print("Normalized SCS", scs_integ)
```



Out:

Normalized SCS 0.02572717419074843

Total running time of the script: (0 minutes 17.270 seconds)

Estimated memory usage: 16 MB

Bibliography

[JC1972] (P. B. Johnson and R. W. Christy. Optical constants of the noble metals, Phys. Rev. B 6, 4370-4379 (1972)).

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