CF-03 Python interface for OOMMF

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Overview – JOOMMF Project

- Towards embedding OOMMF into the Jupyter Notebook (→ J-OOMMF)
- · Step 1: Drive OOMMF through Python interface
- · Step 2: Develop data analysis tools
- Step 3: Interactive documentation, micromagnetic tutorial, reproducibility

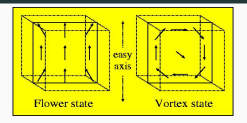
Status

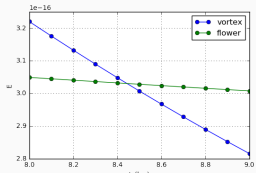
Step 1 Prototype completed

- Python interface for OOMMF:
- "OOMMF Calculator"

 ≡ OOMMFC

Standard Problem 3





Full problem specification:
http://www.ctcms.nist.
gov/~rdm/spec3.html

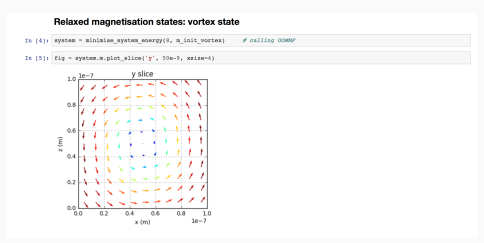
CF-03: Hans Fangohr: Pythoh^(lex)terface for OOMMF

```
In [1]: import commfc as oc  # access to OOMMF Calculator

import discretisedfield as df  # other setup to keep the next slides brief
import numpy as np
from math import sin, cos, pi, sqrt
import matplotlib.pyplot as plt
%matplotlib inline
```

Micromagnetic standard problem 3

```
In [2]: def m init flower(pos):
            """Given a pos vector pos = (x, y, z), return the magnetisation
            vector (mx, my, mz) for that position. """
            x, y, z = pos[0]/1e-9, pos[1]/1e-9, pos[2]/1e-9
            # flower pattern:
            mx = 0
            my = 2 * z - 1
            mz = -2 * y + 1
            norm squared = mx**2 + my**2 + mz**2
            if norm squared <= 0.05:
                return (1, 0, 0)
            else:
                return (mx, mv, mz)
        def m init vortex(pos):
            """Given a pos vector pos = (x, y, z), return the magnetisation
            vector (mx, mv, mz) for that position."""
            x, y, z = pos[0]/1e-9, pos[1]/1e-9, pos[2]/1e-9
            # vortex pattern
            mv = 0
            mv = sin(pi/2 * (x-0.5))
            mz = cos(pi/2 * (x-0.5))
            return (mx, my, mz)
```



Flower state: In [6]: system = minimise system energy(8, m init flower) In [7]: fig = system.m.plot_slice('y', 50e-9, xsize=4) y slice 1.0 0.8 0.6 z (m) 0.4 0.2 0.4 0.8 0.0 0.6 x (m) 1e-7

Create the energy crossing plot

```
In [8]: L array = np.linspace(8, 9, 3) # values of L, from 8 to 9 in 4 steps
        vortex energies = []
        flower energies = []
        for L in L array:
            print("Computing vortex L={} using OOMMF".format(L))
            vortex = minimise system energy(L, m init vortex)
            print("Computing flower L={} using OOMMF".format(L))
            flower = minimise system energy(L, m init flower)
            vortex energies.append(vortex.total energy()) # remember energies for later
            flower energies.append(flower.total energy())
                                                           # plotting
        Computing vortex L=8.0 using OOMMF
        Computing flower L=8.0 using OOMMF
        Computing vortex L=8.5 using OOMMF
        Computing flower L=8.5 using OOMMF
        Computing vortex L=9.0 using OOMMF
        Computing flower L=9.0 using OOMMF
```

```
In [9]: plt.plot(L_array, vortex_energies, 'o-', label='vortex')
        plt.plot(L_array, flower_energies, 'o-', label='flower')
        plt.xlabel('L (lex)')
        plt.ylabel('E')
        plt.xlim([7.9, 9.1])
        plt.grid()
        plt.legend();
               le-16
                                                        vortex
           3.2
                                                         flower
           3.1
           3.0
           2.9
           2.8
                  8.0
                           8.2
                                     8.4
                                              8.6
                                                       8.8
                                                                 9.0
```

L (lex)

Use bisection method to find energy crossing automatically

```
In [10]: from scipy.optimize import bisect
         def energy difference(L):
             print("Computing energy difference at L = {}".format(L))
             vortex = minimise system energy(L, m init vortex)
             flower = minimise system energy(L, m init flower)
             return vortex.total energy() - flower.total energy()
         cross section = bisect(energy difference, 8.3, 8.5, xtol=0.01)
         print("The transition between vortex and flower states occurs approximately at {}*lex".format(cross section))
         Computing energy difference at L = 8.3
         Computing energy difference at L = 8.5
         Computing energy difference at L = 8.4
         Computing energy difference at L = 8.45
         Computing energy difference at L = 8.425
         Computing energy difference at L = 8.4125
         Computing energy difference at L = 8.41875
         The transition between vortex and flower states occurs approximately at 8.41875*lex
```

Benefits

Present

- OOMMF simulation study in single (Python) file
- Multiple simulation runs within the same script
- Exploit existing libraries and tools (root finding)

Future

- Embedding interactive simulation data analysis, and visualisation
- Reproducibility
- · Interactive documenation
- ...

How does the interface to OOMMF work?

Via MIF files

- 1. write MIF file
- 2. execute OOMMF
- 3. read output files

Why?

- · most robust approach
- · see https://arxiv.org/abs/1609.07432 for details

How to install?

- Need OOMMF natively installed
 (and set variable OOMMFTCL to point to oommf.tcl file)
 or
 Docker (http://docker.com)
- 2. Need Python (Suggest Anaconda distribution)
- 3. Install oommfc via\$> pip install oommfc

Is it ready to use?

Software ready to use?

- · Yes(-ish)
- · interface may change, although we try to avoid it
- · beta users and questions welcome

Installation and support workshop for OOMMFC

- Wednesday 17:00 19:00 (today)
- Thursday 17:00 19:00 (tomorrow)

Outside "Galerie 4" on Level 2, drop-in anytime

Summary

Python interface for OOMMF (OOMMF Calculator)

- part of JOOMMF Project
- invite the community to engage
 - · with ideas, questions and bug reports
 - subscribe to joommf-news mailing list
 - come to workshop tonight/tomorrow
- http://joommf.github.io

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