## MPB Demo

## — numerical Maxwell eigensolver on Athena

For more information on MPB, see also the MPB web page at <a href="http://ab-initio.mit.edu/mpb">http://ab-initio.mit.edu/mpb</a> (which includes a link to documentation). All of these files and links can also be found on the course web page.

The following are a sequence of Athena and Matlab commands that I will (hopefully) use in class to perform a simple calculation and analyze the output. Here, we analyze a simple 2d waveguide formed by an  $\varepsilon$ =12 region with thickness 1, surrounded by air.

The MPB input file, 2dwaveguide.ctl, is on the following page. The last page is the input file and commands for a more complicated problem—a 2d waveguide formed by a periodic sequence of dielectric cylinders.

A note on units: Since Maxwell's equations are scale-invariant, there is no need for MPB to settle on any particular units. Instead, you simply pick whatever units of distance you want — typically you select some characteristic lengthscale a in the system and set that equal to 1. Then, you specify a wavevector k to MPB in units of  $2\pi/a$ , and the frequencies  $\omega$  are returned in units of  $2\pi c/a$ . See also the MPB manual for more details (it gets more complicated for 2d-periodic systems, where the lattice or reciprocal lattice vectors are used as a basis for vectors or wavevectors).

```
% foo = unix command
>> foo = Matlab command
% add mpb

% mpb 2dwaveguide.ctl > 2dwaveguide.out
% cat 2dwaveguide.out
% grep tmyevenfreqs: 2dwaveguide.out > foo.dat
```

```
>> foo = dlmread('foo.dat', ',', 1, 1)
>> kx = foo(:,2)
>> f = foo(:,6:end)
>> plot(kx, f, 'ro-')
>> hold on
>> plot(kx, kx, 'k', 'LineWidth', 2)
\rightarrow patch([kx;flipud(kx)], [kx; ones(size(kx))*max(kx)], [0.9,0.9,0.9])
% grep tmyoddfreqs: 2dwaveguide.out > foo.dat
>> foo = dlmread('foo.dat', ',', 1, 1)
>> fo = foo(:,6:end)
>> plot(kx, fo, 'bo-')
>> plot(kx, kx / sqrt(12), 'k')
% h5ls epsilon.h5
>> ep = hdf5read('epsilon.h5', 'data')
>> figure
>> plot(ep, 'k')
% h5ls e.k01.b01.z.tm.h5
>> ez1 = hdf5read('e.k01.b01.z.tm.h5', 'z.r') +
hdf5read('e.k01.b01.z.tm.h5','z.i') * i;
>> plot(real(ez1) * 100, 'r');
>> ez2 = hdf5read('e.k01.b02.z.tm.h5', 'z.r') +
hdf5read('e.k01.b02.z.tm.h5','z.i') * i;
>> plot(real(ez2) * 100, 'b');
>> ez3 = hdf5read('e.k01.b03.z.tm.h5', 'z.r') +
hdf5read('e.k01.b03.z.tm.h5','z.i') * i;
>> plot(real(ez3) * 100, 'm');
>> (ep .* ez1) * ez3'
```

```
; Example MPB input file for 18.325, illustrating a simple 2d dielectric
; waveguide along the x direction. Run it with:
        mpb 2dwaveguide.ctl > 2dwaveguide.out
; to produce an output file 2dwaveguide.out (as well as some .h5 data files).
: As described in the manual, you can extract the eigenfrequencies
; by doing "grep freqs: 2dwaveguide.out" at the Unix shell.
; (Note that anything after a ";" on a line is ignored by MPB.)
......
; First, we will define some parameters describing our structure. Defining
: them symbolically here makes it easier to change them. (e.a. we
; can change the dielectric constant from the command line via
; "mpb eps-hi=13 2dwavequide.ctl".) We then use these parameters below
(define-param eps-hi 12); the waveguide dielectric constant
(define-param eps-lo 1): the surrounding low-dielectric material
(define-param h 1); the thickness of the waveguide (arbitrary units)
(define-param Y 10); the size of the computational cell in the y direction
; Define the structure and the computational cell
; Here we define the size of the computational cell. Since it is 2d,
; it has no-size in the z direction. Because it is a waveguide in the
; x direction, then the eigenproblem at a given k has no-size in the
: x direction as well.
(set! geometry-lattice (make lattice (size no-size Y no-size)))
; the default-material is what fills space where we haven't placed objects
(set! default-material (make dielectric (epsilon eps-lo)))
; a list of geometric objects to create structures in our computational cell:
; (in this case, we only have one object, a block to make the wavequide)
(set! geometry
     (list (make block; a dielectric block (a rectangle)
         (center 0 0 0); centered at origin
         (size infinity h infinity); block is finite only in y direction
         (material (make dielectric (epsilon eps-hi))))))
; MPB discretizes space with a given resolution. Here, we set
; a resolution of 32 pixels per unit distance. Thus, with Y=10
; our comptuational cell will be 320 pixels wide. In general,
; you should make the resolution fine enough so that the pixels
: are much smaller than the wavelenath of the light.
```

```
(set-param! resolution 32)
; Tell MPB what eigenmodes we want to compute.
; Generally, we want omega(k) for a range of k values. MPB
; can automatically interpolate a set of k values between any
; given bounds. Here, we will interpolate 10 k's between 0 and 2.
(define-param kmin 0)
(define-param kmax 2)
(define-param k-interp 10)
: k-points is the list of k values that MPB computes eigenmodes at.
; (vector3 x y z) specifies a vector. (k is in units of 2 pi/distance)
(set! k-points (interpolate k-interp
                        (list (vector3 kmin 0 0) (vector3 kmax 0 0))))
; we also need to specify how many eigenmodes we want to compute, given
; by "num-bands":
(set-param! num-bands 5)
; to compute *all* the modes, we now simply type (run).
; However, it is convenient to compute only one symmetry of mode
; at a time. In particular, we will compute only TM (E in z direction)
; modes, and separately compute even and odd modes with respect to the
; y=0 mirror symmetry plane.
(run-tm-yeven)
(run-tm-yodd)
; (If we don't have y=0 mirror symmetry we should just use run-tm).
; That's it! We're done! However, suppose we now want to get the
; *fields* at a given k. To do this, we'll call the run function
; again, this time giving it an option to output the modes.
(define-param k 1); the k value where we'll output the modes
(set! k-points (list (vector3 k 0 0))); compute only a single k now
; output-efield-z does just what it says. There are also options
; to output any other field component we care to examine.
(run-tm output-efield-z)
```

```
; Example MPB input file for 18.325, for a periodic (period = 1) sequence
                                                                           ; let's just stick with the even modes for now.
; of dielectric cylinders in the x direction. (This file is otherwise
                                                                           (run-tm-yeven)
; very similar to 2dwaveguide.ctl ... refer to that file for more details.)
                                                                           ; uncomment to output some fields at k=0.4:
; Some parameters:
                                                                          (define-param k 0.4); the k value where we'll output the modes
(define-param eps-hi 12); the waveguide dielectric constant
                                                                          (set! k-points (list (vector3 k 0 0))); compute only a single k now
(define-param eps-lo 1); the surrounding low-dielectric material
(define-param h 1); the thickness of the waveguide (arbitrary units)
                                                                           ; (run-tm-yeven output-efield-z)
(define-param r 0.2) : the radius of the cylinders
                                                                          % foo = Unix (Athena) command
                                                                          >> foo = Matlab command
(define-param Y 10); the size of the computational cell in the y direction
                                                                          % add mpb
; Define the structure and the computational cell
                                                                          % mpb 2dwaveguide-periodic.ctl > 2dwaveguide-periodic.out
: note that now the size in the x direction is 1 (one period)
                                                                          % grep tmyevenfreqs: 2dwaveguide.out > foo.dat
(set! geometry-lattice (make lattice (size 1 Y no-size)))
                                                                          >> foo = dlmread('foo.dat', ',', 1, 1)
(set! default-material (make dielectric (epsilon eps-lo)))
                                                                          \gg kx = foo(:,2)
(set! geometry
                                                                          >> fe = foo(:,6:end)
     (list (make cylinder; cylinder oriented along z direction
         (center 0 0 0); centered at origin
                                                                          >> plot(kx, fe, 'ro-')
         (radius r) (height infinity)
                                                                          >> hold on
         (material (make dielectric (epsilon eps-hi))))))
                                                                          >> plot(kx, kx, 'k', 'LineWidth', 2)
                                                                          >> plot(kx + 1, kx, 'k', 'LineWidth', 2)
(set-param! resolution 16)
                                                                          >> plot(-kx + 1, kx, 'k', 'LineWidth', 2)
                                                                          >> plot(-kx + 2, kx, 'k', 'LineWidth', 2)
>> axis([0 2 0 0.6])
; Tell MPB what eigenmodes we want to compute.
                                                                          Run again, uncommenting line to output fields...
(define-param kmin 0)
(define-param kmax 2)
                                                                          >> ez1 = hdf5read('e.k01.b01.z.tmyeven.h5', 'z.r') +
(define-param k-interp 20)
                                                                                  hdf5read('e.k01.b01.z.tmyeven.h5','z.i') * i;
; k-points is the list of k values that MPB computes eigenmodes at.
                                                                          >> pcolor(real(ez1))
; (vector3 x y z) specifies a vector. (k is in units of 2 pi/distance)
                                                                          >> axis imaae
(set! k-points (interpolate k-interp
                      (list (vector3 kmin 0 0) (vector3 kmax 0 0))))
                                                                          % mpb-data -x 5 e.k01.b01.z.tmyeven.h5 epsilon.h5
                                                                          % h5topng -S 4 -Zc bluered -C epsilon.h5:data-new -d z.r-new
; we also need to specify how many eigenmodes we want to compute, given
                                                                                                                  e.k01.b01.z.tmyeven.h5
: by "num-bands":
                                                                          % add graphics
(set-param! num-bands 5)
                                                                          % xv e.k01.b01.z.tmyeven.png &
```