# 18.369 Problem Set 4

Due Friday, 28 March 2008.

#### **Problem 1: Perturbation theory**

- (a) In class, we derived the 1st-order correction in the eigenvalue for an ordinary Hermitian eigenproblem  $\hat{O}\psi = \lambda\psi$  for a small perturbation  $\Delta\hat{O}$ . Now, do the same thing for a *generalized* Hermitian eigenproblem  $\hat{A}\psi = \lambda\hat{B}\psi$ .
  - (i) That is, assume we have the solution  $\hat{A}^{(0)}\psi^{(0)} = \lambda^{(0)}\hat{B}^{(0)}\psi^{(0)}$  to an unperturbed system (where  $\hat{A}^{(0)}$  and  $\hat{B}^{(0)}$  are Hermitian, and  $\hat{B}^{(0)}$  is positive-definite) and find the first-order correction  $\lambda^{(1)}$  when we change *both*  $\hat{A}$  and  $\hat{B}$  by small amounts  $\Delta \hat{A}$  and  $\Delta \hat{B}$ . You may assume that  $\lambda^{(0)}$  is non-degenerate, for simplicity.
  - (ii) Now, apply this solution to the generalized eigenproblem  $\nabla \times \nabla \times \mathbf{E} = \frac{\omega^2}{c^2} \varepsilon \mathbf{E}$  for a small change  $\Delta \varepsilon$ , and show that the first-order correction  $\Delta \omega$  is the same as the one derived in class using the **H** eigenproblem.
- (b) Recall the problem of the modes in an L × L metal box that we solved in class for the H<sub>z</sub> (TE) polarization, and which you solved in problem set 2 for the E<sub>z</sub> (TM) polarization. Originally, this box was filled with air (ε = 1). Now, suppose that we increase ε by some small constant Δε in the lower-left <sup>L</sup>/<sub>2</sub> × <sup>L</sup>/<sub>2</sub> corner of the box. What is the first-order Δω for the first four (lowest ω) modes of the TM polarization? What happens to the degenerate modes (be careful, remember to diagonalize the perturbation explicitly or via symmetry)?
- (c) In class, we calculated the band gap Δω that appeared in a uniform (1d) material ε(x) = ε<sub>1</sub> when *half* of the unit cell a was changed to ε<sub>1</sub> - Δε for some small Δε, by using perturbation theory. In particular, we derived the gap between the first two bands that appears at k = π/a, the edge of the Brillouin zone.
  - (i) Now, you should compute the gap that appears between band 2 and band 3 at k = 0,

the *center* of the Brillouin zone, to first order. However, do it more generally: assume that the  $\varepsilon_1$  region has thickness  $d_1$ and that the  $\varepsilon_1 - \Delta \varepsilon$  region has thickness  $a - d_1$ , repeated with period a, and compute  $\Delta \omega$  as a function of  $d_1$ .

(ii) At what  $d_1$  values does this gap at k = 0disappear ( $\Delta \omega = 0$ )? What is  $\Delta \omega$  for the quarter-wave stack thicknesses?

#### **Problem 2: Band gaps in MPB**

Consider the 1d periodic structure consisting of two alternating layers:  $\varepsilon_1 = 12$  and  $\varepsilon_2 = 1$ , with thicknesses  $d_1$  and  $d_2 = a - d_1$ , respectively. To help you with this, I've created a sample input file *bandgap1d.ctl* that is posted on the course web page.

- (a) Using MPB, compute and plot the fractional TM gap size (of the *first* gap, i.e lowest  $\omega$ ) vs.  $d_1$  for  $d_1$  ranging from 0 to a. What  $d_1$  gives the largest gap? Compare to the "quarter-wave" thicknesses  $d_{1,2} = a\sqrt{\varepsilon_{2,1}}/[\sqrt{\varepsilon_1} + \sqrt{\varepsilon_2}]$ .
- (b) Given the optimal parameters above, what would be the physical thicknesses in order for the mid-gap vacuum wavelength to be  $\lambda = 2\pi c/\omega = 1.55\mu$ m? (This is the wavelength used for most optical telecommunications.)
- (c) Plot the 1d TM band diagram for this structure, with  $d_1$  given by the quarter wave thickness, showing the first five gaps. Also compute it for  $d_1 = 0.12345$  (which I just chose randomly), and superimpose the two plots (plot the quarterwave bands as solid lines and the other bands as dashed). What special features does the quarterwave band diagram have?

### **Problem 3: Bands and supercells**

*Note:* this problem does *not* require you to do any numerical calculations in MPB etcetera—it actually appeared on the spring 2007 midterm.

Calvin Q. Luss, a Harvard student, posts to the MPB mailing list that he has discovered a bug in MPB. He writes:

I'm getting ready to do a 2d-crystal calculation, but first I wanted to do a 1d crystal as a test case since I know the band diagram analytically for that (from

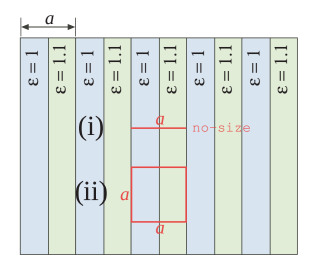


Figure 1: Two MPB unit cells for the band structure of a 1d-periodic quarter-wave stack: (i) a 1d  $a \times no-size$  unit cell (ii) a 2d  $a \times a$  unit cell.

Yeh's book). I used the quarter-wave stack shown in fig. 1(i), with a 1d computational cell of  $a \times no - size \times no - size$ , and plotted the TM band structure  $\omega(k_x)$  (for  $\mathbf{k} = (k_x, 0, 0)$  with  $k_x$  from 0 to 0.5 in MPB units, i.e. from 0 to  $\pi/a$ )—everything works fine! Then I do the same calculation but with a computational cell of  $a \times a \times no - size$ , as shown in fig. 1(ii), and the result is wrong! I get all sorts of extra bands at bogus frequencies; why doesn't the result match the 1d computation, since the structure hasn't changed? I think it must be a bug; you MIT people obviously don't know what you're doing.

Sketch out the plots that Calvin got from his two calculations, and explain why MPB is correctly answering exactly the question that he posed. Sketch at least 4 bands in the 1d calculation, and at least 6 bands in the 2d calculation (not counting degeneracies), and label any bands that are doubly (or more?) degenerate.

(You can use the fact that the  $\varepsilon$  contrast in this case is only 10%—the structure is *nearly* homogeneous to help you sketch out the bands more quantitatively. But no need to be *too* quantitative, however: you don't need to use perturbation theory or anything like that; a reasonable guess is sufficient.)

## **Problem 4: Defect modes in MPB**

In MPB, you will create a (TM polarized) defect mode by increasing the dielectric constant of a single layer by  $\Delta \varepsilon$ , pulling a state down into the gap. The periodic structure will be the same as the one from problem 2, with the quarter-wave thickness  $d_1 = 1/(1 + \sqrt{12})$ . To help you with this, I've created a sample input file *defect1d.ctl* that is posted on the course web page.

- (a) When there is no defect (Δε), plot out the band diagram ω(k) for the N = 5 supercell, and show that it corresponds to the band diagram of problem 2 "folded" as expected.
- (b) Create a defect mode (a mode that lies in the band gap of the periodic structure) by increasing the ε of a single ε<sub>1</sub> layer by Δε = 1, and plot the E<sub>z</sub> field pattern. Do the same thing by increasing a single ε<sub>2</sub> layer. Which mode is even/odd around the mirror plane of the defect? Why?
- (c) Gradually increase the  $\varepsilon$  of a single  $\varepsilon_2$  layer, and plot the defect  $\omega$  as a function of  $\Delta \varepsilon$  as the frequency sweeps across the gap. At what  $\Delta \varepsilon$  do you get two defect modes in the gap? Plot the  $E_z$  of the second defect mode. (Be careful to increase the size of the supercell for modes near the edge of the gap, which are only weakly localized.)
- (d) The mode must decay exponentially far from the defect (multiplied by an e<sup>i π/α x</sup> sign oscillation and the periodic Bloch envelope, of course). From the E<sub>z</sub> field computed by MPB, extract this asymptoic exponential decay rate (i.e. κ if the field decays ~ e<sup>-κx</sup>) and plot this rate as a function of ω, for the first defect mode, as you increase ε<sub>2</sub> as above (vary ε<sub>2</sub> so that ω goes from the top of the gap to the bottom).