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2010 J. Opt. 12 104013

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Theory of disorder-induced coherent scattering and light localization in slow-light photonic crystal waveguides

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Received 22 February 2010, accepted for publication 5 May 2010
Published 24 September 2010
Online at stacks.iop.org/JOpt/12/104013

Abstract

We introduce a theoretical formalism to describe disorder-induced extrinsic scattering in slow light photonic crystal waveguides. This work details and extends the optical scattering theory used in a recent issue of Physics Review Letters (Patterson et al 2009 Phys. Rev. Lett. 102 253903) to describe coherent scattering phenomena and successfully explain related experimental measurements. Our presented theory, which combines Green function and coupled mode methods, allows us to self-consistently account for arbitrary multiple scattering for the propagating electric field and recover experimental features such as resonances near the band edge. The technique is fully three-dimensional and can calculate the effects of disorder on the propagating field over thousands of unit cells. As an application of this theory, we explore various sample lengths and disordered instances, and demonstrate the profound effect of multiple scattering in the waveguide transmission. The spectra yield rich features associated with disorder-induced localization and multiple scattering, which are shown to be exacerbated in the slow light propagation regime.

Keywords: photonic crystals, slow light, disorder-induced scattering, localization

(Some figures in this article are in colour only in the electronic version)
structure. In section 1, we introduce the theoretical formalism. The magnetic Bloch mode 
̂h_k(r) is defined similarly. The electric field mode 
̂e_k(r) is obtained from the magnetic field mode 
̂h_k(r) by the magnetic field relation 
̂h_k(r) = \frac{1}{\sqrt{\varepsilon_0 \mu_0}} \sqrt{\mu_0/\varepsilon_0} \nabla \times \nabla \times \hat{e}_k(r).

1. Theory

1.1. Waveguide Bloch modes

The ideal PC waveguide is periodic along the propagation direction (z) with periodicity a: \varepsilon(r + a\hat{k}) = \varepsilon(r), where \varepsilon(r) is the dielectric constant that we will assume is real and \hat{k} is a unit vector. Consequently, Bloch’s theorem applies and the electric field mode may be written as \nabla \times \nabla \times \hat{e}_k(r) e^{i k x} = 0.

\[ \mathcal{P}_k \mathbf{E} = \int \int_{r=x} d\alpha d\beta \mathbf{e}_k(r) e^{-i k \alpha} \times \mathbf{E}(r) + \mathbf{P}(r; \omega) / \varepsilon_0, \]

where the overbar represents a tensor or dyadic. For convenience, we partition the electric field wave equation, namely

\[ \mathbf{E}(r; \omega) = \mathbf{E}_k(r; \omega) + \int_{\text{all space}} d\alpha d\beta \mathbf{G}(r, r'; \omega) \mathbf{e}_k(r') e^{i k x} = \frac{1}{\sqrt{\varepsilon_0 \mu_0}} \sqrt{\mu_0/\varepsilon_0} \nabla \times \nabla \times \hat{e}_k(r)

For the present work, since we are interested in developing sub-unit-cell propagation equations, we would prefer the integration was over only the plane perpendicular to the propagation direction. Using the electric and magnetic field orthogonality relations, the Maxwell constitutive relations, and the divergence theorem, one can derive

\[ 0 = \int_{r=x} d\alpha d\beta \mathbf{e}_k(r) e^{-i k \alpha} \times \mathbf{E}(r) + \mathbf{P}(r; \omega) / \varepsilon_0, \]

where the integration here is performed over a single plane transverse to the propagation direction. For k \neq k', the term in brackets is non-zero and the integral must evaluate to zero.

For k = k', the integral can be recognized as the power flux at the transverse plane which is clearly non-zero (except for a radiation mode propagating perpendicular to the slab). Thus, a new projection (orthogonality) operator can be defined as

\[ \mathcal{P}_k \mathbf{E} = \int_{r=x} d\alpha d\beta \mathbf{e}_k(r) e^{-i k \alpha} \times \mathbf{E}(r) + \mathbf{P}(r; \omega) / \varepsilon_0, \]

where \mathbf{E}(r) is the field being projected and x = x_0 is an arbitrary plane. This result is in agreement with that of Marcuse [20] and the standard form for overlap integrals [21]. The projection operator \mathcal{P}_k has the useful property that \mathcal{P}_k \mathbf{e}_k(r) e^{i k x} = \delta_{k,k'}.

1.2. Green function approach for the electric field

The electric field properties of the disordered structure can be calculated analytically from the Green function solution to the electric field wave equation, namely

\[ \mathbf{E}(r; \omega) = \mathbf{E}_k(r; \omega) + \int_{\text{all space}} d\alpha d\beta \mathbf{G}(r, r'; \omega) \mathbf{e}_k(r') e^{i k x} = \frac{1}{\sqrt{\varepsilon_0 \mu_0}} \sqrt{\mu_0/\varepsilon_0} \nabla \times \nabla \times \hat{e}_k(r), \]

where the overbar represents a tensor or dyadic. For convenience, we partition the Green function into contributions from the bound waveguide mode, radiation modes, and other modes as

\[ \mathbf{G}(r, r'; \omega) = \mathbf{G}_B(r, r'; \omega) + \mathbf{G}_R(r, r'; \omega) + \mathbf{G}_T(r, r'; \omega). \]

The bound mode Green function is given analytically from properties of the bound mode [6, 19]

\[ \mathbf{G}_B(r, r'; \omega) = \frac{\alpha \omega}{2 \nu_g} \mathbf{e}_k(r) \otimes \mathbf{e}_k(r') e^{i k (t - x)} \Theta(t - x), \]

where the group velocity, \nu_g, is assumed positive (in the case of anomalous dispersion, \nu_g is then negative), \otimes is a tensor

\[ \mathbf{G}(r, r'; \omega) = \mathbf{G}_B(r, r'; \omega) + \mathbf{G}_R(r, r'; \omega) + \mathbf{G}_T(r, r'; \omega). \]
product, $e_{-k}(\mathbf{r}) = e_k^*(\mathbf{r})$, and $\Theta(x)$ is the Heaviside step function, equal to one if $x > 0$ and 0 if $x < 0$. The mode properties can be calculated with any mode solving technique; for example, we use a freely available plane wave expansion code [22].

The radiation Green function, $\mathbf{G}_R(\mathbf{r}, \mathbf{r}'; \omega)$, contains contributions from the continuum of radiation modes above the light line that are not confined to the slab by total internal reflection. The radiation Green function, whose contribution is significantly smaller than the dominant bound mode, is rather featureless and is well approximated by using a homogeneous dielectric slab with an effective permittivity determined through numerical finite-difference time-domain (FDTD) [23] simulations. We compute the radiation Green function efficiently by using the method of [24] (see also [19] for more details of our specific implementation).

The remainder of the contributions to the Green function are contained in $\mathbf{G}_0(\mathbf{r}, \mathbf{r}'; \omega)$ (‘O’ represents others), such as the possibility of having other modes (bound or leaky), and the divergence contribution of the real part of the Green function as $\mathbf{r} \to \mathbf{r}'$. Since we consider a waveguide with one bound mode in the frequency range of interest, we can safely neglect other bound modes. For the divergent contribution to $\mathbf{G}_0(\mathbf{r}, \mathbf{r}'; \omega)$, we shall neglect its contribution in this work; the dominant effect is to cause a frequency shift [25] and introduce local field corrections [12, 26].

1.3. Forward wave envelope equation

The electric field in the ideal waveguide can be decomposed into the complete Bloch mode basis consisting of the target bound waveguide modes $e_{\pm k}(\mathbf{r})$, and the set of radiation modes $\{\mathbf{q}(\mathbf{r})\}$ as

$$\mathbf{E}(\mathbf{r}; \omega) = \mathcal{E}_0 \left[ e_k(\mathbf{r}) e^{i k_z x} \psi_1(x) + e_k^*(\mathbf{r}) e^{-i k_z x} \psi_b(x) + \sum_q \mathbf{q}(\mathbf{r}) e^{i q z} \psi_q(x) \right],$$

(8)

where $\mathcal{E}_0$ is an amplitude and $\psi_1(x)$, $\psi_b(x)$, and $\{\psi_q(x)\}$ are the envelopes for the forward, backward, and radiation modes. We stress that we use envelopes only for convenience and do not require that they are slowly varying. Although we are only interested in the envelopes for the bound waveguide modes, we initially track the radiation modes to include radiation scattering.

The field in a disordered waveguide can be calculated analytically from equation (4), using the effective PC waveguide Green function and the disorder polarization density $\mathbf{P}(\mathbf{r}; \omega) = \epsilon_0 \Delta \varepsilon(\mathbf{r}) \mathbf{E}(\mathbf{r}; \omega)$, as

$$\mathbf{E}(\mathbf{r}; \omega) \simeq \mathcal{E}_0 \left[ e_k(\mathbf{r}) e^{i k_z x} \psi_1(x) + e_k^*(\mathbf{r}) e^{-i k_z x} \psi_b(x) + \int d\mathbf{r}' \left[ \mathbf{G}_B(\mathbf{r}, \mathbf{r}'; \omega) + \mathbf{G}_R(\mathbf{r}, \mathbf{r}'; \omega) \right] \cdot \left[ \Delta \varepsilon(\mathbf{r}') \mathbf{E}(\mathbf{r}'; \omega) \right] \right],$$

(9)

where $\Delta \varepsilon(\mathbf{r}) = \varepsilon(\mathbf{r}) - \varepsilon_i(\mathbf{r})$ is the disorder function and $\varepsilon_i(\mathbf{r})$ is the dielectric constant for the ideal structure. We assume an initial electric field $\mathbf{E}_0(\mathbf{r}; \omega) = \mathcal{E}_0 e_k(\mathbf{r}) e^{i k_z x}$, and a total field including scattering $\mathbf{E}(\mathbf{r}; \omega)$ given by equation (8). The explicit notation of the integration limits has been dropped; unless stated otherwise the integration is performed over the entire domain of the variable of integration (all space).

We begin by projecting equation (9) onto a forward propagating wave by operating with $\mathbf{P}_F$. We then multiply by $e_k^{\pm 1}$ and differentiate with respect to $x$. The left-hand side becomes simply $d\psi_1(x)/dx$. The projection of $\mathbf{E}_0(\mathbf{r}; \omega)$ equals one and differentiating eliminates the contribution of the field in the ideal structure. This derivation will transform the integral description of the total electric field into a set of coupled propagation equations and the electric field in the ideal structure will be included as a wave injected from the input port. Equation (9) for the forward wave becomes

$$\frac{d}{dx} \psi_1(x) = \frac{i}{v_g} \left[ c_{b1}(x) \psi_1(x) + c_{b1}(x) e^{-2ikx} \psi_b(x) + \sum_q c_{q1}(x) \psi_q(x) \right].$$

(10)

The terms on the right-hand side all arise from the projection of the $\mathbf{G}_B(\mathbf{r}, \mathbf{r}'; \omega)$ term; the projection of the $\mathbf{G}_0(\mathbf{r}, \mathbf{r}'; \omega)$ term is 0 since the constituent radiation modes are orthogonal to the chosen bound mode. The volume integral has been converted to an integral over the transverse plane by the derivative of the Heaviside function in $\mathbf{G}_B(\mathbf{r}, \mathbf{r}'; \omega)$. The scattering coefficients, corresponding to forward-forward, forward-backward, and forward-radiation scatter, are

$$c_{b1}(x) = \frac{a \omega}{2} \int dy dz e_k^*(\mathbf{r}) \cdot e_k(\mathbf{r}) \Delta \varepsilon(\mathbf{r}),$$

(11)

$$c_{b2}(x) = \frac{a \omega}{2} \int dy dz e_k^*(\mathbf{r}) \cdot e_k(\mathbf{r}) \Delta \varepsilon(\mathbf{r}),$$

(12)

$$c_{q1}(x) = \frac{a \omega}{2} \int dy dz e_k^*(\mathbf{r}) e^{-ikx} \cdot \mathbf{q}(\mathbf{r}) e^{ikx} \Delta \varepsilon(\mathbf{r}).$$

(13)

The $c_{q1}(x)$ coefficients are cumbersome to work with since they require knowledge of all the radiation modes of the system. Later we will combine the sum over these coefficients into a single effective radiative scatter coefficient, $c_{b1}(x)$.

An analogous equation to equation (10) for $d\psi_1(x)/dx$ is formed by projecting equation (9) onto a backward propagating wave. One has

$$\frac{d}{dx} \psi_b(x) = \frac{-i}{v_g} \left[ c_{b1}(x) \psi_1(x) + c_{b2}(x) e^{2ikx} \psi_1(x) + \sum_q c_{q2}(x) \psi_q(x) \right].$$

(14)

where the negative sign arises from the Heaviside function in equation (7), $c_{b2}(x) = c_{f1}(x)$, $c_{q2}(x) = c_{b1}(x)$, and

$$c_{q2}(x) = \frac{a \omega}{2} \int dy dz e_k^*(\mathbf{r}) e^{ikx} \cdot \mathbf{q}(\mathbf{r}) e^{-ikx} \Delta \varepsilon(\mathbf{r}).$$

1.4. Disorder-mediated coupled mode equations

Next, we seek to eliminate the $\psi_q(x)$ from the equation since there are a large (infinite) number of radiation modes, and we would rather not have to solve for all the $\psi_q(x)$. We project equation (9) onto any one of the radiation modes to derive a radiation mode envelope equation. The left-hand side becomes

$$\frac{d}{dx} \psi_q(x) = \frac{a \omega}{2} \int dy dz e_k^*(\mathbf{r}) e^{ikx} \cdot \mathbf{q}(\mathbf{r}) e^{-ikx} \Delta \varepsilon(\mathbf{r}).$$

simply \( \psi_q(x) \). Only the \( \mathbf{G}_{\mathbf{r}}(\mathbf{r}, \mathbf{r}'; \omega) \) term on the right-hand side will have a non-zero projection since any chosen radiation mode will be orthogonal to the bound waveguide modes. Thus we obtain a set of equations, one for each of the radiation modes \( q \).

\[
\psi_q(x) = E_0^{−1} P_q \int d\mathbf{r}' \mathbf{G}_{\mathbf{r}}(\mathbf{r}, \mathbf{r}'; \omega) \cdot [E(\mathbf{r}'; \omega) \Delta \varepsilon(\mathbf{r}')] \\
+ P_q \int d\mathbf{r}' \mathbf{G}_{\mathbf{r}}(\mathbf{r}, \mathbf{r}'; \omega) \cdot \mathbf{e}_q(\mathbf{r}') e^{-i k x} \psi_i(x) \Delta \varepsilon(\mathbf{r}').
\]

This is equation (15) with only term (15)

In equation (10), the projected Green function (in wave equation is interpreted as a matrix multiplication by the basis vector (in identity transform of \( \mathbf{G}_{\mathbf{r}}(\mathbf{r}, \mathbf{r}'; \omega) \)) are \( \mathbf{e}_q(\mathbf{r}') \) multiplied by the basis vector (in \( \psi_{\mathbf{r}}(\mathbf{r}; \omega) \)) and then the forward mode. These assumptions are reasonable because the radiation modes quickly leak from the slab and so do not interact with the scattering regions for very long. This leaves only (15a) which accounts for loss from the forward mode into the radiation modes. The \( P_q \) prefix in equation (15) is a projection operator acting on the radiation Green function. In equation (10), the projected Green function (in \( \psi_q(x) \)) is multiplied by the basis vector (in \( \mathbf{e}_q(\mathbf{r}') \)). Since the set \( \{q(\mathbf{r})\} \) spans all radiation modes included in \( \mathbf{G}_{\mathbf{r}}(\mathbf{r}, \mathbf{r}'; \omega) \), this is an identity transform of \( \mathbf{G}_{\mathbf{r}}(\mathbf{r}, \mathbf{r}'; \omega) \) and equation (10), under substitution by equation (15), becomes

\[
\psi(x) = i c_{fi}(x) \psi_i(x) + i c_{fi}(x) e^{-i k x} \psi_b(x) + i c_{bi}(x) \psi_i(x),
\]

where the effective radiation loss coefficient \( c_{fi} \) (given in equation (18)) includes the sum over the individual radiation mode wave amplitudes \( \psi_q \) and scattering coefficients \( c_{eq} \) (equation (13)). Note that we have conveniently eliminated the sum over \( q \).

For the backward wave, equation (14) is transformed using equation (15) with only term (15b) retained. The backward wave equation is

\[
\frac{d\psi_b(x)}{dx} = i c_{fb}(x) \psi_b(x) + i c_{fb}(x) e^{i k x} \psi_i(x) + i c_{bi}(x) \psi_b(x).
\]

The final coupled mode equations are equations (16) and (17). The coupling coefficients can be physically interpreted as \( c_{fb} = c_{bh} (11) \) driving scattering from a mode into itself, \( c_{bi} = c_{hb} (12) \) driving scattering into the counter-propagating mode, and \( c_{fi} \) and \( c_{be} \) driving scattering from the waveguide mode into radiation modes above the light line. With the elimination of the radiation mode envelopes, the coupling coefficients into radiation modes become

\[
c_{fi}(x) = \frac{\alpha_0}{2} \int \int dy dz \int_{all space} d\mathbf{r}' \Delta \varepsilon(\mathbf{r}) \Delta \varepsilon(\mathbf{r}') e^{-i k x} \psi_i(x) \psi_b(x),
\]

for the backward wave, equation (14) is transformed using equation (15), and scattering from a mode into itself, \( c_{fi} = c_{bi} \) (11) driving scattering from a mode into itself, \( c_{bi} = c_{hb} \) (12) driving scattering into the counter-propagating mode, and \( c_{fi} \) and \( c_{be} \) driving scattering from the waveguide mode into radiation modes above the light line. With the elimination of the radiation mode envelopes, the coupling coefficients into radiation modes become

\[
c_{fi}(x) = \frac{\alpha_0}{2} \int \int dy dz \int_{all space} d\mathbf{r}' \Delta \varepsilon(\mathbf{r}) \Delta \varepsilon(\mathbf{r}') e^{-i k x} \psi_i(x) \psi_b(x),
\]

Importantly, this theory incorporates the full 3D structure of the waveguide, Bloch modes, and disorder functions in calculating the scattering.

The radiation scattering coefficients of equations (18) and (19) are difficult to evaluate due to the integral over the entire waveguide. Although we assume disorder between holes is uncorrelated in the expectation sense, for any instance of disorder, there may be a non-zero correlation between holes mediated by radiation modes. However, we are primarily interested in coherent scattering that is contained within the waveguide, and can reasonably assume that any field scattered out of a bound mode will not be scattered back into a bound mode; this justifies as the bound mode scattering channel is by far the dominant one. Therefore, we can simplify the radiation loss by using \( c_{fi} = i (\langle \alpha_{rad} \rangle / 2 \alpha) \). Figure (18) is the incoherent average radiation loss [6]

\[
\langle \alpha_{rad} \rangle = \frac{\alpha_0}{2} \int \int dy dz \int_{all space} d\mathbf{r}' \Delta \varepsilon(\mathbf{r}) \Delta \varepsilon(\mathbf{r}') e^{-i k x} \psi_i(x) \psi_b(x),
\]

Here the integrations are over a single unit cell. Comparing equations (18) and (18), the former is just the expectation value of the imaginary part of the latter, integrated along \( x \). The factor of two is necessary to convert from a power loss to an amplitude loss.

For modelling an incident field at one end of the waveguide, the boundary conditions for a wave injected into the waveguide (and consistent with \( E_i(\mathbf{r}; \omega) \)) are

\[
\psi(x_{start}) = 1,
\]

\[
\psi_b(x_{end}) = 0,
\]

where \( x_{start} \) and \( x_{end} \) are the positions of the input and output ports. The propagating envelopes are then computed at all spatial position within the waveguide using the presented coupled mode equations (equations (16) and (17)).

2. Disorder model

The equations can now be used with any disorder model. In our experience [4, 13] and in agreement with the analysis of images of PC slabs [27], we have found that disorder in PC slab structures is dominated by perturbations of the perimeter of the holes, as shown in figure 2. We take the radial perturbation \( \Delta r \) to be a Gaussian random variable with a mean of 0 and a standard deviation of \( \sigma \). Two radial perturbations are correlated by

\[
\langle \Delta r_i(\phi_i) \Delta r_j(\phi_j) \rangle = \sigma^2 e^{-R(\phi_i−\phi_j)/\Delta r} \delta_{ij},
\]
where the subscript indexes the holes, \( \phi_i \) is the angular position of the point measured about the centre of the hole, \( R \) is the ideal hole radius, and \( L_p \) is the correlation length measured around the circumference.

The change in dielectric constant about a single hole \( i \) is given exactly by

\[
\Delta \varepsilon_i (r_i, \phi_i) = (\varepsilon_2 - \varepsilon_1) \left[ \Theta(r_i - R) - \Theta(r_i - R - \Delta r_i(\phi_i)) \right],
\]

where \((r_i, \phi_i)\) are cylindrical coordinates centred about hole \( i \). This form holds for both positive and negative values of \( \Delta r_i(\phi_i) \). The disorder \( \Delta \varepsilon \) appears in the formalism in spatial integrals where it is multiplied by functions of the electric fields and Green function. We consider such an integration, where \( f(r_i, \phi_i) \) represents one of the scalar fields and is slowly varying over the relevant length scale (e.g., \( f(r_i, \phi_i) = \mathbf{e}_i(r) \cdot \mathbf{e}_i \)). The field \( f(r_i, \phi_i) \) can be expanded in a Taylor series along the radial coordinate to evaluate the integral as

\[
\int dr_i \Delta \varepsilon_i (r_i, \phi_i) f(r_i, \phi_i) = \int dr_i \Delta \varepsilon_i (r_i, \phi_i) \times (f(R, \phi_i) + f'(R, \phi_i)(r_i - R) + O((r_i - R)^2))
\]

\[
= f(R, \phi_i) \int dr_i \Delta \varepsilon_i (r_i, \phi_i)
\]

\[
+ f'(R, \phi_i) \int dr_i \Delta \varepsilon_i (r_i, \phi_i)(r_i - R) + O((r_i - R)^2)
\]

\[
= f(R, \phi_i) (\varepsilon_2 - \varepsilon_1) \Delta r_i(\phi_i)
\]

\[
+ f'(R, \phi_i) (\varepsilon_2 - \varepsilon_1) \frac{\Delta r_i(\phi_i)^2}{2} + O(\Delta r_i(\phi_i)^3).
\]

(25)

To include the disorder to first order in \( \Delta r_i(\phi_i) \), it is sufficient to take the field at the ideal hole radius \( f(R, \phi_i) \). For convenience of notation, we then rewrite equation (24) as

\[
\Delta \varepsilon_i (r_i, \phi_i) = (\varepsilon_2 - \varepsilon_1) \delta(r_i - R) \Delta r_i(\phi_i),
\]

which agrees with equation (25) to first order.

3. Implementation

3.1. Ideal structure

This calculation requires, as inputs, the ideal waveguide mode dispersion and spatial field distribution. As a representative example we consider a W1 semiconductor waveguide with pitch \( a = 480 \text{ nm} \) and index of refraction \( n = 3.18 \). The dispersion of the waveguide mode is shown in figure 3(a) (blue, solid, left scale) along with the group index (green, dashed, right scale). Near the band edge \( (k = 2\pi/a) \), the group index is large, increasing scattering as the light slows down. The spatial distribution of the electric field in the centre of the slab is shown in figure 3(b).
3.2. Numerical implementation

To solve equations (16) and (17) numerically, the coupling coefficients are assumed to be constant over a short ($\Delta x \ll a$) interval in $x$ and are integrated analytically. This yields a pair of transfer equations linking the envelopes on either side of the chosen interval. In this way, a set of transfer equations that span the entire waveguide length can be built, and then solved using linear algebra techniques. This approach is particularly amenable to adding reflective facets and other features by simply including an appropriate transfer matrix.

The average coupling constants for each interval are calculated by, for each hole, generating an instance of a disordered profile from the statistical distribution of equation (23). The coupling coefficients are calculated at multiple points within the interval, and then averaged. Typically, there are 20 intervals per unit cell to satisfy the assumption that the coefficients are relatively constant. As shown in figure 4, if the discretization of the unit cell is too coarse, the loss is underestimated. Thus, one must include sub-unit-cell propagation effects.

We highlight that the calculation is orders of magnitude more efficient than standard brute-force numerical techniques, e.g. FDTD. We also note that we only need to calculate the coupled mode coefficients wherever disorder has an influence, namely at the hole interfaces. However, the final computation, though efficient, is not instantaneous. Producing a high resolution transmission spectrum (1000 frequency points) for a 1 mm waveguide (2500 unit cells and 50000 grid points) takes approximately 1 cpu day (on a 2.4 GHz AMD Opteron processor). However, the calculations at each frequency are independent and the total calculation can also be greatly accelerated by exploiting parallelism. In contrast, we estimate that a minimum of about 40 GBytes of memory and 5800 cpu days are required to perform the simulation using FDTD. Clearly, this semi-analytic treatment is a significant advantage.

4. Computed transmission spectra

Figure 5 shows transmission spectra for four disordered waveguides calculated by solving equations (16) and (17) (blue, solid). For reference, previous incoherent scattering results, computed within a second-order Born approximation [6], are also shown (red, dashed); we also note that extensions to the incoherent scattering theory to account for multiple scattering have been introduced recently [13]. Each row of plots is for a different waveguide with the left plot showing a broad frequency range and the right plot showing a narrow frequency range near the band edge. Plots (a) and (b) are for a disordered 1.5 mm waveguide. Plots (c) and (d) are for a different disorder instance of the same 1.5 mm waveguide. Plots ((e) and (f)) and plots ((g) and (h)) are for the same disorder instance as ((c) and (d)) but with the length reduced to 1.0 mm and 0.5 mm respectively. The calculation uses an RMS roughness of $\sigma = 3$ nm, and a disorder correlation length of $l_p = 40$ nm. The forward wave intensity as a function of position is given in figure 6 for the two points marked with crosses in (d).
1.0 mm and 0.5 mm respectively. Here the qualitative roll off changes due to the length reduction but disordered resonances can be found at similar frequencies across the three lengths, especially between the 1.5 and 1.0 mm cases.

We can examine the position-dependent distribution of energy in the waveguide under continuous wave illumination. In the second row, right column of figure 5, a neighbouring transmission minimum and maximum are marked with red crosses. The forward wave intensity at these frequencies is plotted in figure 6. Although the points are very close in frequency, the minute difference in group index \(n_g = 25.11\) compared to \(n_g = 24.96\) creates a difference in the accumulated phase and a dramatic change in the transmission.

By including multiple, coherent scattering we reproduce the experimental phenomenon of sharp spectral resonances near the band edge. Although initially unexpected, these features are just disorder-induced Fabry–Pérot-like fringes between extrinsic scattering sites. The slow group velocity enhances scattering to create the scattering sites and also increases the effective cavity length between sites, narrowing the resonance line-width. It may be tempting to think of these resonances as an example of Anderson localization [28, 29], but by using the established 3D criterion for Anderson localization [30, 31], we are actually far from the criterion for true strong localization [15].

5. Conclusions

We have described and applied a theory for self-consistently modelling coherent scattering and light localization in a disordered PC waveguide instance, allowing one to map directly onto a realistic experimental situation. Slow light propagation enhances back scattering (and, to a lesser extent, radiation scattering) leading to high losses near the band edge. The formation of sharp spectral resonances and localization near the band edge is shown and is mediated by Fabry–Pérot-like resonances between disorder sites. This theory is computationally efficient, making the analysis of very long waveguides (thousands of periods using the full 3D structure) feasible on a desktop computer. Although the presented model may not be quantitatively exact (e.g. it neglects local field effects), the qualitative results such as the formation of sharp resonances near the band edge certainly can, and already have been, used to explain a rich range of experimental features without introducing any fitting parameters [15]. The role of local field effects will be reported in future work, and the effects on incoherent frequency shifts are described elsewhere [32].

Acknowledgments

This work was supported by the National Sciences and Engineering Research Council of Canada, and the Canadian Foundation for Innovation. We thanks S Combré and A De Rossi for many useful discussions.

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