Sample-efficient Search for Satisfactory OLED Designs

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Abstract

Higher efficiency OLEDs can operate with less power, but identifying efficient designs can be a slow process because the cost of fabricating proposed designs is extreme. Computational simulations are more expedient, but they provide only an approximation to the eventual manufactured outcome. Moreover, blind search for the optimal structure with simulations is not a favorable choice due to the number of design parameters and high computational costs. We apply the Constraint Active Search methodology to identify a diverse set of designs which all have suitably high efficiency for eventual consideration for physical fabrication.

Introduction

Organic light emitting diodes (OLEDs) are a fundamental part of modern electronics; cell phones, monitors and television screens use OLEDs as a power-efficient mechanism for facilitating their displays. An OLED device is made of multiple layers to emit the light. The most generic OLED device is made of a cathode layer, an organic layer which the light is generated in, a transparent anode layer and a glass substrate. Although the modern manufactured OLEDs are much more complicated and have many different layers and multiple doped organic layers to increase the internal quantum efficiency, we consider a less complex model to simulate for the sake of simplicity and because most of the additional layers, mainly organic layers, have very close refractive index and this does not affect the out-coupling efficiency that we are investigating. See Figure 1 for a depiction of the OLEDs we consider here.

In the presented model, to increase the extraction efficiency, an anti-reflection layer is introduced between the Anode (ITO) and the glass substrate. In general, a middle layer can be used to reduce the reflection of light at the contact surface of two materials with different refractive indexes. The additional layer gives the best performance if its refractive index follows this equation: $$n_3 = \sqrt{n_1 \cdot n_2}.$$ Refractive indexes of the ITO and glass are 1.94 and 1.46 respectively at the wavelength of 520nm (green light) which the simulations are performed at. The best material to use as anti-reflection layer should have refractive index of $$n = \sqrt{1.94 \cdot 1.46} = 1.68.$$ Figure 1: This graphic representation of the cross-section of an OLED showing the elements of OLED design under consideration during our experimentation. Text in red denotes parameters we consider tuning, and text in black denotes design choices we have fixed for this article.

During the last two decades, many successful works have been done to increase the internal quantum efficiency to 100% (Adachi et al. 2001; Xiao et al. 2009; Zhang et al. 2015; Williams et al. 2007) and today’s problem with low efficiencies of OLED devices is mainly the out-coupling efficiency and the trapped light within the OLED structure that keeps the EQE low. Department of Energy has set the target extraction efficiency for OLED devices to 75% for

Efficiency of an OLED device is represented by External Quantum Efficiency (EQE) which encompasses two distinct factors: internal quantum efficiency and out-coupling efficiency. EQE is the ratio of the number of the emitted photons to the number of the injected electron charges (Glowacki et al. 2012). Equation 1 defines the OLED efficiency:

$$EQE = \gamma \cdot \eta_{\text{rad, eff}} \cdot \eta_{\text{S/T}} \cdot \eta_{\text{out}} \quad (1)$$

in which $$\eta_{\text{int}} = \gamma \cdot \eta_{\text{rad, eff}} \cdot \eta_{\text{S/T}}$$ is defined as internal quantum efficiency and $$\eta_{\text{out}}$$ is the out-coupling efficiency. Internal quantum efficiency encircles efficiencies pertaining to the used materials and manufacturing quality such as electrical efficiency ($\gamma$), effectiveness of the emitter ($\eta_{\text{rad, eff}}$), and the portion of excitons turning into emitted photons ($\eta_{\text{S/T}}$), while the out-coupling efficiency ($\eta_{\text{out}}$) represents the efficiency of the optical environment design.

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2035 and 60% for interim 2025 target (Pattison et al. 2020). Finite-difference time-domain (FDTD) simulations helps us to understand the out-coupling efficiency of different OLED structures and enables us to search for optimal designs. To perform FDTD simulations, Lumerical software has been used.

Many different OLED structures were simulated and manufactured to increase the extraction efficiencies. Micro-lens arrays were casted on top of the glass substrate by (Wei and Su 2004; Ho et al. 2013) and showed upto 56% and 74% improvement in luminance efficiency respectively. Also, OLED devices with cylindrical micro-lens arrays was manufactured by (Lee et al. 2008) but comprised the image quality to increase the outcoupling efficiency. A corrugated oled model was proposed and manufactured by (Schwab et al. 2014) and their measurements showed 13.5% increase in EQE. In their model corrugation are in all the OLED layers similar to the conformal structure proposed in this article.

In this article, we focus on improving the OLED extraction efficiency (out-coupling efficiency) through a conformal OLED structure. Previous research has shown that tempering with flat structures can be a promising method to achieve higher light extraction values or lower reflections, see (Haghanifar et al. 2020) for an application of aditively manufactured cones on top of the glass to reduce the reflection of a piece of glass.

**Numerical Simulation**

Lumerical FDTD is a numerical solver of Maxwell’s equations for complex geometries and is usable for all electromagnetics and photonics problems. For more information please see (Lumerical Inc. 2021a).

The simulated model is composed of the cathode layer: Aluminum, organic layer(s): Ir(ppy)$_3$, Anode: ITO, anti-reflection layer: Al$_2$O$_3$, and glass substrate. In practice, emission occurs at all points in the organic layer; though, to be able to model the emission of light in simulations we should consider enough number of dipoles within the organic layer to cover the whole area properly and average the results over all dipoles to obtain the final result. The vertical location of the dipoles within the organic layer is assumed to be half of the layer thickness (the red dashed line in Figure 1). Also due to the symmetry, we only consider half of the device pitch to place the dipoles.

FDTD is inherently a coherent simulation method. To achieve incoherent results which we are interested in, we should understand and consider coherency in FDTD simulations. For comprehensive explanation please see (Lumerical Inc. 2021b). For our 2D OLED simulations, we should consider spatial incoherence meaning to avoid including more than one dipole in each simulation and also avoid periodic boundary conditions when electromagnetic filed is not close to zero at boundaries, and also consider polarization incoherence meaning to average the results over three orthogonal dipoles at each dipole location to obtain the results for an unpolarized beam. Simulation domain or width of the OLED device should be large enough to get an accurate result. Photons traveling between the OLED layers can eventually find their way out of the structure after traveling through a couple of conformal structures. Therefore, it is necessary to make sure the simulation width has been chosen large enough.

Thickness of the layers in the simulated OLED devices in this paper are shown in Table 1. Although for further optimization, layer thicknesses can be chosen as optimization variables, in all simulations in the current study these parameters are fixed and are not included for optimization.

Also note that the initially defined pitch value will be modified during the simulation according to the layer thicknesses by the factor of $2 \cdot t_1 \cdot \tan(\theta/2)$ to ensure each layer has the same thickness around the cones and also each cone has properly spaced by their adjacent structures. See Figure 1 for details.

Table 1: Thickness of OLED layers for the simulated device.

<table>
<thead>
<tr>
<th>Layer</th>
<th>Thickness</th>
</tr>
</thead>
<tbody>
<tr>
<td>Organic Layer(s)</td>
<td>200 nm</td>
</tr>
<tr>
<td>Anode</td>
<td>100 nm</td>
</tr>
<tr>
<td>Anti-reflection</td>
<td>200 nm</td>
</tr>
<tr>
<td>Glass</td>
<td>200 nm</td>
</tr>
</tbody>
</table>

To ensure that performed FDTD simulations are valid and accurate, three certain parameters that directly affects the computational accuracy was investigated: mesh size, simulation width, and number of dipoles.

**Mesh size.** FDTD method solves Maxwell’s equations by discretization of the simulation domain and time. Similar to the all other discretized computational methods, usually, smaller mesh size means higher accuracy but also means higher computational times. To find the best uniform mesh size, multiple simulations with different mesh size for two different structures, one with smaller geometries and one with larger geometries, was performed. Table 2 shows the structures used in convergence tests.

Table 2: Size of the structures investigated for convergence tests.

<table>
<thead>
<tr>
<th>Structure</th>
<th>Height</th>
<th>Pitch</th>
<th>$D_{out}$</th>
<th>$D_{inp}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Small</td>
<td>800 nm</td>
<td>400 nm</td>
<td>300 nm</td>
<td>100 nm</td>
</tr>
<tr>
<td>Large</td>
<td>700 nm</td>
<td>2500 nm</td>
<td>2000 nm</td>
<td>500 nm</td>
</tr>
</tbody>
</table>

Figure 2 shows that by choosing mesh size smaller than or equal to 10nm, we can have reliable results for OLED extraction efficiency values. For faster but a slightly inaccurate results 10nm mesh size is favorable to use. To have high accuracy 5nm mesh size should be used in the simulations.

**Simulation width.** As discussed before, the simulation domain should be chosen large enough so that we make sure the obtained extraction efficiency has converged properly to the actual value. If a small simulation domain is selected then the result value for the extraction efficiency would be lower. shows how extraction efficiency changes for different initial simulation domains.
Number of dipoles. One important parameter in the OLED simulations is the number of dipoles. The red dashed line in Figure 1 shows all the possible locations for the dipoles. To have the most accurate result, unlimited number of dipoles should be averaged on this line, but it is obviously impossible. To have an insight of how efficiency changes for different dipole locations, Figure 4 shows the variation of the out-coupling efficiency for 50 uniformly distributed dipoles along half the pitch for each of the two structures. Red lines denotes the average values of these 50 dipoles which we can safely consider as the actual out-coupling efficiency of the OLED device.

Before delving into the proposed optimization procedure in this paper, we represent two simple approaches for dipole selection to show their difference and accuracy: uniform selection and random selection.

For uniform selection, we can consider selecting e.g. 10 dipoles uniformly distributed in half the pitch (red dashed line in Figure 1) and average efficiencies over those 10 dipoles. For random selection, we can randomly choose e.g. 10 dipoles and average the results. Figure 5 and Figure 6 show the results for these two approaches. In random selection approach, for each sample size (e.g. 10 randomly selected dipoles) the sampling was repeated 100 times and the mean and standard deviation of the extraction efficiencies of these 100 samples were calculated and plotted.

Our goal in this research is to maximize the OLED extraction efficiency, denoted $\eta_{out}$, as measured as the fraction of the emitted photons form the OLED through the glass substrate.

Identifying Satisfactory Designs

As indicated earlier, we only consider the implications of the additive manufacturing on the efficiency – the other OLED elements are fixed throughout our experimentation. Our search domain (see Figure 1 for reference) is defined as:

- Simulation Domain: $a \in [100, 2000]$ nm
- Cone Top: $d_{top} \in [100, 2000]$ nm
- Cone Bottom: $d_{bot} \in [100, 2000]$ nm
- Height: $h \in [100, 2000]$ nm

To enforce the cone shape during manufacturing we enforce the domain constraints:

- Simulation Domain > Cone Bottom, and
- Cone Bottom > Cone Top.

We denote this constrained search space as $\mathcal{X}$ and each individual design as $x \in \mathcal{X}$.

As described earlier, our goal in this project is to identify an OLED design involving these 4 parameters which can surpass 75% extraction efficiency. We use the numerical simulation strategy described earlier to search for this
design, but the numerical simulation is only an approximation to the performance of the ultimate fabrication. The mesh size, simulation width and number of dipoles involved in the simulation all affect the accuracy; additionally. Furthermore, as discussed in (Haghanifar et al. 2020), the physical manufacturing process incurs its own imprecisions in actually producing the prescribed design (the maskless reactive ion etching fabrication process (Haghanifar et al. 2017) will not be precise at 1 nm length scales).

**Constraint Active Search**

To account for the gap between the problem we are studying (the efficiency estimated in our numerical simulation) and the actual problem we want to study (the efficiency of the physical OLED), we utilize Constraint Active Search (Malkomes et al. 2021). This strategy replaces the optimization of efficiency with a search for a diverse set of designs satisfying efficiency, the final choice, or choices, of fabrication can be made from this set based on other criteria (such as fabrication speed, consistency, durability, etc). We choose 70% as our satisfactory value, under the belief that the gap between simulation and fabricated performance may be 5%.

Constraint active search (CAS) requires an acquisition function, and we use the probability of success acquisition function

\[
g(x) = P(Y_x > \tau|D),
\]

where \(Y_x|D\) denotes a Gaussian process (Fasshauer and Mc-Court 2015) \(Y\) considered at the location \(x\) conditioned on \(D\), the data observed thus far (\(x_i\) and \(\text{EQE}(x_i)\) values) and \(\tau\) represents our definition of success (which we stated will be 70%).

In (Malkomes et al. 2021), the authors use a resolution parameter to define a desire to search away from previously observed \(x\) values – in this setting, this has the effect of accounting for our limited manufacturing precision by not over-resolving a small region of the design space. We use a resolution parameter of 150 here, which is incorporated into our acquisition function optimization process by only optimizing over points which are at least 150 nm away in design space, i.e., the next design to test satisfies 

\[
\max_{x \in \mathcal{X}} g(x), \text{ such that } \forall x_i \in D, \|x - x_i\|_2 > 150.
\]

In the random search (RS) settings, we also account for the limited manufacturing precision by restricting all parameters to 10 nm increments, i.e., Cone Bottom can only take values 100, 110, 120, . . . , 2000 nm; this matches the strategy used in (Haghanifar et al. 2020).

**Search Implementation Considerations**

For each design \(x \in \mathcal{X}\), the resulting efficiency evaluation \(\text{EQE}(x)\) is the sum of three integrals: the \(x\), \(y\) and \(z\)-axis emissions integrated over the possible dipole locations as depicted with the dashed red line in Figure 1. Without loss of generality, the dipole locations can be parametrized as being within a domain \([0, 1]\), thus the efficiency values may be written as

\[
\text{EQE}(x) = \int_0^1 \text{EQE}_x(x; p) \, dp \\
+ \int_0^1 \text{EQE}_y(x; p) \, dp \\
+ \int_0^1 \text{EQE}_z(x; p) \, dp,
\]

where \(\text{EQE}_x(x; p)\), \(\text{EQE}_y(x; p)\) and \(\text{EQE}_z(x; p)\) are the efficiencies along the \(x\), \(y\) and \(z\) axes for a single dipole \(p \in [0, 1]\).

These integrals can be computed in any fashion, but we can leverage the structure of our optimization algorithm to estimate the integrals at low accuracy before improving the quality of only those with high efficiency. We estimate the integral with Quasi Monte Carlo (Choi et al. 2020+), allowing up to 33 total points in the estimate; for example, in the \(x\)-axis, this would look like:

\[
\int_0^1 \text{EQE}_x(x; p) \, dp = \frac{1}{n} \sum_{i=1}^{n} \text{EQE}_x(x; p_i)
\]

where \(p_i\) come from the Halton sequence and \(n \leq 33\).

After a new design \(x\) has been chosen (whether through RS or CAS) We use the first 5 \(p_i\) in the sequence and study the mean and variance of the integral estimator in (3). As the optimization/search progresses, if the integral for a given design \(x\) has too much variance, another more dipoles are included in the integral estimate (which reduces the variance), up to 33 total dipoles. This strategy is analogous to that of early stopping in machine learning (Caruana, Lawrence, and Giles 2001) – it has also been used in a medical imaging setting (McCourt, Dewancker, and Ganci 2016).

In RS, we consider a design to have too much variance if its 2 standard deviation confidence interval contains the mean of the design with the highest mean seen thus far (including the highest mean seen thus far). For CAS, a design has too much variance if its 2 standard deviation confidence interval contains the efficiency threshold 70%. To seed the optimization/search, we initialize with 12 designs drawn randomly from \(\mathcal{X}\) before conducting any termination.
To balance between decreasing variance and searching \( \chi \), we keep a global max dipole count, starting at 5 and increasing by 4 (up to 33) each time a new \( x \) is considered. After the new \( x \) has computed its dipoles up to the global max (allowing for early termination), all the previously considered designs are also given the next 4 dipoles in the Halton sequence if their efficiency estimator is considered to have too much variance.

For CAS, we use the Gaussian process estimates of mean and variance, and for RS we use the standard Monte Carlo estimate. After each new dipole is computed, we refit the Gaussian process hyperparameters using maximum likelihood estimation. The covariance kernel for the Gaussian processes is the \( C^4 \) Matérn kernel, and we use a constant polynomial mean fit with generalized least squares.

**Experimental Results**

In our experiment, we gave both RS and CAS 500 dipoles worth of computations to consider and studied the resulting designs. Table 3 shows that CAS was able to produce 10 extra satisfactory designs despite only creating roughly the same number of unique designs. This is partly possible through more intelligently choosing the sampling locations, and also because the Gaussian process model used in CAS provides more accurate estimates of the fully accurate integrals in (3) (though the covariance incorporating data from other designs in the efficiency estimates).

<table>
<thead>
<tr>
<th>Strategy</th>
<th># Designs</th>
<th># Satis. Designs</th>
<th>Avg. # Dipoles per Unsatis.</th>
</tr>
</thead>
<tbody>
<tr>
<td>RS</td>
<td>67</td>
<td>11</td>
<td>6.875</td>
</tr>
<tr>
<td>CAS</td>
<td>68</td>
<td>21</td>
<td>5.071</td>
</tr>
</tbody>
</table>

Table 3: Constraint active search is able to produce many more satisfactory outcomes for consideration in eventual fabrication. It does this, in part, by being able to more quickly terminate unsatisfactory outcomes through use of the Gaussian process model.

Figure 7 provides two sets of plots: the parallel axes graphs which show that for nearly all satisfactory outcomes the Diameter Top should be quite small; and the scatter plots, which show how the other parameters define the satisfactory region. Of greatest note for eventual fabrication, satisfactory cones seems to be of the shorter, fatter variety, with Height consistently less than Diameter Bottom. The best results also seems to prefer some space between cones, with Box Size greater than Diameter Bottom as opposed to roughly equal to Diameter Bottom (as helped minimize reflection in (Haghanifar et al. 2020)). Cone Angle is an auxiliary shape description, provided here to show that there is a specific range of Cone Angle values which is required to see satis-
factory efficiency (within [0.7, 1.0] radians).

**Conclusion**

We have adapted Constraint Active Search to incorporate estimates of uncertainty generated by the iterative approximation of the emission in each dimension. In doing so, we were able to improve our computational efficiency while still identify a diverse set of satisfactory manufacturing designs.

In future work, we hope to conduct 3D simulations which lead to actual fabrications; doing so may require some multifidelity computations involving lower quality computational grids or more cheaply simulated approximations to the in-fidelity computations involving lower quality computational lead to actual fabrications; doing so may require some multi-fidelity approximations using Matlab rather than through the QMC strategy.

We also plan to compare the performance of CAS to Bayesian optimization to understand how the iterative approximation strategy performs in that setting. We should also rerun these experiments multiple times to produce some sense of statistical certainty regarding the results.

**References**


