

# Lumencor's Novel Light Engine Approach to Life Science Illumination

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Among the trends redefining the life sciences in the 21st century are (1) the rapidly expanding use of genomics for diagnostics and (2) the design of low-cost, portable analyzers. Life science instruments propel these trends using light as a powerful, analytical tool. Although high intensity, low cost light sources are essential to the design and proliferation of new bioanalytical instruments, until now this key subsystem has limited instrument performance and hindered substantial system cost reduction. As large instruments are reengineered into miniaturized analyzers and ultimately point-of-care devices, the availability of low cost, high performance, miniaturized Light Engines remains an elusive goal. Lumencor offers an inexpensive lighting solution that no competitive technology provides for today.

Life science instruments present unique challenges for light generation: inexpensive, high intensity, uniform, monochromatic light across a broad UV-Vis-NIR spectrum is required. Currently available commercial light sources have significant performance limitations including insufficient power, spectral drift and short lifetimes which may only be overcome with prohibitively large investments. Each of today's available light sources fall short of satisfying all these requirements. Their limitations compromise throughput and sensitivity, lead to frequent instrument calibration and generate high ownership cost. Consequently, instrument performance is constrained and often excessively priced with no obvious near term solution.

Lumencor is uniquely positioned to provide new Light Engines to satisfy this growing market. Lumencor Light Engines (patents pending) utilize solid state luminescent materials, excitation source and optimized light management techniques. A variety of light source technologies is employed to optimize the output of each color. Commercially available components are utilized to solve long standing technical problems and are contributing to the growth of an established instrument market. Initial single color products are configured to directly replace existing light subsystems and sell for less than \$3000. A working prototype has been tested and validated by a third party. Comparable prototypes are being shown to prospective customers. These are miniature, high power, cyan Light Engines designed as alternatives to the biotech workhorse, the Argon ion laser.

## Objective

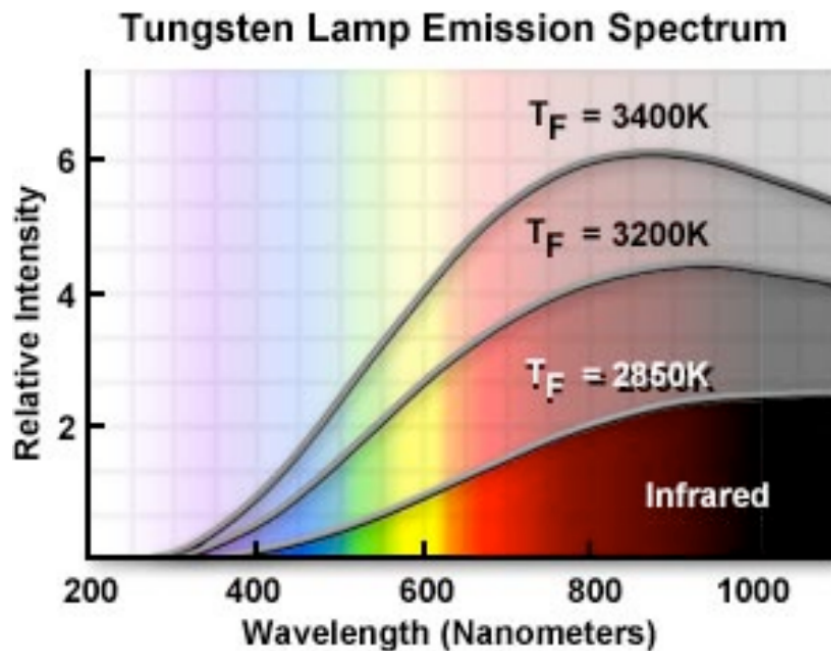
This white paper provides a technical overview of commercially available lighting technologies employed in the life sciences (their merits and shortcomings), a description of Lumencor's novel Light Engine approach, a product family map highlighting Lumencor's development plan and a summary of Lumencor's proprietary position.

## The Competitive Landscape

Today's commercial light sources include lamps, lasers and LEDs. Arc and tungsten lamps offer white light for a broad range of uses. Lasers are the brightest and most spectrally pure light sources. While LEDs offer by far the lowest cost light source option. A more detailed examination of the competitive technical and business landscape follows.

## Lamps

The primary light sources for PCR and Sanger sequencing instruments are tungsten halogen lamps, arc lamps and lasers. The tungsten halogen lamp is a mature product having been repeatedly optimized for display and lighting applications. Thus, improvements in its low light-collection efficiency or mitigation of high thermal emissions are unlikely. Durability and stability issues are also inherent and difficult to resolve. Its cost could come down but only if driven by volume applications. This white light source with a width of over 1000 nm must be heavily filtered to create the 10 nm band required in bioanalytical instruments. Figure 1 provides the typical emission curves as a function of drive current or operating temperature. As such, the light output to the sample volume is quite low, on the order of mWs, which relegates use of this lamp to low throughput instruments.



**Figure 2.** Typical tungsten lamp spectrum as function of drive current or operating temperature.

To overcome this limitation, instrument designs employ arc lamps. These high power lamps generate large light levels, however, they also generate very broad white light that must be heavily filtered. Additionally, they create significant heat loads which must be carefully managed to avoid compromising the bioanalytical chemistry and instrument performance. Power instability due to the changing arc over its lifetime is an notorious problem. The arc lamp has been significantly improved due to its proliferation in newer microdisplay based projection televisions and business projectors. However, the availability of low cost LEDs have diffused efforts to improve the arc lamp. Typically arc lamps, with their associated filters and spectral wheels, mirrors, controllers and shutters can require the addition of \$10,000 to the cost of a \$3,000 to \$5,000 bulb.

## Lasers

A common alternative to broadband lamps is the laser. One of the most frequently employed in bioanalyses is an air cooled Argon ion laser. A typical laser, for capillary based sequencing for example, provides 10 to 25 mW in a collimated, multi-line beam. However, the laser color-balance is inherently unstable; the power output of individual wavelengths varies over time and from product to product. Furthermore, lasers are limited in the number of wavelengths available. They are nonlinear devices and as such are built to be run in a continuous mode. They are not readily modulated except by external mechanics. The failure mechanism of lasers is a catastrophic event and as a consequence require replacement every 6 to 8 months. Replacement

costs average \$5,000 and, because they are typically replaced twice in the course of a year, result in high annual maintenance costs. Serious safety concerns exist about the use of these light sources by inexperienced users and in non-traditional testing environments.

As a consequence of the above shortcomings competition from improved light sources will likely come from diode lasers and LEDs. Significant improvements in these semiconductor products have already resulted in enhanced brightness, wavelength range and cost reductions. However, diode lasers are primarily targeted for telecommunications. They do not readily provide wavelengths in the visible and UV ranges of the spectrum, those required for bioanalyses. Techniques typically employed to generate these wavelengths, including frequency doubling, are unlikely to generate sufficient light for low cost Light Engines. Difficulties with beam shape, power stability and reliability are other issues yet to be resolved.

One exception receiving significant attention in newer instrument applications are the up-converted IR diode lasers, the so called DPSSs (diode pumped solid state lasers). Users will find them less expensive, more powerful and longer lived than their predecessors, however they retain certain limitations: namely, a limited number of emission lines, the need for external modulation and they remain prohibitively expensive for many applications.

## LEDs

LEDs are now available in a relatively wide range of wavelengths. However their output is significantly broad so as to require filtering. Additionally, output in the visible spectrum is profoundly reduced in the green. This gap in usable wavelength range compromises their utility for bioanalyses which require multicolor excitation throughout the visible. Because sufficient light intensity cannot be obtained from individual units, arrays of LEDs are employed, leading to uniformity, stability and durability issues. The spatially broad output leads to lower collection efficiencies and contrast ratios. Their main advantage, low unit cost, is often offset by the cost and complexity resulting from the needed additional components for their implementation.

Most importantly, these light technologies cannot be readily improved for bioanalytical applications. The associated Light Engine market simply does not justify the large investment necessary to overcome fundamental performance limitations. As a result, analytical instrument performance and price is constrained by the light source with no clear solution in sight. Moreover the numerous manufacturers of lamps and lasers provide only a source, not an integrated Light Engine. Such companies include ILC Technology, Lumileds, Spectra-Physics and Sylvania. Lumencor is in a unique position to provide a highly integrated, cost effective subsystem that bests our direct competitors.

## Lumencor Light Engines

Lumencor's Light Engine outperforms competing technologies in all figures of merit, including cost. Table 1 provides a general comparison of competing Light Engines. As an example, a Lumencor Light Engine can emit narrowband light exceeding 500 mW with intensities above 100 W/cm<sup>2</sup>. Bandwidths are as narrow as 10 nm. Lumencor Light Engines also have unique temporal and spatial properties that should enable new instrument designs and capabilities.

Light Engine	Useable Light	Uniformity	Temporal Response	Heating	Durability	Engine BOM
Arc Lamp	med	poor	none	high	low	high
Laser	high	poor	none	low	low	very high
LED	low	poor	fast	low	high	medium
Tungsten	low	poor	none	medium	low	medium
Lumencor	high	high	fast	low	high	low

**Table 1:** General comparison of various Light Engine used in bioanalyses.

A more detailed comparison among present day Light Engines employed in Sanger sequencers and QPCR tools with that of Lumencor's Light Engine is shown in Table 2. Such detailed comparisons are difficult. One must compare proprietary subsystems, not published specifications for the light sources. Based on discussions with customers, as a general rule Lumencor's Light Engine products demonstrate at least twice the performance of the current Light Engines at half the price.

Specification	Sanger Sequencing		Q-PCR	
Light Engine	Lumencor	Ar Ion Laser	Lumencor	Arc Lamp
Power	> 20 mW	< 10 mW	> 2 mW/cm <sup>2</sup>	< 1 mW/cm <sup>2</sup>
Wavelength	505 nm	488, 514 nm	4 visible colors	
Bandwidth	< 10 nm	~ 26 nm	< 10 nm	~ 15 nm
Stability	0.1%	> 1%	0.1%	> 1%
Duty Cycle	10 - 100%	50 - 100%	10 - 100%	100%
MTBF	> 10,000 hrs	< 4,000 hrs	> 10,000 hrs	< 1,000 hrs
Price Point	< \$3,000	> \$5,000	< \$5,000	> \$10,000

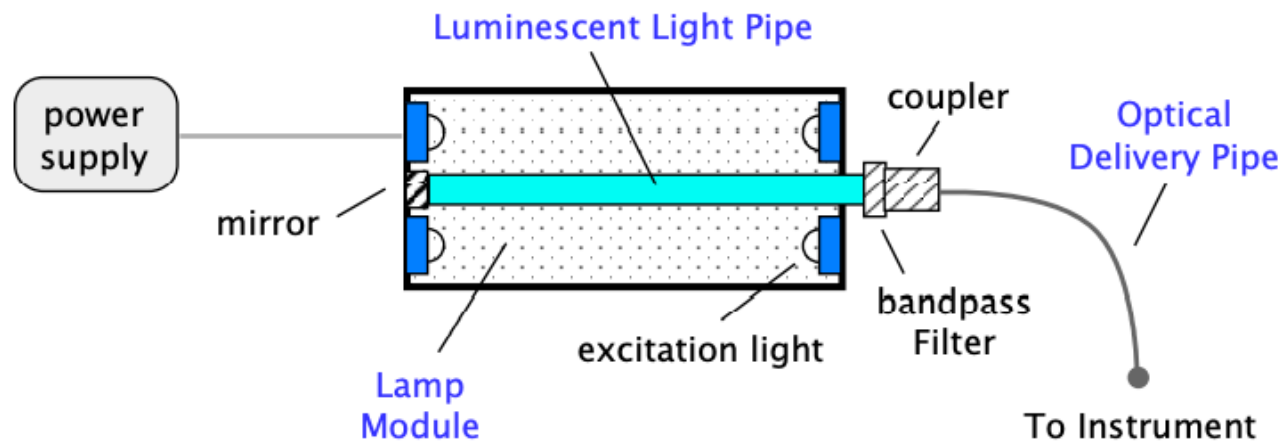
**Table 2:** Lumencor Light Engine output vs. the competition in two bioanalytical instruments.

## Solution

Lumencor achieves its cost and performance advantages utilizing a hybrid of illumination technologies including its own, proprietary light pipe technology (patents pending). Light Engines are based on solid state luminescent materials and optimized light management tools. Readily available components are employed to solve long standing technical problems. **As a result, Lumencor can deliver high performance, low cost solutions that meet all of biotech's requirements.**

## Light Engine Subsystem

Each Light Engine consists of a lamp and delivery optics. The lamp subsystem embodies multiple modules optimized to each produce a precise set of wavelengths. One such module is shown in Figure 2. Here Lumencor's light pipe technology is the source. Light is generated by a luminescent light pipe which is driven by an e-beam, LEDs or UV lamps. The pipe, which may consist of any of a variety of luminescent materials (e.g. rare earth doped glass, organic doped plastic, single crystal fiber) is tailored to emit light in the UV through the NIR. In the case of the rare earths, emission is inherently narrow due to their atomic-like energy structure. Any unwanted light is removed by a simple bandpass filter. The light output from the pipe is fed into an optical delivery pipe and into the instrument. The fast luminescence decay results in rapid switching with no appreciable warm up time.



**Figure 2:** Schematic of a typical Light Engine consisting of a lamp module and delivery optics.

The light pipe geometry integrates a significant fraction of the light leading to high external efficiencies. These efficiencies are optimized by the innovative design of the lamp module including the excitation source and the unique geometric shape of the pipe. Increased power levels, exceeding those typical for lasers in these applications, can readily be obtained by scaling the light pipe and associated excitation.

Each of the colors produced by Lumencor Light Engines requires a unique combination of light pipe and excitation source. Each instrument application requires one of several different approaches to coupling light from the delivery pipe into sample volumes. As an example, the light in a Sanger sequencer is coupled to a row of capillaries, each with a bore diameter of 50  $\mu\text{m}$ . A Q-PCR tool requires uniform excitation in an array of wells within a microtiter plate with no illumination of the interstitial spaces. To satisfy the range of instrument architectures, imaging and non-imaging optics are part of the Lumencor design portfolio.

## Product Family Map

Lumencor's product families are based on a common set of single color wavelengths and a set of light delivery systems. The colors and instrument geometries have been defined by commercial instrument users and instrument manufacturers. Six wavelengths have been identified based on the most common fluorescent dyes used today as well as the common sets of dyes most often employed in sets of two, three or four. These dyes and dye sets are used in microscopy, whole cell analysis, DNA sequencing, PCR, as well as across numerous other instrument applications throughout the life sciences.

PERFORMANCE CRITERIA	VIOLET	BLUE	CYAN	GREEN	YELLOW	RED
Central $\lambda$ , nm	395	445	480	550	575	650
$\lambda$ Range, nm	379 - 399	436 - 455	460 - 492	527 - 556	560-589	640 - 657
$\lambda$ Width, nm	20	19	32	29	29	17
LLG Power, mW	200	250	200	200	200	75
Stability, %	> 99.5	> 99.5	> 99.5	> 99.5	> 99.5	> 99.5
Lifetime, hrs	>5,000	>5,000	>5,000	>5,000	>5,000	>5,000
Modulation v, kHz	Up to 1	Up to 1	Up to 1	Up to 1	Up to 1	Up to 1
Warm Up, ms	1	1	1	1	1	1

**Table 3:** Product Specification Table.

Many light delivery systems share common geometries. As such three optical systems have been identified upon which to base Light Engine product families. They are direct illumination; light piping using an optical fiber or liquid light guide; and flood illumination. Direct illumination is an option for microscopes; piping is important for microscopy, sequencing, sensors, endoscopy; and flood illumination should be implemented for microarrays, microtiter plates, bioanalytical chips and any systems in which a larger areas must be irradiated.

Lumencor's first product family is a set of Light Engines for fluorescence microscopy. The important wavelengths upon which the Light Engine products are based are listed in Table 3. These colors are the basis for the development of all subsequent Lumencor product families. In microscopy applications, there is significant demand for single, dual, triple and four color Light Engines. A list of specification is outlined in table 3.

### Intellectual Property

Lumencor has core patent applications that cover Light Engines for use in the broad arena of life sciences. Specifically, the company has filed US and PCT applications covering all life science instruments based on Lumencor Light Engine technology. This strategy is key in that it secures Lumencor's position in the life sciences preventing competitors from building similar Light Engines for use in this sector. As well, Lumencor has several disclosures for proprietary materials that satisfy a subset of unique technical requirements; these clearly separate Lumencor from the extensive prior art on luminescent glass, plastic and/or crystalline fiber.