



Cytek® Northern Lights

Say Hello to a New Reality



Meet Northern Lights:



A new flow cytometry system that shifts the paradigm in what scientists expect to see in performance from an affordable three laser system.

Cytek Northern Lights incorporates the same groundbreaking technologies as its older sister, the Cytek® Aurora. Like Aurora, its optical design and unmixing algorithm give scientists remarkable flexibility, enabling the use of a wide array of new fluorochrome combinations without reconfiguring the system for each application. The state-of-the-art optics and low-noise electronics provide excellent sensitivity and resolution. Flat-top laser beam profiles, combined with a uniquely designed fluidics system, translate to outstanding performance at high sample flow rates.

The end result is a system that sits in a sweet spot for scientists that have budgeted for a one to three laser system, but desire the ability to run panels of higher complexity.

SpectroFlo® software offers an intuitive workflow from QC to data analysis with technology-enabling tools that simplify running applications.

The Cytek team has reimagined what you should expect from an affordable cytometer and has delivered an instrument that brings the benefits of full spectrum cytometry to more scientists.

High Value

Upgradeable from one laser and nine colors to three lasers and 24 colors, there is a Northern Lights configuration to fit your needs.

Remarkable Sensitivity

Sensitivity redefined using state-of-the-art optics and low-noise electronics.

Superb resolution of dim and rare populations, even in high complexity panels and high flow rates.

Easy, Flexible, and Intuitive

One configuration for all assays - no need to change optical filters.

Use any commercially available fluorochrome excited by the onboard lasers.

Intuitive software with familiar workflow.

Low Cost of Ownership

Fewer lasers to run more colors.

Low maintenance lasers, more fluorochrome choice, one configuration, and up to 24 colors per sample equates to **greater cost savings and less setup time between experiments.**



Application Flexibility For More Users

The revolutionary technologies on board Northern Lights enable capabilities usually seen in much pricier systems. With its onboard 100mW 405nm laser (available with three laser configuration only) and highly sensitive violet side scatter detector, particles nearing 100nm in size can be analyzed. Northern Lights opens the door to a wide variety of small particle applications. For those challenging applications involving highly autofluorescent particles, let the software's autofluorescence extraction tool bring new levels of resolution.

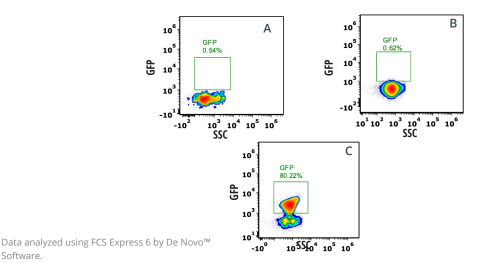
Small Particle Detection Example: ViroFlow

Murine Leukemia Virus (MLV-124 nm \pm 14 nm) genetically engineered to express superfolder GFP (sfGFP) as a fusion protein with the viral envelope glycoprotein.

The plots on the right show:

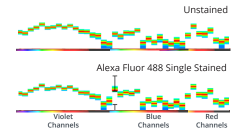
- A) Buffer only
- B) MLV with no sfGFP (MV-M-Zero)
- C) MLV with sfGFP-Env (MV-M-sfGFP)

All samples were run on a three laser Aurora using violet SSC as a threshold trigger. Virus reference particles were provided by ViroFlow Technologies (www.viroflowtechnologies.com).

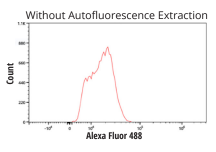


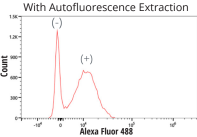
Autofluorescence Extraction Example: PrimeFlow™ RNA Assay

Human U937 cells were subjected to the PrimeFlow™ RNA Assay. The cells underwent a series of hybridization steps to label mRNA for HMBS, a low expressed gene (~10 copies/cell), with Alexa Fluor® 488. The sample was run on Northern Lights and analyzed using SpectroFlo® software with two different strategies, one with autofluorescence extraction and one without.



Spectrum plots of unstained and Alexa Fluor 488 stained cells acquired on Northern Lights. Note that the two spectra heavily overlap.





Due to high autofluorescence, separation of negative and positive signals was marginal (upper histogram). Autofluorescence extraction greatly improved the resolution of the two cell populations (lower histogram).

 ${\sf PrimeFlow^{\rm TM}} \ {\sf is} \ {\sf a} \ {\sf trademark} \ {\sf of} \ {\sf Thermo} \ {\sf Fisher} \ {\sf Scientific}.$



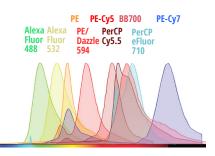
The Power of Full Spectrum Cytometry

Full spectrum cytometry opens a wide range of new possibilities. Northern Lights allows scientists to run complex multicolor experiments with as few as one or two lasers. The unique optical system enables the use of dyes with highly overlapping peak emissions without sacrificing resolution, translating to more flexibility in dye choice. Only one configuration is used for all applications, saving time in experimental setup, and minimizing the chance for experimental error. Experiments on the following pages are examples of what is possible with Northern Lights:

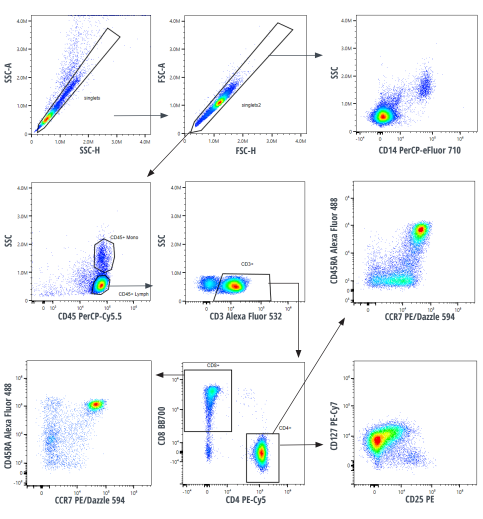
9-Color Blue Laser Panel

Peripheral blood mononuclear cells (PBMCs) were thawed, stained, washed, and analyzed on a one-laser Northern Lights. In this nine color blue laser excitable dyes panel, monocytes and several CD4 T cell and CD8 T cell subsets were easily identified. Markers and fluorochromes used in this assay are summarized in the table below.

SPECIFICITY	FLUOROCHROME		
CD45RA	Alexa Fluor® 488		
CD3	Alexa Fluor® 532 PE		
CD25			
CCR7	PE/Dazzle™ 594		
CD4	PE-Cy™5		
CD45	PerCP-Cy™5.5		
CD8	BD Horizon™ BB700		
CD14	PerCP-eFluor® 710		
CD127	PE-Cy™7		



9-Color Panel Dyes Emission Spectra





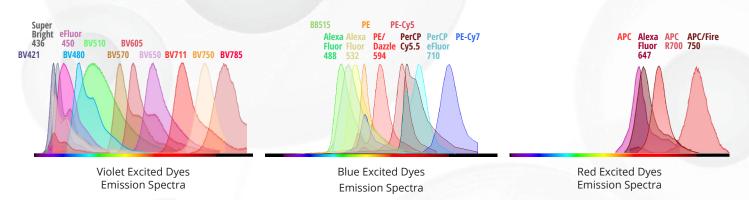
24 Colors With Three Lasers...Is it Possible?

The optical design combined with the unmixing capability in SpectroFlo® software allows greater fluorochrome choice, panel flexibility, and easy setup without having to change filters. The three laser configuration provides outstanding multi-parametric data for a wide array of applications. Markers and fluorochromes in a 24-color panel designed for identification of circulating cell subsets in human peripheral blood are summarized in the table below:

SPECIFICITY	FLUOROCHROME	SPECIFICITY	FLUOROCHROME	SPECIFICITY	FLUOROCHROME
CCR7	Brilliant Violet 421™	CD11c	BD Horizon™ BB515	CD27	APC
CD19	Super Bright 436	CD45RA	Alexa Fluor® 488	CD123	Alexa Fluor® 647
CD16	eFluor® 450	CD3	Alexa Fluor® 532	CD127	BD Horizon™ APC R700
TCR γ/δ	BD Horizon™ BV480	CD25	PE	HLA DR	APC/Fire™ 750
CD14	Brilliant Violet 510™	IgD	PE/Dazzle™ 594		
CD8	Brilliant Violet 570™	CD95	PE-Cy™5		
CD1c	Brilliant Violet 605™	CD11b	PerCP-Cy™5.5		
PD-1	Brilliant Violet 650™	CD38	PerCP-eFluor® 710		
CD56	Brilliant Violet 711™	CD57	PE-Cy™7		
CD4	Brilliant Violet 750™				
CD28	Brilliant Violet 785™			24-COL	OR DATA

On the next page, this 24-color panel is demonstrated in a healthy donor using a whole blood lyse wash sample preparation.

The 24-Color Panel Includes Many Highly Overlapping Dyes:

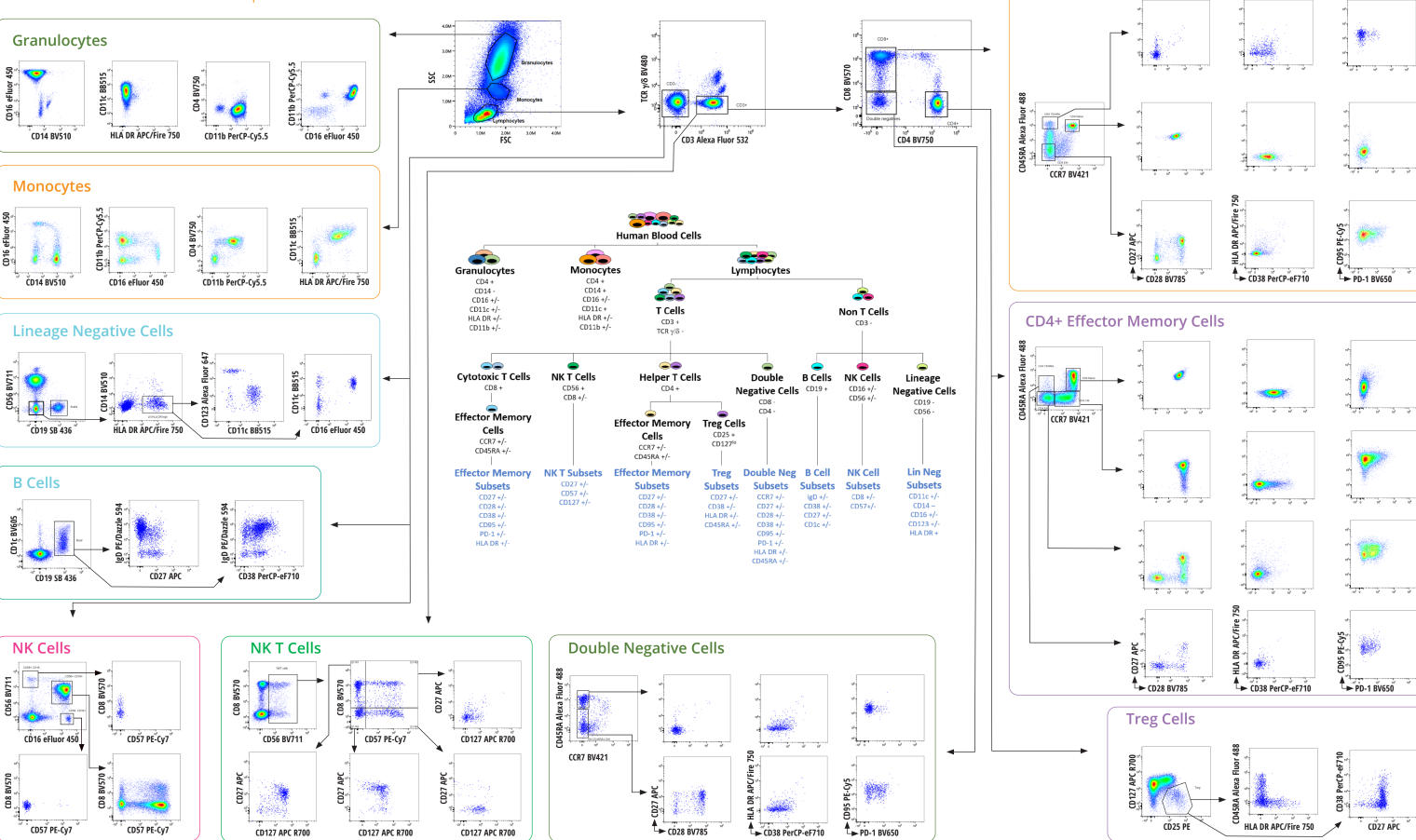


APC/Fire™ and PE/Dazzle™ are the trademarks and property of BioLegend,Inc.
Brilliant Violet™ is a trademark of Sirigen Group Ltd.
BD Horizon™ and Brilliant Blue (BB) are trademarks of BD Biosciences.
Alexa Fluor®, eFluor®, and Super Bright are trademarks of Thermo Fisher Scientific.
Cy® and CyDye® are registered trademarks of GE Healthcare

Allophycocyanin (APC) conjugates: US Patent No. 5,714,386 PE-Cy7: US Patent Number 4,542,104. APC-Cy7: US Patent Number 5,714,386. Trademarks are the property of their respective owners.

A New Reality:

3 Lasers, 24 Colors, Unparalleled Resolution



Northern Lights Makes It Possible

CD8+ Effector Memory Cells



Fluorescent Proteins and Challenging Dye Combinations

The detection of some fluorescent protein or fluorochrome combinations by conventional flow cytometry presents a challenge due to high amounts of spectral overlap (Figures 1, 4). Northern Lights addresses this challenge by using differences in full emission spectra signatures across all lasers to clearly resolve these combinations, even if the populations are co-expressed (Figures 2, 3, 5 and 6).

Example 1: GFP and YFP

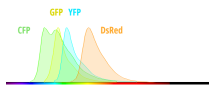


Figure 1: Spectrum plots from a conventional spectrum viewer shows heavy overlap between GFP and YFP.

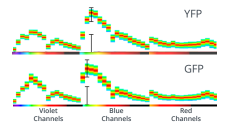


Figure 2: Spectrum plots from Northern Lights show distinct signatures across three lasers.

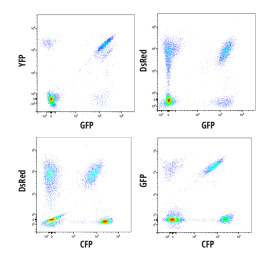


Figure 3: Sp2/0 cells were transfected with GFP, YFP, CFP and/or DsRed (alone or in combination) and run on Northern Lights (plots are gated on FSC vs SSC). Each population is clearly identified.

Example 2: Qdot 705 and BV711



Figure 4: Spectrum plots from a conventional spectrum viewer shows heavy overlap between Qdot 705 and BV711.

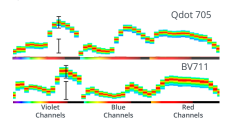


Figure 5: Spectrum plots from Northern Lights show distinct signatures for Qdot 705 and BV711.

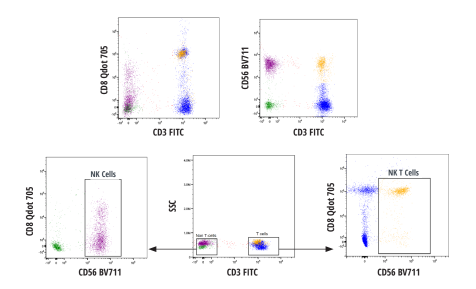


Figure 6: Normal human whole blood was stained, lysed, washed, and analyzed on Northern Lights. Subsets of NK and NK T cells that co-express CD56 BV711 and CD8 Qdot 705 were easily identified.



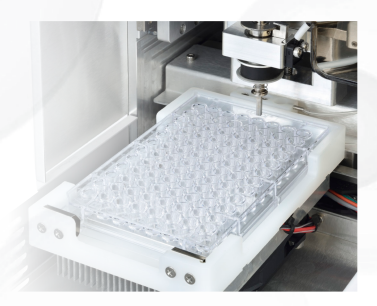
Get to Know Our Automatic Micro-Sampling System (AMS)



Meet the New AMS

The new AMS offers preset and user adjustable settings that allows the loader to be fine tuned to your experimental requirements. The AMS is specifically designed to streamline experimental workflow and seamlessly integrates into Northern Lights. The AMS also offers ease of use, low carry over, and minimal dead volume.

- Reliable and easy
 - Reliable 96 well plate acquisition maximizes productivity.
 - **Easily change** between plates and tubes in a matter of seconds.
- Three throughput modes
 - **Optimized** acquisition speeds, from low carry over to high throughput.
- () User customizable modes
 - **Fully customizable** with a suite of user modes to fit a variety of applications and workflows.



Say Hello to a New Reality

SpectroFlo® Software Guided Workflows 🧳





The new SpectroFlo software offers an intuitive workflow from QC to data analysis with technology-enabling tools that simplify running any application.

QC and Setup:

Run Daily QC to monitor instrument performance and add reference controls.

Library:

Add or remove experiment templates, worksheet templates, fluorochrome information, QC bead information, and more.



Extra Tools:

Unmix data using controls from different experiments or apply virtual filters to your data.

Users:

For administrative controls.

Preferences:

Customize the software appearance. Set default plot sizes, text sizes and fonts, gate colors, print layout, statistics table options, and more.

Acquisition:



Experiment Menu



Worksheet Menu



Fluidics Menu

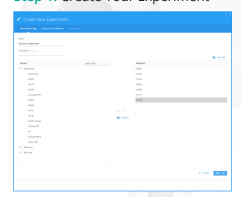


Plate Calibration

Experiment Workflow:

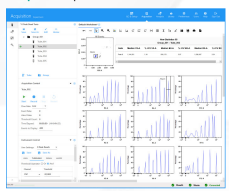
From the Acquisition menu, you can start a new experiment and get to your data in three simple guided steps.

Step 1: Create Your Experiment



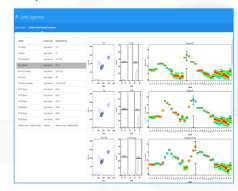
Create your experiment, choose fluorochromes, and add labels, tubes, worksheets, and stopping criteria in this guided workflow.

Step 2: Acquire Your Tubes



Load and acquire your samples.

Step 3: Unmix Your Data



Visualize your reference controls spectra using our unmixing wizard.



Specifications

Optics

EXCITATION OPTICS

OPTICAL PLATFORM

Northern Lights contains a fixed optical assembly configured with one to three spatially separated laser beams. Laser delays are automatically adjusted during instrument QC.

LASERS

One laser configuration: 488 nm: 50 mW Two laser configuration: 488 nm: 50 mW, 640 nm: 80 mW

Three laser configuration: 405 nm: 100 mW, 488 nm: 50 mW, 640 nm: 80 mW

BEAM GEOMETRY

Flat-Top laser beam profile with narrow vertical beam height optimized for small particle detection.

EMISSION OPTICS

EMISSION COLLECTION

Fused silica cuvette coupled to high NA lens for optimum collection efficiency to optical fibers.

FORWARD AND SIDE SCATTER DETECTION

FSC: high-performance semiconductor detector with 488 nm bandpass filter.

SSC: Two high-performance semiconductor detectors with 405 nm and 488 nm bandpass filter. Note: 405 nm side scatter applies to three laser configurations only.

FLUORESCENCE DETECTORS

Proprietary high sensitivity Coarse Wavelength Division Multiplexing (CWDM) semiconductor array per laser enabling more efficient spectrum capture in the 420-829 nm range. No filter changes required for any fluorochrome excited by the 405 nm, 488 nm, and 640 nm lasers.

STANDARD OPTICAL CONFIGURATION

Violet detector module (only available in three laser configurations): 16 channels uneven spaced bandwidth from 420-829 nm.

Blue detector module: 14 channels uneven spaced bandwidth from 498-829 nm.

Red detector module: 8 channels uneven spaced bandwidth from 652-829 nm.

Fluidics

SAMPLE FLOW RATES

Low: 15 μ L/min, Medium: 30 μ L/min, High: 60 μ L/min, Plate high-throughput mode: 100 μ L/min

FLUIDIC MODES

Long clean, SIT flush, Purge filter, Clean flow cell

MANUAL SAMPLE INPUT FORMATS

12x75mm polystyrene and polypropylene tubes

STANDARD FLUIDIC RESERVOIRS

4L fluid container set with level-sensing provided. Compatible with 20L sheath and waste cubitainers.

VOLUMETRIC SENSOR

Volumetric measurement during sample recording enables calculation of counts per µL for any gated population.

PLATE LOADER OPTION

96-well microtiter plate capability

Throughput time 35 minutes at High Throughput mode sampling 7 µL/well

Plate stage temperature: 4-30°C

PLATE LOADER CARRYOVER

Default mode: ≤0.3%, Low Carryover mode: ≤0.1%, High Throughput mode: ≤1%

Performance

FLUORESCENCE LINEARITY

FITC R² ≥0.995 / PE R² ≥0.995

FORWARD AND SIDE SCATTER

Performance is optimized for resolving lymphocytes, monocytes, and granulocytes.

SIDE SCATTER RESOLUTION

Capable of resolving 0.2µm beads from noise.

CARRYOVER

<0.1%

DATA ACQUISITION RATE

35,000 events/s*

*Three-laser system

Software

SPECTROFLO® SOFTWARE

Live unmixing during acquisition

Developed specifically to streamline assay setup, data acquisition, and file export

Automated QC module

Autofluorescence extraction

Raw and Unmixed FCS 3.1 files

Electronics

SIGNAL PROCESSING

Digital signal processing with automatic window gate adjustment.

22-bit 6.5 log decades.

Threshold using any single parameter or combination of parameters.

PULSE SHAPE PARAMETERS

Pulse Area and Height for every parameter. Width for scatter parameters and one fluorescence parameter for each laser.

Workstation

OPERATING SYSTEM

Windows® 10 Pro 64-bit

PROCESSOR

Intel® Core™ i7 processor

RAM

16GB

HARD DRIVE

500GB SSD and 1TB SATA

VIDEO PROCESSOR

NVIDIA® HD GeForce

MONITOR

32" UHD 4K Monitor

Installation Requirements

Dimensions (W x D X H)

INSTRUMENT DIMENSIONS

Without loader: 54 x 52 x 52 cm With loader: 58 x 62 x 52 cm

INSTRUMENT WEIGHT

Instrument weight: 61 kg Loader weight: 13 kg

RECOMMENDED WORKSPACE

165 x 76 x 132 cm

Room Requirements

POWER

100-140 VAC, 15A or 200-250 VAC, 10A

HEAT DISSIPATION

500W with all solid-state lasers

TEMPERATURE

15-28°C

HUMIDITY

20%-85% relative non-condensing

AIR FILTERING

No excessive dust or smoke

LIGHTING

No special requirements

Regulatory Status

For Research Use Only. Not for use in diagnostic or therapeutic procedures.





Cytek Biosciences is dedicated to enhancing our customers' user experience. The Northern Lights system is backed by our world-class service and support team that can provide phone-or field based assistance. Various levels of maintenance options are available to meet your needs now, and in the future.

For more information, email us at: sales@cytekbio.com or call 1-877-922-9835

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