Seventeen-colour flow cytometry: unravelling the immune system

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The increasing need for polychromatic approaches to flow cytometry, coupled with rapid technological advances, has pushed the frontiers of flow cytometry beyond 12-colour systems. Recent breakthroughs have allowed the design and implementation of instruments that measure 19 parameters (17 fluorescent colours and 2 physical parameters). This article describes the instrumentation and considers the reagents, analysis and applications for this powerful, new extension of flow-cytometric technology.

Fluorescence-activated flow cytometry is one of the most powerful technologies that is routinely used in immunology. It allows not only for the rapid multiparametric analysis of cells on a cell-by-cell basis, but also for the viable separation (sorting) of highly purified populations of cells. Flow cytometry was born in the Herzenberg laboratory at Stanford University (Stanford, United States) in the late 1960s1–4. Early cytometers measured three parameters (one fluorescence signal and two scattered-light signals); the power of multiparametric analysis to discriminate functionally distinct subsets of leukocytes drove the technology to the commonly used four-colour machines that can distinguish six parameters for each cell. This four-colour analysis was first performed in the mid-1980s. In the 1990s, further technological advances were made necessary by the growing appreciation of researchers for the complexity of the immune system; a systematic series of developments of hardware, software and chemistry were spearheaded by the Herzenberg laboratory to create machines capable of measuring more than 19 parameters (4–5 fluorescent, 10–15 physical)5.

The ability to discriminate between these cell types is crucial to our understanding of cellular immunity and disease pathogenesis. This can be illustrated by recent studies of HIV, which show considerable changes to specific T-cell subpopulations beyond ‘bulk’ CD4+ T cells. For example, disease caused by HIV has been associated with a depletion of naïve6–7 and newly produced8 T cells, viral infection of memory9 and HIV-specific10 CD4+ T cells, and clonal expansion of activated11 and CD8+ (REF. 12) T-cell subsets. These are specialized subsets of T cells that carry out distinct programmed functions; the selective change in their representation reflects a marked alteration in the composition of the immune system, even when overall numbers of T cells might not be significantly altered. For each of these studies, three- or four-colour flow cytometry techniques were sufficient. However, future studies will require the simultaneous measurement of more phenotypic and functional parameters. Technology that can measure six or more colours — polychromatic flow cytometry — is well suited to such endeavours; however, investigators need to determine how many colours are required to answer immunological questions.

For simple discrimination between peripheral blood cells, as few as two physical parameters (forward and side scatter) and two fluorescence parameters are sufficient. Forward- and side-scatter characteristics can be used to distinguish monocytes and granulocytes (high side scatter and high forward scatter) from platelets and erythrocytes (low side scatter and low forward scatter). Lymphocytes and circulating dendritic cells show intermediate forward and side scatter and can be further segregated by lineage (that is, B cells, T cells or natural killer (NK) cells) using one or more fluorescence parameters. For example, T cells can be identified using fluorescently labelled antibodies specific for the CD3 cell-surface marker and either the CD4 or CD8 cell-surface markers (two-colour flow cytometry).

Within T-cell populations, various markers have been proposed to distinguish naïve, effector- and memory-cell populations. These markers are often paired (for example, CD45RA and CD62 ligand (CD62L), co-expression of which identifies naïve T cells) to create the simple phenotyping schemes commonly used in four-colour experiments. However, attempts to describe the complex T-cell compartment using these phenotyping schemes have raised many questions. For example, most of the common markers that are used to identify the stages of T-cell development.
The use of additional colours also has distinct advantages for improving the accuracy of measurements of rare cell populations. Specifically, irrelevant cell populations can be excluded from analysis using a 'dump channel', which contains all of the antibodies specific for unwanted cells in one fluorescence channel, such as the Cy5PE (cyanine-5–phycoerythrin) channel. Cy5PE is a useful channel for 'dump' antibodies, because dyes that selectively label dead cells, such as propidium iodide and 7AAD, fluoresce in this channel; therefore, any fluorescence in this channel can be used to eliminate dead or unwanted cells. It should be noted that the differential compensation required by the viability dyes and Cy5PE is not important, because all cells that fluoresce in this channel will be eliminated from further analysis. Studies of T-cell populations could include a dump channel for CD19+ cells (B cells), CD56+ cells (NK cells) and CD14+ cells (monocytes), with all of the antibodies conjugated to just one fluorochrome. For better detection and discrimination of low-frequency populations, markers shared between the cells of interest and irrelevant cells could be assigned to additional fluorochromes. So, using more colours in an experiment allows more-accurate and more-sensitive measurements of low-frequency cell populations. This benefit is particularly important in vaccine development, in which the cells that provide immune protection can comprise a small proportion of the overall T-cell compartment.

Over time, the utility of polychromatic flow cytometry is also likely to become apparent in other settings, such as oncology and haematology. Because the requirements for developing and implementing this technology have now been defined, many of the main hurdles to its widespread adoption have been cleared. This article describes the sophisticated instrumentation, computer algorithms, and advances in reagent technology that now allow use of a 17-colour system.

Developing 17-colour cytometry

Ten-parameter instrumentation, which measures eight fluorescent colours and two physical parameters, was first described in 1997. Subsequently, the development of a 13-parameter (11-colour) flow cytometer was reported in 2001 (REVS.14–16). Although these instruments provided considerable advances over previous generations of cytometers, they needed high-power lasers, were extensively customized and required highly skilled operators. Recently, however, numerous technological and computing advances have produced a new generation of technology that incorporates more than ten fluorescent colours.
of instruments that are cheaper, require less technical skill to operate and are adaptable for applications that require as many as 19 parameters.

An example of this technology is the LSR II flow cytometer (Becton, Dickinson and Company Biosciences, San Jose, United States). As illustrated in FIG. 1a, our 19-parameter system, the LSR II, uses three solid-state lasers, which emit in the green (532 nm), blue (488 nm) and violet (407 nm) ranges. In addition, a fourth (helium–neon) laser emits in the red (633 nm) range. These four lasers provide the excitation for the fluorochrome-conjugated antibodies. After excitation, each fluorochrome emits photons of light with a unique spectrum; this emission is collected in an octagon or trigon detection system specific to each laser. The octagon detection system for the green-laser-excited fluorochromes (that is, phycoerythrin (PE), Texas Red–PE (TRPE), Cy5PE, Cy5.5PE and Cy7PE) is shown in FIG. 1b,c. At each corner of the assembly, a photomultiplier tube detects light in a particular wavelength range, as controlled by the filters in the centre of the assembly; in this way, each detector measures mainly (but not solely) the emission from a particular fluorochrome. This process occurs in two steps. First, the emitted photons encounter one or more dichroic mirrors, which either reflect or transmit photons, depending on the wavelength characteristic of the filter. In this example (FIG. 1c, red line), a 685 long-pass (685LP) dichroic filter reflects wavelengths shorter than 685 nm and transmits photons with wavelengths longer than 685 nm. Then, the transmitted light is further ‘purified’ by a band-pass filter before it reaches the detector (using, for example, a 710/50 band-pass filter, which eliminates photons outside the range 685–735 nm). So, the signature excitation and emission curve of each fluorochrome produces photons that are selectively measured by a series of dichroic mirrors and band-pass filters. The main objective of this system is to have the narrowest filter range possible, while still capturing the maximum amount of light. In the 17-colour detection system that we built, the green laser is configured with five detectors (for the fluorochromes PE, TRPE, Cy5PE, Cy5.5PE and Cy7PE). If additional fluorochromes become available that have distinct emission spectra after excitation by the green laser, the detection capacity could be increased even further. The blue laser is configured with a single fluorescence detector (for fluorescein isothiocyanate, FITC); the red laser is configured with three detectors (for allophycocyanin (APC), Cy7APC and Alexa Fluor 680); and the violet laser is configured with eight detectors (for quantum dots (also known as Q dots or QDs) -545, QD565, QD585, QD605, QD655, QD705, AmCyan and Cascade Blue). As more fluorescent dyes become available, the detection capacity can be increased through the incorporation of additional lasers or more-complex detection assemblies; there is no theoretical limit on the number of fluorochromes that can be simultaneously measured.

Developing complex reagent panels
The development of reagent panels for immunophenotyping begins with the selection of surface or intracellular proteins for study. Ideal markers for immunophenotyping are proteins that are likely to be relevant in the life cycle of a cell type, such as the lymph node homing molecules CD62L and CCR7 for naive T cells17–20. In addition, the ideal marker (or combination of markers) would be differentially expressed across the cell subsets to be investigated. Finally, the best markers for phenotyping cell subsets are unaltered in three situations: first, in vitro stimulation, which is required for many functional assays; second, viral infection, which might alter...
expression of cell proteins as an immune-evasion mechanism; and third, cell cryopreservation, which can, for example, strip CD62L from the membrane of frozen cells. The latter is a particularly important consideration for immunogenicity analysis in large-scale vaccine trials, in which it is untenable to carry out all assays on freshly isolated cells.

After antibodies specific for the markers of interest have been selected, the most suitable fluorochromes for detection need to be chosen. The availability of many fluorescent dyes, with distinct excitation or emission spectra, allows for several different fluorochrome–antibody pairings, which can be combined to produce a large number of possible staining panels. For example, in an 11-colour flow-cytometry experiment, there are nearly 40 million different staining panels that can be assembled by assigning antibodies to different fluorochromes, and for an 11-colour experiment with 17 available channels, there are more than $5 \times 10^{13}$ possibilities. Although the best combination of antibodies and fluorochromes for a particular application can as yet only be ascertained empirically, there are considerations that can guide users in their design of staining panels.

The assignment of antibodies to fluorochromes is determined mainly by the background staining of, and spectral overlap between, the chosen fluorescent dyes. Some background staining is generated by the natural autofluorescence of organic molecules in cells (such as NADH); the emission spectrum of autofluorescence overlaps that of dyes such as Cascade Blue, FITC and PE. High background staining can also result from the non-specific binding of a fluorescent conjugate to cellular elements, which occurs when the antibody concentration is too high. In either case, the problems caused by high background staining can be avoided by the titration of antibody reagents before their use. From titration data, the antibody concentration that provides the maximal brightness of the positive population with the least amount of background staining should be chosen for use in the panel$^{21}$.

The spectral overlap between fluorescent dyes also needs to be considered in the assignment of antibodies to fluorochrome channels. This overlap needs to be mathematically eliminated using a process known as compensation. Compensation is the mathematical deconvolution of the outputs from multiple detectors to yield measurement values proportional to each of the fluorescent dyes used. The need for compensation arises from the spillover of light between detectors and the relatively wide excitation energies of most fluorochromes. As illustrated in Figure 2, signals emitted by one fluorochrome (y-axis) can contaminate neighbouring detectors (x-axis), which results in 16 potential sources of contaminating light. For example, the TRPE fluorochrome shows a large percentage spillover into the PE, Cy5PE and Cy5.5PE detectors. By contrast, light from FITC and Cascade Blue have little spillover into other detectors.

However, even with computer-assisted compensation, inescapable measurement errors lead to a broadening of the distribution of the fluorescence of cells that have low or no fluorescence, particularly in the far-red channels (APC, Alexa Fluor 680 and Cy7APC)$^{14,22}$. This phenomenon can adversely affect the resolution between dimly staining cell populations and negative cell populations (that is, populations for which there is no detectable cell-surface expression of the marker). So, markers that stain brightly and provide good separation between positive and negative cell populations should be chosen for the far-red channels (for example, antibodies specific for CD3, CD4 and CD8). Bright fluorochromes, such as PE or Cy5PE, should be used for markers with unknown staining patterns or for markers that require clear discrimination between dimly staining and negative cell populations (for example, CD28 and CD127).

In this sample experiment, numerous dead cells are apparent in the light-scatter gate (a); however, not all of these can be effectively removed solely by gating on forward scatter (FSC) and side scatter (SSC). These cells can be effectively removed from the analysis by adding ethidium monoazide (EMA) to the cell sample and then gating on the EMA+ cells (live-cell gate) (b). Further analysis uses the light-scatter gate after this ‘EMA gating’ (c). Data are shown for cells stained with CD4- and CD8-specific antibodies before (d) and after (e) EMA gating. A double-positive population (arrow) can be observed as a result of the dead cells present in the light-scatter gate. Until now, the identification of dead cells has not been crucial in many experiments. However, because polychromatic flow cytometry can identify hundreds of cell subsets, many of these will be present at frequencies below the fraction of dead cells (typically a few percent of all cells). Therefore, it is crucial to eliminate even this small percentage of dead cells from the analysis, to ensure that they do not artefactually comprise most of the subset of interest.

**Figure 3 | Dead cells can be an important artefact.** In this sample experiment, numerous dead cells are apparent in the light-scatter gate (a); however, not all of these can be effectively removed solely by gating on forward scatter (FSC) and side scatter (SSC). These cells can be effectively removed from the analysis by adding ethidium monoazide (EMA) to the cell sample and then gating on the EMA+ cells (live-cell gate) (b). Further analysis uses the light-scatter gate after this ‘EMA gating’ (c). Data are shown for cells stained with CD4- and CD8-specific antibodies before (d) and after (e) EMA gating. A double-positive population (arrow) can be observed as a result of the dead cells present in the light-scatter gate. Until now, the identification of dead cells has not been crucial in many experiments. However, because polychromatic flow cytometry can identify hundreds of cell subsets, many of these will be present at frequencies below the fraction of dead cells (typically a few percent of all cells). Therefore, it is crucial to eliminate even this small percentage of dead cells from the analysis, to ensure that they do not artefactually comprise most of the subset of interest.
antibodies or streptavidin. By varying the amount of cadmium and selenium, QDs can emit light in a broad range (525–705 nm); therefore, after conjugation to monoclonal antibodies, these reagents increase the colour capabilities of the 407 nm violet-laser detection system from two to eight parameters23. Another recent addition to the violet-laser repertoire is the organic fluorochrome AmCyan. This reagent is brighter than older reagents that are detected in the violet laser channel, such as Alexa Fluor 430. Both QD and AmCyan are attractive choices for low-power lasers, such as the violet solid-state laser.

Although these new reagents make it possible to envisage 17-colour panels, new studies need to begin with preliminary experiments to assess the staining patterns of smaller panels that use four or five well-defined markers. The staining patterns observed in these smaller panels then need to be compared with those previously reported or previously observed in simple two- or three-colour experiments. If the results are not comparable, then new combinations of fluorochromes need to be tested. After comparable results are obtained, the investigator could add markers of interest to the panel, but care should be taken to ensure that the staining patterns of the original markers are not changed by the addition of new markers.

Two crucial considerations in the development of new staining panels are the use of viability markers to exclude dead cells and the incorporation of controls to help identify cell subsets based on their fluorescence measurements. Viability markers are often required in experiments in which the cells of interest are rare (less than 1% of the total population) or have been activated or thawed. In these types of sample, cell death can result in large errors because of the propensity of dead cells to bind nonspecifically to monoclonal antibody conjugates. For example, FIG. 3 shows how the viability marker ethidium monoazide (EMA) can eliminate a substantial number of dead cells from the light-scatter dot plot (FIG. 3a), allowing more-precise identification of lymphocytes (FIG. 3c). In FIG. 3d, the way in which staining artefacts, such as the double-positive (CD4+CD8+) staining of peripheral blood cells (indicated by the arrow), can be attributed to dead cells in the lymphocyte gate is shown. It should be noted that these double-positive cells are not present in the sample after the EMA+ cells have been gated out (FIG. 3e).

The second important aspect to panel design is the inclusion of a ‘fluorescence-minus-one’ (FMO) control. This is a sample that has been stained with all of the reagents except one; the analysis of this sample allows for the precise definition of cells that have fluorescence above background levels (FIG. 4). In this example, cells stained with anti-CD3–FITC (that is, FITC-conjugated CD3-specific monoclonal antibodies), anti-CD4–PE, anti-CD8–Cy5PE and anti-CD45RO–Cy7PE are compared with cells stained with all monoclonal-antibody conjugates except the CD4-specific antibodies (FMO control) (FIG. 4a). It should be noted that by using the unstained sample to determine the cut-off between positive and negative cell populations (‘unstained gate’ line), the population gate would be too low to accurately distinguish CD4+ and CD4– T cells (FIG. 4b). However, the cut-off determined by the FMO control (‘FMO gate’ line) can correctly separate these populations, thereby giving a more accurate representation of the percentage of the population that comprises CD4+ cells21.

Analysis tools for 19-parameter data

Data analysis is by far the most time-consuming aspect of polychromatic flow-cytometry experiments; therefore, there is considerable demand for tools that can organize analyses into databases, as well as assist in the exploration of complex data sets.

Recently, automated data-analysis techniques were developed to identify subpopulations of cells in complex multivariate data. These automated techniques are intended to complement the commonly used technique of dividing data by manually segregating events into regions two parameters at a time (known as bivariate gating). The automated approach is similar to cluster algorithms, which group individual datum points together based on similarity or disparity; the computation of similarity is the core of such algorithms and varies markedly between different approaches. The algorithms are based on published techniques that quantify the difference between multivariate distributions; these are known as probability-binning comparison algorithms24-28. Automated subset-identification algorithms might also be useful in settings in which cells respond in complex, multivariate patterns or for the rapid identification of immunophenotyping patterns that correlate with pathogenic states27.

Examples of cluster- and bivariate-gating analyses are shown in FIG. 5. The antibodies that were used for staining are indicated in the figure. It should be noted that cluster analysis identifies many different cell populations and assigns a different colour to each figure (FIG. 5a). The expression level of each marker can then be visualized for the various clusters by using dot plots and histograms. So, the orange cluster mainly contains CD4+ T cells (arrow 1), whereas the green and yellow clusters are mainly CD8+ T cells (arrow 2). The main advantage of this analysis is the ability to rapidly (and automatically) explore populations without the constraints of defined gates. For example, cluster analysis can quickly identify a population of CD4+CD45RO+ T cells that express both interleukin-2 (IL-2) (arrow 3) and interferon-γ (IFN-γ) (arrow 4). Similarly,
Peripheral-blood mononuclear cells were stimulated with staphylococcal enterotoxin B (for 6 hours in the presence of brefeldin A) then permeabilized and stained with antibodies specific for various cell-surface markers and intracellular cytokines. Data are gated on CD3+ T cells. a | Cluster analysis identifies multiple cell populations, each of which is assigned a particular colour. In this example, orange clusters consist of CD4+ T cells (arrow 1), whereas green and yellow clusters consist of CD8+ T cells (arrow 2). Cluster analysis can quickly identify CD4+ T cells that express CD45RO and interleukin-2 (IL-2) (arrow 3), together with interferon-γ (IFN-γ) (arrow 4). Similarly, heterogeneous expression of CD45RO (arrow 5), yellow population) is quickly recognized in CD8+ T cells that lack IL-2 expression (arrow 6). So, automatic identification and visualization of clusters can be a powerful tool. However, because overlaid clusters can be difficult to distinguish, the level of IFN-γ expression in the CD8+IL-2– cell population is unclear (arrow 7). When bivariate-gating analysis is used (Fig. 5b), clearer more-quantitative results are obtained, albeit at the expense of considerable analyst time. For instance, bivariate analysis shows the presence of CD8+ IFN-γ+ cells that do not express IL-2. Moreover, the relationship between cytokine production by CD8+CD45RO– cells (arrow 8) and CD8+CD45RO+ cells (arrow 9) can be directly assessed. To summarize, cluster analysis automatically provides a visual display that enables rapid determination of the level of expression of multiple markers across multiple dimensions; however, it may not provide the clear, structured and quantitative analysis that bivariate gating does. At present, cluster analysis is recommended as a tool for rapid, qualitative (not quantitative) data exploration. Certainly, more tools that aid data exploration are needed.

As indicated earlier, the multidimensional nature of data from polychromatic experiments can make cluster analyses unwieldy. Therefore, there is a need for other visualization tools that simultaneously allow data exploration, comparison and quantification with maximum flexibility. In this context, flexibility indicates the ability to rapidly rearrange the way in which data are presented, by choosing a new parent phenotype to anchor the analysis or by hiding certain markers. Representing data in a form similar to a hierarchical tree, such as the one presented in Fig. 6, might offer some of these advantages. In this type of graphical representation, each marker that is measured (such as CD11a) is assigned to a particular level of the tree, and different branches represent various expression levels of the marker (for example, positive versus negative) in the immediate parent population. Phenotypes that do not occur in a cell population are not displayed on the tree, allowing rapid comparison of samples according to the ‘shape’ of the tree. To further explore the data, the order of the markers can be rearranged until branches with similar functional characteristics are brought together, thereby identifying phenotypes that are similar in function. For example, high IFN-γ production is shown to be confined to CD11abright cells in Fig. 6. Similarly, the overlap between the markers that are examined could be determined by comparing the full phenotypic tree to a tree in which one or more markers are hidden. Finally, as indicated in Fig. 6, quantitative elements could be added to tree graphics. In the simplest case, a frequency histogram underneath the tree can show the proportion of cells that belong to a particular branch. Alternatively, the histogram could indicate the frequency of a subset that shows a certain function. Such tree-based graphics offer many options for data exploration, comparison and quantification.

Concluding remarks
As immunologists further dissect the heterogeneity of the cell populations that coordinate defences against various pathogens, polychromatic flow cytometry will be an invaluable tool. Until recently, it has been limited by instrument complexity, reagent availability and a lack of powerful analysis software; however, all of these areas are now progressing, thereby enabling this technology to advance rapidly.

Nonetheless, there remain considerable hurdles before this application becomes widely used. First, we need flow cytometers...
Figure 6 | *Visualization of polychromatic data using hierarchical trees.* The data shown in this hierarchical tree were obtained from peripheral-blood mononuclear cells stimulated with staphylococcal enterotoxin B (for 6 hours in the presence of brefeldin A). The cells were subsequently permeabilized and stained for various cell-surface markers and intracellular cytokines. Branches of the tree segregate and stained for various cell-surface markers and intracellular cytokines. Branches of the tree segregate based on the expression level (for example, positive versus negative or bright versus dim) of a particular marker, and each terminal branch represents one fully gated subset. So, reading from the top, the branch below arrow 1 represents cells that are CD3+CD4+CD27–CD45RA–CCR7+ cell population indicated by arrow 1 expresses only IL-2, which is constant with a naive phenotype. Interestingly, IL-2 is expressed by a greater fraction of a naive-like subset (CD11a+CD27–CD45RA–CCR7+ cells), in which cells have lost expression of CD27 that each configuration need not be separately optimized. Second, more-sophisticated systems are needed for software-controlled data acquisition, to help guide the user through the still relatively complex set-up and optimization procedure. Third, manufacturers need to devise a new model for reagent distribution so that laboratories can test numerous conjugated monoclonal antibodies at minimal cost before committing to purchase larger amounts of the final optimized panels. This will require manufacturers to have a large inventory, because the number of reagent–fluorochrome combinations will be an order of magnitude greater than what is required at present. Last, we desperately need new tools to aid in the exploration of these complex data sets, as well as new techniques for the presentation of data. Journals will need to accommodate the requirement for the presentation of far more complex graphics and analyses, and the inclusion of such information as online supporting documentation will become ubiquitous.

Undoubtedly, polychromatic flow-cytometry technology such as we describe here will become more common in immunology and clinical-research laboratories worldwide. In our laboratory, we use this technology to identify correlates of immune protection, therapeutic efficacy and disease morbidity for disease caused by HIV and for vaccination against infectious diseases. Although the identification of such correlates might not require more than six–to eight-colour flow cytometry, the process will be markedly accelerated by the ability to make many more measurements simultaneously.

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