

SPECIALIST IN CYTOMETRY, SCYM(ASCP) INTERNATIONAL SPECIALIST IN CYTOMETRY, SCYM(ASCPⁱ) EXAMINATION CONTENT GUIDELINE

EXAMINATION MODEL

The SCYM(ASCP) and SCYM(ASCPⁱ) certification examination is composed of 100 questions given in a 2 hour 30 minute time frame. All exam questions are multiple-choice with one best answer.

Examination questions may be both theoretical and/or procedural. Theoretical questions measure skills necessary to apply knowledge, calculate results, and correlate patient results to disease states. Procedural questions measure skills necessary to perform laboratory techniques and follow quality assurance protocols. Additionally, regulatory questions are based on U.S. sources (e.g., AABB, FDA, CLIA, etc.).

EXAMINATION CONTENT AREAS

The SCYM exam questions encompass the following content areas within Cytometry: Instrumentation, Panel/Experiment Design, Applications, Data, and Laboratory Operations. Each of these content areas comprises a specific percentage of the overall 100-question exam. The content areas and percentages are described below:

CONTENT AREAS	DESCRIPTION	EXAM PERCENTAGES
INSTRUMENTATION	Principles of fluidic, optical, and electronic instrumentation including troubleshooting	15 – 20%
PANEL/EXPERIMENT DESIGN	Sample source, sample integrity, sample preparation and staining, cell enrichment, and assay development (including target, sample state for functional studies, probe types, fluorochrome selection, spectral overlap and compensation, assay controls, and assay optimization)	25 – 30%
APPLICATIONS	Immunophenotyping, functional assays, multiplex bead assays, solid organ transplant, stem cell analysis, cell cycle / DNA ploidy, rare event analysis, small particle analysis, fetal hemoglobin assay, cell sorting, imaging cytometry, and mass cytometry	25 – 30%
DATA	Data standards, signal processing, data display, gating, statistical methods, common data modeling techniques, quantitative cytometry, and troubleshooting	15 – 20%
LABORATORY OPERATIONS	Quality control, assay validation, safety, and laboratory administration	10 – 15%

For a more specific overview of the SCYM exam, please refer to the [CONTENT OUTLINE](#) starting on page 2.

SPECIALIST IN CYTOMETRY, SCYM(ASCP) INTERNATIONAL SPECIALIST IN CYTOMETRY, SCYM(ASCPⁱ) EXAMINATION CONTENT OUTLINE

I. INSTRUMENTATION (15 – 20%)

- A. Fluidics
 - 1. Hydrodynamic focusing and properties of sheath fluids
 - 2. Sample delivery (e.g., syringe pump, pressure based, vacuum, acoustic)
- B. Optics
 - 1. Optical filters (e.g., long pass, band pass, short pass, dichroics, neutral density, polarizing)
 - 2. Light source (e.g., laser type, laser line, arc lamp, led)
 - 3. Lenses (e.g., beam shape, collecting, focusing, objective)
 - 4. Optical pathway (e.g., transmission, reflection, interrogation point, collinear, spatial separation, light scatter)
- C. Electronics
 - 1. Amplifiers (e.g., linear, logarithmic)
 - 2. Detectors (e.g., photomultiplier tube, photodiode, CCD camera, avalanche photon detector)
 - 3. Digital vs. analog systems
 - 4. Noise
 - 5. Pulse measurement (e.g., time delay, window extension, area, width, Coulter impedance)
 - 6. Threshold/discriminator
- D. Troubleshooting

II. PANEL/EXPERIMENT DESIGN (25 – 30%)

- A. Sample
 - 1. Sample source (e.g., beads, blood, bone marrow, solid tissue, body fluids, subcellular components, cultured cells, microorganisms, plants, whole organisms)
 - 2. Sample integrity (e.g., collection, handling, storage viability)
 - 3. Sample preparation and staining (e.g., disaggregation, lysing agents, aggregates, filtering, fixation, permeabilization)
 - 4. Cell enrichment (e.g., cell sorting, density gradient isolation, magnetic beads)

B. Assay Development

- 1. Target (e.g., cell type, subcellular location, molecule)
- 2. Sample state for functional studies (e.g., activated, resting, proliferating)
- 3. Probe types (e.g., antibodies, viability/DNA dyes, physiological, tracking dyes, fluorescent proteins)
- 4. Fluorochrome selection (e.g., antigen density, protein coexpression, optimal combination, photostability, F/P ratio, quenching, signal to noise)
- 5. Spectral overlap and compensation
- 6. Assay controls (e.g., fluorescence minus one [FMO], autofluorescence, biological systems control, background measurement controls)
- 7. Assay optimization (e.g., appropriate use of limited sample, frequency of target, cell concentration, kinetics, scalability, blocking, statistical design)

III. APPLICATIONS (25 – 30%)

- A. Immunophenotyping (e.g., immunologic evaluations, hematologic disorders)
- B. Functional Assays (e.g., cytokines, chronic granulomatous disease, calcium flux, phospho flow)
- C. Multiplex Bead Assays (e.g., cytokines, proteins, chemokines)
- D. Solid Organ Transplant (e.g., HLA crossmatch)
- E. Stem Cell Analysis (e.g., CD34 absolute counts)
- F. Cell Cycle / DNA Ploidy
- G. Rare Event Analysis (e.g., circulating tumor cells, measurable [minimal] residual disease [MRD], circulating endothelial cells)
- H. Small Particle Analysis
 - I. Fetal Hemoglobin Assay
 - J. Cell Sorting
 - K. Imaging Cytometry
 - L. Mass Cytometry

IV. DATA (15 – 20%)

- A. Data Standards (e.g., image file format, FCS format, listmode, MIFlowCyt checklist, storage requirements)
 - B. Signal Processing (e.g., binning, compensation, pulse processing, baseline restoration, background correction)
 - C. Data Display (e.g., types of displays, transformations)
 - D. Gating (e.g., hierarchical vs. Boolean gating, gates, regions)
 - E. Statistical Methods (e.g., central tendency, standard deviation, CV, KS statistics, cluster analysis, principal component analysis, discriminant analysis)
 - F. Common Data Modeling Techniques (e.g., cell cycle analysis, proliferation, phenotyping, ratiometric, high dimensional)
 - G. Quantitative Cytometry (e.g., molecules of equivalent soluble fluorochrome [MESF], absolute counts)
 - H. Troubleshooting (data quality assessment)
- 3. Chemical safety (e.g., mutagenic agents, cytotoxic agents)
 - 4. Environmental safety (e.g., waste disposal)
- D. Laboratory Administration
 - 1. Financial (e.g., capital equipment acquisition, cost analysis, reimbursement, purchasing, inventory)
 - 2. Operations (e.g., customer service, facility management, information technology)
 - 3. Personnel (e.g., staffing and productivity, performance standards, training and evaluation)
 - 4. Quality management (e.g., continuous quality improvement, risk management/medical-legal issues)

Examples provided (as indicated by e.g.) are not limited to those listed.

All Board of Certification examinations use conventional and SI units for results and reference ranges.

END OF CONTENT GUIDELINE

V. LABORATORY OPERATIONS (10 – 15%)

- A. Quality Control
 - 1. Instrument quality control (e.g., optical alignment, detector calibration)
 - 2. Instrument optimization
 - 3. Reagent quality control (e.g., panel verification, titration, lot to lot variation, storage, handling)
 - 4. Sample integrity
 - 5. Appropriate sample quality controls selection (internal, external)
 - 6. Trend analysis and interpretation
- B. Assay Validation
 - 1. Method validation (e.g., accuracy, reproducibility/precision, sensitivity, specificity, linearity, reference range, robustness)
 - 2. Method calibration (e.g., standards, controls)
- C. Safety
 - 1. Biosafety procedures (e.g., biosafety categories, personal protective equipment [PPE], specimen transport and preparation precautions, aerosols, decontamination)
 - 2. Instrument safety (e.g., lasers, electronics)

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Suggested Reading for Examination Preparation

This list is intended only as a partial reference source. Its distribution does not indicate endorsement by the American Society for Clinical Pathology Board of Certification (ASCP BOC), nor does the BOC wish to imply that the content of the examination will be drawn solely from these publications.

JOURNALS

- Barsky, et al. (2016). [International Society for Advancement of Cytometry \(ISAC\) Flow Cytometry Shared Resource Laboratory \(SRL\) Best Practices](#). *Cytometry*, 89(11), 1017-1030. <https://doi.org/10.1002/cyto.a.23016>
- Cossarizza, A., et al. (2019). [Guidelines for the Use of Flow Cytometry and Cell Sorting in Immunological Studies](#) (2nd ed.). *European Journal of Immunology*, 49(10), 1457-1973. <https://doi.org/10.1002/eji.201970107>
- Davis B.H., et al. (2013). [Validation of Cell-Based Fluorescence Assays: Practice Guidelines from the International Council for Standardization of Haematology and the International Clinical Cytometry Society](#). *Cytometry Part B: Clinical Cytometry*, 84(5), 279-357.
- Lee J.A., et al. (2008). [MIFlowCyt: The Minimum Information about a Flow Cytometry Experiment](#). *Cytometry. Part A: the Journal of the International Society for Analytical Cytology*, 73(10), 926-930. <https://doi.org/10.1002/cyto.a.20623>
- McFarlin, B.K. (Ed.). (2017). [Flow Cytometry](#). *Methods*, 112, pages 1-220.
- Snow, C. (2004). [Flow Cytometry Electronics](#). *Cytometry*, 57A, 63-69. <https://doi.org/10.1002/cyto.a.10120>
- Tung, J.W., et al. (2007). [Modern Flow Cytometry: A Practical Approach](#). *Clinics in Laboratory Medicine*, 27(3), 453-468. <https://doi.org/10.1016/j.cll.2007.05.001>
- Welsh, J.A., et al. (2020). [MIFlowCyt-EV: A Framework for Standardized Reporting of Extracellular Vesicle Flow Cytometry Experiments](#). *Journal of Extracellular Vesicles*, 9(1), 1713526. <https://doi.org/10.1080/20013078.2020.1713526>

TEXTS

GENERAL FLOW CYTOMETRY

- Givan, A. L. (2001). [Flow Cytometry: First Principles](#) (2nd ed.). New York: Wiley-Liss.
- Hawley, T.S. & Hawley, R.G. (Eds.). (2018). [Flow Cytometry Protocols](#) (4th ed.). New York: Humana Press.
- McCoy Jr., J.P. (Ed.). (2019). [Immunophenotyping Methods and Protocols](#). Humana Press.
- Ormerod, M.G. (2008). [Flow Cytometry: A Basic Introduction](#). Redhill: M. G. Ormerod.

CLINICAL FLOW CYTOMETRY

- Carey, J.L., McCoy, J.P., & Keren, D.F. (2007). [Flow Cytometry in Clinical Diagnosis](#) (4th ed.). Chicago: ASCP Press.
- CLSI. (2021). [Validation of Assays Performed by Flow Cytometry](#) (1st ed.). CLSI Guideline H62. Clinical and Laboratory Standards Institute.
- Dorfman, D.M., Karlson, W.J., & Linden, M.A. (2023). [Color Atlas of Flow Cytometry](#). College of American Pathologists.
- McPherson, R.A. & Pincus, M.R. (2021). [Henry's Clinical Diagnosis and Management by Laboratory Methods: Chapters 35 and 46](#) (24th ed.). Philadelphia: Elsevier.
- Nguyen, D.T., Diamond L.W., & Braylan, R.C. (2007). [Flow Cytometry in Hematopathology: A Visual Approach to Data Analysis and Interpretation](#) (2nd ed.). Totowa, NJ: Humana Press.
- Stevenson, M. (2007). [Clinical Flow Cytometric Analysis of Neoplastic Hematolymphoid Cells: Approved Guideline](#) (2nd ed.). Wayne, PA: Clinical and Laboratory Standards Institute.
- Stewart, C.C. & Nicholson, J.K. (Eds.). (2001). [Immunophenotyping: Chapters 5 – 12](#). New York: Wiley-Liss.
- Swerdlow, S.H., et al. (2017). [World Health Organization Classification of Tumour: Pathology and Genetics: Tumours of Haematopoietic and Lymphoid Tissues](#) (Revised 4th ed.). Lyon, France: IARC Press.

RESEARCH FLOW CYTOMETRY

Barteneva, N.S. & Vorobjev, I.A. (Eds.). (2016). [*Imaging Flow Cytometry: Methods and Protocols*](#). New York: Springer.

Shapiro, H.M. (2003). [*Practical Flow Cytometry*](#) (4th ed.). New York: Wiley-Liss.

Sobti R.C. & Krishan, A. (Eds.). (2003). [*Advanced Flow Cytometry: Applications in Biological Research*](#). Norwell, MA: Kluwer Academic Publishers.

LABORATORY ADMINISTRATION

U.S. Department of Health and Human Services (2020). [*Biosafety in Microbiological and Biomedical Laboratories*](#) (6th ed.). HHS. Publication No. (CDC) 300859.

Garcia, L.S. (2013). [*Clinical Laboratory Management*](#) (2nd ed.). Washington D.C.: ASM Press.

Harmening, D.M. (2020). [*Laboratory Management: Principles and Processes*](#) (4th ed.). St. Petersburg: D.H. Pub. & Consulting, Inc.

ONLINE

[The Single Cell Omics Group](#)

[Centers for Medicare and Medicaid Services](#) (includes CLIA regulations)

[Clinical and Laboratory Standards Institute \(CLSI\)](#)

[Clinical Cytometry Education Network \(CCEN\)](#)

[Clinical Laboratory Management Association \(CLMA\)](#)

[College of American Pathologists \(CAP\)](#)

[International Society for Advancement of Cytometry \(ISAC\) Cyto University](#)

[International Clinical Cytometry Society \(ICCS\)](#)

- [International Clinical Cytometry Society \(ICCS\) Quality and Standards Committee](#)

[Occupational Safety & Health Administration \(OSHA\)](#)

Specialist in Cytometry, SCYM

Credential Maintenance Program (CMP)

CMP Requirements

Total Number of Points Required: **36**

Point Distribution:

- **1** point in laboratory or patient safety (i.e., quality control, quality assurance)
- **1** point in medical ethics (e.g., HIPAA, DEI (Diversity, Equity, Inclusion), sexual harassment/microaggressions, compliance, confidentiality, IRB, etc.)
- **10** points in cytometry
- **Remaining** points in area(s) of lab specialty, management, education, or other related laboratory areas of interest

Credential Renewal Notification

Upon successful completion of the program, you will receive a **three-year renewal notification** for your credential.

Audit Provisions

To ensure the integrity of the Credential Maintenance Program, the ASCP Board of Certification will review and audit a percentage of declaration forms each year.

If your declaration is selected for audit, you will be notified and requested to submit supporting documentation of the completed continuing education credits submitted.

CMP Application Fee

\$95

CMP Reinstatement Fee

\$125 (required fee for expired certifications in addition to the application fee).

Reduced Fee for Multiple Certifications

\$15 (for multiple certifications with the same expiration date or within three months of each other, you must submit multiple declarations separately on the same business day to receive the reduced fee). Note: CMP fees are non-refundable.