# vi. Antibody isolation and purification, ELISA etc.

# vi. Antibody Isolation and Purification, ELISA, Immunoblotting, Immunohistochemistry, Immunoprecipitation, Immune Cell Isolation, Flow Cytometry, and Immunotherapy

Immunological research and diagnostics heavily rely on (1) isolating and characterizing antibodies, (2) assaying antigens and immune complexes (ELISA, immunoblotting, immunohistochemistry, immunoprecipitation), (3) isolating and analyzing immune cells (flow cytometry), and (4) applying immunotherapy for disease control. Below is a comprehensive look at each technique and concept, elucidating their principles, applications, and impact on modern biomedicine (plus some integrative insights from Ayurvedic immunomodulation).

# **Antibody Isolation and Purification**

### **Generation of Antibodies**

### 1. Polyclonal Antibodies

- Produced by **immunizing** an animal (e.g., rabbit, goat) with an antigen; serum is harvested, containing a
  mixture of IgGs recognizing multiple epitopes.
- o Pros: Broad reactivity, cost-effective. Cons: Batch variability, cross-reactivity.

### 2. Monoclonal Antibodies (mAbs)

- Derived from a single B-cell clone; generated via **hybridoma** technology (fusion of splenic B cells with myeloma cells).
- o Pros: High specificity, uniform batch production. Cons: More expensive initial production.

# **Isolation from Serum or Culture Supernatant**

### 1. Salt Precipitation

- (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> precipitation to fractionate immunoglobulins from serum.
- o An older method, simple but less precise.

# 2. Affinity Chromatography

- **Protein A** or **Protein G** columns binding Fc region of IgG.
- Highly selective, yields high-purity antibody.
- Alternatively, antigen-specific columns can isolate desired antigen-binding fraction (purifying a specific subset of polyclonal or a single mAb).

### 3. Ion-Exchange or Size-Exclusion Chromatography

Used for further polishing steps, removing aggregates or contaminants.

# **Applications**

• Purified antibodies used in **diagnostics** (ELISA, immunoblot), **therapeutics** (monoclonals in cancer, autoimmune diseases), **research** (cell labeling, immunoprecipitation, immunofluorescence).

# **Immunological Assays**

# **ELISA (Enzyme-Linked Immunosorbent Assay)**

# 1. Principle

- Detects antigen-antibody interactions using an enzyme-linked secondary antibody that produces a colorimetric (or chemiluminescent) readout.
- Variants: Indirect, Sandwich, Competitive.

### 2. Steps

- Coating of antigen or capture antibody on plates → blocking → sample addition → detection antibody → substrate for color development.
- $\circ \ \ Quantifiable \ via \ spectrophotometer.$

### 3. Applications

 $\circ \ \textbf{Serological} \ \ \text{screening for infectious diseases (HIV, hepatitis), hormone level assays, cytokine}$ 

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measurements.

• High throughput, relatively inexpensive, widely used in clinical/industrial labs.

# **Immunoblotting (Western Blot)**

### 1. Overview

- Separates proteins by SDS-PAGE → transfer onto membrane → probing with specific antibody.
- Visualizes **molecular weight** and presence of target proteins.

### 2. Procedure

- Protein sample loaded → electrophoresis → membrane transfer → blocking → primary antibody → secondary antibody (HRP/AP conjugate) → detection (chemiluminescence, colorimetric).
- Confirms protein identity, post-translational modifications, or expression changes.

### 3. Clinical Use

- o Confirmatory test for HIV (historically), verifying presence of viral proteins.
- Research applications in analyzing protein expression or immune reactivity.

# Immunohistochemistry (IHC)

#### 1. **Definition**

• Uses antigen-antibody detection in tissue sections, visualizing distribution of proteins or markers in situ.

#### 2. Method

 Tissue embedding (paraffin or frozen) → sectioning → blocking → primary antibody → detection system (enzyme-labeled or fluorescent-labeled) → microscope visualization.

# 3. Diagnostic Relevance

- Identifying tumor markers (HER2 in breast cancer, p53, Ki-67), localizing pathogens, or analyzing immune cell infiltrates.
- o Essential in histopathology for guiding clinical decisions.

# Immunoprecipitation (IP)

# 1. Concept

 Precipitating a specific antigen-antibody complex out of solution, often used to **pull down** proteins or protein complexes.

# 2. Procedure

- Mix cell lysate with antibody, add protein A/G beads → wash → analyze bound fraction (Western blot, mass spectrometry).
- o Co-immunoprecipitation (Co-IP) reveals protein-protein interactions.

# 3. Applications

o Identifying binding partners of a protein, verifying post-translational modifications, mapping signal transduction pathways.

# Immune Cell Isolation and Flow Cytometry

### **Immune Cell Isolation**

### 1. Density Gradient Centrifugation

- **Ficoll-Hypaque** or **Percoll** gradients separate mononuclear cells (lymphocytes, monocytes) from RBCs and granulocytes.
- Standard procedure for PBMC extraction from peripheral blood.

# 2. Magnetic-Activated Cell Sorting (MACS)

- Antibody-labeled microbeads selectively bind a cell type, magnetically separated.
- o Rapid, gentle, preserves viability for downstream functional assays.

### 3. Fluorescence-Activated Cell Sorting (FACS)

Flow cytometry with cell sorting function. Allows multi-parameter gating for pure subsets (CD4+ T cells, B cells, etc.).

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# Flow Cytometry

### 1. Principle

- Single-cell suspension flows past lasers; fluorophore-labeled antibodies to cell-surface or intracellular antigens produce distinct signals.
- o Measures forward scatter (size), side scatter (granularity), and multiple fluorescent channels.

# 2. Applications

- Immunophenotyping (e.g., T, B, NK cell proportions), intracellular cytokine staining, cell cycle analysis, viability checks.
- · High-throughput, multi-dimensional data on immune cell subsets, activation states, or functional responses.

### 3. Clinical Utility

- Diagnosing immunodeficiencies (e.g., counting CD4+ T cells in HIV), leukemia/lymphoma subtyping, minimal residual disease detection.
- In research, crucial for evaluating vaccine-induced immune changes or sorting cells for adoptive immunotherapy (CAR T-cell manufacturing).

# **Immunotherapy**

# **Definition and Scope**

### 1. Concept

• Manipulation of immune responses for **therapeutic** benefit: augmenting anti-tumor activity, reducing autoimmunity or transplant rejection, controlling allergies, etc.

# 2. Main Categories

- o Monoclonal Antibodies: e.g., anti-TNF, PD-1 inhibitors.
- Cell-based Therapies: CAR T-cells, adoptive TIL therapy.
- $\circ$  **Cytokine Therapy**: IL-2, IFN- $\alpha$  in certain cancers.
- **Vaccines**: prophylactic or therapeutic (HPV vaccine, personalized tumor vaccines).

# **Key Examples**

### 1. Cancer Immunotherapy

- o Checkpoint Inhibitors (CTLA-4, PD-1, PD-L1 blockade) releasing T-cell responses against tumors.
- **CAR T-Cell Therapy**: Engineering patient T cells to target specific tumor antigens (CD19 in B-cell malignancies).

### 2. Autoimmune Disease Treatments

- Biologics (e.g., anti-IL-6, anti-IL-17, anti-TNF) dampen hyperactive pathways in RA, psoriasis, Crohn's.
- Regulatory T Cell (Treg) expansion or infusion considered for severe autoimmunity.

# 3. Allergy and Infectious Disease

- Allergen immunotherapy (subcutaneous or sublingual) modulates Th2 responses.
- Passive immunization with high-titer immunoglobulins for prophylaxis or post-exposure (e.g., rabies IG, antivenom).

# 4. Ayurvedic/Herbal Immunomodulators

- Guduchi (Tinospora cordifolia) or Ashwagandha (Withania somnifera) believed to enhance immune resilience.
- o Potential synergy with mainstream immunotherapies, though requiring clinical validation.

# **Integrative Summary and Applications**

# 1. Laboratory Diagnostics

- **Immunoassays** (ELISA, Western blot, immunohistochemistry, IP) plus advanced **flow cytometry** crucial for characterizing immune responses.
- **Cell isolation** techniques ensure robust experimental or therapeutic cell populations.

# 2. Therapeutic Advances

- Immunotherapy revolutionizing oncology, autoimmune disease management, vaccine development.
- o Antibody isolation and purification underpins monoclonal antibody production—foundation of many

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"biologicals."

### 3. Ayurvedic Parallels

- While classical Ayurveda does not detail "ELISA" or "FACS," it acknowledges the principle of discerning internal vs. external substances (e.g., kṛmi) and balancing doṣas.
- Emerging integrative research might combine herbal immunomodulators with conventional immunotherapies for comprehensive immune regulation.

# **Conclusion**

Modern immunological techniques—antibody isolation, ELISA, immunoblotting, immunohistochemistry, immunoprecipitation, flow cytometry, immune cell isolation—provide powerful tools for studying and harnessing the immune system. These methods illuminate disease mechanisms (autoimmune, cancer immunopathology), guide diagnostic strategies (e.g., flow-based immunophenotyping), and fuel immunotherapy breakthroughs (monoclonal antibodies, adoptive cell transfers). Through synergy with traditional prophylactic strategies and lifestyle-based immunity from Ayurvedic thought, science continues to refine precision treatments, bridging fundamental research with transformative clinical applications.

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