
Contents

Preface

xxi

CHAPTER 1

Antibody Production by the Immune System

1

Stefanie Sarantopoulos

INTRODUCTION

1

Immune Responses Combat Microbial Invasion

1

Innate and Adaptive Immunity Are Linked Together by Cell–Cell Interactions

2

B Cells Differentiate into Antibody-Secreting Cells

3

Memory B Cells Are Produced after an Initial Encounter with Foreign Antigens

4

Adaptive Immunity Relies on Clonal Selection

4

B Cells and Antibodies Can Distinguish Foreign Organisms and Molecules from Self

6

Antibody Cross-Reactivity

6

Antibodies Mediate Effector Functions

6

CHAPTER 2

The Antibody Molecule

9

Stefanie Sarantopoulos

INTRODUCTION

9

Antibody Structure

9

Antibodies from the IgG Class Have Two Identical Antigen-Binding Sites

10

In Addition to the IgG Molecules, Serum Contains Other Classes of
Antibody Molecules

11

Comparison of the Primary Amino Acid Sequences of Light Chains Reveals
a Constant and a Variable Region

12

Comparison of the Sequences of Heavy Chains Also Reveals Variable and
Constant Regions

12

The Variable Regions of the Heavy and Light Chains Form the Antigen-Binding Site

14

Antibody Diversity

14

DNA Rearrangements Are Required for the Formation of a Functional κ Gene

14

DNA Rearrangements Are Also Required to Form a Functional λ Gene

15

Forming a Functional Heavy-Chain Gene Requires Two DNA Rearrangements

15

These Rearrangements and Additional Special Mechanisms Create a Vast Number of Antigen-Binding Sites	17
Allelic Exclusion Ensures That Only One Rearranged Light-Chain Gene and One Rearranged Heavy-Chain Gene Are Expressed in Any One B Cell	17
Somatic Mutation and Affinity Maturation: The Hypervariable CDRs	18
Additional Recombination Events Are Used to Generate the Different Classes and Subclasses of Antibodies	19

CHAPTER 3

Antibody–Antigen Interactions

Stefanie Sarantopoulos

21

INTRODUCTION	21
Structure of the Antibody–Antigen Complex	21
Affinity	24
Avidity	24

CHAPTER 4

Antibody Responses

Stefanie Sarantopoulos

31

INTRODUCTION	31
B-Cell Receptor (BCR) and BAFF Receptor Signaling Is Required for B-Cell Survival	31
Primary and Secondary Antibody Responses to Vaccination Occur with Stereotypical Kinetics	32
At the End of the Primary Response, the Antigen Is Cleared, Leaving Primed Memory Cells	32
Subsequent Injections of Antigen Induce a More Potent “Secondary” Antibody Response	33
The Type of Antigen and Its Method of Delivery Can Determine the Type of Immune Response	33
Antigen Is Presented to T Cells after Phagocytosis and Antigen Processing by APCs	34
Characteristics of the T-Cell Receptor Genes and Protein Account for Their Capacity to Recognize Antigens	35
Properties of the Interactions between Class II and Peptide and between MHC–Peptide Complex and the TCR Explain Why Only Some Compounds Make Good Antigens	35
High-Affinity Antibody Production Requires Both TCR Binding to an Antigen–Fragment–Class II Protein Complex on an APC and BCR Recognition of That Antigen	36
B-Cell Activation Requires Second Signal Delivery That Occurs with T-Cell–B-Cell Synapse	36
Additional Signals and Cytokines Lead to B-Cell Activation and Differentiation	37
Immune Tolerance Can Result in Difficult-to-Produce Antibodies	37
High-Affinity, Functional Antibody Production Occurs in the Germinal Center Reaction	39
Binding of T_{FH} Cells to B Cells Leads to Proliferation and Differentiation of B Cells	39

CHAPTER 5**Selecting the Antigen****43***Edward A. Greenfield, James DeCaprio, and Mohan Brahmandam*

INTRODUCTION	44
Immunogenicity	44
Sources of Immunogens	49
Making Weak Antigens Strong	61
PROTOCOLS	66
1 Modifying Antigens by Dinitrophenol Coupling	66
2 Modifying Antigens by Arsnyl Coupling	67
3 Modifying Protein Antigens by Denaturation	69
4 Preparing Immune Complexes for Injection	70
5 Coupling Antigens to Red Blood Cells	72
6 Coomassie Brilliant Blue Staining	73
7 Sodium Acetate Staining	74
8 Copper Chloride Staining	75
9 Side-Strip Method	76
10 Fragmenting a Wet Gel Slice	77
11 Lyophilization of a Gel Slice	78
12 Electroelution of Protein Antigens from Polyacrylamide Gel Slices	79
13 Electrophoretic Transfer to Nitrocellulose Membranes	81
14 Autoradiography	83
15 Cross-Linking Peptides to KLH with Maleimide	85
16 Preparing GST-Fusion Proteins from Bacteria	87
17 Preparing His-Fusion Proteins from Bacteria	89
18 Preparing MBP-Fusion Proteins from Bacteria	92
19 Sarkosyl Preparation of Antigens from Bacterial Inclusion Bodies	95
20 Preparing mFc- and hFc-Fusion Proteins from Mammalian Cells	97
21 Preparing Live Cells for Immunization	100
22 Preparing Antigens Using a Baculovirus Expression System	101
23 Transfected Dendritic Cell Immunizations	104

CHAPTER 6**Immunizing Animals****107***Edward A. Greenfield*

INTRODUCTION	108
Anesthesia	112
Adjuvants	113
Form of the Immunogen	118
Routes of Injection	120
Sampling Serum	127

Serum Preparation	129
Exsanguination	129
Inducing Ascites Fluid in Mice	129
Closing Comments	130
PROTOCOLS	136
<i>Anesthesia and Adjuvants</i>	136
1 Administering Anesthesia to Mice, Rats, and Hamsters	136
2 Administering Anesthesia to Rabbits	139
3 Preparing Freund's Adjuvant	142
4 Using Ribi Adjuvant	144
5 Using Hunter's TiterMax Adjuvant	145
6 Using Magic Mouse Adjuvant	146
7 Preparing Aluminum Hydroxide (Alum) Adjuvant	148
<i>Routes of Antigen Injection</i>	150
8 Injecting Rabbits Subcutaneously	150
9 Subcutaneous Injections with Adjuvant into Mice and Rats	152
10 Subcutaneous Injections without Adjuvant into Mice and Rats	153
11 Immunizing Mice and Rats with Nitrocellulose-Bound Antigen	154
12 Injecting Rabbits Intramuscularly	157
13 Injecting Rabbits Intradermally	159
14 Injecting Rabbits Intravenously	161
15 Injecting Mice Intravenously	163
16 Intraperitoneal Injections with Adjuvant into Mice and Rats	165
17 Intraperitoneal Injections without Adjuvant into Mice and Rats	166
18 Immunizing Mice, Rats, and Hamsters in the Footpad or Hock	168
<i>Serum Sampling and Preparation</i>	170
19 Sampling Rabbit Serum from the Marginal Ear Vein	170
20 Sampling Mouse Serum from the Tail Vein	172
21 Sampling Rat Serum from the Tail Vein	174
22 Sampling Mouse and Rat Serum from the Retro-Orbital Sinus	175
23 Sampling Mouse and Rat Serum from the Submandibular Vein	177
24 Sampling Mouse and Rat Serum from the Saphenous Vein	179
25 Serum Preparation	181
<i>Immunization Protocols</i>	182
26 Induction of Ascites Using Freund's Adjuvant	182
27 Induction of Ascites in BALB/c Mice Using Myeloma Cells	183
28 Standard Immunization of Mice, Rats, and Hamsters	184
29 Standard Immunization of Rabbits	186
30 Repetitive Immunization at Multiple Sites (RIMMS) of Mice, Rats, and Hamsters	188
31 Subtraction Immunization for Mice, Rats, and Hamsters	189
32 Decoy Immunization for Mice, Rats, and Hamsters	190
33 Adoptive Transfer Immunization of Mice	191
34 cDNA Immunization of Mice, Rats, and Hamsters	192

<i>Harvesting Tissue</i>	194
35 Euthanizing Mice, Rats, and Hamsters Using CO ₂ Asphyxiation	194
36 Harvesting Spleens from Mice, Rats, and Hamsters	196
37 Harvesting Lymph Nodes from Mice, Rats, and Hamsters	198

CHAPTER 7

Generating Monoclonal Antibodies

201

Edward A. Greenfield

INTRODUCTION	202
Characteristics of Monoclonal Antibodies	203
Production of Monoclonal Antibodies	207
Developing the Screening Method	207
Generating Hybridomas	216
Preparation for Fusions	217
Fusions	220
Plating Strategies	222
Feeding Hybridomas	224
Supernatant Collection Strategies for Screening	224
Screening	225
Expanding and Freezing Positive Clones	225
Dealing with Contamination	229
Isotyping and Subclassing of Monoclonal Antibodies	231
Interspecies Hybridomas	234
Human Hybridomas	234
Future Trends	235
PROTOCOLS	238
1 Antibody Capture in Polyvinyl Chloride Wells: Enzyme-Linked Detection (Indirect ELISA)	238
2 Antibody Capture in Polyvinyl Chloride Wells: Enzyme-Linked Detection When Immunogen Is an Immunoglobulin Fusion Protein (Indirect ELISA to Detect Ig Fusion Proteins)	240
3 Antibody Capture on Nitrocellulose Membrane: Dot Blot	242
4 Antibody Capture on Nitrocellulose Membrane: High-Throughput Western Blot Assay for Hybridoma Screening	244
5 Antibody Capture on Whole Cells: Cell-Surface Binding (Surface Staining by Flow Cytometry/FACS)	246
6 Antibody Capture on Permeabilized Whole Cells Binding (Intracellular Staining by Flow Cytometry/FACS)	249
7 Antibody Capture on Whole Cells: Cell-Surface Binding (Surface Staining by Immunofluorescence)	251
8 Antibody Capture on Permeabilized Whole Cells (Immunofluorescence)	253
9 Antibody Capture on Tissue Sections (Immunohistochemistry)	255
10 Antigen Capture in 96-Well Plates (Capture or Sandwich ELISA)	258

8 Isolation of IgY from Chicken Eggs: Polyethylene Glycol Method	399
9 Conjugation of Peptides to Thiol-Reactive Gel	401
10 Peptide Affinity Purification of Antibodies	403

CHAPTER 11

Engineering Antibodies 405

James R. Dasch and Amy L. Dasch

INTRODUCTION	405
PROTOCOLS	407
1 Creation of Recombinant Antibodies Using Degenerate Oligonucleotides	407
2 Creation of Recombinant Antibodies Using 5'-RACE	410
3 Using Phage Display to Create Recombinant Antibodies	412
4 Modification of Antibody Function by Mutagenesis	422
5 Rescue Strategy for Nonviable Hybridomas	424
CONCLUSION	428

CHAPTER 12

Labeling Antibodies 429

Eric A. Berg and Jordan B. Fishman

INTRODUCTION	429
Choosing a Labeling Target	430
Selecting Appropriate Cross-Linkers	431
PROTOCOLS	435
<i>Labeling Antibodies by Biotinylation</i>	435
1 Labeling Antibodies with NHS-LC-Biotin	436
2 Biotinylating Antibodies Using Biotin Polyethylene Oxide (PEO) Iodoacetamide	438
3 Biotinylating Antibodies Using Biotin-LC Hydrazide	440
<i>Labeling Antibodies with Fluorophores</i>	442
4 Labeling Antibodies Using NHS-Fluorescein	444
5 Labeling Antibodies Using a Maleimido Dye	446
<i>Forming Protein-Antibody Conjugates</i>	448
6 Conjugation of Antibodies to Horseradish Peroxidase	450
7 Labeling Antibodies with Cy5-Phycoerythrin	452
<i>Labeling Antibodies with Metals and Elements</i>	455
8 Labeling Antibodies Using Europium	457
9 Labeling Antibodies Using Colloidal Gold	459
<i>Radiolabeling Antibodies</i>	461
10 Iodination of Antibodies with Immobilized Iodogen	462

<i>Cleanup, Purification, and Storage of Antibody Conjugates</i>	464
11 Desalting or Buffer Exchange Using Size-Exclusion Chromatography	467

CHAPTER 13

Immunoblotting 469

Larisa Litovchick

INTRODUCTION	470
Choice of Antibody	471
Detection Methods	473
Experimental Strategies	473
PROTOCOLS	479
<i>Sample Preparation</i>	479
1 Preparing Whole-Cell Lysates for Immunoblotting	480
2 Preparing Protein Solutions for Immunoblotting	484
3 Preparing Immunoprecipitations for Immunoblotting	487
<i>Resolution of the Proteins in the Sample</i>	490
4 Resolving Proteins for Immunoblotting by Gel Electrophoresis	491
<i>Transfer of Proteins from Gels to Membranes</i>	497
5 Semi-Dry Electrophoretic Transfer	499
6 Wet Electrophoretic Transfer	503
7 Staining the Blot for Total Protein with Ponceau S	507
<i>Blocking and Incubation with Antibodies</i>	509
8 Blocking and Incubation with Antibodies: Immunoblots Prepared with Whole-Cell Lysates and Purified Proteins (Straight Western Blotting)	510
9 Blocking and Incubation with Antibodies of Immunoblots Prepared with Immunoprecipitated Protein Antigens (Immunoprecipitation/Western Blotting)	515
<i>Detection</i>	519
10 Detection with Enzyme-Labeled Reagents	521
11 Detection with Fluorochromes	525
<i>Reprobing the Membrane after Immunoblotting Using Chemiluminescence or Fluorochromes</i>	526
12 Stripping of the Blot for Reprobing	527

CHAPTER 14

Immunoprecipitation 531

James DeCaprio and Thomas O. Kohl

INTRODUCTION	532
Cell Preparation	534
Cell Lysis	534
Antibodies	537
Forming an Antibody-Antigen Complex	538
Epitope Tags	539

Immunoadsorbants	539
Visualizing Immune Complexes	541
Optimization	542
Background Problems	543
Controls	543
PROTOCOLS	545
<i>Radiolabeling Protein Antigens</i>	545
1 Metabolic Labeling of Antigens with [³⁵ S]Methionine	546
2 Pulse-Chase Labeling of Antigens with [³⁵ S]Methionine	552
3 Metabolic Labeling of Antigens with [³² P]Orthophosphate	558
<i>Cell Lysis</i>	563
4 Freezing Cell Pellets for Large-Scale Immunoprecipitation	563
5 Detergent Lysis of Tissue Culture Cells	565
6 Detergent Lysis of Animal Tissues	568
7 Lysis Using Dounce Homogenization	571
8 Differential Detergent Lysis of Cellular Fractions	575
9 Lysing Yeast Cells with Glass Beads	579
• Alternative Protocol: Using Lysis Buffer 2 without Detergents	582
10 Lysing Yeast Cells Using a Coffee Grinder	585
11 Denaturing Lysis	588
<i>Cross-Linking Antibodies to Beads</i>	591
12 Cross-Linking Antibodies to Beads Using Dimethyl Pimelimidate (DMP)	593
13 Cross-Linking Antibodies to Beads with Disuccinimidyl Suberate (DSS)	597
<i>Immunoprecipitation</i>	601
14 Immunoprecipitation	602
15 Tandem Immunoaffinity Purification Using Anti-FLAG and Anti-HA Antibodies	607
<i>Chromatin Immunoprecipitation</i>	614
16 Chromatin Immunoprecipitation	617
17 Dual Cross-Linking for Chromatin Immunoprecipitation	625
CHAPTER 15	
Immunoassays	629
<i>Thomas O. Kohl and Carl A. Ascoli</i>	
INTRODUCTION	629
Deciding Where to Start	630
Detecting and Quantitating Antigens	630
Detecting and Quantifying Antibodies	632
Selecting the Appropriate Microtiter Plate for Your Assay	632
ELISA Blocking Buffers	635
ELISA Wash Buffers	636
Reporter-Labeled Secondary Antibodies	636
Immunoassay Substrates	637

ELISA Parameters	639
ELISA Validation Parameters	640
Types of Immunoassays	642
PROTOCOLS	644
1 Indirect Immunometric ELISA	644
2 Immunometric Antibody Sandwich ELISA	651
3 Immunometric Double-Antibody Sandwich ELISA	658
4 Direct and Indirect Cell-Based ELISA	661
5 Direct Competitive ELISA	666
6 Indirect Competitive ELISA	670
 CHAPTER 16	
Cell Staining	675
<i>Scott J. Rodig</i>	
 INTRODUCTION	676
Major Constraints	676
Choice of Antibody	677
Protocols for Cell Staining	679
Troubleshooting Background Problems	683
 PROTOCOLS	686
1 Growing Adherent Cells on Coverslips or Multiwell Slides	686
2 Growing Adherent Cells on Tissue Culture Dishes	687
3 Attaching Suspension Cells to Slides Using the Cytocentrifuge	688
4 Attaching Suspension Cells to Slides Using Poly-L-Lysine	689
5 Preparing Cell Smears	690
6 Attaching Yeast Cells to Slides Using Poly-L-Lysine	691
7 Preparing Frozen Tissue Sections	693
8 Preparing Paraffin Tissue Sections	696
• Additional Protocol: Heat-Induced Epitope Retrieval	698
9 Preparing Cell Smears from Tissue Samples or Cell Cultures	700
10 Embedding Cultured Cells in Matrigel	701
11 Fixing Attached Cells in Organic Solvents	703
12 Fixing Attached Cells in Paraformaldehyde or Glutaraldehyde	705
13 Fixing Suspension Cells with Paraformaldehyde	707
14 Lysing Yeast	708
15 Binding Antibodies to Attached Cells or Tissues	709
16 Binding Antibodies to Cells in Suspension	711
17 Detecting Horseradish Peroxidase-Labeled Cells Using Diaminobenzidine	713
18 Detecting Horseradish Peroxidase-Labeled Cells Using Diaminobenzidine and Metal Salts	715
19 Detecting Horseradish Peroxidase-Labeled Cells Using Chloronaphthol	716
20 Detecting Horseradish Peroxidase-Labeled Cells Using Aminoethylcarbazole	717
21 Detecting Alkaline Phosphatase-Labeled Cells Using NABP-NF	718

22	Detecting Alkaline Phosphatase-Labeled Cells Using BCIP-NBT	719
23	Detecting β -Galactosidase-Labeled Cells Using X-Gal	721
24	Detecting Fluorochrome-Labeled Reagents	722
25	Detecting Gold-Labeled Reagents	725
26	Detecting Iodine-Labeled Reagents	727
27	Counterstaining Cells	729
28	Mounting Cell or Tissue Samples in DPX	730
29	Mounting Cell or Tissue Samples in Gelvatol or Mowiol	731
30	Photographing the Samples	733
CHAPTER 17		
Antibody Screening Using High-Throughput Flow Cytometry		735
<i>Thomas D. Duensing and Susan R. Watson</i>		
	INTRODUCTION	735
	Overview of Flow Cytometry	735
	Plate-Based Flow Cytometry	738
	Designing and Optimizing Flow Cytometry-Based Screening Assays	738
	PROTOCOLS	742
1	Simple Multiplexed Antibody-Binding Assay	742
	<i>High-Throughput Functional Assays for Further Assessment of Antibodies</i>	746
2	Complement-Dependent Cytotoxicity Assay	747
3	Assessment of Apoptosis (Programmed Cell Death)	749
4	Antibody-Dependent Cellular Cytotoxicity (ADCC) by Flow Cytometry	751
APPENDIX I		
Electrophoresis		755
	SDS-Polyacrylamide Gel Electrophoresis	756
	Solutions for Tris/Glycine SDS-Polyacrylamide Gel Electrophoresis	758
	Alternative Buffer Systems for Discontinuous Polyacrylamide Gel Electrophoresis	759
	Partial Proteolytic Peptide Maps—V8 Protease	760
	Two-Dimensional Isoelectric Focusing/SDS-Polyacrylamide Gel Electrophoresis	762
	Alkylation of Proteins before Running on a Gel	765
	PPO Fluorography	766
	Counting ^{35}S , ^{14}C , or ^3H from Polyacrylamide Gel Slices	767
	Counting ^{32}P or ^{125}I from Polyacrylamide Gel Slices	767
	Coomassie Blue Staining—Standard Method	768
	Coomassie Blue Staining—Quick Method	768
	Coomassie Blue Staining—Maximal Sensitivity	769
	Silver Staining of Gels—Ammoniacal Silver Staining	770
	Silver Staining of Gels—Neutral Silver Staining	771
	Copper Staining of Gels	772

Fixing Gels	773
Drying Gels	773
Rehydrating Dried Polyacrylamide Gels	774
Autoradiography and Fluorography	775
Sensitivities of Autoradiography versus Fluorography	775

APPENDIX II

Protein Techniques 777

Ammonium Sulfate Saturation Tables	778
Proteins Used as Molecular Weight Standards	779
Amino Acids	780
Genetic Code	781
Mitochondrial Genetic Code	782
Amino Acid and Codon Usage	783
Log-Odds Matrix for Relationships between Protein Sequences (MDM ₇₈)	785
Accepted Amino Acid Substitutions	786
Common Protein Sequence Motifs	787
David's Life Chart II	788
Protein Quantitation—Bradford	789
Protein Quantitation—Bradford Spot Test	790
Protein Quantitation—Coomassie Spot Test	790
Protein Quantitation—UV Detection	791
Protein Quantitation—Bicinchoninic Acid	792
Protein Quantitation—Lowry	792
Protein Quantitation—NanoDrop	793
Concentration of Protein Samples by Ultrafiltration	793
Magnetic Beads—Immunoprecipitation	793
Proteases	794
Protease Inhibitors	794
Preparing Dialysis Tubing	795
TCA Precipitation—Filtration	795
TCA Precipitation—Spotting	796
Chromogenic Substrates Yielding Water-Soluble Products	797
Chromogenic Substrates Yielding Water-Insoluble Products	797

APPENDIX III

General Information 799

Commonly Used Buffers	800
Preparation of Stock Solutions	801
Concentrations of Commercial Liquids	803
Strains of Laboratory Mice	803
Detergents	804

Detergent Removal	805
Properties of Many Commonly Used Detergents	806
 APPENDIX IV	
Bacterial Expression	809
Screening for Expression in Bacteria—SDS Lysis	810
Screening for Expression in Bacteria—Chloroform Lysis	811
λ Maps	812
Plasmid Maps	816
Gateway Expression Vectors	817
 APPENDIX V	
General Safety and Hazardous Material Information	819
Index	825

General Safety and Hazardous Material Information

This manual should be used by laboratory personnel with experience in laboratory and chemical safety or students under the supervision of such trained personnel. The procedures, chemicals, and equipment referenced in this manual are hazardous and can cause serious injury unless performed, handled, and used with care and in a manner consistent with safe laboratory practices. Students and researchers using the procedures in this manual do so at their own risk. It is essential for your safety that you consult the appropriate Material Safety Data Sheets, the manufacturers' manuals accompanying equipment, and your institution's Environmental Health and Safety Office, as well as the General Safety and Hazardous Material Information in Appendix V for proper handling of hazardous materials described in this manual. Cold Spring Harbor Laboratory makes no representations or warranties with respect to the material set forth in this manual and has no liability in connection with the use of these materials.

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