LASER FUNDAMENTALS

SECOND EDITION

WILLIAM T. SILFVAST

School of Optics / CREOL University of Central Florida



PUBLISHED BY THE PRESS SYNDICATE OF THE UNIVERSITY OF CAMBRIDGE The Pitt Building, Trumpington Street, Cambridge, United Kingdom

CAMBRIDGE UNIVERSITY PRESS
The Edinburgh Building, Cambridge CB2 2RU, UK
40 West 20th Street, New York, NY 10011-4211, USA
477 Williamstown Road, Port Melbourne, VIC 3207, Australia
Ruiz de Alarcón 13, 28014 Madrid, Spain

Dock House, The Waterfront, Cape Town 8001, South Africa

http://www.cambridge.org

First published 1996 Reprinted 1999, 2000, 2003

First edition © Cambridge University Press Second edition © William T. Silfvast 2004

This book is in copyright. Subject to statutory exception and to the provisions of relevant collective licensing agreements, no reproduction of any part may take place without the written permission of Cambridge University Press.

First published 2004

Printed in the United States of America

Typeface Times 10.5/13.5 and Avenir System AMS-TEX [FH]

A catalog record for this book is available from the British Library.

Library of Congress Cataloging in Publication data Silfvast, William Thomas, 1937– Laser fundamentals / William T. Silfvast. – 2nd ed. p. cm. Includes bibliographical references and index. ISBN 0-521-83345-0 1. Lasers. I. Title.

TA1675.S52 2004 621.36'6 - dc21

2003055352

ISBN 0 521 83345 0 hardback

Contents

P	reface to the Second Edition	page xix
P	reface to the First Edition	xxi
A	cknowledgments	xxiii
1	INTRODUCTION	1
	OVERVIEW	1
	Introduction	1
	Definition of the Laser	1
	Simplicity of a Laser	1 2 2 3
	Unique Properties of a Laser	2
	The Laser Spectrum and Wavelengths	3
	A Brief History of the Laser	4
	Overview of the Book	5
5	SECTION 1. FUNDAMENTAL WAVE PROPERTIES OF LIGHT	
2	WAVE NATURE OF LIGHT - THE INTERACTION OF LIGHT	
	WITH MATERIALS	9
	OVERVIEW	9
	2.1 Maxwell's Equations	9
	2.2 Maxwell's Wave Equations	12
	Maxwell's Wave Equations for a Vacuum	12
	Solution of the General Wave Equation – Equivalence of Light and	
	Electromagnetic Radiation	13
	Wave Velocity – Phase and Group Velocities	17
	Generalized Solution of the Wave Equation	20
	Transverse Electromagnetic Waves and Polarized Light	21
	Flow of Electromagnetic Energy	21
	Radiation from a Point Source (Electric Dipole Radiation)	22
	2.3 Interaction of Electromagnetic Radiation (Light) with Matter	23
	Speed of Light in a Medium	23
	Maxwell's Equations in a Medium	24
	Application of Maxwell's Equations to Dielectric Materials – Laser Gain Media	25
		28
	Complex Index of Refraction – Optical Constants Absorption and Dispersion	28
	AUSOLUUUI AIIU DISUCISIUII	29

viii CONTENTS

	Estimating Particle Densities of Materials for Use in the	
	Dispersion Equations	34
	2.4 Coherence	36
	Temporal Coherence	37
	Spatial Coherence	38
	REFERENCES	39
	PROBLEMS	39
S	ECTION 2. FUNDAMENTAL QUANTUM PROPERTIES OF LIGHT	
3	PARTICLE NATURE OF LIGHT - DISCRETE ENERGY LEVELS	45
	OVERVIEW	45
	3.1 Bohr Theory of the Hydrogen Atom	45
	Historical Development of the Concept of Discrete Energy Levels	45
	Energy Levels of the Hydrogen Atom	46
	Frequency and Wavelength of Emission Lines	49
	Ionization Energies and Energy Levels of Ions	51
	Photons	54
	3.2 Quantum Theory of Atomic Energy Levels	54
	Wave Nature of Particles	54
	Heisenberg Uncertainty Principle	56
	Wave Theory	56
	Wave Functions	57
	Quantum States	57
	The Schrödinger Wave Equation	59
	Energy and Wave Function for the Ground State of the	
	Hydrogen Atom	61
	Excited States of Hydrogen	63
	Allowed Quantum Numbers for Hydrogen Atom Wave Functions	66
	3.3 Angular Momentum of Atoms	67
	Orbital Angular Momentum	67
	Spin Angular Momentum	68
	Total Angular Momentum	69
	3.4 Energy Levels Associated with One-Electron Atoms	70
	Fine Structure of Spectral Lines	70
	Pauli Exclusion Principle	72
	3.5 Periodic Table of the Elements	72
	Quantum Conditions Associated with Multiple Electrons Attached	70
	to Nuclei	72
	Shorthand Notation for Electronic Configurations of Atoms Having	76
	More Than One Electron 3.6 Energy Levels of Multi-Electron Atoms	76 77
		77
	Energy-Level Designation for Multi-Electron States	
	Russell–Saunders or LS Coupling – Notation for Energy Levels Energy Levels Associated with Two Electrons in Unfilled Shells	78 79
	Rules for Obtaining S, L, and J for LS Coupling	82
	Degeneracy and Statistical Weights	84
	j-j Coupling	85
	Isoelectronic Scaling	85

CONTENTS

	REFERENCES	86
	PROBLEMS	86
4	RADIATIVE TRANSITIONS AND EMISSION LINEWIDTH	89
	OVERVIEW	89
	4.1 Decay of Excited States	90
	Radiative Decay of Excited States of Isolated Atoms -	
	Spontaneous Emission	90
	Spontaneous Emission Decay Rate – Radiative Transition	0.4
	Probability	94
	Lifetime of a Radiating Electron – The Electron as a Classical Radiating Harmonic Oscillator	95
	Nonradiative Decay of the Excited States – Collisional Decay	98
	4.2 Emission Broadening and Linewidth Due to Radiative Decay	101
	Classical Emission Linewidth of a Radiating Electron	101
	Natural Emission Linewidth as Deduced by Quantum Mechanics	
	(Minimum Linewidth)	103
	4.3 Additional Emission-Broadening Processes	105
	Broadening Due to Nonradiative (Collisional) Decay	106
	Broadening Due to Dephasing Collisions	107
	Amorphous Crystal Broadening	109
	Doppler Broadening in Gases	109
	Voigt Lineshape Profile	114
	Broadening in Gases Due to Isotope Shifts	115
	Comparison of Various Types of Emission Broadening	118
	4.4 Quantum Mechanical Description of Radiating Atoms	121
	Electric Dipole Radiation Electric Dipole Matrix Element	122 123
	Electric Dipole Madrix Element Electric Dipole Transition Probability	123
	Oscillator Strength	124
	Selection Rules for Electric Dipole Transitions Involving Atoms	124
	with a Single Electron in an Unfilled Subshell	125
	Selection Rules for Radiative Transitions Involving Atoms with	
	More Than One Electron in an Unfilled Subshell	129
	Parity Selection Rule	130
	Inefficient Radiative Transitions - Electric Quadrupole and Other	
	Higher-Order Transitions	131
	REFERENCES	131
	PROBLEMS	131
5	ENERGY LEVELS AND RADIATIVE PROPERTIES OF MOLECULES,	
	LIQUIDS, AND SOLIDS	135
	OVERVIEW	135
	5.1 Molecular Energy Levels and Spectra	135
	Energy Levels of Molecules	135
	Classification of Simple Molecules	138
	Rotational Energy Levels of Linear Molecules	139
	Rotational Energy Levels of Symmetric-Top Molecules Selection Rules for Rotational Transitions	141 141
	OCICCUON RUICS FOR ROTAUDITAL FRANKLUOUS	141

x CONTENTS

Vibrational Energy Levels	143
Selection Rule for Vibrational Transitions	143
Rotational-Vibrational Transitions	144
Probabilities of Rotational and Vibrational Transitions	148
Electronic Energy Levels of Molecules	149
Electronic Transitions and Associated Selection Rules of	
Molecules	150
Emission Linewidth of Molecular Transitions	150
The Franck-Condon Principle	151
Excimer Energy Levels	152
5.2 Liquid Energy Levels and Their Radiation Properties	153
Structure of Dye Molecules	153
Energy Levels of Dye Molecules	155
Excitation and Emission of Dye Molecules	156
Detrimental Triplet States of Dye Molecules	157
5.3 Energy Levels in Solids – Dielectric Laser Materials	158
Host Materials	158
Laser Species – Dopant Ions	159
Narrow-Linewidth Laser Materials	161
Broadband Tunable Laser Materials	
	166 168
Broadening Mechanism for Solid-State Lasers	
5.4 Energy Levels in Solids – Semiconductor Laser Materials	168
Energy Bands in Crystalline Solids	168
Energy Levels in Periodic Structures	170
Energy Levels of Conductors, Insulators, and Semiconductors	172
Excitation and Decay of Excited Energy Levels – Recombination	172
Radiation	173
Direct and Indirect Bandgap Semiconductors	174
Electron Distribution Function and Density of States in	177
Semiconductors	175
Intrinsic Semiconductor Materials	179
Extrinsic Semiconductor Materials – Doping	179
p-n Junctions - Recombination Radiation Due to Electrical	100
Excitation	182
Heterojunction Semiconductor Materials	184
Quantum Wells	186
Variation of Bandgap Energy and Radiation Wavelength with	101
Alloy Composition	191
Recombination Radiation Transition Probability and Linewidth	195
REFERENCES	195
PROBLEMS	195
RADIATION AND THERMAL EQUILIBRIUM - ABSORPTION AND	
STIMULATED EMISSION	199
OVERVIEW	199
6.1 Equilibrium	199
Thermal Equilibrium	199
Thermal Equilibrium via Conduction and Convection	200
Thermal Equilibrium via Radiation	200
	200

CONTENTS

	6.2 Radiating Bodies	201
	Stefan-Boltzmann Law	204
	Wien's Law	205
	Irradiance and Radiance	206
	6.3 Cavity Radiation	207
	Counting the Number of Cavity Modes	208
	Rayleigh–Jeans Formula	209
	Planck's Law for Cavity Radiation	210
	Relationship between Cavity Radiation and Blackbody	
	Radiation	211
	Wavelength Dependence of Blackbody Emission	214
	6.4 Absorption and Stimulated Emission	215
	The Principle of Detailed Balance	216
	Absorption and Stimulated Emission Coefficients	217
	REFERENCES	221
	PROBLEMS	221
S	ECTION 3. LASER AMPLIFIERS	
7	CONDITIONS FOR PRODUCING A LASER - POPULATION	
	INVERSIONS, GAIN, AND GAIN SATURATION	225
	OVERVIEW	225
	7.1 Absorption and Gain	225
	Absorption and Gain on a Homogeneously Broadened Radiative	
	Transition (Lorentzian Frequency Distribution)	225
	Gain Coefficient and Stimulated Emission Cross Section for	
	Homogeneous Broadening	229
	Absorption and Gain on an Inhomogeneously Broadened Radiative	
	Transition (Doppler Broadening with a Gaussian Distribution)	230
	Gain Coefficient and Stimulated Emission Cross Section for	
	Doppler Broadening	231
	Statistical Weights and the Gain Equation	232
	Relationship of Gain Coefficient and Stimulated Emission	
	Cross Section to Absorption Coefficient and Absorption	
	Cross Section	233
	7.2 Population Inversion (Necessary Condition for a Laser)	234
	7.3 Saturation Intensity (Sufficient Condition for a Laser)	235
	7.4 Development and Growth of a Laser Beam	238
	Growth of Beam for a Gain Medium with Homogeneous	32732
	Broadening	238
	Shape or Geometry of Amplifying Medium	241
	Growth of Beam for Doppler Broadening	244
	7.5 Exponential Growth Factor (Gain)	245
	7.6 Threshold Requirements for a Laser	247
	Laser with No Mirrors	247
	Laser with One Mirror	248
	Laser with Two Mirrors	249
	REFERENCES	253
	PROBLEMS	253

xii CONTENTS

8	LASER OSCILLATION ABOVE THRESHOLD	255
	OVERVIEW	255
	8.1 Laser Gain Saturation	255
	Rate Equations of the Laser Levels That Include Stimulated	
	Emission	255
	Population Densities of Upper and Lower Laser Levels with	
	Beam Present	250
	Small-Signal Gain Coefficient	257
	Saturation of the Laser Gain above Threshold	257
	8.2 Laser Beam Growth beyond the Saturation Intensity	258
	Change from Exponential Growth to Linear Growth	258
	Steady-State Laser Intensity	261
	8.3 Optimization of Laser Output Power	26
	Optimum Output Mirror Transmission	26
	Optimum Laser Output Intensity	264
	Estimating Optimum Laser Output Power	264
	8.4 Energy Exchange between Upper Laser Level Population and	
	Laser Photons	260
	Decay Time of a Laser Beam within an Optical Cavity	267
	Basic Laser Cavity Rate Equations	268
	Steady-State Solutions below Laser Threshold	270
	Steady-State Operation above Laser Threshold	272
	8.5 Laser Output Fluctuations	273
	Laser Spiking	273
	Relaxation Oscillations	270
	8.6 Laser Amplifiers	279
	Basic Amplifier Uses	279
	Propagation of a High-Power, Short-Duration Optical Pulse through	
	an Amplifier	280
	Saturation Energy Fluence	282
	Amplifying Long Laser Pulses	284
	Amplifying Short Laser Pulses	284
	Comparison of Efficient Laser Amplifiers Based upon Fundamental	
	Saturation Limits	285
	Mirror Array and Resonator (Regenerative) Amplifiers	285
	REFERENCES	288
	PROBLEMS	288
9	REQUIREMENTS FOR OBTAINING POPULATION INVERSIONS	290
	OVERVIEW	290
	9.1 Inversions and Two-Level Systems	290
	9.2 Relative Decay Rates – Radiative versus Collisional	292
	9.3 Steady-State Inversions in Three- and Four-Level Systems	293
	Three-Level Laser with the Intermediate Level as the Upper Laser	
	Level	295
	Three-Level Laser with the Upper Laser Level as the Highest Level	298
	Four-Level Laser	301
	9.4 Transient Population Inversions	304

CONTENTS

	9.5 Processes That Inhibit or Destroy Inversions	307
	Radiation Trapping in Atoms and Ions	308
	Electron Collisional Thermalization of the Laser Levels in Atoms	
	and Ions	311
	Comparison of Radiation Trapping and Electron Collisional Mixing	
	in a Gas Laser	315
	Absorption within the Gain Medium	316
	REFERENCES	319
	PROBLEMS	319
10	LASER PUMPING REQUIREMENTS AND TECHNIQUES	322
10	OVERVIEW	322
	10.1 Excitation or Pumping Threshold Requirements	322
		324
	10.2 Pumping Pathways Excitation by Direct Pumping	324
	Excitation by Direct Pumping Excitation by Indirect Pumping (Pump and Transfer)	324
		330
	Specific Pump-and-Transfer Processes	330
	10.3 Specific Excitation Parameters Associated with	220
	Optical Pumping	339
	Pumping Geometries	339
	Pumping Requirements	342 344
	A Simplified Optical Pumping Approximation	
	Transverse Pumping	346
	End Pumping	348
	Diode Pumping of Solid-State Lasers	350
	Characterization of a Laser Gain Medium with Optical Pumping	252
	(Slope Efficiency)	352
	10.4 Specific Excitation Parameters Associated with	
	Particle Pumping	355
	Electron Collisional Pumping	355
	Heavy Particle Pumping	359
	A More Accurate Description of Electron Excitation Rate to a	
	Specific Energy Level in a Gas Discharge	359
	Electrical Pumping of Semiconductors	361
	REFERENCES	363
	PROBLEMS	364
SE	CTION 4. LASER RESONATORS	
11	LASER CAVITY MODES	371
	OVERVIEW	371
	11.1 Introduction	371
	11.2 Longitudinal Laser Cavity Modes	372
	Fabry-Perot Resonator	372
	Fabry–Perot Cavity Modes	379
	Longitudinal Laser Cavity Modes	380
	Longitudinal Mode Number	380
	Requirements for the Development of Longitudinal	230
	Laser Modes	382

xiv CONTENTS

	11.3 Transverse Laser Cavity Modes	384
	Fresnel-Kirchhoff Diffraction Integral Formula	385
	Development of Transverse Modes in a Cavity with Plane-Parallel	
	Mirrors	386
	Transverse Modes Using Curved Mirrors	390
	Transverse Mode Spatial Distributions	391
	Transverse Mode Frequencies	392
	Gaussian-Shaped Transverse Modes within and beyond the	
	Laser Cavity	393
	11.4 Properties of Laser Modes	396
	Mode Characteristics	396
	Effect of Modes on the Gain Medium Profile	397
	REFERENCES	399
	PROBLEMS	399
12	STABLE LASER RESONATORS AND GAUSSIAN BEAMS	402
	OVERVIEW	402
	12.1 Stable Curved Mirror Cavities	402
	Curved Mirror Cavities	402
	ABCD Matrices	404
	Cavity Stability Criteria	406
	12.2 Properties of Gaussian Beams	410
	Propagation of a Gaussian Beam	411
	Gaussian Beam Properties of Two-Mirror Laser Cavities	412
	Properties of Specific Two-Mirror Laser Cavities	417
	Mode Volume of a Hermite-Gaussian Mode	421
	12.3 Properties of Real Laser Beams	423
	12.4 Propagation of Gaussian Beams Using ABCD Matrices –	
	Complex Beam Parameter	425
	Complex Beam Parameter Applied to a Two-Mirror Laser Cavity	428
	REFERENCES	432
	PROBLEMS	432
13	SPECIAL LASER CAVITIES AND CAVITY EFFECTS	434
	OVERVIEW	434
	13.1 Unstable Resonators	434
	13.2 Q-Switching	439
	General Description	439
	Theory	441
	Methods of Producing Q-Switching within a Laser Cavity	446
	13.3 Gain-Switching	450
	13.4 Mode-Locking	451
	General Description	451
	Theory	451
	Techniques for Producing Mode-Locking	456
	13.5 Pulse Shortening Techniques	462
	Self-Phase Modulation	463
	Pulse Shortening or Lengthening Using Group Velocity Dispersion	464
	Pulse Compression (Shortening) with Gratings or Prisms	465 467
	Ultrashort-Pulse Laser and Amplifer System	40/

CONTENTS

	13.6 Ring Lasers	468
	Monolithic Unidirectional Single-Mode Nd: YAG Ring Laser	469
	Two-Mirror Ring Laser	470
	13.7 Complex Beam Parameter Analysis Applied to Multi-Mirror	
	Laser Cavities	470
	Three-Mirror Ring Laser Cavity	470
	Three- or Four-Mirror Focused Cavity	473
	13.8 Cavities for Producing Spectral Narrowing of	
	Laser Output	478
	Cavity with Additional Fabry-Perot Etalon for Narrow-Frequency	
	Selection	478
	Tunable Cavity	478
	Broadband Tunable cw Ring Lasers	480
	Tunable Cavity for Ultranarrow-Frequency Output	480
	Distributed Feedback (DFB) Lasers	481
	Distributed Bragg Reflection Lasers	484
	13.9 Laser Cavities Requiring Small-Diameter Gain Regions -	
	Astigmatically Compensated Cavities	484
	13.10 Waveguide Cavities for Gas Lasers	485
	REFERENCES	486
	PROBLEMS	488
SE	ECTION 5. SPECIFIC LASER SYSTEMS	
4	LASER SYSTEMS INVOLVING LOW-DENSITY GAIN MEDIA	491
	OVERVIEW	491
	14.1 Atomic Gas Lasers	491
	Introduction	491
	Helium-Neon Laser	492
	General Description	492
	Laser Structure	493
	Excitation Mechanism	494
	Applications	497
	Argon Ion Laser	497
	General Description	497
	Laser Structure	498
	Excitation Mechanism	499
	Krypton Ion Laser	500
	Applications	501
	Helium-Cadmium Laser	501
	General Description	501
	Laser Structure	502
	Excitation Mechanism	504
	Applications	505
	Copper Vapor Laser	505
	General Description	505
	Laser Structure	507
	Excitation Mechanism	507
	Applications	509

xvi CONTENTS

	14.2 Molecular Gas Lasers	510
	Introduction	510
	Carbon Dioxide Laser	511
	General Description	511
	Laser Structure	511
	Excitation Mechanism	515
	Applications	515
	Excimer Lasers	516
	General Description	516
	Laser Structure	517
	Excitation Mechanism	518
	Applications	520
	Nitrogen Laser	520
	General Description	520
	Laser Structure and Excitation Mechanism	521
	Applications	522
	Far-Infrared Gas Lasers	522
	General Description	522
	Laser Structure	523
	Excitation Mechanism	523
	Applications	524
	Chemical Lasers	524
	General Description	524
	Laser Structure	524
	Excitation Mechanism	524
	Applications	525
	14.3 X-Ray Plasma Lasers	525
	Introduction	525
	Pumping Energy Requirements	525
	Excitation Mechanism	528
	Optical Cavities	532
	X-Ray Laser Transitions	532
	Applications	532
	14.4 Free-Electron Lasers	535
	Introduction	535
	Laser Structure	536
	Applications	537
	REFERENCES	537
15	LASER SYSTEMS INVOLVING HIGH-DENSITY GAIN MEDIA	539
	OVERVIEW	539
	15.1 Organic Dye Lasers	539
	Introduction	539
	Laser Structure	540
	Excitation Mechanism	543
	Applications	544
	15.2 Solid-State Lasers	545
	Introduction	545

CONTENTS xviii

Ruby Laser	547
General Description	547
Laser Structure	548
Excitation Mechanism	548
Applications	549
Neodymium YAG and Glass Lasers	550
General Description	550
Laser Structure	551
Excitation Mechanism	553
Applications	554
Neodymium:YLF Lasers	555
General Description	555
Laser Structure	556
Excitation Mechanism	556
Applications	557
Neodymium:Yttrium Vanadate (Nd:YVO ₄) Lasers	557
General Description	557
Laser Structure	557
Excitation Mechanism	558
Applications	558
Ytterbium: YAG Lasers	559
General Description	559
Laser Structure	560
Excitation Mechanism	560
Applications	561
Alexandrite Laser	562
General Description	562
Laser Structure	563
Excitation Mechanism	563
Applications	564
Titanium Sapphire Laser	565
General Description	565
Laser Structure	566
Excitation Mechanism	567
Applications	568
Chromium LiSAF and LiCAF Lasers	568
General Description	568
Laser Structure	568
Excitation Mechanism	569
Applications	570
Fiber Lasers	570
General Description	570
Laser Structure	571
Excitation Mechanism	571
Applications	572
Color Center Lasers	573
General Description	573
Laser Structure	574

xviii CONTENTS

	Excitation Mechanism	574
	Applications	576
	15.3 Semiconductor Diode Lasers	576
	Introduction	576
	Four Basic Types of Laser Materials	579
	Laser Structure	581
	Frequency Control of Laser Output	591
	Quantum Cascade Lasers	592
	p-Doped Germanium Lasers	594
	Excitation Mechanism	594
	Applications	596
	REFERENCES	597
	REFERENCES	
	ECTION 6. FREQUENCY MULTIPLICATION OF LASER BEAMS	
16	FREQUENCY MULTIPLICATION OF LASERS AND OTHER	
	NONLINEAR OPTICAL EFFECTS	601
	OVERVIEW	601
	16.1 Wave Propagation in an Anisotropic Crystal	601
	16.2 Polarization Response of Materials to Light	603
	16.3 Second-Order Nonlinear Optical Processes	604
	Second Harmonic Generation	604
	Sum and Difference Frequency Generation	605
	Optical Parametric Oscillation	607
	16.4 Third-Order Nonlinear Optical Processes	607
	Third Harmonic Generation	608
	Intensity-Dependent Refractive Index – Self-Focusing	609
	16.5 Nonlinear Optical Materials	610
	16.6 Phase Matching	610
	Description of Phase Matching	610
	Achieving Phase Matching	613
	Types of Phase Matching	615
	16.7 Saturable Absorption	615
	16.8 Two-Photon Absorption	617
	16.9 Stimulated Raman Scattering	618
	16.10 Harmonic Generation in Gases	619
	REFERENCES	619
App	pendix	621
Ind	lex	625
27104	The state of the s	020

Introduction

OVERVIEW A laser is a device that amplifies light and produces a highly directional, high-intensity beam that most often has a very pure frequency or wavelength. It comes in sizes ranging from approximately one tenth the diameter of a human hair to the size of a very large building, in powers ranging from 10^{-9} to 10^{20} W, and in wavelengths ranging from the microwave to the soft–X-ray spectral regions with corresponding frequencies from 10^{11} to 10^{17} Hz. Lasers have pulse energies as high as 10^4 J and pulse durations as short as 5×10^{-15} s. They can easily drill holes in the most durable of materials and can

weld detached retinas within the human eye. They are a key component of some of our most modern communication systems and are the "phonograph needle" of our compact disc players. They perform heat treatment of high-strength materials, such as the pistons of our automobile engines, and provide a special surgical knife for many types of medical procedures. They act as target designators for military weapons and provide for the rapid check-out we have come to expect at the supermarket. What a remarkable range of characteristics for a device that is in only its fifth decade of existence!

INTRODUCTION

There is nothing magical about a laser. It can be thought of as just another type of light source. It certainly has many unique properties that make it a special light source, but these properties can be understood without knowledge of sophisticated mathematical techniques or complex ideas. It is the objective of this text to explain the operation of the laser in a simple, logical approach that builds from one concept to the next as the chapters evolve. The concepts, as they are developed, will be applied to all classes of laser materials, so that the reader will develop a sense of the broad field of lasers while still acquiring the capability to study, design, or simply understand a specific type of laser system in detail.

DEFINITION OF THE LASER

The word *laser* is an acronym for Light Amplification by Stimulated Emission of Radiation. The laser makes use of processes that increase or amplify light signals after those signals have been generated by other means. These processes include (1) stimulated emission, a natural effect that was deduced by considerations relating to thermodynamic equilibrium, and (2) optical feedback (present in most

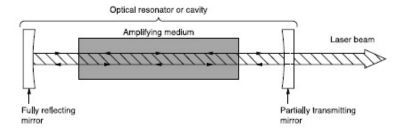


Figure 1-1 Simplified schematic of typical laser

lasers) that is usually provided by mirrors. Thus, in its simplest form, a laser consists of a gain or amplifying medium (where stimulated emission occurs), and a set of mirrors to feed the light back into the amplifier for continued growth of the developing beam, as seen in Figure 1-1.

SIMPLICITY OF A LASER

The simplicity of a laser can be understood by considering the light from a candle. Normally, a burning candle radiates light in all directions, and therefore illuminates various objects equally if they are equidistant from the candle. A laser takes light that would normally be emitted in all directions, such as from a candle, and concentrates that light into a single direction. Thus, if the light radiating in all directions from a candle were concentrated into a single beam of the diameter of the pupil of your eye (approximately 3 mm), and if you were standing a distance of 1 m from the candle, then the light intensity would be 1,000,000 times as bright as the light that you normally see radiating from the candle! That is essentially the underlying concept of the operation of a laser. However, a candle is not the kind of medium that produces amplification, and thus there are no candle lasers. It takes relatively special conditions within the laser medium for amplification to occur, but it is that capability of taking light that would normally radiate from a source in all directions - and concentrating that light into a beam traveling in a single direction - that is involved in making a laser. These special conditions, and the media within which they are produced, will be described in some detail in this book.

UNIQUE PROPERTIES OF A LASER

The beam of light generated by a typical laser can have many properties that are unique. When comparing laser properties to those of other light sources, it can be readily recognized that the values of various parameters for laser light either greatly exceed or are much more restrictive than the values for many common light sources. We never use lasers for street illumination, or for illumination within our houses. We don't use them for searchlights or flashlights or as headlights in

INTRODUCTION 3

our cars. Lasers generally have a narrower frequency distribution, or much higher intensity, or a much greater degree of collimation, or much shorter pulse duration, than that available from more common types of light sources. Therefore, we do use them in compact disc players, in supermarket check-out scanners, in surveying instruments, and in medical applications as a surgical knife or for welding detached retinas. We also use them in communications systems and in radar and military targeting applications, as well as many other areas. A laser is a specialized light source that should be used only when its unique properties are required.

THE LASER SPECTRUM AND WAVELENGTHS

A portion of the electromagnetic radiation spectrum is shown in Figure 1-2 for the region covered by currently existing lasers. Such lasers span the wavelength range from the far infrared part of the spectrum ($\lambda=1,000~\mu\text{m}$) to the soft–X-ray region ($\lambda=3$ nm), thereby covering a range of wavelengths of almost six orders of magnitude. There are several types of units that are used to define laser wavelengths. These range from micrometers or microns (μ m) in the infrared to nanometers (nm) and angstroms (Å) in the visible, ultraviolet (UV), vacuum ultraviolet (VUV), extreme ultraviolet (EUV or XUV), and soft–X-ray (SXR) spectral regions.

WAVELENGTH UNITS

$$1 \mu m = 10^{-6} \text{ m};$$

 $1 \text{ Å} = 10^{-10} \text{ m};$
 $1 \text{ nm} = 10^{-9} \text{ m}.$

Consequently, 1 micron (μ m) = 10,000 angstroms (Å) = 1,000 nanometers (nm). For example, green light has a wavelength of 5 × 10⁻⁷ m = 0.5 μ m = 5,000 Å = 500 nm.

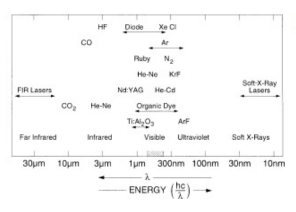


Figure 1-2 Wavelength range of various lasers

WAVELENGTH REGIONS

Far infrared: 10 to 1,000 μ m; middle infrared: 1 to 10 μ m; near infrared: 0.7 to 1 μ m;

visible: 0.4 to 0.7 μ m, or 400 to 700 nm; ultraviolet: 0.2 to 0.4 μ m, or 200 to 400 nm; vacuum ultraviolet: 0.1 to 0.2 μ m, or 100 to 200 nm;

extreme ultraviolet: 10 to 100 nm;

soft X-rays: 1 nm to approximately 20-30 nm (some overlap with EUV).

A BRIEF HISTORY OF THE LASER

Charles Townes took advantage of the stimulated emission process to construct a microwave amplifier, referred to as a *maser*. This device produced a coherent beam of microwaves to be used for communications. The first maser was produced in ammonia vapor with the inversion between two energy levels that produced gain at a wavelength of 1.25 cm. The wavelengths produced in the maser were comparable to the dimensions of the device, so extrapolation to the optical regime – where wavelengths were five orders of magnitude smaller – was not an obvious extension of that work.

In 1958, Townes and Schawlow published a paper concerning their ideas about extending the maser concept to optical frequencies. They developed the concept of an optical amplifier surrounded by an optical mirror resonant cavity to allow for growth of the beam. Townes and Schawlow each received a Nobel Prize for his work in this field.

In 1960, Theodore Maiman of Hughes Research Laboratories produced the first laser using a ruby crystal as the amplifier and a flashlamp as the energy source. The helical flashlamp surrounded a rod-shaped ruby crystal, and the optical cavity was formed by coating the flattened ends of the ruby rod with a highly reflecting material. An intense red beam was observed to emerge from the end of the rod when the flashlamp was fired!

The first gas laser was developed in 1961 by A. Javan, W. Bennett, and D. Harriott of Bell Laboratories, using a mixture of helium and neon gases. At the same laboratories, L. F. Johnson and K. Nassau demonstrated the first neodymium laser, which has since become one of the most reliable lasers available. This was followed in 1962 by the first semiconductor laser, demonstrated by R. Hall at the General Electric Research Laboratories. In 1963, C. K. N. Patel of Bell Laboratories discovered the infrared carbon dioxide laser, which is one of the most efficient and powerful lasers available today. Later that same year, E. Bell of Spectra Physics discovered the first ion laser, in mercury vapor. In 1964 W. Bridges of Hughes Research Laboratories discovered the argon ion laser, and in 1966 W. Silfvast, G. R. Fowles, and B. D. Hopkins produced the first blue helium—cadmium metal vapor

INTRODUCTION 5

laser. During that same year, P. P. Sorokin and J. R. Lankard of the IBM Research Laboratories developed the first liquid laser using an organic dye dissolved in a solvent, thereby leading to the category of broadly tunable lasers. Also at that time, W. Walter and co-workers at TRG reported the first copper vapor laser.

The first vacuum ultraviolet laser was reported to occur in molecular hydrogen by R. Hodgson of IBM and independently by R. Waynant et al. of the Naval Research Laboratories in 1970. The first of the well-known rare-gas-halide excimer lasers was observed in xenon fluoride by J. J. Ewing and C. Brau of the Avco-Everett Research Laboratory in 1975. In that same year, the first quantumwell laser was made in a gallium arsenide semiconductor by J. van der Ziel and co-workers at Bell Laboratories. In 1976, J. M. J. Madey and co-workers at Stanford University demonstrated the first free-electron laser amplifier operating in the infrared at the CO2 laser wavelength. In 1979, Walling and co-workers at Allied Chemical Corporation obtained broadly tunable laser output from a solid-state laser material called alexandrite, and in 1985 the first soft-X-ray laser was successfully demonstrated in a highly ionized selenium plasma by D. Matthews and a large number of co-workers at the Lawrence Livermore Laboratories. In 1986, P. Moulton discovered the titanium sapphire laser. In 1991, M. Hasse and co-workers developed the first blue-green diode laser in ZnSe. In 1994, F. Capasso and co-workers developed the quantum cascade laser. In 1996, S. Nakamura developed the first blue diode laser in GaN-based materials.

In 1961, Fox and Li described the existence of resonant transverse modes in a laser cavity. That same year, Boyd and Gordon obtained solutions of the wave equation for confocal resonator modes. Unstable resonators were demonstrated in 1969 by Krupke and Sooy and were described theoretically by Siegman. Qswitching was first obtained by McClung and Hellwarth in 1962 and described later by Wagner and Lengyel. The first mode-locking was obtained by Hargrove, Fork, and Pollack in 1964. Since then, many special cavity arrangements, feedback schemes, and other devices have been developed to improve the control, operation, and reliability of lasers.

OVERVIEW OF THE BOOK

Isaac Newton described light as small bodies emitted from shining substances. This view was no doubt influenced by the fact that light appears to propagate in a straight line. Christian Huygens, on the other hand, described light as a wave motion in which a small source spreads out in all directions; most observed effects – including diffraction, reflection, and refraction – can be attributed to the expansion of primary waves and of secondary wavelets. The dual nature of light is still a useful concept, whereby the choice of particle or wave explanation depends upon the effect to be considered.

Section One of this book deals with the fundamental wave properties of light, including Maxwell's equations, the interaction of electromagnetic radiation with matter, absorption and dispersion, and coherence. Section Two deals with the fundamental quantum properties of light. Chapter 3 describes the concept of discrete energy levels in atomic laser species and also how the periodic table of the elements evolved. Chapter 4 deals with radiative transitions and emission linewidths and the probability of making transitions between energy levels. Chapter 5 considers energy levels of lasers in molecules, liquids, and solids – both dielectric solids and semiconductors. Chapter 6 then considers radiation in equilibrium and the concepts of absorption and stimulated emission of radiation. At this point the student has the basic tools to begin building a laser.

Section Three considers laser amplifiers. Chapter 7 describes the theoretical basis for producing population inversions and gain. Chapter 8 examines laser gain and operation above threshold, Chapter 9 describes how population inversions are produced, and Chapter 10 considers how sufficient amplification is achieved to make an intense laser beam. Section Four deals with laser resonators. Chapter 11 considers both longitudinal and transverse modes within a laser cavity, and Chapter 12 investigates the properties of stable resonators and Gaussian beams. Chapter 13 considers a variety of special laser cavities and effects, including unstable resonators, *Q*-switching, mode-locking, pulse narrowing, ring lasers, and spectral narrowing.

Section Five covers specific laser systems. Chapter 14 describes eleven of the most well-known gas and plasma laser systems. Chapter 15 considers twelve well-known dye lasers and solid-state lasers, including both dielectric solid-state lasers and semiconductor lasers. The book concludes with Section Six (Chapter 16), which provides a brief overview of frequency muliplication with lasers and other nonlinear effects.

Radiation and Thermal Equilibrium

Absorption and Stimulated Emission

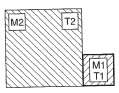
OVERVIEW Chapters 3 through 5 dealt with energy levels of gaseous, liquid, and solid materials and discussed the spontaneous radiative properties associated with transitions occurring between those levels. The rate of radiation occurring from energy levels was described in terms of transition probabilities and relative oscillator strengths of the transitions. So far, no attempt has been made to consider the collective properties of a large number of radiating atoms or

molecules. Such considerations will be dealt with in this chapter and will lead to the concept of radiation in thermodynamic equilibrium. Planck's radiation law evolved from the analysis of equilibrium radiation from dense bodies. From this law and other principles, Einstein was able to deduce the concept of stimulated emission, which is the underlying principle leading to the development of the laser.

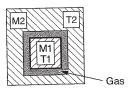
6.1 EQUILIBRIUM

Thermal Equilibrium

We will consider the processes that bring various masses into equilibrium when they are located near each other. In the study of thermodynamics, thermal equilibrium describes the case where all of the individual masses within a closed system have the same temperature. If a substance of small mass M1 and temperature T1 is placed near to or in contact with a much larger mass M2 at a higher temperature T2, then the state of thermal equilibrium is achieved when the mass M1 reaches the same temperature as that of M2. The duration over which this occurs could be a period as short as picoseconds or as long as minutes or even hours, depending upon the situation. In fact, the final temperature will be less than T2, since M2 will also be cooled slightly as it transfers some of its energy to M1. The temperature decrease of M2 will depend upon the relative masses of M1 and M2, but the relevant factor, when a new equilibrium is reached, is that the total energy will be divided equally among all of the atoms of the combined system. Thermal equilibrium can be achieved between the two masses by one or more possible heat transfer processes: conduction, convection, and radiation.

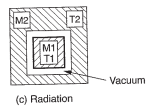


(a) Conduction



(b) Convection

Figure 6-1 Examples of energy transfer via (a) conduction, (b) convection, and (c) radiation



Thermal Equilibrium via Conduction and Convection

If M1 is placed in direct contact with M2, as shown in Figure 6-1(a), this will generally bring the masses to equilibrium in the shortest period of time. In this case, the method of heat transfer is referred to as *conduction*. If M1 and M2 are composed of two metals, for example, the rapid heat transfer by the conduction electrons of the metals would bring the two masses quickly to an equilibrium temperature.

If the bodies are placed as shown in Figure 6-1(b), with M1 located in a large cavity inside M2, and if a gas (such as air) at atmospheric pressure fills the cavity between M1 and M2, then the molecules of the gas would provide the energy transport from M2 to M1 and so bring M1 and M2 to a final equilibrium temperature somewhere between T1 and T2. This case of energy transport leading to thermal equilibrium is referred to as *convection*. A small portion of the energy would also end up in the gas, which would arrive at the same final equilibrium temperature as that of the two masses. Since the mass density of gas at atmospheric pressure is approximately three orders of magnitude lower than that of solids, the convection process takes a much longer time to reach equilibrium than does the conduction process.

Thermal Equilibrium via Radiation

If all of the air is evacuated from the space between M1 and M2 as in Figure 6-1(c), so that there is no material contact between them, there is still another process that

will bring the two masses into equilibrium – namely, *radiation*. For such a process to occur, two effects must take place: the masses must be radiating energy, and they must be capable of absorbing radiation from the other body.

6.2 RADIATING BODIES

We will first consider the question of whether or not the masses are radiating. If a collection of atoms is at temperature T then, according to the Boltzmann equation, the probability distribution function f_i that any atom has a discrete energy E_i is given by

$$f_i(E_i) = C_1 g_i e^{-E_i/kT}.$$
 (6.1)

In this equation, Boltzmann's constant k is 8.6164×10^{-5} eV/K and g_i is the statistical weight of level i. The C_1 term is a normalizing constant that is the same for all energy levels and is subject to the constraint that

$$\sum_{i} f_{i} = \sum_{i} C_{1} g_{i} e^{-E_{i}/kT} = 1, \tag{6.2}$$

which suggests that the electron must exist in one of the i energy levels. If N is the total number of atoms per unit volume of this species and N_i is the population density occupying a specific energy level i, then

$$\sum_{i} N_i = N,\tag{6.3}$$

where N_i could then be expressed as

$$N_i = f_i N = C_1 g_i e^{-E_i/kT} N. (6.4)$$

For a high-density material such as a solid, the energy levels are usually continuously distributed (with some exceptions, including insulators such as solid-state laser host materials and their dopant ions). Thus the distribution function would be expressed as a probability per unit energy g(E) such that the probability of finding a fraction of that material excited to a specific energy E within an energy width dE would be given by

$$g(E) dE = C_2 e^{-E/kT} dE, (6.5)$$

where we have ignored the statistical weights. This probability would also be subject to the normalizing constraint that

$$\int_0^\infty g(E) dE = \int_0^\infty C_2 e^{-E/kT} dE = 1.$$
 (6.6)

From this equation we can readily show that $C_2 = 1/kT$, and thus g(E) can be expressed as

$$g(E) = \frac{1}{kT}e^{-E/kT}. (6.7)$$

Again, if N is the total number of atoms per unit volume in the solid and if we refer to the number of atoms per unit volume within a specific energy range dE as N(E), then the normalizing condition requires that

$$N = \int_0^\infty N(E) dE. \tag{6.8}$$

The number of atoms at energy E within a specific energy range dE can thus be given as

 $N(E) dE = \frac{N}{kT} e^{-E/kT} dE.$ (6.9)

We can compute the ratio of the populations that exists at two specific energies, either for the case of discrete energy levels such as those for isolated atoms or for high-density materials such as solids. For discrete energy levels, according to (6.4) the ratio of the population densities N_u and N_l (number of particles per unit volume) of atoms with electrons occupying energy levels u and l (with corresponding energies E_u and E_l) would be expressed as

$$\frac{N_u}{N_l} = \frac{g_u}{g_l} e^{-(E_u - E_l)/kT} = \frac{g_u}{g_l} e^{-\Delta E_{ul}/kT},$$
(6.10)

where it is assumed that E_u is higher than E_l and that $\Delta E_{ul} = E_u - E_l$. Similarly, the ratio of population densities of a dense material (such as a solid) at energies E_u and E_l within an energy interval dE can be expressed from (6.9) as

$$\frac{N(E_u) dE}{N(E_l) dE} = \frac{N(E_u)}{N(E_l)} = e^{-\Delta E_{ul}/kT}.$$
 (6.11)

In dense materials there are so many sublevels within small ranges of energy that the statistical weights for most levels are essentially the same; hence they would effectively cancel in an expression such as (6.11). It can thus be seen that (6.10) and (6.11) are identical except for the statistical weight factor, which we have ignored for the high-density material. Therefore, when a collection of atoms – whether in the form of a gas, a liquid, or a solid – are assembled together and reach equilibrium, not only the kinetic energies related to their motion will be in thermal equilibrium; the distribution of their internal energies associated with the specific energy levels they occupy will also be in thermal equilibrium, according to (6.10) and (6.11).

EXAMPLE

Determine the temperature required to excite electrons of atoms within a solid to energies sufficient to produce radiation in the visible portion of the electromagnetic spectrum when the electrons decay from those excited levels.

We assume a typical solid material with a density of approximately $N=5\times 10^{28}$ atoms/m³ in the ground state. For several different temperatures we will

6.2 RADIATING BODIES 203

compute how many of those atoms would occupy energy levels high enough to radiate that energy as visible light. Visible radiation comprises wavelengths ranging from 700 nm in the red to 400 nm in the violet, or photons with energies ranging from approximately 1.7 to 3.1 eV. We would thus be interested in energy levels above 1.7 eV that are populated within the solid, since any electron having an energy higher than 1.7 eV will have the potential of radiating a visible photon. We will calculate the number of species that have an electron in an excited energy level that lies higher than 1.7 eV above the ground state for several different temperatures by taking the integral of N(E) dE from (6.9) over the energy range from 1.7 eV to infinity. We must recognize that, for a typical solid, most of the atoms will decay nonradiatively from these excited levels, but a certain portion could emit visible radiation depending upon the radiation efficiency of the material.

At room temperature, $T\cong 300~{\rm K}$ and $kT=0.026~{\rm eV}$. Thus $N_{\rm vis}$ can be expressed as

$$N_{\text{vis}} \cong \frac{N}{kT} \int_{1.7 \text{ eV}}^{\infty} e^{-E/kT} dE = -N[e^{-E/kT}]_{1.7 \text{ eV}}^{\infty}$$
$$= (-5 \times 10^{28})[0 - 4 \times 10^{-29}] \cong \frac{0}{\text{m}^3}.$$
 (6.12)

Thus there are essentially no atoms in this energy range from which visible photons could radiate. This is, of course, why we can see nothing when we enter a room that has no illumination, even though the human eye is very sensitive and can detect as little as only a few photons. No thermal radiation in the visible spectrum could be emitted from the walls, floors, ceiling, or furniture when those various masses are at room temperature.

We next consider a temperature of 1,000 K or 0.086 eV. At this temperature we determine the population able to emit visible photons as

$$N_{\text{vis}} = \frac{N}{kT} \int_{1.7 \text{ eV}}^{\infty} e^{-E/kT} dE = -N[e^{-E/kT}]_{1.7 \text{ eV}}^{\infty}$$
$$= (-5 \times 10^{28})[0 - 2.6 \times 10^{-9}] = \frac{1.3 \times 10^{20}}{\text{m}^3}.$$
 (6.13)

Thus, increasing the temperature by little more than a factor of 3, we have gone from essentially no atoms in those excited levels to an appreciable number in those levels. We can and do see such radiation: in the glowing coals of a campfire, in the glowing briquettes of a barbecue fire, or from the heating elements of an electric stove; all are at temperatures of approximately 1,000 K.

A temperature of 5,000 K or 0.43 eV (the temperature of the sun) can also be considered for comparison:

$$N_{\text{vis}} = \frac{N}{kT} \int_{1.7 \text{ eV}}^{\infty} e^{-E/kT} dE = -N[e^{-E/kT}]_{1.7 \text{ eV}}^{\infty}$$
$$= (-5 \times 10^{28})[0 - 1.9 \times 10^{-2}] = \frac{9.5 \times 10^{26}}{\text{m}^3}.$$
 (6.14)

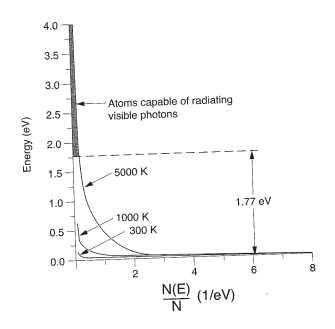


Figure 6-2 Population distribution of occupied states versus energy for temperatures of 300 K, 1,000 K, and 5,000 K

At this temperature nearly 10% of the atoms would be excited to an energy of 1.7 eV or higher, and the material would be radiating with an intensity that is too bright to look at.

A graph of N(E)/N, which is the distribution of population versus energy as taken from (6.9), is plotted in Figure 6-2 for 300 K, 1,000 K, and 5,000 K. The graph clearly shows the rapid increase in the population at higher energies as the temperature is increased. It can be seen that the energy levels that are high enough to produce visible radiation are populated only when the temperature is significantly above room temperature, as discussed in the preceding examples. Such radiation emitted from masses at those various temperatures would emit a continuous spectrum of frequencies over a certain frequency range. The radiation is referred to as thermal radiation, since it is emitted from objects in thermal equilibrium.

There are two effects we should consider when observing thermal radiation such as that of glowing coals. First, more energy is radiated from the object as the temperature is increased; this is described by the Stefan-Boltzmann law. Second, if the spectral content of the radiation from the glowing coals is analyzed, it will be found that the radiation increases with decreasing wavelength to a maximum value at a specific wavelength, and then it decreases relatively rapidly at even shorter wavelengths. The wavelength at which the maximum value occurs can be obtained from Wien's law.

Stefan-Boltzmann Law

The Stefan-Boltzmann law is an empirical relationship obtained by Stefan and later derived theoretically by Boltzmann. It states that the total radiated intensity (W/m^2) emitted from a body at temperature T is proportional to the fourth power of the temperature, T^4 . This can be written as

$$I = e_M \sigma T^4, \tag{6.15}$$

where $\sigma = 5.67 \times 10^{-8}$ W/m²-K⁴ and e_M is the emissivity that is specific for a given material. The emissivity, a dimensionless quantity that varies between zero and unity, represents the ability of a body to radiate efficiently and is also associated with its ability to absorb radiation, as will be discussed in more detail in Section 6.3. Equation (6.15) describes an extremely rapidly increasing function with temperature and accounts for the tremendous flux increase we estimated to be radiating in the visible from our simple system at 300 K, 1,000 K, and 5,000 K. The total radiation from such a mass would increase in going from 300 K to 1,000 K by a factor of 123, and in going from 300 K to 5,000 K by a factor of over 77,000!

Wien's Law

A graph of the spectrum of radiation emitted versus frequency from a heated mass is shown for three different temperatures in Figure 6-3. The specific wavelength at which the radiation is a maximum was found to vary inversely with temperature. The wavelength λ_m at which the maximum emission occurs for any given temperature is described by Wien's law as follows:

$$\lambda_m T = 2.898 \times 10^{-3} \text{ m-K}.$$
 (6.16)

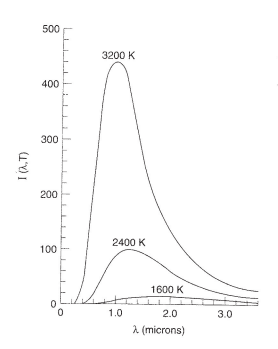


Figure 6-3 Spectrum of radiation versus wavelength of a heated mass (blackbody) for several temperatures

TABLE 6-1

Total Emissivity e_M at Low

Temperatures

Highly polished silver	0.02
Aluminum	0.08
Copper	0.15
Cast iron	0.25
Polished brass	0.60
Black gloss paint	0.90
Lampblack	0.95

Irradiance and Radiance

Recall the case depicted in Figure 6-1(c). Because M2 is initially at a higher temperature than M1, we surmise that M1 will eventually be raised to equilibrium with M2, at a temperature T3 such that T1 < T3 < T2, by absorbing the radiation from M2. However, when M1 reaches the same temperature as M2, it must radiate as much energy as it absorbs. Otherwise, it would continue to heat up to a higher temperature than T2, which defies the laws of thermodynamics. Thus the relationship – in total power per unit area – between the *irradiance* I_1 incident upon M1 and the radiance H_1 leaving M1 must be proportional:

$$H_1 = b_1 I_1$$
 or $I_1 = \frac{H_1}{b_1}$, (6.17)

where $b_1 \le 1$ is the proportionality constant that represents the fraction of the power absorbed by M1. If instead of just M1 there were several masses (M1, M3, and M4) all inside the cavity of M2, we could express their radiative characteristics with respect to the incident flux in the same way as that of (6.17) for M1. We would obtain the relationships $H_1 = b_1 I_1$, $H_3 = b_3 I_3$, and $H_4 = b_4 I_4$. Because I (the power per unit area) arriving at each body would be the same, we can write

$$I = \frac{H_1}{b_1} = \frac{H_3}{b_3} = \frac{H_4}{b_4} = \cdots, \tag{6.18}$$

which indicates that the ratio of the power radiated from a body to the fraction of radiation absorbed is a constant, independent of the material. Thus strong absorbers are also strong radiators, and since b represents the fraction of radiation absorbed we can conclude that $b = e_M$. This relationship is known as *Kirchhoff's law*. Examples of emissivity are shown in Table 6-1. It can be seen that emissivities range from very small numbers for highly reflecting materials to values near unity for highly absorbing materials. Lampblack, the soot deposited from burning candles, is perhaps the blackest material we ever observe; it has an emissivity of $e_M = 0.95$.

The ideal case of a perfect absorber ($b = \varepsilon_M = 1$) is also a perfect emitter in that it would radiate as much energy as is incident upon it. Such a perfectly absorbing body is known as a blackbody (since it appears to be very black in color)

6.3 CAVITY RADIATION 207

and is the best emitter of thermal radiation. The radiation it emits is consequently referred to as blackbody radiation. If such a blackbody is placed within a cavity in thermal equilibrium – say, M1 in Figure 6-1(c) – then it would radiate an intensity and frequency distribution that is characteristic of the thermal radiation within the medium because the radiance would be equal to the thermal radiation (irradiance) within the cavity. Hence blackbody radiation is also referred to as cavity radiation.

6.3 CAVITY RADIATION

We have developed the idea that ideal blackbodies radiate with the same spectral power as that occurring within a cavity in thermal equilibrium such as the radiation within the cavity depicted in Figure 6-1(c). It will be useful for us to obtain the quantity and wavelength distribution of the radiated flux within such a cavity in order to later obtain the stimulated emission coefficient, which is the fundamental concept leading to amplification in a laser medium. We must allow for the possibility of all frequencies of radiation within the cavity and would like to obtain an expression describing that frequency distribution for any specific temperature. Toward this end, we begin by considering the properties of radiation within a cavity. The boundary conditions – as deduced from electromagnetic theory – suggest that, in order for electromagnetic waves to be supported or enhanced (or at least not rapidly die away), the value of the electric vector must be zero at the boundary of the cavity. This involves the development of distinct "standing waves" within the cavity - that is, waves whose functions exhibit no time dependence. Such standing waves, which have a value of zero for the electric field at the boundaries, are referred to as cavity modes. The existence of each mode implies that the wave has an integral number of half-wavelengths occurring along the wave direction within the cavity. For example, three distinct cavity modes oriented in a specific direction are shown in Figure 6-4: waves of one half-wavelength, two half-wavelengths (one complete cycle), and three half-wavelengths. If we can calculate the number of such modes at each frequency or wavelength within a cavity and multiply by the average energy of each mode at those wavelengths, we can obtain the frequency

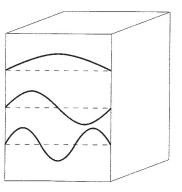


Figure 6-4 Several modes of electromagnetic radiation within a confined cavity

distribution of the emission spectrum for cavity radiation. Hence we will now calculate the number of modes that exist within a cavity of a given size.

Counting the Number of Cavity Modes

We will carry out a counting procedure for these modes in a way similar to that used in Section 5.4 for counting the number of modes available for electrons in the conduction band of a semiconductor. Assume we have a cavity that is rectangular in shape with dimensions L_x , L_y , and L_z , as shown in Figure 6-1(c). Because we are considering only standing waves within the cavity, in counting the number of modes we will include only the spatial dependence of the electromagnetic waves, which is described by an oscillatory function of the form $e^{-i(k_x L_x + k_y L_y + k_z L_z)}$. In order to satisfy the boundary condition that the field be periodic in L and also zero at the boundaries, the exponential phase factor must be an integral multiple of π . This can be achieved if we specify individual modes in the directions x, y, z such that

$$k_x L_x = n_x \pi, \quad k_y L_y = n_y \pi, \quad k_z L_z = n_z \pi,$$

 $n_x, n_y, n_z = 0, 1, 2, \dots$ (6.19)

Any mode in this cavity will have a specific value of k; its mode number can be identified by specifying mode numbers n_x , n_y , n_z since

$$k^{2} = (k_{x}^{2} + k_{y}^{2} + k_{z}^{2}) = \left[\left(\frac{n_{x}\pi}{L_{x}} \right)^{2} + \left(\frac{n_{y}\pi}{L_{y}} \right)^{2} + \left(\frac{n_{z}\pi}{L_{z}} \right)^{2} \right].$$
 (6.20)

The total number of modes in any volume $V = L_x L_y L_z$ up to a given value of k can then be counted by using a three-dimensional space whose axes are defined as the number of modes. For example, the number of modes in the x direction would be the length of the cavity in that direction L_x divided by one half-wavelength or $n_x = L_x/(\lambda/2)$, which is equivalent to the mode number in that direction. Consequently, the number of modes along each axis of the volume for a specific wavelength λ is obtained as $n_x = 2L_x/\lambda$, $n_y = 2L_y/\lambda$, and $n_z = 2L_z/\lambda$. We will choose one octant of an ellipsoid to describe the mode volume, since this is a convenient way of counting the modes. (We choose only one octant with the foregoing dimensions for the major axes because all of the values of n are positive.) Each mode – say $n_x = 1$, $n_y = 2$, and $n_z = 1$, for example – is specified in our mode volume as the intersection of those specific mode numbers within that octant, as shown in Figure 6-5. Such a mode can be specified, for the purposes of this calculation, by assuming that it is represented by a cube of unit dimensions within that volume. This approximation is very good for volumes that are significantly larger than the wavelength λ , and since we are dealing with relatively small (short) wavelengths, the volume calculation for counting the number of modes is quite accurate.

The volume of the octant can be written as

$$\frac{1}{8} \cdot \frac{4\pi}{3} \cdot n_x n_y n_z = \frac{1}{8} \cdot \frac{4\pi}{3} \left(\frac{2L_x}{\lambda} \cdot \frac{2L_y}{\lambda} \cdot \frac{2L_z}{\lambda} \right); \tag{6.21}$$

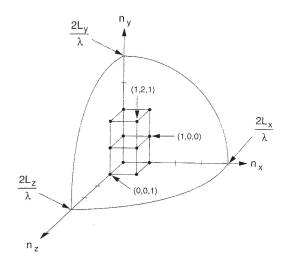


Figure 6-5
Three-dimensional diagram of the cavity mode volume

likewise, the number of modes M can be expressed as

$$M = \frac{1}{8} \cdot \frac{4\pi}{3} \left(\frac{2L_x 2L_y 2L_z}{\lambda^3} \right) = \frac{4\pi\eta^3 v^3}{3c^3} V, \tag{6.22}$$

where $V = L_x L_y L_z$ and we have used the relationship $\lambda \nu = c/\eta$. This calculation defines the number of modes for *all frequencies up to and including the frequency* ν within a volume V. This number must be doubled to allow for the fact that two orthogonal polarizations of the electromagnetic waves must be represented for each spatial mode. Higher frequencies would be outside of this volume and thus would not be counted. For frequencies up to the value ν , the mode density ρ (number of modes per unit volume) is then given by

$$\rho(\nu) = \frac{2M}{V} = \frac{8\pi\eta^3\nu^3}{3c^3}.$$
 (6.23)

This equation can be differentiated to obtain the number of modes per unit volume within a given frequency interval between ν and $\nu + d\nu$,

$$\frac{d\rho(v)}{dv} = \frac{8\pi\eta^3 v^2}{c^3}.$$
 (6.24)

We have used a rectangular cavity to derive the mode density as well as the mode density per unit frequency for the number of modes up to frequency ν , but it can be shown that (6.23) and (6.24) are generally applicable formulas for any shape of cavity.

Rayleigh-Jeans Formula

If we assume that the average energy per mode is of the order of kT, then we can obtain the Rayleigh–Jeans formula for the energy density u(v) of radiation per unit

volume within the frequency v to v + dv. We do this by multiplying the mode density per unit frequency by the average energy kT per mode. Therefore, using kTas the value of the energy per mode, for the energy density per unit frequency we have

 $u(v) = \frac{d\rho(v)}{dv}kT = \frac{8\pi\eta^3 v^2}{c^3}kT.$ (6.25)

This result suggests that there is a continuous increase in energy density with frequency for a given temperature T. This expression is referred to as the Rayleigh-Jeans formula for the energy density of cavity radiation. It agrees with experiments for lower frequencies but does not predict the experimentally observed maximum value for a given temperature at higher frequencies as shown in Figure 6.3. Instead, the Rayleigh-Jeans expression suggests that energy density approaches infinity as the frequency is increased.

Planck's Law for Cavity Radiation

Planck was disturbed by the invalidity of the Rayleigh-Jeans formula for the intensity of cavity radiation at higher frequencies and so he questioned the basic assumption of assigning a value of kT for the average energy per cavity mode. As an alternative, Planck explored the possibility of quantizing the mode energy, postulating that an oscillator of frequency v could have only discrete values mhv of energy, where $m = 0, 1, 2, 3, \dots$ He referred to this unit of energy hv as a quanturn that could not be further divided. We can apply this condition to obtain the energy of each cavity mode in thermal equilibrium. We will use the Boltzmann distribution function (eqn. 6.1) to describe the energy distribution of the modes; however, we will assign discrete values for the energy $E_m = mhv$ in the function instead of a continuous variable E, and we will ignore the statistical weight factor since we are talking about very dense materials. Thus, for a given temperature T, we can express the distribution function as

$$f_m = Ce^{-E_m/kT} = Ce^{-mh\nu/kT}.$$
 (6.26)

The normalizing condition for the distribution function requires that

$$\sum_{m=0}^{\infty} f_m = \sum_{m=0}^{\infty} Ce^{-mh\nu/kT} = C \sum_{m=0}^{\infty} e^{-mh\nu/kT} = 1,$$
 (6.27)

which can be solved for C to obtain

$$C = 1 - e^{-h\nu/kT}. (6.28)$$

We can then compute the average mode energy \bar{E} of those oscillators in the usual manner:

$$\bar{E} = \frac{\sum_{m=0}^{\infty} E_m f_m}{\sum_{m=0}^{\infty} f_m} = \frac{C \sum_{m=0}^{\infty} (mh\nu) e^{-mh\nu/kT}}{C \sum_{m=0}^{\infty} e^{-mh\nu/kT}}$$

$$= \frac{[1 - e^{-h\nu/kT}] \sum_{m=0}^{\infty} (mh\nu) e^{-mh\nu/kT}}{1}.$$
(6.29)

The value of the summation in the numerator on the right-hand side of (6.29) is

$$\sum_{m=0}^{\infty} (mh\nu) e^{-mh\nu/kT} = h\nu \sum_{m=0}^{\infty} me^{-mh\nu/kT} = \frac{h\nu e^{-h\nu/kT}}{(1 - e^{-h\nu/kT})^2}.$$
 (6.30)

This leads to an expression for the average energy \bar{E} of

$$\bar{E} = \frac{h\nu}{e^{h\nu/kT} - 1}.\tag{6.31}$$

Using this expression for the average energy per mode instead of the kT value used in the Rayleigh–Jeans law to obtain (6.25), we arrive at the following relationship for the energy density per unit frequency:

$$u(\nu) = \frac{8\pi h \eta^3 \nu^3}{c^3 (e^{h\nu/kT} - 1)}.$$
 (6.32)

This relationship has a maximum at a specific frequency for a given temperature, and both the location of the maximum and the shape of the distribution agree very well with experimental observations. This expression became known as Planck's law for cavity radiation.

The expression $u(\nu)$ describes the energy density per unit frequency ν for radiation anywhere within an enclosed cavity in thermal equilibrium at temperature T. It consists of waves traveling in all directions within the cavity. If we wanted to compute the total energy density u emitted at all frequencies, we could of course simply integrate the energy density $u(\nu) d\nu$ over the frequencies:

$$u = \int_0^\infty u(v) \, dv. \tag{6.33}$$

Carrying out this integration leads to the Stefan-Boltzmann law.

Relationship between Cavity Radiation and Blackbody Radiation

If we were able to make a small hole of unit area through mass M2 into the cavity within M2 of Figure 6-1(c), we would observe a small amount of radiation of intensity $I(\nu)$ at any specific frequency ν emerging from the hole; that intensity is related to the energy density $u(\nu)$ within the cavity. We could then calculate the total radiation flux emerging from that cavity at frequency ν , through the hole,

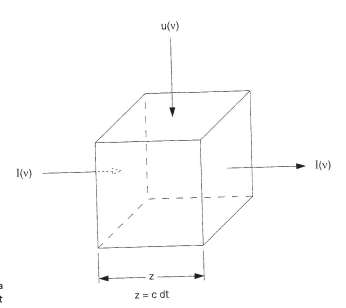


Figure 6-6 Beam of light intensity l(v) incident upon a volume of thickness z = c dt

traveling in all directions within a solid angle of 2π (a hemisphere). This value for the radiation flux at frequency ν would describe the irradiance of a blackbody, since we deduced earlier that the spectral density within a cavity has the same energy density as that at the surface of a blackbody. We will therefore refer to this radiation as the blackbody radiation intensity per unit frequency, or $I_{\rm BB}(\nu)$.

Before performing this calculation, we need to obtain a relationship between the energy density u(v) (energy per unit volume-frequency) of electromagnetic radiation within a volume element and the flux I(v) (energy per time-area-frequency) passing through a surface enclosing part of that volume element and traveling in a specific direction, as shown in Figure 6-6. Let us consider a beam of light of intensity per unit frequency I(v) at frequency v, with a cross-sectional area dA and traveling in a direction z that passes through that area dA in a time dt. The energy density of radiation would be the product of the intensity of the beam, the cross-sectional area, and the time duration, or I(v) dA dt, which has units of energy per unit frequency. This would be equivalent to considering the energy density u(v) of a beam existing within a volume $dV = dA \cdot z$ if an instantaneous photograph were taken of the beam within that volume dV of length z = v dt, where v is the velocity of the beam. This is described by the expression

$$I(v) dA dt = u(v) dV = u(v) dA \cdot z = u(v) dA \cdot v dt.$$
 (6.34)

If the volume element is in a vacuum then $v \equiv c$, which leads to the relationship

$$I(v) = u(v)c \tag{6.35}$$

for a beam traveling in a specific direction.

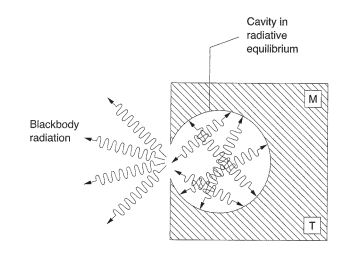


Figure 6-7(a) Blackbody radiation escaping from a cavity within mass *M* at equilibrium temperature *T*

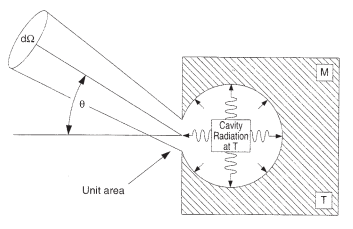


Figure 6-7(b) Coordinates for analyzing blackbody radiation escaping from a confined cavity

When the volume element is in a medium whose index of refraction is η , the velocity is expressed as $v = c/\eta$; this leads to

$$I(v) = u(v)\frac{c}{\eta}. (6.36)$$

We can now convert the expression for the energy density of radiation within a cavity of temperature T to an expression for blackbody radiation emerging from a hole of unit surface area accessing that cavity and radiating into a hemisphere, as indicated in Figure 6-7(a). Radiation from any point within the cavity is traveling in all directions – that is, within a 4π solid angle. We consider the coordinates shown in Figure 6-7(b), where the fraction of radiation traveling within a solid angle $d\Omega/4\pi$ at a particular angle θ with respect to the normal to the plane of the hole is described by $I(\nu)\cos\theta \,d\Omega/4\pi$. In this expression we have assumed that the intensity emitted from the source is independent of angle, but the factor $\cos\theta$ represents the decrease in the effective area of the hole with increasing angle from the normal as seen by the observer. Expressing the solid angle as $d\Omega = \sin\theta \,d\theta \,d\phi$,

the radiation flux in that direction can be written as $I(\nu)\cos\theta\sin\theta \,d\theta\,d\phi/4\pi$. The total blackbody flux $I_{\rm BB}(\nu)$ is obtained by integrating this component of the flux over the entire hemisphere (2π solid angle):

$$I_{\rm BB}(\nu) = \int_0^{2\pi} \int_0^{\pi/2} \frac{I(\nu) \cos \theta \sin \theta \, d\theta \, d\phi}{4\pi}.$$
 (6.37)

Replacing I(v) by u(v)c from (6.35) gives the relationship between the blackbody radiation intensity per unit frequency, $I_{BB}(v)$, and the energy density per unit frequency, u(v), of the cavity:

$$I_{\text{BB}}(\nu) = \int_0^{2\pi} \int_0^{\pi/2} u(\nu)c \cdot \frac{\cos\theta \sin\theta \, d\theta \, d\phi}{4\pi} = \frac{u(\nu)c}{4}.$$
 (6.38)

Using (6.38) in conjunction with the value of u(v) indicated in (6.32), we can obtain the spectral radiance of a blackbody as a function of v and T as follows:

$$I_{BB}(\nu) = \frac{2\pi h \nu^3}{c^2} \frac{1}{e^{h\nu/kT} - 1}.$$
 (6.39)

Wavelength Dependence of Blackbody Emission

The total radiance of $I_{\rm BB}(\nu)$ from a frequency interval $d\nu$ is given as

$$I_{BB}(v) dv = \frac{2\pi h v^3}{c^2} \frac{dv}{e^{hv/kT} - 1}.$$
 (6.40)

Because $\lambda \nu = c$ (i.e., $\nu = c/\lambda$), it follows that

$$|dv| = \frac{c}{\lambda^2} \, d\lambda. \tag{6.41}$$

We can therefore express the intensity per unit wavelength at temperature T as

$$I_{\rm BB}(\lambda, T) = \frac{2\pi c^2 h}{\lambda^5 (e^{ch/\lambda kT} - 1)}.$$
(6.42)

The total radiance emitted from the blackbody surface within a specific wavelength interval $\Delta\lambda$ would then be expressed as

$$I_{\rm BB}(\lambda, \Delta\lambda, T) = I_{\rm BB}(\lambda, T)\Delta\lambda.$$
 (6.43)

Specific values of the radiance $I_{BB}(\lambda, \Delta\lambda, T)$ as given by (6.43), in units of power per unit area (intensity) over the wavelength interval $\Delta\lambda$, can be obtained from the following empirical expression for $I_{BB}(\lambda, T)$:

$$I_{\text{BB}}(\lambda, T) = \frac{3.75 \times 10^{-22}}{\lambda^5 (e^{0.0144/\lambda T} - 1)} \text{ W/m}^2 - \mu \text{m}, \tag{6.44}$$

where λ must be given in meters, $\Delta\lambda$ in micrometers (μ m), and T in degrees Kelvin. Alternatively,

$$I_{\rm BB}(\lambda, T) = \frac{3.75 \times 10^{-25}}{\lambda^5 (e^{0.0144/\lambda T} - 1)} \,\text{W/m}^2\text{-nm}$$
 (6.45)

for λ in meters, $\Delta\lambda$ in nanometers, and T in degrees Kelvin. Remember that these expressions are for the radiation into a 2π steradian solid angle.

EXAMPLE

Compute the radiation flux or power in watts coming from a surface of temperature 300 K (near room temperature) and area 0.02 m² over a wavelength interval of 0.1 μ m at a wavelength of 1.0 μ m.

Use (6.43) and (6.44) to obtain the intensity (W/m^2) and multiply it by the area ΔA (m^2) to obtain the power P in watts coming from the surface:

$$\begin{split} P &= I_{\rm BB}(\lambda,T) \Delta \lambda \Delta A \\ &= \frac{3.75 \times 10^{-22}}{(1 \times 10^{-6})^5 (e^{0.0144/(1 \times 10^{-6})300} - 1)} (0.1)(0.02) = 1.06 \times 10^{-15} \, \rm W. \end{split}$$

We can see that at room temperature the power radiated from a blackbody within this wavelength region is almost too small to be measurable.

6.4 ABSORPTION AND STIMULATED EMISSION

We have examined the issues relating to the decay of electrons from a higher energy level to a lower level. We have also investigated the natural radiative decay process, which is inherent in all excited states of all materials and is referred to as *spontaneous emission*. However, we saw in Chapter 4 that such emission is not always the dominant decay process. We described how collisions with other particles (in the case of gases) or phonons (in the case of solids) can depopulate a level faster than the normal radiative process. Such collisions can also populate or *excite* energy levels.

Excitation or de-excitation can also occur by way of photons – "light particles" that have specific energies. The phenomenon of light producing excitation, referred to as *absorption*, has been known for well over 100 years. Such a process

could also have been referred to as "stimulated" absorption, since it requires electromagnetic energy to stimulate the electron and thereby produce the excitation. There is no reason to suppose that the inverse of that process would not also occur, but it was never seriously considered until Einstein suggested the concept of stimulated emission in 1917.

The Principle of Detailed Balance

Einstein was considering the recently developed Planck law for cavity radiation, the expression we just derived. He began questioning how the principle of detailed balance must apply in the case of radiation in equilibrium, a situation associated with cavity radiation. This principle states that, *in equilibrium*, the total number of particles leaving a certain quantum state per unit time equals the number arriving in that state per unit time. It also states that in equilibrium the number leaving by a particular pathway equals the number arriving by that pathway.

We tacitly used this principle in the first part of this chapter when arguing that the mass M1 could not continue to receive excess energy from M2 once it had reached the new equilibrium temperature. The principle describes why the population of a specific energy state cannot increase indefinitely. This principle has also been called the principle of *microscopic reversibility* and was originally applied to considerations of thermodynamic equilibrium.

The principle of detailed balance suggests that if a photon can stimulate an electron to move from a lower energy state l to higher energy state u by means of absorption, then a photon should also be able to stimulate an electron from the same upper state u to the lower state l. In the case of absorption, the photon disappears, with the energy being transferred to the absorbing species. In the case of stimulation, or stimulated emission, the species would have to radiate an additional photon to conserve energy. Such a stimulated emission process must occur in order to keep the population of the two energy levels in thermal equilibrium, if that equilibrium is determined by cavity radiation as described earlier in this chapter. Thus, the relationships between absorption and stimulated emission must be associated in some way with the Planck law for radiation in thermal equilibrium.

We have identified the three radiative processes producing interactions between two bound levels in a material: spontaneous emission, absorption (stimulated absorption), and stimulated emission. These processes are diagrammed in Figure 6-8. In the case of both of the stimulated processes (absorption and stimulated emission) occurring between bound (discrete) states, the energy (related to the frequency) of the light must correspond exactly to that of the energy difference between the two energy states. For stimulated emission, an additional photon is emitted at exactly the same energy (or frequency) as that of the incident photon (conservation of energy), and in exactly the same direction in phase with the incident photon (conservation of momentum).

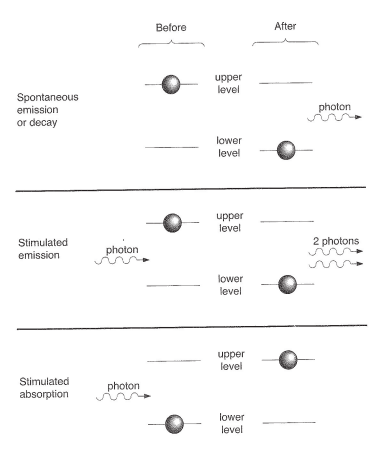


Figure 6-8 The three fundamental radiation processes associated with the interaction of light with matter: spontaneous emission, stimulated emission, and absorption

Absorption and Stimulated Emission Coefficients

We will now derive the absorption and stimulated emission coefficients associated with these processes by considering radiation in thermal equilibrium. We will consider a group of atoms having electrons occupying either energy levels u or l with population densities N_u and N_l (number of atoms per unit volume). We assume the atoms are in thermal equilibrium with each other and must therefore be related by the Boltzmann distribution function given in (6.10),

$$\frac{N_u}{N_l} = \frac{g_u}{g_l} e^{-(E_u - E_l)/kT} = \frac{g_u}{g_l} e^{-\Delta E_{ul}/kT},$$
 (6.46)

where g_u and g_l are the statistical weights of levels u and l and where $E_u - E_l = \Delta E_{ul}$.

We will consider photons interacting with such a collection of atoms. The photons will be assumed to have energies ΔE_{ul} such that $\Delta E_{ul} = h \nu_{ul}$, corresponding to the exact difference in energy between the levels u and l. We have defined A_{ul} as the spontaneous transition probability, the rate at which spontaneous transitions

occur from level u to level l (number per unit time). Thus, the number of spontaneous transitions from u to l per unit time per unit volume is simply $N_u A_{ul}$.

We have also suggested that stimulated processes should occur. These processes would be proportional to the photon energy density u(v) at frequency v_{ul} as well as to the population in the appropriate level. If we assume that the proportionality constant for such stimulated transitions is B, then the upward flux – the number of stimulated upward transitions per unit volume per unit time per unit frequency – would be $N_l B_{lu} u(v)$. Similarly, the downward flux would be $N_u B_{ul} u(v)$. The constants A_{ul} , B_{ul} , and B_{lu} are referred to as the Einstein A and B coefficients.

For the populations N_u and N_l to be in *radiative thermal equilibrium* (as described by eqn. 6.46) and for the principle of detailed balance to apply, the downward radiative flux should equal the upward radiative flux between the two levels:

$$N_u A_{ul} + N_u B_{ul} u(v) = N_l B_{lu} u(v). (6.47)$$

From this equation we can solve for u(v) as follows:

$$u(v) = \frac{N_u A_{ul}}{N_l B_{lu} - N_u B_{ul}}. (6.48)$$

Dividing the top and bottom terms in the right-hand side of (6.48) by N_u and then using (6.46) for the ratio N_u/N_l , we are led to the expression

$$u(v) = \frac{A_{ul}}{B_{ul}} \left(\left[\frac{g_l B_{lu}}{g_u B_{ul}} \right] e^{hv_{ul}/kT} - 1 \right)^{-1}, \tag{6.49}$$

where we have used the relationship $\Delta E_{ul} = h \nu_{ul}$.

Equation (6.49) has a familiar form if we compare it to (6.32). Since both equations concern radiation in thermal equilibrium, if true then they must be equivalent. The equivalence follows if

$$\frac{g_l B_{lu}}{g_u B_{ul}} = 1$$
 or $g_l B_{lu} = g_u B_{ul}$ (6.50)

and

$$\frac{A_{ul}}{B_{ul}} = \frac{8\pi h \eta^3 v^3}{c^3}. (6.51)$$

We have thus derived the relationship between the stimulated emission and absorption coefficients B_{ul} and B_{lu} (respectively), along with their relationship to the spontaneous emission coefficient A_{ul} . We can rewrite (6.51) to obtain

$$B_{ul} = \frac{c^3}{8\pi h \eta^3 v^3} A_{ul}.$$
 (6.52)

We can now substitute the expression for A_{ul} in terms of $A_{ul}(v)$ from (4.64) to obtain

 $B_{ul} = \frac{c^3}{8\pi h \eta^3 v^3} \frac{(v - v_0)^2 + (\gamma_{ul}^T / 4\pi)^2}{\gamma_{ul}^T / 4\pi^2} A_{ul}(v).$ (6.53)

If we now define

$$B_{ul}(\nu) = \frac{\gamma_{ul}^T / 4\pi^2}{(\nu - \nu_0)^2 + (\gamma_{ul}^T / 4\pi)^2} B_{ul}, \tag{6.54}$$

which describes the frequency dependence of B_{ul} , then (6.53) may be rewritten as

$$\frac{A_{ul}(v)}{B_{ul}(v)} = \frac{8\pi h \eta^3 v^3}{c^3} \equiv \frac{A_{ul}}{B_{ul}}.$$
 (6.55)

Hence the frequency-dependent expressions for $B_{ul}(\nu)$ and $B_{lu}(\nu)$ will satisfy (6.49) if

$$\frac{g_l B_{lu}(\nu)}{g_u B_{ul}(\nu)} = 1 \quad \text{or} \quad g_l B_{lu}(\nu) = g_u B_{ul}(\nu). \tag{6.56}$$

We have thus derived the stimulated emission coefficients B_{ul} and B_{lu} , as well as their frequency-dependent counterparts $B_{ul}(\nu)$ and $B_{lu}(\nu)$, that define the way a photon beam interacts with a two-level system of atoms that was obtained by considering radiation in thermal equilibrium. These relationships provide the fundamental concepts necessary for producing a laser.

It is interesting to examine the ratio of stimulated to spontaneous emission rates from level u. This ratio can be obtained from (6.32) and (6.51) as

$$\frac{B_{ul}u(v)}{A_{ul}} = \frac{1}{e^{hv_{ul}/kT} - 1}. (6.57)$$

Thus, stimulated emission plays a significant role only for temperatures in which kT is of, or greater than, the order of the photon energy hv_{ul} . The ratio is unity when $hv_{ul}/kT = \ln 2 = 0.693$. For visible transitions in the green portion of the spectrum (photons of the order of 2.5 eV), such a relationship would be achieved for a temperature of 33,500 K. Thus, in the visible spectrum, the dominance of stimulated emission over spontaneous emission normally happens only in stars, in high-temperature and -density laboratory plasmas such as laser-produced plasmas, or in lasers. In low-pressure plasmas the radiation can readily escape, so there is no opportunity for the radiation density to build up to a value where the stimulated decay rate is comparable to the radiative decay rate. In lasers, however, the ratio of (6.57) can be significantly greater than unity.

EXAMPLE

A helium–neon laser operating at 632.8 nm has an output power of 1.0 mW with a 1-mm beam diameter. The beam passes through a mirror that has 99% reflectivity and 1% transmission at the laser wavelength. What is the ratio of $B_{ul}u(v)/A_{ul}$

for this laser? What is the effective blackbody temperature of the laser beam as it emerges from the laser output mirror? Assume the beam diameter is also 1 mm inside the laser cavity and that the power is uniform over the beam cross section (this is only an approximation, as we will learn in Chapter 11). Assume also that the laser linewidth is approximately one tenth of the Doppler width for the transition.

The laser frequency is determined by

$$v = \frac{c}{\lambda} = \frac{3 \times 10^8 \text{ m/s}}{6.328 \times 10^{-7} \text{ m}} = 4.74 \times 10^{14} \text{ Hz}.$$

From (6.51) we can now compute the ratio of A_{ul}/B_{ul} for $\eta=1$ as

$$\frac{A_{ul}}{B_{ul}} = \frac{8\pi h v^3}{c^3} = \frac{(8\pi)(6.63 \times 10^{-34} \text{ J-s})(4.74 \times 10^{14} \text{ Hz})^3}{(3 \times 10^8 \text{ m/s})^3}$$
$$= 6.57 \times 10^{-14} \text{ J-s/m}^3;$$

hence

$$\frac{B_{ul}}{A_{ul}} = 1.52 \times 10^{13} \text{ m}^3/\text{J-s}.$$

We must now compute the energy density u(v), which from (6.35) is related to the intensity per unit frequency I(v) as u(v) = I(v)/c. We can compute I(v) by dividing the laser beam power in the cavity by the beam cross-sectional area and the frequency width of the beam. The power of the beam within the cavity traveling toward the output mirror must be 100 mW and that reflected would be 99 mW (1 mW passes through the mirror). Thus, the total power in the cavity is 199 mW. The Doppler width of the helium–neon 632.8-nm transition (see Table 4-1) is 1.5×10^9 Hz. Thus

$$u(v) = \frac{I(v)}{c}$$

$$= \frac{[(199 \times 1.0 \text{ mW})/(\pi \cdot (5 \times 10^{-4} \text{ m})^2)]/(0.1)(1.5 \times 10^9 \text{ Hz})}{3 \times 10^8 \text{ m/s}}$$

$$= 5.63 \times 10^{-12} \text{ J-s/m}^3,$$

so the ratio is

$$\frac{B_{ul}u(v)}{A_{ul}} = (1.52 \times 10^{13} \text{ m}^3/\text{J-s})(5.63 \times 10^{-12} \text{ J-s/m}^3) = 85.6.$$

The stimulated emission rate is therefore almost 86 times the spontaneous emission rate on transitions from the upper to the lower laser level at 632.8 nm.

Using (6.57), the preceding ratio can be rewritten as

$$\frac{B_{ul}u(v)}{A_{ul}} = \frac{1}{e^{hv_{ul}/kT} - 1} = 85.6,$$

which yields

$$e^{h\nu_{ul}/kT} - 1 = 1/85.6 = 0.0117$$

or

$$h\nu_{ul}/kT = \ln(1.0117) = 1.16 \times 10^{-2}$$
.

We can thus solve for T as follows:

$$T = \frac{hv_{ul}}{(1.16 \times 10^{-2})k} = \frac{(6.63 \times 10^{-34} \text{ J-s})(4.74 \times 10^{14} \text{ Hz})}{(1.16 \times 10^{-2})(1.38 \times 10^{-23} \text{ J/K})}$$
$$= 1.96 \times 10^6 \text{ K} = 1,960,000 \text{ K}.$$

This calculation indicates that the radiation intensity of the laser beam inside the laser cavity has a value equivalent to that of a nearly 2,000,000-K blackbody – if we consider only the radiation emitted from the blackbody in the frequency (or wavelength interval) over which the laser operates.

REFERENCES

A. Corney (1977), Atomic and Laser Spectroscopy. Oxford: Clarendon Press, Chapter 9.

R. B. Leighton (1959), Principles of Modern Physics. New York: McGraw-Hill, Chapter 2.

R. Loudon (1973), The Quantum Theory of Light. Oxford: Clarendon Press, Chapter 1.

PROBLEMS

- 1. Calculate the number of radiation modes in a cube 1 mm on a side for a spread of 0.001 nm centered at 514.5 nm and a spread of 0.01 μ m centered at 10.6 μ m.
- 2. Consider a 1-mm—diameter surface area of carbon (graphite). Calculate how many atoms would exist in energy levels from which they could emit radiation at wavelengths shorter than 700 nm (visible light and shorter wavelengths) for surface temperatures of 300 K, 1,000 K, and 5,000 K when the solid is in thermal equilibrium at those temperatures. Assume that only those atoms within a depth of 10 nm of the material surface can emit observable radiation.
- 3. In Problem 2, if the excited atoms that emit visible radiation decay in 10^{-13} s and if only 0.002% of them decay radiatively (quantum yield of 2%), how much power would be radiated from that surface at the aforementioned temperatures? Assume that half of the atoms that radiate emit into the 2π solid angle that would result in their leaving the surface of the material, and assume an average visible photon energy of 2.5 eV. Also compute the total amount of power that could be radiated (over all wavelengths) from the surface at the given temperatures using the Stefan–Boltzmann law (eqn. 6.15). Speculate as to why the two approaches for computing the radiated power are inconsistent at a temperature of 5,000 K.
- **4.** How much power is radiated from a 1-mm² surface of a body at temperature T when the peak measured wavelength is that of green light at 500 nm?
- 5. Determine the number of modes in a 1-cm³ box for frequencies in the visible spectrum between 400 and 700 nm. Compare that value to the number of modes in a

- sodium streetlamp that emits over a wavelength interval of 3 nm at a center wavelength of 589 nm. Assume that the streetlamp is a cylinder of radius 0.5 cm and length 10 cm.
- **6.** Estimate the number of photons in both the box and the streetlamp of Problem 5 for temperatures of 300 K, 1,000 K, and 5,000 K.
- 7. A 100-W incandescent lamp has a tungsten filament composed of a wire, 0.05 cm in diameter and 10 cm in length, that is coiled up to fit within the light bulb. Assume the filament is heated to a temperature of 3,000 K when the light bulb is turned on. How much power (watts) is emitted within the visible spectrum from the filament, assuming that it is emitting as a blackbody? As an approximation, you could divide the visible spectral region into several segments and compute the average contribution from each segment. Then simply add the averages together (instead of trying to integrate the blackbody function over the entire visible spectral range).
- **8.** Show that Planck's radiation law of (6.39) will lead to the Stefan–Boltzmann relationship of (6.15) if the power radiated over all wavelengths is considered. Determine the coefficients of the Stefan–Boltzmann constant. *Hint*:

$$\int_0^\infty \frac{x^3 \, dx}{e^x - 1} = \frac{\pi^4}{15}.$$

- 9. An argon ion laser emits 2 W of power at 488.0 nm in a 2-mm-diameter beam. What would be the effective blackbody temperature of the output beam of that laser radiating over the frequency width of the laser transition, given that the laser linewidth is approximately one fifth of the Doppler linewidth? Assume that the laser is operating at an argon gas temperature of 1,500 K and that the laser output is uniform over the width of the beam.
- 10. For the laser in Problem 9, how much power would be required for the stimulated emission rate to equal the radiative decay rate?
- 11. A pulsed and Q-switched Nd:YAG laser is focused to a 200-μm-diameter spot size on a solid flat metal target in an attempt to produce a bright plasma source for microlithography applications. The plasma is observed to radiate uniformly into 2π steradians with a wavelength distribution that is approximately that of a blackbody. The intensity of the Nd:YAG laser is adjusted so that the peak of the blackbody emission occurs at a wavelength of 13.5 nm, the optimum wavelength as a source for EUV microlithography. What is the temperature of the blackbody? What fraction of the total energy radiated by the blackbody would be radiated within the useful bandwidth for microlithography of 0.4 nm centered around the emission maximum? Assume that the plasma doesn't expand significantly during the 10-ns emission time (the duration of the Nd:YAG laser pulses).
- 12. The blackbody spectral distribution curve has a maximum wavelength λ_m that is dependent upon the temperature T of the radiating body. Show that the product $\lambda_m T$ is a constant for any temperature (Wien's law, eqn. 6.16). *Hint*: Use the frequency version of the blackbody radiation formula instead of the wavelength version to show this.
- 13. A blue argon laser beam at 488 nm is propagating around a coliseum as part of a laser light show. The power is measured to be 10 W cw at a specific location with a beam diameter of 5 mm. What is the energy density per unit frequency u(v) of the beam at that location?

15.2 SOLID-STATE LASERS 565

are self-selected by the beam. The use of a short laser pulse allows heat to destroy the bad cells without the heat spreading to the surrounding cells. In the case of tattoo removal, the tattoo itself is absorbing and hence requires no tagging. Other applications include pollution detection and removal of kidney stones.

TITANIUM SAPPHIRE LASER

General Description

The titanium sapphire laser (Ti:Al₂O₃) is the most widely used tunable solid-state laser. It can be operated over a wavelength range of 660-1,180 nm and thus has the broadest gain bandwidth of any laser. It also has a relatively large stimulated emission cross section for a tunable laser. Titanium ions are typically doped into a sapphire (aluminum oxide) crystal at a concentration of 0.1% by weight. This laser has achieved cw outputs of nearly 50 W as well as terawatts of peak power from 100-fs-duration mode-locked pulses. The laser can be flashlamp pumped, but the technique is not efficient owing to the unusually short upper laser level lifetime of the laser crystal (3.8 μ s at room temperature), which does not match well with the longer pulse duration of typical flashlamps. Commercial titanium sapphire lasers are therefore typically pumped with either argon ion lasers (for cw operation) or frequency-doubled Nd:YAG or Nd:YLF lasers (for pulsed operation). The pump absorption band covers the range from less than 400 nm to just beyond 630 nm and peaks around 490 nm, as can be seen in Figure 15-13. The titanium sapphire crystal has high thermal conductivity, good chemical inertness, good mechanical rigidity, and high hardness.

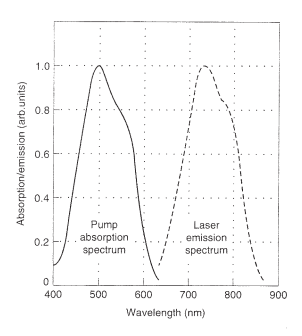
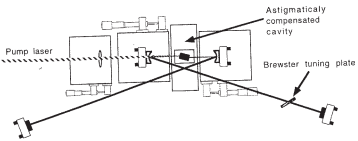


Figure 15-13 Absorption and emission spectra of titanium:sapphire laser amplifier rod



(a) cw Ti:sapphire - X fold configuration

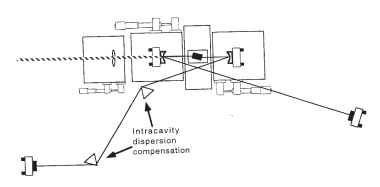


Figure 15-14 Diagrams of (a) a cw titanium sapphire laser and (b) a femtosecond mode-locked titanium sapphire laser (courtesy of CREOL)

(b) Femtosecond mode-locked Ti:sapphire

Laser Structure

An example of a cw Ti:Al₂O₃ laser using an X-cavity design is shown in Figure 15-14(a). It uses an astigmatically compensated cavity for the $Ti:Al_2O_3$ laser crystal (see Section 13.9). In such a cavity design, the crystal typically ranges from 2 to 10 mm in length, depending upon the dopant level, and is arranged with the output faces of the crystal at Brewster's angle. The longer-length crystal lengths with lower doping concentrations are used with higher pumping flux intensities in order to obtain higher power output. Generally, either a cw argon ion laser or a doubled Nd:YAG laser is used as the pumping source. The pump beam enters the cavity from the left, as shown in the figure. A birefringent filter, installed within the cavity at Brewster's angle, can be rotated for wavelength tuning. A modified version of this cavity, shown in Figure 15-14(b), is used to produce mode-locked pulses. It includes two prisms for intracavity dispersion compensation and uses the Kerr lens mode-locking (KLM) technique described in Section 13.4. The necessary aperture within the Ti:Al₂O₃ crystal to produce KLM is provided by a separate aperture located next to the crystal (not shown), or simply by the aperturing effect associated with the small diameter of the pump beam. Extremely precise adjustments and alignment of the mirrors and cavity dimensions are essential to maintain a stable mode-locked output for this laser.

Excitation Mechanism

The energy-level structure of this laser is similar to that of a dye laser. The ground state, a 2T_2 state, has a broad sequence of overlapping vibrational or vibronic levels extending upward from the lowest level, as shown in Figure 5-16. The first excited state is a 2E state that also extends upward with a series of overlapping vibronic levels. This energy-level structure is unique for laser crystals in that there are no d-state energy levels above the upper laser level. Thus, the simple energy-level structure involving a 3d electron eliminates the possibility of excited-state absorption, an effect that reduces the tuning range of other tunable solid-state lasers (see Section 9.5).

Excitation therefore occurs from the lowest vibronic levels of the 2T_2 ground state (those that are sufficiently populated at room temperature) to the broad range of excited vibronic levels of the 2E excited state. The population pumped to all of the vibrational levels of the broadband excited state rapidly relaxes to the lowest levels of that state. It then decays back to any one of the vibronic levels of the ground state in a manner that is similar to a dye laser, but with a much lower radiative rate. When the population reaches the excited vibronic levels of the ground state, it very rapidly relaxes to the lowest-lying levels, leaving a distribution dictated by the Boltzmann relationship of (6.11).

Laser wavelengths (λ_{ul})	660–1,180 nm
Laser transition probability (A_{ul})	2.6×10^{5} /s
Upper laser level lifetime (τ_u)	$3.8~\mu s$
Stimulated emission cross section (σ_{ul})	$3.4 \times 10^{-23} \text{ m}^2$
Spontaneous emission linewidth and	
gain bandwidth, FWHM ($\Delta \nu_{ul}$)	1.0×10^{14} /s ($\Delta \lambda_{ul} = 180 \text{ nm}$)
Inversion density (ΔN_{ul})	$6 \times 10^{23} / \text{m}^3$
Small-signal gain coefficient (g_0)	20/m
Laser gain-medium length (L)	0.1 m
Single-pass gain $(e^{\sigma_{ul}\Delta N_{ul}L})$	7–10
Doping density	$3.3 \times 10^{25} / \text{m}^3$
Index of refraction of gain medium	1.76
Operating temperature	300 K
Thermal conductivity of laser rod	3.55 W/m-K
Thermal expansion coefficient of laser rod	$5 \times 10^{-6} / \text{K}$
Pumping method	optical (flashlamp or laser)
Pumping bands	380-620 nm
Output power	up to 50 W (cw),
• •	10^{12} W for 100-fs pulse
Mode	single-mode or multi-mode

The laser energy-level arrangement is effectively a four-level system, as in a dye laser, in which all of the higher-lying vibronic levels of the ${}^{2}E$ state serve as level i of the four-level system described in Section 9.3. The lowest vibrational levels of the ${}^{2}E$ state serve as the upper laser level u. These levels decay to any of the excited vibrational levels of the ground state ${}^{2}T_{2}$, any of which can be considered as the lower laser level l. These levels then rapidly relax to the lowest levels of ${}^{2}T_{2}$ serving as the ground state 0.

Applications

Titanium sapphire lasers are used in infrared spectroscopy of semiconductors and in laser radar, rangefinders, and remote sensing. They are used in medical applications such as photodynamic therapy. They are also used to produce short pulses of X-rays by focusing the mode-locked pulses onto solid targets from which high-density and high-temperature radiating plasmas are produced, plasmas that in turn emit large fluxes of X-rays.

CHROMIUM LISAF AND LICAF LASERS

General Description

The chromium-doped lithium strontium aluminum fluoride (Cr:LiSAF) and lithium calcium aluminum fluoride (Cr:LiCAF) lasers are broadband tunable lasers in the same category as the alexandrite and titanium sapphire lasers. The Cr:LiSAF laser can be tuned over a wavelength ranging from 780 to 1,010 nm and the Cr:LiCAF laser can be tuned from 720 to 840 nm. Both cw and pulsed output have been obtained from both lasers: a cw output of up to 1.2 W and a pulsed output of over 10 J with a slope efficiency of 5%. These lasers have relatively long upper-level lifetimes of 67 μ s for Cr:LiSAF and 170 μ s for Cr:LiCAF, so both can be effectively flashlamp pumped. They have also been laser pumped with AlGaAs diode lasers and argon ion lasers. Both of these fluoride laser materials can be doped to very high concentrations (up to 15% Cr) without affecting the upper laser level lifetime, which results in more uniform flashlamp pumping as indicated in Figure 10-19(c). These lasers have also been used as regenerative amplifiers leading to very short-pulse amplification. The laser crystals are chemically stable when treated properly. Their thermal properties are closer to those of Nd:glass than to Nd:YAG. They are durable but not as hard as YAG. Their average power-handling capabilities are not as good as Ti:Al₂O₃.

Laser Structure

A diagram of a flashlamp-pumped Cr:LiSAF laser is shown in Figure 15-15. This laser uses a LiSAF rod of 6-mm diameter and 0.1-m length. A birefringent filter is used to tune the wavelength from 750 to 1,000 nm, and a saturable absorber is