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Preface

The three volumes VIII/1A, B, C document the state of the art of "Laser Physics and Applications". Scientific trends and related technological aspects are considered by compiling results and conclusions from phenomenology, observation and experience. Reliable data, physical fundamentals and detailed references are presented.

In the recent decades the laser source matured to a universal tool common to scientific research as well as to industrial use. Today a technical goal is the generation of optical power towards shorter wavelengths, shorter pulses and higher power for application in science and industry. Tailoring the optical energy in wavelength, space and time is a requirement for the investigation of laser-induced processes, i.e. excitation, non-linear amplification, storage of optical energy, etc. According to the actual trends in laser research and development, Vol. VIII/1 is split into three parts: Vol. VIII/1A with its two subvolumes 1A1 and 1A2 covers laser fundamentals, Vol. VIII/1B deals with laser systems and Vol. VIII/1C gives an overview on laser applications.

In Vol. VIII/1A1 the following topics are treated in detail:

Part 1: Fundamentals of light-matter interaction

This part compiles the basic elements of classical electromagnetic wave theory, non-relativistic quantum mechanics of the two-level system and its interaction with the non-quantized radiation field. The relevant relations with their approximations and range of validity are discussed. It starts with Maxwell's equations, wave equation and SVE-approximations, presents the Schrödinger equations, the field/atom interaction including the Einstein coefficients and cross-sections. The main parameters characterizing the two-level system with typical numbers are given in several tables. Finally, the coherent interaction is briefly discussed. This semiclassical approach is sufficient for most applications in laser technology. The fully quantized theory is offered in Vol. VIII/1A2, Chap. 5.

Part 2: Radiometry

In the first section the definitions of the radiometric quantities and their measurement are summarized. In the second part the main elements of laser beam characterization are compiled with a detailed discussion of the theoretical background. The experimental determination of the essential quantities according to the ISO-normalizations is given.

Part 3: Linear optics

The design of optical resonators and beam handling requires a broad knowledge in optics. In this part the fundamentals of beam propagation, Gaussian beams, diffraction, refraction, lens design and crystal optics are presented. The extensive references give access to detailed information.

VIII Preface

Part 4: Nonlinear optics

Nonlinear effects are widely used in laser technology to generate new wavelengths or to improve beam quality. In four sections the essential nonlinear optical effects are discussed: frequency conversion in crystals, frequency conversion in gases and liquids, stimulated scattering and phase conjugation. In extensive tables the coefficients of the nonlinear processes are compiled.

August 2005 The Editors

Contents

Part 1 Fu	Part 1 Fundamentals of light-matter interaction		
1.1	Fundamentals of the semiclassical laser theory V.A. LOPOTA, H. WEBER	3	
1.1.1	The laser oscillator	3	
1.1.2 1.1.2.1 1.1.2.2 1.1.2.2.1 1.1.2.2.2 1.1.2.2.3 1.1.2.2.4 1.1.2.3	The electromagnetic field . Maxwell's equations . Homogeneous, isotropic, linear dielectrics . The plane wave . The spherical wave . The slowly varying envelope (SVE) approximation . The SVE-approximation for diffraction . Propagation in doped media .	5 5 6 7 8 9 9	
1.1.3 1.1.3.1 1.1.3.2 1.1.3.2.1 1.1.3.2.2 1.1.3.3 1.1.3.3.1 1.1.3.3.1.1 1.1.3.3.1.2 1.1.3.3.1.3 1.1.3.3.2	The two-level system The dipole approximation Inversion density and polarization The interaction with a monochromatic field The Maxwell–Bloch equations Decay time T_1 of the upper level (energy relaxation) Spontaneous emission Interaction with the host material Pumping process	11 11 12 12 14 15 15 15 16 16	
1.1.4 1.1.4.1 1.1.4.2 1.1.4.3	Inversion density and polarization	17 17 19 19	
1.1.5 1.1.5.1 1.1.5.2 1.1.5.3 1.1.5.3.1 1.1.5.3.2 1.1.5.4 1.1.5.5	Adiabatic equations Rate equations Thermodynamic considerations Pumping schemes and complete rate equations The three-level system The four-level system Adiabatic pulse amplification Rate equations for steady-state laser oscillators.	20 20 21 22 23 24 25 26	
1.1.6 1.1.6.1 1.1.6.1.1 1.1.6.1.2	Line shape and line broadening Normalized shape functions Lorentzian line shape Gaussian line shape		

X Contents

1.1.6.1.3 1.1.6.2 1.1.6.2.1 1.1.6.2.2 1.1.6.2.3 1.1.6.2.4 1.1.6.3 1.1.6.3.1 1.1.6.3.2 1.1.6.4	Normalization of line shapes Mechanisms of line broadening Spontaneous emission Doppler broadening. Collision or pressure broadening Saturation broadening Types of broadening Homogeneous broadening. Inhomogeneous broadening Time constants	28 28 28 29 29 29 30
1.1.0.4 1.1.7 1.1.7.1 1.1.7.2 1.1.7.3 1.1.7.3.1 1.1.7.3.1.1 1.1.7.3.1.2 1.1.7.3.2 1.1.8	Coherent interaction The Feynman representation of interaction Constant local electric field Propagation of resonant coherent pulses Steady-state propagation of $n\pi$ -pulses 2π -pulse in a loss-free medium π -pulse in an amplifying medium Superradiance Notations References for 1.1	31 32 33 34 35 36 37 37
Part 2 Ra	adiometry	
2.1	Definition and measurement of radiometric quantities B. Wende, J. Fischer	45
2.1.1	Introduction	45
2.1.2	Definition of radiometric quantities	45
2.1.2 2.1.3 2.1.3.1 2.1.3.2	Definition of radiometric quantities Radiometric standards Primary standards Secondary standards	47 47
2.1.3 2.1.3.1	Radiometric standards	47 47 48
2.1.3 2.1.3.1 2.1.3.2	Radiometric standards	47 47 48 50
2.1.3 2.1.3.1 2.1.3.2	Radiometric standards Primary standards Secondary standards Outlook – State of the art and trends	47 47 48 50 51
2.1.3 2.1.3.1 2.1.3.2 2.1.4	Radiometric standards Primary standards Secondary standards Outlook – State of the art and trends References for 2.1 Beam characterization B. Eppich	477 478 488 500 513
2.1.3 2.1.3.1 2.1.3.2 2.1.4 2.2	Radiometric standards Primary standards Secondary standards Outlook – State of the art and trends References for 2.1 Beam characterization B. Eppich Introduction	47 47 48 50 51 53 55
2.1.3 2.1.3.1 2.1.3.2 2.1.4 2.2 2.2.1 2.2.2	Radiometric standards Primary standards Secondary standards Outlook – State of the art and trends References for 2.1 Beam characterization B. Eppich Introduction The Wigner distribution	47 47 48 50 51 53 53
2.1.3 2.1.3.1 2.1.3.2 2.1.4 2.2 2.2.1 2.2.2 2.2.3	Radiometric standards Primary standards Secondary standards Outlook – State of the art and trends References for 2.1 Beam characterization B. EPPICH Introduction The Wigner distribution The second-order moments of the Wigner distribution	47 47 48 50 51 53 53 55 55
2.1.3 2.1.3.1 2.1.3.2 2.1.4 2.2 2.2.1 2.2.2	Radiometric standards Primary standards Secondary standards Outlook – State of the art and trends References for 2.1 Beam characterization B. Eppich Introduction The Wigner distribution	477 447 488 500 511 533 535 545 546 546 546 546 546 546 546 546 54

Contents	XI

2.2.5.3 2.2.5.4 2.2.5.5	General astigmatic beams Pseudo-symmetric beams Intrinsic astigmatism and beam conversion	64
2.2.6	Measurement procedures	
2.2.7	Beam positional stability	
2.2.7.1 2.2.7.2 2.2.7.3	Absolute fluctuations	67 69
	References for 2.2	70
Part 3 Li	near optics	
3.1	Linear optics	
0.1	R. GÜTHER	73
3.1.1	Wave equations	73
3.1.2	Polarization	75
3.1.3	Solutions of the wave equation in free space	78
3.1.3.1	Wave equation	78
3.1.3.1.1 3.1.3.1.2	Monochromatic plane wave	
3.1.3.1.3	Spherical vector wave	
3.1.3.2	Helmholtz equation	79
3.1.3.2.1 3.1.3.2.2	Plane wave	
3.1.3.2.3	Cylindrical wave	
3.1.3.2.4	Diffraction-free beams	79
3.1.3.2.4.1 3.1.3.2.4.2	Diffraction-free Bessel beams	
3.1.3.2.4.2	Vectorial Bessel beams	
3.1.3.3	Solutions of the slowly varying envelope equation	
3.1.3.3.1	Gauss-Hermite beams (rectangular symmetry)	
3.1.3.3.2 3.1.3.3.3	Gauss-Laguerre beams (circular symmetry)	
3.1.4	Diffraction	
3.1.4.1	Vector theory of diffraction	
3.1.4.2	Scalar diffraction theory	
3.1.4.3 3.1.4.4	Time-dependent diffraction theory	
3.1.4.4.1	Fraunhofer diffraction patterns	
3.1.4.4.2	Circular aperture with radius a	90
3.1.4.4.2.1	Applications	92
3.1.4.4.3 3.1.4.5	Gratings	92 93
3.1.4.5.1	Fresnel's diffraction on a slit	93
3.1.4.5.2	Fresnel's diffraction through lens systems (paraxial diffraction)	94
3.1.4.6	Fourier optics and diffractive optics	94
3.1.5	±	
3.1.5.1 3.1.5.2	Dielectric media	96 97

XII Contents

3.1.5.3	Dispersion characteristics for short-pulse propagation	. 97
3.1.5.4	Optics of metals and semiconductors	. 98
3.1.5.5	Fresnel's formulae	. 98
3.1.5.6	Special cases of refraction	. 101
3.1.5.6.1	Two dielectric isotropic homogeneous media (\hat{n} and \hat{n}' are real)	
3.1.5.6.2	Variation of the angle of incidence	
3.1.5.6.2.1	External reflection $(n < n')$	
3.1.5.6.2.2	Internal reflection $(n < n')$	
3.1.5.6.3	Reflection at media with complex refractive index	. 101
3.1.3.0.3	(Case $\hat{n} = 1$ and $\hat{n}' = n' + i k'$)	109
3.1.5.7	Crystal optics	
	v i	
3.1.5.7.1	Classification	
3.1.5.7.2	Birefringence (example: uniaxial crystals)	
3.1.5.8	Photonic crystals	
3.1.5.9	Negative-refractive-index materials	
3.1.5.10	References to data of linear optics	. 108
3.1.6	Geometrical optics.	. 108
3.1.6.1	Gaussian imaging (paraxial range)	
3.1.6.1.1	Single spherical interface	
3.1.6.1.2	Imaging with a thick lens.	
3.1.6.2	Gaussian matrix (ABCD-matrix, ray-transfer matrix) formalism for paraxial	. 110
5.1.0.2	optics	111
3.1.6.2.1	Simple interfaces and optical elements with rotational symmetry	
-		
3.1.6.2.2	Non-symmetrical optical systems	
3.1.6.2.3	Properties of a system	
3.1.6.2.4	General parabolic systems without rotational symmetry	
3.1.6.2.5	General astigmatic system	
3.1.6.2.6	Symplectic optical system	
3.1.6.2.7	Misalignments	
3.1.6.3	Lens aberrations	. 117
3.1.7	Beam propagation in optical systems	. 120
3.1.7.1	Beam classification	
3.1.7.2	Gaussian beam: complex q-parameter and its ABCD-transformation	
3.1.7.2.1	Stigmatic and simple astigmatic beams	
3.1.7.2.1.1	Fundamental Mode	
3.1.7.2.1.2	Higher-order Hermite-Gaussian beams in simple astigmatic beams	
3.1.7.2.2	General astigmatic beam	
3.1.7.3	Waist transformation	
3.1.7.3.1	General system (fundamental mode)	
	Thin lens (fundamental mode)	
3.1.7.3.2		
3.1.7.4	Collins integral	
3.1.7.4.1	Two-dimensional propagation	
3.1.7.4.2	Three-dimensional propagation	. 127
3.1.7.5	Gaussian beams in optical systems with stops, aberrations, and waveguide	
	coupling	. 127
3.1.7.5.1	Field distributions in the waist region of Gaussian beams including stops and	
	wave aberrations by optical system	
3.1.7.5.2	Mode matching for beam coupling into waveguides	
3.1.7.5.3	Free-space coupling of Gaussian modes	
3.1.7.5.4	Laser fiber coupling	. 129
	References for 3.1	. 131

Part 4 Nonlinear optics		
4.1	Frequency conversion in crystals G.G. Gurzadyan	. 141
4.1.1 4.1.1.1 4.1.1.1.2 4.1.1.3 4.1.1.2	Introduction	. 141 . 141 . 142 . 142
4.1.2 4.1.2.1 4.1.2.2 4.1.2.3 4.1.2.4 4.1.2.5 4.1.2.5.1 4.1.2.5.2 4.1.2.5.3	Fundamentals Three-wave interactions Uniaxial crystals Biaxial crystals Effective nonlinearity Frequency conversion efficiency General approach Plane-wave fixed-field approximation SHG in "nonlinear regime" (fundamental wave depletion)	. 144 . 145 . 145 . 155 . 155 . 154
4.1.3	Selection of data	. 154
4.1.4	Harmonic generation (second, third, fourth, fifth, and sixth)	. 156
4.1.5	Sum frequency generation	. 167
4.1.6	Difference frequency generation	. 172
4.1.7	Optical parametric oscillation	. 176
4.1.8	Picosecond continuum generation	. 186
	References for 4.1	
4.2	Frequency conversion in gases and liquids C.R. Vidal	201
4.2.1 4.2.1.1 4.2.1.2 4.2.1.3 4.2.1.4 4.2.1.5	Fundamentals of nonlinear optics in gases and liquids Linear and nonlinear susceptibilities Third-order nonlinear susceptibilities Fundamental equations of nonlinear optics Small-signal limit Phase-matching condition	. 205 . 205 . 206 . 207
4.2.2 4.2.2.1 4.2.2.2 4.2.2.3	Frequency conversion in gases Metal-vapor inert gas mixtures Mixtures of different metal vapors Mixtures of gaseous media References for 4.2	. 209 . 209 . 209
4.3	Stimulated scattering A. LAUBEREAU	
4.3.1 4.3.1.1 4.3.1.2	Introduction	. 217

XIV Contents

4.3.2	General properties of stimulated scattering	219
4.3.2.1	Exponential gain by stimulated Stokes scattering	
4.3.2.2	Experimental observation	220
4.3.2.2.1	Generator setup	220
4.3.2.2.2	Oscillator setup	220
4.3.2.2.3	Stimulated amplification setup	221
4.3.2.3	Four-wave interactions	221
4.3.2.3.1	Third-order nonlinear susceptibility	221
4.3.2.3.2	Stokes—anti-Stokes coupling	222
4.3.2.3.3	Higher-order Stokes and anti-Stokes emission	
4.3.2.4	Transient stimulated scattering	222
4.3.3	Individual scattering processes	223
4.3.3.1	Stimulated Raman scattering (SRS)	223
4.3.3.2	Stimulated Brillouin scattering (SBS) and stimulated thermal Brillouin	
	scattering (STBS)	227
4.3.3.3	Stimulated Rayleigh scattering processes, SRLS, STRS, and SRWS	
	References for 4.3	232
4.4	Phase conjugation	
	H.J. EICHLER, A. HERMERSCHMIDT, O. MEHL	235
4.4.1	Introduction	235
4.4.2	Basic mathematical description	236
4.4.3	Phase conjugation by degenerate four-wave mixing	236
4.4.4	Self-pumped phase conjugation	237
4.4.5	Applications of SBS phase conjugation	240
4.4.6	Photorefraction	242
	References for 4.4	245
Index		247

1.1 Fundamentals of the semiclassical laser theory

V.A. LOPOTA, H. WEBER

A rigorous description of light–matter interaction requires a fully quantized system of field equations, which is the content of quantum optics [70Hak, 95Wal, 97Scu, 95Man, 01Vog]. This theory is well developed and the results are confirmed perfectly by many experiments (see Chap. 5.1). But most problems of laser design and laser technology can be solved in a satisfactory way by applying the semiclassical theory. This means a non-relativistic quantum-mechanical approach for the electronic system and a non-quantized, classical electromagnetic field.

Non-relativistic means that the velocity of the interacting electrons is small compared with the velocity of light. This holds for the outer shell electrons of the atoms and molecules, which are relevant in laser physics. It is not true for the free-electron laser and for the interaction of strong fields with plasmas, which demand a relativistic treatment.

A non-quantized electromagnetic field implies that the photon is neglected. In laser technology the photon flux in most applications is extremely high and the granulation of light beams is of no importance. It is of significance for metrology, where the lower limit of detectability is partly given by photon statistics. There are some other effects, which are not covered by the semiclassical theory:

- Planck's law, related to photon statistics,
- squeezed states,
- entangled photons,
- zero-point energy effects,
- spontaneous emission,

and some spectral line shifts (Lamb-shift [47Lam]), of minor importance for laser technology, although of great experimental interest for the confirmation of the fundamental theory. The spontaneous emission of excited atoms/molecules is responsible for the lower limit of laser line width [74Sar, 95Man] and for the on-set of laser oscillation. Therefore, spontaneous emission has to be included in the semiclassical theory by a phenomenological term as shown in Fig. 1.1.1.

It is the intention of this chapter to compile the relevant relations of laser dynamics, their application in laser design and to discuss the limitations and approximations. The mathematical derivations can be taken from the references.

1.1.1 The laser oscillator

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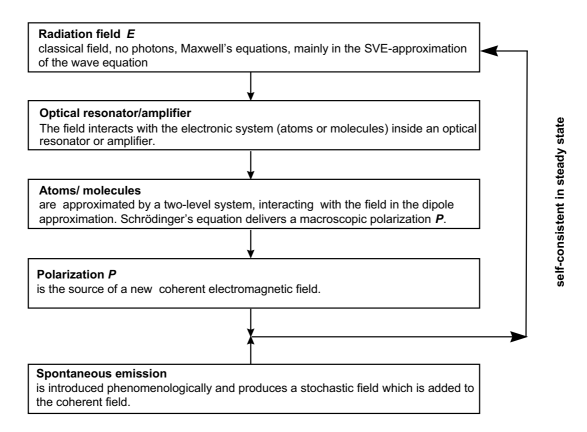


Fig. 1.1.1. The semiclassical laser theory (SVE-approximation: Slowly Varying Envelope approximation, see Sect. 1.1.2.2.3).

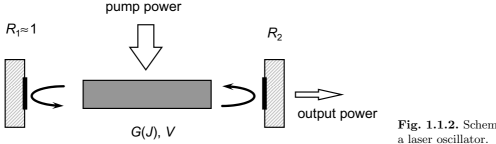


Fig. 1.1.2. Schematic set-up of

The principle set-up of a laser oscillator is plotted in Fig. 1.1.2. Light is amplified by induced emission in an active medium (gas discharge, doped crystals or liquids, pn-transitions). The active medium is characterized by an intensity- and frequency-dependent gain factor G(J) (with J: intensity). The beam bounces forth and back between the two mirrors of an optical resonator. On-set of laser oscillation requires a gain factor exceeding the total losses per round trip:

$$G_0RV > 1$$
 (threshold condition) (1.1.1)

with

 G_0 : small-signal gain factor for the intensities,

 $R = \sqrt{R_1 R_2}$: average reflection factor of the mirrors,

V: internal loss factor of the resonator.

With increasing intensity J the gain decreases due to saturation of the amplifier

$$G(J) \leq G_0$$
.

In steady state the gain has to compensate the losses:

$$G(J)RV = 1$$
 (steady-state condition). (1.1.2)

If the relation G(J) is known, depending on the specific amplifier, (1.1.2) gives the internal intensity of the laser system in steady state.

The wavelength of the field is determined by the resonance condition. After one round trip the phase shift $\Delta \varphi$ of the field must be

$$\Delta \varphi = 2\pi p$$
, $p = 1, 2, 3, \dots$ (resonance condition), (1.1.3)

otherwise the field would be reduced by destructive interference. The resonator is mainly responsible for the mode structure of the output field and can be described by a non-quantized field. Details are given in Chap. 8.1. For the interaction field–amplifier a plane wave is assumed and diffraction is neglected.

1.1.2 The electromagnetic field

1.1.2.1 Maxwell's equations

י אין איני און איני אין איני איני און איני אינון איני אינון איני אינון איני אינון איני אינון איני אינון איני א איני איני איני אינון אינון אינון אינון איני איני איני איני אינון אינון אינון אינון אינון אינון איני איני איני איני אינון איני אינון אינו

The classical electromagnetic field is completely described by Maxwell's equations:

$$\operatorname{curl} \mathbf{E} = -\frac{\partial \mathbf{B}}{\partial t} , \qquad (1.1.4)$$

$$\operatorname{curl} \boldsymbol{H} = \frac{\partial \boldsymbol{D}}{\partial t} + \boldsymbol{j} , \qquad (1.1.5)$$

$$\operatorname{div} \mathbf{D} = \rho \,, \tag{1.1.6}$$

$$\operatorname{div} \boldsymbol{B} = 0 \tag{1.1.7}$$

with

E: electric field (SI-unit: V/m),

H: magnetic field (SI-unit: A/m),

D: electric displacement (SI-unit: As/m²),

B: magnetic induction (SI-unit: Vs/m^2),

j: current density (SI-unit: A/m²),

 ρ : density of electric charges (SI-unit: As/m³).

For all quantities the complex notation is used [99Bor], the real quantities are $Q_{\text{real}} = \frac{1}{2}(Q + Q^*)$. The relations between D, E and B, H are given by the material equations. Under the action of an external electric/magnetic field atomic or molecular electric/magnetic dipoles are generated in matter. The dipole moment per unit volume is called the electric or magnetic polarization P(E, H) or J(E, H), respectively. The resulting material quantities are the electric displacement D and the magnetic induction B given as:

$$D = \varepsilon_0 E + P(E, H) = \varepsilon_0 \varepsilon(E, H) \cdot E , \qquad (1.1.8)$$

$$B = \mu_0 H + J(E, H) = \mu_0 \mu(E, H) \cdot H \tag{1.1.9}$$

with

 $P = \varepsilon_0 \chi_e(E, H) E$: electric polarization (SI-unit: As/m²), $J = \mu_0 \chi_m(E, H) H$: magnetic polarization (SI-unit: Vs/m²),

 $\chi_{\rm e}(\boldsymbol{E},\boldsymbol{H}), \, \chi_{\rm m}(\boldsymbol{E},\boldsymbol{H})$: electric/magnetic susceptibility, in general a tensor and a function of the fields,

 $\varepsilon = 1 + \chi_e$, $\mu = 1 + \chi_m$: permittivity/permeability number, in general tensors, 1: unit tensor,

 $\varepsilon_0 = 8.8542 \times 10^{-12}$ As/Vm: electric constant, $\mu_0 = 4\pi \times 10^{-7}$ Vs/Am: magnetic constant.

The current inside a medium is caused by the electric field and Ohm's law holds

$$j = \sigma_{\rm e} E \tag{1.1.10}$$

with

 $\sigma_{\rm e}$: electric conductivity, in general a tensor and function of the field, (SI-unit: A/Vm).

Electric and magnetic polarization depend in general on both generating fields, E and H. In many cases this relation is linear, but quite often a very complicated relation occurs, as in non-linear optics, ferro-magnetism or ferro-electricity. The material equations can only be evaluated by quantum mechanics. In the following non-conducting ($\sigma_e = 0$), charge-free ($\rho = 0$) and non-magnetic ($\chi_m = 0, \mu = 1$) media are assumed, which holds for dielectrics. The magnetic field can be eliminated and a wave equation results from Maxwell's equations:

grad div
$$\mathbf{E} - \Delta \mathbf{E} + \frac{1}{c_0^2} \frac{\partial^2}{\partial t^2} \left(\mathbf{E} + \frac{1}{\varepsilon_0} \mathbf{P} \right) = 0$$
, (1.1.11)

$$\operatorname{div} \mathbf{D} = 0 \tag{1.1.12}$$

with

$$c_0 = \frac{1}{\sqrt{\varepsilon_0 \mu_0}} = 2.99792458 \times 10^8 \text{ m/s}$$
: vacuum velocity of light.

Equation (1.1.11) is the fundamental equation, describing the propagation of optical fields. It includes diffraction as well as amplification of light and non-linear effects. It has now to be adapted and simplified for the different applications in optics and laser technology.

1.1.2.2 Homogeneous, isotropic, linear dielectrics

$$\frac{1}{1} \cdot \frac{1}{1} \cdot \frac{1}$$

$$\Delta \mathbf{E} - \frac{\varepsilon}{c_0^2} \frac{\partial^2 \mathbf{E}}{\partial t^2} = 0 , \qquad (1.1.13)$$

$$\operatorname{div} \mathbf{E} = 0. \tag{1.1.14}$$

Simple solutions are the plane and the spherical waves.

1.1.2.2.1 The plane wave

The infinite, monochromatic wave with a plane phase front and constant amplitude reads:

$$\boldsymbol{E} = \boldsymbol{E}_0 \, \exp[\mathrm{i}(\omega t - n\boldsymbol{k}_0 \boldsymbol{r})] \,, \tag{1.1.15}$$

$$\boldsymbol{H} = \boldsymbol{H}_0 \exp[\mathrm{i}(\omega t - n\boldsymbol{k}_0 \boldsymbol{r})]; \qquad (1.1.16)$$

$$\boldsymbol{H}_0 = rac{\left[\boldsymbol{k}_0 \times \boldsymbol{E}_0
ight]}{k_0 Z} \ .$$

It is a transversely polarized field with $E \perp H \perp k_0$, as plotted in Fig. 1.1.3.

$$n = \sqrt{\varepsilon} = \sqrt{1 + \chi_e}$$
: the refractive index of the medium, in general complex, (1.1.17)

 $k_0 = 2\pi/\lambda_0$: wave number in vacuum,

 k_0 : wave vector, direction of propagation,

 λ_0 : wavelength in vacuum,

$$Z = \sqrt{\frac{\mu\mu_0}{\varepsilon\varepsilon_0}}$$
: impedance, $Z_0 = \sqrt{\frac{\mu_0}{\varepsilon_0}} = 376.7~\Omega$: vacuum impedance.

The Poynting vector or energy flux is a real quantity with

$$S = [E_{\text{real}} \times H_{\text{real}}]$$
 (SI-unit: W/m²).

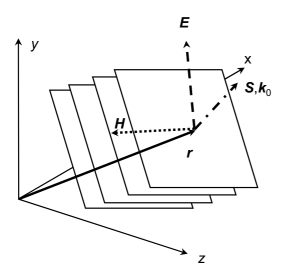


Fig. 1.1.3. The plane wave in a homogeneous, isotropic medium.

Table 1.1.1. Values of refractive index n_r and absorption coefficient α at wavelength λ_0 [85Pal, 82Gra, 78Dri].

Material	$\lambda_0 \; [\mu \mathrm{m}]$	$n_{ m r}$	$\alpha [\mathrm{m}^{-1}]$
Fused quartz	0.54	1.46	very small
Sapphire	0.50	1.765/1.764	very small
Water	0.54	1.332	0.8
Water	1	1.328	80
Copper	0.54	0.7	11.6×10^{6}
Gold	0.54	0.3	11.1×10^{6}
Iron	0.54	2.4	16.4×10^{6}

The intensity is the time average over one period $T = 2\pi/\omega$ and results in:

$$J = \langle \mathbf{S} \rangle_T = \frac{1}{4} \left(\frac{1}{Z} + \frac{1}{Z^*} \right) \langle \mathbf{E}_0 \mathbf{E}_0^* \rangle . \tag{1.1.18}$$

For dielectrics without losses ($\mu = 1$, $n = n_{\rm r}$ is real), (1.1.18) reduces to

$$J = \frac{1}{2}c_0 n_r \varepsilon_0 |E_0|^2 \tag{1.1.19}$$

with both quantities, E_0 and J, inside the medium. For vacuum applies

$$J_{\rm W/m^2} = 1.33 \times 10^{-3} |E_{\rm 0,V/m}|^2$$
, $|E_{\rm 0,V/m}| = 27.4 \sqrt{J_{\rm W/m^2}}$.

For a homogeneous dielectric, low-absorbing medium the complex refractive index is given by [99Bor, p. 739]:

$$\hat{n} = n_{\rm r} - i\frac{\alpha}{2k_0} \,, \quad \alpha \ll k_0 \tag{1.1.20}$$

with

 $n_{\rm r}$: real part of the refractive index,

 α : absorption coefficient, in general the non-resonant broad-band absorption.

For a field propagating in z-direction (1.1.15)/(1.1.20) deliver an exponentially damped amplitude:

$$\mathbf{E}(z,t) = \mathbf{E}_0 \exp\left[\mathrm{i}(\omega t - n_{\rm r}k_0 z) - \frac{\alpha z}{2}\right] . \tag{1.1.21}$$

Some numbers of $n_{\rm r}$, α are compiled in Table 1.1.1.

1.1.2.2.2 The spherical wave

One solution of the wave equation (1.1.13) in spherical coordinates is the quasi-spherical wave, generated by an oscillating dipole (Hertz's dipole), see Fig. 1.1.4. The far field reads [99Jac]:

$$\boldsymbol{E}\left(r,\vartheta,t\right) = \frac{\lambda_0 \boldsymbol{E}_{\vartheta}}{r} \exp\left[\mathrm{i}\left(\omega t - \hat{n}\boldsymbol{k}_0\boldsymbol{r}\right)\right] \sin\vartheta \;, \quad |\boldsymbol{E}_{\vartheta}| = \frac{|\boldsymbol{\mu}| \; 4\pi^2 k_0^3}{\varepsilon_0} \;, \quad r \gg \lambda_0$$

with μ the dipole moment and ϑ the angle between the dipole axis and beam propagation k_0 . In the paraxial approach ($\vartheta \cong \pi/2$, $\theta \ll 1$) the well-known spherical wave, useful for applying Huygens' principle, results:

$$\boldsymbol{E}(z,t) \cong \frac{\lambda_0}{r} \, \boldsymbol{E}_0 \exp\left[\mathrm{i} \left(\omega t - \hat{n} \boldsymbol{k}_0 \boldsymbol{r}\right)\right] , \quad \theta \ll 1 , \qquad (1.1.22)$$

where E is approximately parallel to the dipole axis.

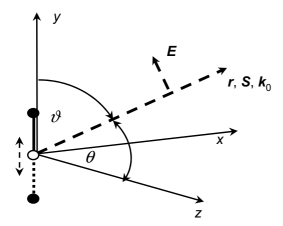


Fig. 1.1.4. A quasi-spherical wave, emitted by an oscillating dipole.

1.1.2.2.3 The slowly varying envelope (SVE) approximation

In the Slowly Varying Envelope approximation (1.1.11) is solved approximately with the ansatz of a quasi-monochromatic, quasi-plane wave

$$E = E_0(x, y, z, t) \exp[i(\omega t - n_r k_0 z)], \quad P = P_0(x, y, z, t) \exp[i(\omega t - n_r k_0 z)].$$
 (1.1.23)

The wave propagates mainly in z-direction and the amplitude is slowly varying with x, y, z, t, which means:

- slowly varying in time (quasi-monochromatic): $\partial |\mathbf{E}_0|/\partial t \ll \omega |\mathbf{E}_0|$, or spectral bandwidth $\Delta \omega \ll \omega$,
- slowly varying in space (quasi-plane wave): $\partial |\mathbf{E}_0|/\partial z \ll k_0|\mathbf{E}_0|$, which means low divergence of the beam $\Delta\theta \ll 1$ (paraxial approach), and a smooth transverse profile,
- slowly varying polarization $\partial |\mathbf{P}_0|/\partial t \ll \omega |\mathbf{P}_0|$,
- slowly varying electric susceptibility $\partial |\chi_{\rm e}|/\partial t \ll \omega |\chi_{\rm e}|$ and $|{\rm grad}~\chi_{\rm e}| \ll k_0 |\chi_{\rm e}|$.

Then second order terms can be neglected and the SVE-approximations are obtained [84She, p. 47], [66War, 86Sie].

1.1.2.2.4 The SVE-approximation for diffraction

$$\left(\Delta_{\rm tr} - 2ik_0 \frac{\partial}{\partial z}\right) \boldsymbol{E}_0 = 0 , \quad \text{div } \boldsymbol{E} = 0 . \tag{1.1.24a}$$

 Δ_{tr} is the transverse delta-operator, which in rectangular coordinates reads

$$\Delta_{\rm tr} = \frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2} \; .$$

The field in (1.1.24a) is a vector field, and the Δ -operator in cylinder coordinates is rather complicated, because the unit-vectors are no longer constant [99Jac], especially for non-uniform polarization in circular birefringent media [82Fer, 93Wit]. In most cases (except birefringence) the

nates

$$\left(\frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2} - 2ik_0 \frac{\partial}{\partial z}\right) E_0 = 0 , \qquad (1.1.24b)$$

$$\left(\frac{\partial^2}{\partial r^2} + \frac{1}{r}\frac{\partial}{\partial r} + \frac{1}{r^2}\frac{\partial^2}{\partial \varphi^2} - 2ik_0\frac{\partial}{\partial z}\right)E_0 = 0.$$
(1.1.24c)

This is the fundamental equation in paraxial diffraction optics. It gives the Fresnel-integral and the eigenmodes of free propagation (Gauss-Hermite/Gauss-Laguerre polynomials, see Chaps. 3.1 and 8.1). Equations (1.1.24a)/(1.1.24b)/(1.1.24c) hold for a homogeneous medium, but can be extended to quadratic index media [86Sie].

1.1.2.3 Propagation in doped media

A plane wave without transverse structure interacts with active atoms or molecules and induces a polarization P_A . In most cases the active atoms are embedded in a host medium (glass, crystal, liquid, gas), which is also polarized by the field, generating an additional polarization P_H . The total polarization is:

$$P = P_{A} + P_{H} = (P_{A0} + P_{H0}) \exp[i(\omega t - n_{r}k_{0}z)].$$
(1.1.25)

The response of the host medium is in most cases very fast $(10^{-12} \dots 10^{-14} \text{ s})$, no transient behavior occurs and nonlinear effects are assumed to be small. Then the host polarization is proportional to the applied field:

$$\boldsymbol{P}_{\mathrm{H}} = \varepsilon_0 \chi_{\mathrm{H}} \boldsymbol{E}$$
.

 $\chi_{\rm H}$ is the complex susceptibility of the host material and is related to the refractive index $n_{\rm r}$ and the loss coefficient α according to (1.1.17)/(1.1.20) [99Ber]:

$$\chi_{\rm H} = (n_{\rm r}^2 - 1) - i \frac{n_{\rm r} \alpha}{k_0} , \quad \alpha \ll k_0 .$$
 (1.1.26)

The imaginary part of $\chi_{\rm H}$ is called extinction coefficient. Some values of refractive indices $n_{\rm r}$ and absorption coefficients α are given in Table 1.1.1. For the polarization of the active atoms one has

$$\mathbf{P}_{\mathbf{A}} = \varepsilon_0 \chi_{\mathbf{A}}(\mathbf{E}_0) \mathbf{E} \,, \tag{1.1.27}$$

where $\chi_{\rm A}$ depends on the field and has to be evaluated quantum-mechanically. Neglecting first and second order derivations of $\boldsymbol{P}_{\rm A0}$ and second order derivations of $\boldsymbol{E}_{\rm 0}$, the SVE-approximation for the interaction is obtained, assuming a plane wave without transverse structure:

$$\left(\frac{\partial}{\partial z} + \frac{1}{c}\frac{\partial}{\partial t} + \frac{\alpha}{2}\right)E_0 = -i\frac{k_0}{2\varepsilon_0 n_r} \mathbf{P}_{A0}\left(\mathbf{E}_0\right) , \quad \text{div } \mathbf{E} = 0$$
(1.1.28)

(SVE-approximation for the amplitude of a plane wave in an active medium)

with $c = c_0/n_r$ the phase velocity of the wave in the host medium. The above equation describes the amplification/attenuation of cw-fields and pulsed radiation by an active medium. It provides also the widely used rate-equation approach, as will be shown in Sect. 1.1.5.1. It fails for fields

with amplitudes varying very rapidly in time or space (fs-pulses). If the intensity J (1.1.19) and the susceptibility of the active medium (1.1.27) are introduced, (1.1.28) reduces to:

$$\left(\frac{\partial}{\partial z} + \frac{1}{c}\frac{\partial}{\partial t}\right)J + \left(\alpha - \frac{k_0}{n_r}\operatorname{Im}\chi_A\right)J = 0.$$
(1.1.29)

The active atoms enhance or reduce the losses of the medium, depending on the sign of the imaginary part $\operatorname{Im} \chi_A$ of the susceptibility, which is a function of the intensity. In steady state and for constant χ_A , which holds for low intensities, (1.1.29) can be integrated and delivers for the intensity

$$J(z) = J(0) \exp \left[-\alpha + \frac{k_0}{n_r} \operatorname{Im} \left(\chi_{\mathcal{A}} \right) \right] z .$$

The amplifying factor is called the small-signal gain factor G_0 of the medium and the exponent the small-signal gain coefficient g_0 :

$$G_0 = \exp\left[\frac{k_0}{n_r}\operatorname{Im}(\chi_A)z\right] = \exp\left[g_0z\right] , \quad g_0 = \frac{k_0}{n_r}\operatorname{Im}(\chi_A) .$$
 (1.1.30)

Some typical values of g_0 are compiled in Table 1.1.4.

1.1.3 Interaction with two-level systems

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1.1.3.1 The two-level system

$$\Delta E_1$$
, μ_{r+1} ,

The two-level system can be part of an atom, ion, molecule, or something more complicated. A monochromatic electric field E of frequency ω in the SVE-approximation according to (1.1.23) acts via the Coulomb force on the bound electrons of the active medium. In linear systems (parabolic potential) the negative electrons will oscillate sinusoidally, whereas the heavy positive nucleus remains more or less at rest. An oscillating dipole is induced with a dipole moment $\mu(t)$, which is given by

$$\mu = -ex \tag{1.1.31}$$

with

e: electron charge,

x: displacement of the electron.

The dipole moment per volume is the macroscopic polarization $P_{\rm A}$ of the active medium. As all single dipoles are aligned by the electric field, the resulting polarization reads:

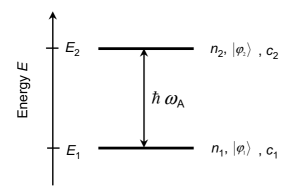


Fig. 1.1.5. The two-level system.

$$\boldsymbol{P}_{\mathbf{A}} = n_0 \boldsymbol{\mu} \tag{1.1.32}$$

with

 n_0 : dipole density (m⁻³), μ : expectation value of the dipole moment (Asm).

In this section the induced dipole moment will be evaluated quantum-mechanically, which requires some simplifications. It is not the intention to discuss in detail the mathematics, but only to summarize briefly the main results of interest for laser technology and to emphasize the approximations and the range of validity. A consistent presentation of the interaction light–matter, starting from first principles, is given in many textbooks [61Mes, 68Sch, 77Coh, 95Man].

From the infinite number of energy levels of an electronic system only two, E_1 and E_2 , are taken into account for the interaction [75All, 89Yar, 69Are], see Fig. 1.1.5. This is a reasonable approach if the field is nearly resonant with the transition from E_1 to E_2 . In this case the other levels of the system will not or only very weakly interact with the field.

It applies

$$|\omega_{\rm A} - \omega| \ll \Delta \omega_{\rm A}$$

with

 $\omega_{\rm A}$: resonance frequency of the transition, $\Delta\omega_{\rm A}$: bandwidth of the transition, ω : frequency of the radiation field, $\hbar=1.0546\times 10^{-34}~{\rm Ws^2}$: Planck's constant.

1.1.3.2 The dipole approximation

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E , which is a E , which is a E , which is a substitution of E .
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1.1.3.2.1 Inversion density and polarization

The interaction of an electromagnetic field with a two-level system was first investigated by Bloch [46Blo] and extensively discussed by Allen and Eberly [75All]. It is characterized by its dipole moment and the population densities in the two levels:

 n_1, n_2 : density of states (atoms, molecules) in the lower/upper level,

 $\Delta n = n_2 - n_1$: inversion density, $n_0 = n_1 + n_2$: total density, const.

The following assumptions are made:

- Non-relativistic interaction. The velocity of the electrons is small compared with the velocity
 of light. This does not hold for inner-shell electrons, hot plasmas and free-electron lasers.
- The wavelength of the light is large compared with the diameter of the atoms/molecules. It means that in the domain of the atomic wave function the electromagnetic field is locally constant. Bohr's radius with $r_{\rm B} = 5.3 \times 10^{-5}$ µm is a typical atomic dimension. The wavelength in the visible range of the spectrum is about 0.5 µm, thus this condition is fulfilled in the visible and UV-part of the spectrum. It is called the dipole approximation [97Scu].
- The permanent dipole moments of the two-level system $\mu_{11} = \mu_{22}$ are zero. Even if larger molecules have a permanent dipole moment, their response to the high-frequency field is small. Only for very strong fields are the permanent dipole moments of importance (see Part 4 on nonlinear optics). A dipole moment exists only for the transition from level 1 to 2 and vice versa. Non-degenerated levels are assumed with $\mu = \mu_{12} = \mu_{21}$.

The two-level system is completely described by its state vector $|\varphi\rangle$, which in general is time-dependent:

$$|\varphi\rangle = c_1(t) |\varphi_1\rangle \exp\left(-i\frac{E_1 t}{\hbar}\right) + c_2(t) |\varphi_2\rangle \exp\left(-i\frac{E_2 t}{\hbar}\right) ,$$
 (1.1.33)

with $|\varphi_1\rangle$, $|\varphi_2\rangle$ the eigenfunctions and E_1 , E_2 the energy eigenstates. The eigenfunctions are normalized, orthogonal and depend on the position vector \mathbf{r} :

$$\int \varphi_i^* \varphi_j dr = \langle \varphi_1 \varphi_2 \rangle = \delta_{ij} . \tag{1.1.34}$$

The state vector has to fulfill the time-dependent Schrödinger equation:

$$i\hbar \frac{\partial |\varphi\rangle}{\partial t} = (H_0 + H_{\rm int}) |\varphi\rangle ,$$
 (1.1.35)

with H_0 the Hamilton operator of the undisturbed system ($H_{\text{int}} = 0$) and H_{int} the interaction energy. For the undisturbed system holds [89Yar]:

$$H_0 |\varphi_i\rangle = E_i |\varphi_i\rangle , \quad i = 1, 2,$$
 (1.1.36)

which follows directly from (1.1.35) by replacing $|\varphi\rangle$ by $|\varphi_i\rangle \exp(-iE_it/\hbar)$. The parameters of interest, the inversion density $\Delta n = n_2 - n_1$ and the macroscopic polarization

$$P_{\mathbf{A}} = n_0 \,\boldsymbol{\mu} \tag{1.1.37}$$

are determined by the coefficients c_1 , c_2 . The probability of the system to be in the lower/upper state is given by $|c_1|^2$, $|c_2|^2$, respectively, which requires:

$$|c_1|^2 + |c_2|^2 = 1$$
. (1.1.38)

The number of atoms in the lower/upper level is then given by:

$$n_1 = n_0 |c_1|^2$$
, $n_2 = n_0 |c_2|^2$, $n_1 + n_2 = n_0$

and hence the inversion density:

$$\Delta n = n_0 \left(|c_2|^2 - |c_1|^2 \right) . \tag{1.1.39}$$

The expectation value of the dipole moment $\langle \mu \rangle = -e \langle \varphi r \varphi \rangle$ is obtained from (1.1.33). Using the afore mentioned assumptions:

$$\langle \boldsymbol{\mu}_{11} \rangle = -e \langle \varphi_1 \boldsymbol{r} \varphi_1 \rangle = 0 , \quad \langle \boldsymbol{\mu}_{22} \rangle = -e \langle \varphi_2 \boldsymbol{r} \varphi_2 \rangle = 0$$

one obtains finally for the polarization from (1.1.33), (1.1.34), (1.1.38)

$$P_{A} = n_{0} \{ \langle \mu_{12} \rangle \ c_{1}^{*} c_{2} \exp(-i\omega_{A} t) + \langle \mu_{21} \rangle \ c_{1} c_{2}^{*} \exp(+i\omega_{A} t) \}$$
(1.1.40)

with $\langle \mu_{12} \rangle$, $\langle \mu_{21} \rangle$ the dipole moment of the transition $E_1 \leftrightarrow E_2$ and vice versa. For non-degenerated transitions one has $\langle \mu_{12} \rangle = \langle \mu_{21} \rangle = \mu_A$. In the following only μ_A will be used, which is a characteristic parameter of the specific transition:

$$\mu_{\rm A} = -e \left\langle \varphi_1 \mathbf{r} \varphi_2 \right\rangle . \tag{1.1.41}$$

1.1.3.2.2 The interaction with a monochromatic field

The interaction Hamiltonian for a non-quantized real field E_{real} corresponds to the classical energy of an electric dipole in an electric field. It reads [97Scu]:

$$H_{\text{int}} = \mu_{\text{A}} E_{\text{real}} = \mu_{\text{A}} \frac{(E + E^*)}{2}. \tag{1.1.42}$$

Substitution of (1.1.42) into (1.1.35), using the orthogonality (1.1.34) and (1.1.41) provides two differential equations for the coefficients c_1 , c_2 of the state vector:

$$\frac{\mathrm{d}c_1}{\mathrm{d}t} = \frac{\mathrm{i}}{\hbar} c_2 \exp\left(-\mathrm{i}\omega_{\mathrm{A}}t\right) \boldsymbol{\mu}_{\mathrm{A}} \frac{(\boldsymbol{E} + \boldsymbol{E}^*)}{2} ,$$

$$\frac{\mathrm{d}c_2}{\mathrm{d}t} = \frac{\mathrm{i}}{\hbar} c_1 \exp\left(+\mathrm{i}\omega_{\mathrm{A}}t\right) \boldsymbol{\mu}_{\mathrm{A}} \frac{(\boldsymbol{E} + \boldsymbol{E}^*)}{2} .$$
(1.1.43)

The time dependence of inversion density and polarization is obtained from (1.1.39), (1.1.40) by differentiating and applying (1.1.43). After some simple mathematics the following two equations for the macroscopic parameters of the two-level-system result are obtained:

$$\frac{\partial \Delta n}{\partial t} = \frac{\mathrm{i}}{\hbar} \left\{ (\boldsymbol{E} + \boldsymbol{E}^*) (\boldsymbol{P}_{\mathrm{A}} - \boldsymbol{P}_{\mathrm{A}}^*) \right\} , \qquad (1.1.44a)$$

$$\frac{\partial \mathbf{P}_{A}}{\partial t} = i \left\{ \omega_{A} \mathbf{P}_{A} + \frac{\mu_{A}}{\hbar} \left\langle \mathbf{\mu}_{A} \left(\mathbf{E} + \mathbf{E}^{*} \right) \right\rangle \Delta n \right\}. \tag{1.1.44b}$$

For E and P_A the SVE-approximations of (1.1.23), (1.1.25) are used. Then in (1.1.44a), (1.1.44b) terms with the frequency 2ω appear, which are neglected. This approach is called the rotating-wave approximation [97Scu, 72Cou]. The above equations simplify to

$$\frac{\partial \Delta n}{\partial t} = \frac{\mathrm{i}}{2\hbar} \left\{ \boldsymbol{E}_0^* \boldsymbol{P}_{\mathrm{A0}} - \boldsymbol{E}_0 \boldsymbol{P}_{\mathrm{A0}}^* \right\} , \qquad (1.1.45a)$$

$$\frac{\partial \mathbf{P}_{A0}}{\partial t} = -i \, \delta \mathbf{P}_{A0} + \frac{i \, \mathbf{\mu}_{A}}{\hbar} \, \langle \mathbf{\mu}_{A} \mathbf{E}_{0} \rangle \, \Delta n \,, \quad \delta = \omega - \omega_{A}$$
(1.1.45b)

(rotating-wave approximation)

with

 $\mu_{\rm A}$: electric dipole moment of the transition,

 ω : frequency of the interacting field,

 $\omega_{\rm A}$: resonance frequency of the two-level system,

 $\hbar = 1.0546 \times 10^{-34} \text{ Ws}^2$: Planck's constant.

Some typical values of dipole moments are given in Table 1.1.2.

Table 1.1.2. Typical values of dipole moments [01Men].

Transition			$ \mu_{ m A} \; [{ m As \; m}]$
Bohr's radius × electron of	charge		10^{-29}
Hydrogen	$\begin{array}{c} 1s-2p\\ 4f-5g \end{array}$	$\lambda_0 = 121 \text{ nm}$ $\lambda_0 = 4053 \text{ nm}$	$0.8 \times 10^{-29} \\ 8.3 \times 10^{-29}$
Chromium ions in ruby	$4A_2(3/2) - E$ levels	$\lambda_0 = 694 \text{ nm}$	10^{-29}

1.1.3.3 The Maxwell-Bloch equations

So far the interaction of the two-level system with the electromagnetic field is purely coherent, no perturbations by external influences on the system are considered. Stochastic processes will modify the interaction considerably. Here only a very basic description is presented. A detailed analysis of these statistical processes is given in [70Hak, 97Scu].

1.1.3.3.1 Decay time T_1 of the upper level (energy relaxation)

Three incoherent processes reduce or increase the upper-level population and have to be considered in (1.1.45a), (1.1.45b):

- spontaneous emission,
- interaction with the host material (collisions, lattice vibrations),
- increase of the population by pumping (light, electron collisions, or other processes).

1.1.3.3.1.1 Spontaneous emission

The two-level system is coupled to the modes of the optical resonator or to the free-space modes. Spontaneous emission into these modes reduces the upper-level population. Moreover, by each spontaneous emission process the phase relation between the field and the two-level eigenfunction is destroyed. If the dimensions of the resonator are large compared with the wavelength, the decay is given by $\partial n_2/\partial t = -n_2/T_{\rm sp}$, with $A_{21} = 1/T_{\rm sp}$, the Einstein coefficient of spontaneous emission. If the resonator dimensions are comparable with the wavelength, spontaneous emission is strongly influenced by the resonator geometry, it can be enhanced or reduced (see Chap. 8.1).

1.1.3.3.1.2 Interaction with the host material

This interaction reduces the population density. Energy is transferred to the host material and converted into heat. A simple approach for this decay is again an exponential ansatz $\partial n_2/\partial t = -n_2/T_{\rm H}$. This decay time together with the spontaneous decay time delivers a resulting decay T_1 of the upper-level population, also called energy relaxation time or longitudinal relaxation time.

1.1.3.3.1.3 Pumping process

The dynamics of upper-level excitation depend on the special pumping scheme and are discussed in Sect. 1.1.5.3 and in Vol. VIII/1B, "Solid-state laser systems". In any case the pump produces in steady state and without a coherent field ($E_0 = 0$) an inversion density Δn_0 .

These three processes are included into (1.1.45a) by the term:

$$\frac{\partial \Delta n}{\partial t} = -\frac{\Delta n - \Delta n_0}{T_1} \tag{1.1.46}$$

with

 T_1 : the resulting time constant.

1.1.3.3.2 Decay time T_2 of the polarization (entropy relaxation)

An external field E induces dipoles, which generate the macroscopic polarization P_A . If the external field is switched off, the polarization will disappear for several reasons:

The energy of the two-level system decays with T_1 , which means that the polarization disappears at least with the same time constant.

Due to incoherent interaction with the host material (collisions), the single dipoles are disoriented in their direction or dephased. The resulting polarization becomes zero, although the single dipole still exists. This process can be much faster than T_1 (see Table 1.1.6) and is characterized by a time constant T_2 . This decay strongly depends on the interaction process. The simplest approach is:

$$\frac{\partial \mathbf{P}_{A0}}{\partial t} = -\frac{\mathbf{P}_{A0}}{T_2} \,, \tag{1.1.47}$$

and (1.1.45b) has to be completed by (1.1.47). T_2 is called the transverse relaxation time, the entropy time constant or the dephasing time. Finally, the two-level equations together with the SVE-approximation, (1.1.28), of the wave equation read:

$$\frac{\partial \Delta n}{\partial t} = -\frac{\mathrm{i}}{2\hbar} \left(\boldsymbol{E}_0^* \boldsymbol{P}_{\mathrm{A0}} - \boldsymbol{E}_0 \boldsymbol{P}_{\mathrm{A0}}^* \right) - \frac{\Delta n - \Delta n_0}{T_1} , \qquad (1.1.48a)$$

$$\frac{\partial \mathbf{P}_{A0}}{\partial t} = -\left(i\,\delta + \frac{1}{T_2}\right)\mathbf{P}_{A0} + i\frac{\mu_A \langle \mu_A \mathbf{E}_0 \rangle}{\hbar} \Delta n \;, \quad \delta = \omega - \omega_A \;, \tag{1.1.48b}$$

$$\left(\frac{\partial}{\partial z} + \frac{1}{c}\frac{\partial}{\partial t} + \frac{\alpha}{2}\right)\boldsymbol{E}_0 = -i\frac{k_0}{2\varepsilon_0 n_r}\boldsymbol{P}_{A0}$$
(1.1.48c)

(Maxwell–Bloch equations).

They describe the propagation of radiation in two-level systems and are called Maxwell–Bloch equations. Equation (1.1.48c) holds, if the transition frequency $\omega_{\rm A}$ for all two-level atoms is the same (homogeneous system). In inhomogeneous systems (see Sect. 1.1.6.3, Fig. 1.1.13) different groups of atoms exist with center frequencies $\omega_{\rm A}$ of each group and a center frequency $\omega_{\rm R}$ of the ensemble. Therefore (1.1.48c) has to be replaced by [81Ver]:

$$\left(\frac{\partial}{\partial z} + \frac{1}{c}\frac{\partial}{\partial t}\right)\boldsymbol{E}_{0} = -i\frac{k_{0}}{2\varepsilon_{0}n_{r}}\int h(\omega_{A},\omega_{R})\boldsymbol{P}_{A0}(\boldsymbol{E}_{0},\omega_{A})d\omega_{A}.$$
(1.1.48d)

 $h(\omega, \omega_{\rm A})$ is the spectral density of atoms with the transition frequency $\omega_{\rm A}$ according to (1.1.92)/(1.1.93). For the solution of these equations, three different regimes are distinguished:

- 1 - 11-1 - - 1111

The temporal variations of the radiation field are slow compared with T_1 .

$$\frac{\partial \Delta n}{\partial t} = 0 \qquad \frac{\partial \boldsymbol{P}_{A0}}{\partial t} = 0$$

no transient effects of the atom, $T_2 \ll T_1$.

$$\frac{\partial \Delta n}{\partial t} \neq 0 \qquad \frac{\partial \mathbf{P}_{A0}}{\partial t} = 0$$

The width τ of the interacting pulses is short compared with T_1, T_2 ; (1.1.45a), (1.1.45b) can be applied.

$$\frac{\partial \Delta n}{\partial t} \neq 0 \qquad \frac{\partial \mathbf{P}_{A0}}{\partial t} \neq 0$$

1.1.4 Steady-state solutions

$$m{r}_1$$
 , $m{r}_2$, $m{r}_3$, $m{r}_4$, $m{r}_4$

1.1.4.1 Inversion density and polarization

The stationary solutions of (1.1.48a), (1.1.48b) are obtained immediately:

$$\Delta n = \frac{\Delta n_0}{1 + (J/J_s) f(\omega)} \quad (1.1.49)$$

$$\chi_{\rm A} = \frac{n_{\rm r}\sigma}{k_0} \left[\frac{\omega - \omega_{\rm A}}{\Delta \omega_{\rm A}/2} + i \right] \Delta n \quad (1.1.50)$$

$$\mathbf{P}_{A0} = \varepsilon_0 \chi_A \mathbf{E}_0 \quad (\mathbf{I}_{A}, \mathbf{I}_{A}, \mathbf{I}_{A}, \mathbf{I}_{A}) \tag{1.1.51}$$

with

$$J = \frac{1}{2} \varepsilon_0 c_0 n_{\rm r} |\boldsymbol{E}_0|^2 \quad \text{(intensity of the field)}, \tag{1.1.52}$$

$$J_{\rm s} = \frac{\hbar \,\omega_{\rm A}}{2\sigma_0 T_1} \quad \text{(saturation intensity of the two-level transition)}, \tag{1.1.53}$$

$$\sigma = \sigma_0 f(\omega, \omega_{\rm A})$$
 (frequency-dependent cross section of the transition), (1.1.54)

$$\sigma_0 = \frac{|\boldsymbol{\mu}_{\rm A}|^2 \,\omega_{\rm A} T_2}{\varepsilon_0 c_0 n_{\rm r} \hbar} \quad \text{(cross section in resonance)},\tag{1.1.55}$$

$$f_{\rm L}(\omega, \omega_{\rm A}) = \frac{(\Delta \omega_{\rm A}/2)^2}{(\omega_{\rm A} - \omega)^2 + (\Delta \omega_{\rm A}/2)^2} \quad \text{(spectral line shape, Lorentzian)}, \tag{1.1.56}$$

$$\Delta\omega_{\rm A} = 2/T_2$$
 (line width of the transition), (1.1.57)

$$g_{\rm h}(\omega, \omega_{\rm A}) = \Delta n \, \sigma = \frac{\Delta n_0 \sigma_0 f(\omega, \omega_{\rm A})}{1 + (J/J_{\rm s}) f(\omega, \omega_{\rm A})}$$
 (gain coefficient, homogeneously broadened),

$$g_{\rm inh}(\omega, \omega_{\rm R}) = \frac{\Delta n_0 \sigma_0}{\sqrt{1 + J/J_{\rm s}}} h(\omega, \omega_{\rm R}) \frac{\Delta \omega_{\rm A}}{2}$$
 (gain coefficient, inhomogeneously (1.1.58b)

broadened, see Sect. 1.1.6.3).

In Table 1.1.3 some numbers of relevant laser transitions are compiled, in Table 1.1.4 some typical values of the small-signal gain coefficient in resonance are given. The susceptibility strongly depends on the frequency as shown in Fig. 1.1.6. According to (1.1.26) the real part of χ_A produces an additional refractive index, and the imaginary part absorption or amplification:

Re
$$\chi_{\rm A} = n_{\rm r}^2 - 1 = \frac{n_{\rm r}\sigma}{k_0} \left(\frac{\omega - \omega_{\rm A}}{\Delta\omega_{\rm A}/2}\right) \Delta n$$
, (1.1.59a)
Im $\chi_{\rm A} = -n_{\rm r}\alpha k_0 = \frac{n_{\rm r}\sigma}{k_0} \Delta n$. (1.1.59b)

$$\operatorname{Im} \chi_{A} = -n_{r}\alpha k_{0} = \frac{n_{r}\sigma}{k_{0}}\Delta n. \tag{1.1.59b}$$

The steady-state propagation of the electric field is obtained from (1.1.48c):

$$\frac{\mathrm{d}\boldsymbol{E}_0}{\mathrm{d}z} = \left[-\frac{\alpha}{2} + \frac{\sigma\Delta n}{2} + \mathrm{i}\sigma\Delta n \frac{\omega - \omega_{\mathrm{A}}}{\Delta\omega_{\mathrm{A}}} \right] \boldsymbol{E}_0 , \qquad (1.1.60)$$

where Δn is a function of the field or the intensity.

Table 1.1.3. Examples of resonance wavelength λ_0 , resonance cross section σ_0 , upper-level lifetime T_1 and saturation intensity J_s . The simple relation (1.1.53) for the saturation intensity holds for two-level systems only and is not applicable in general [01Men].

	λ_0	σ_0	T_1	$J_{ m s}$
	$[\mu \mathrm{m}]$	$[m^2]$	[s]	$[\mathrm{W/m}^2]$
Amplifiers				
CO_2 -gas (1300 Pa)	10.6	10^{-20}	10^{-5}	2×10^5
Neodymium-ion in glass	1.06	4×10^{-24}	3×10^{-4}	$8 \dots 12 \times 10^7$
Neodymium-ion in YAG	1.06	5×10^{-23}	2×10^{-4}	2×10^{7}
Chromium-ion in Al ₂ O ₃	0.69	2×10^{-24}	3×10^{-3}	2.4×10^{7}
(ruby, T = 300 K)				
Neon (25 Pa)	0.63	3×10^{-17}	10^{-8}	5.3×10^{5}
Rhodamine 6G in ethanol	0.57	4×10^{-20}	5×10^{-9}	10^{9}
Absorbers				
SF_6	10.6	8×10^{-22}	4×10^{-4}	2.5×10^{5}
KODAK dye 9860	1.06	4×10^{-20}	$\sim 10^{-11}$	5.6×10^{11}
KODAK dye 9740	1.06	6×10^{-20}	$\sim 10^{-11}$	4×10^{11}
Cryptocyanine-dye in methanol	0.7	5×10^{-20}	5×10^{-10}	2×10^{10}

Table 1.1.4. Typical values of the small-signal gain coefficient $g_0 = \Delta n_0 \sigma_0$ in resonance. The exact values depend on pumping, doping, and other parameters of operation [01Men].

System	$\lambda_0 \; [\mathrm{nm}]$	$g_0 \ [{\rm m}^{-1}]$
He/Ne laser	632.8	0.1
Nd-doped glass	1060	5
Nd-doped YAG	1060	50
GaAs-diode	880	4×10^3

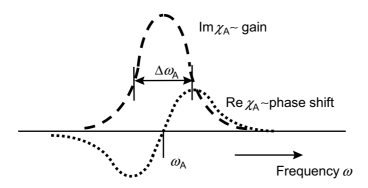


Fig. 1.1.6. Real and imaginary part of the susceptibility vs. frequency.

1.1.4.2 Small-signal solutions

$$J_{s_1,\ldots,s_1,\ldots,s_{s_1,\ldots,s_1,\ldots,s_{s_1,$$

At low intensities $J \ll J_s$, the inversion density is not affected by the intensity,

$$\Delta n = \Delta n_0 ,$$

and (1.1.60) can be integrated. Together with (1.1.23), the complete field is obtained:

$$\boldsymbol{E}(z) = \boldsymbol{E}_0(0) \exp[\mathrm{i}(\omega t - n_t k_0 z) - \frac{1}{2}(\alpha - \Delta n_0 \sigma) z]$$
(1.1.61)

with a total refractive index $n_{\rm t}$

$$n_{\rm t} = n_{\rm r} \left(1 + \frac{\sigma \Delta n_0}{n_{\rm r} k_0} \frac{\omega - \omega_{\rm A}}{\Delta \omega_{\rm A}} \right) . \tag{1.1.62}$$

The active atoms of the two-level system cause an additional phase shift or refractive index and an additional absorption or amplification, depending on the sign of Δn_0 . The small-signal gain factor according to (1.1.30)/(1.1.50) is:

$$G_0 = \exp[\sigma(\omega)\Delta n_0 z] . \tag{1.1.63}$$

Amplification, $G_0 > 1$, requires inversion $\Delta n_0 > 0$. The complex amplitude transmission factor A is defined as the ratio of the monochromatic field amplitudes and can be written:

$$A = \frac{E_0(z)}{E_0(0)} = \exp\left[i\frac{\sigma_0 \Delta n_0}{2} \frac{\Delta \omega_A/2}{(\omega - \omega_A) + i\Delta \omega_A/2}\right] z.$$
 (1.1.64)

It depends on the frequency of the field, which means dispersion. Time-dependent fields and especially short pulses are distorted by the amplifying system, pulse broadening and chirping occur.

1.1.4.3 Strong-signal solutions



The inversion now depends on the intensity. For the propagation of the intensity, (1.1.48c) gives in steady state

$$\frac{\mathrm{d}J}{\mathrm{d}z} = (g(J) - \alpha) J , \qquad (1.1.65)$$

where g(J) is the saturated gain coefficient of (1.1.58a), (1.1.58b). For a homogeneously broadened transition and without losses ($\alpha = 0$) this equation can be can be integrated and provides a transcendental relation for the gain factor G:

$$\frac{G_0}{G} = \exp\left[\frac{J(0)}{J_s}f(\omega) (G-1)\right] \tag{1.1.66}$$

with G_0 the small-signal gain factor of (1.1.62) and G the ratio of output/input intensities

$$G = J(z)/J(0) .$$

For inhomogeneously broadened transitions a more complicated relation is obtained [81Ver].

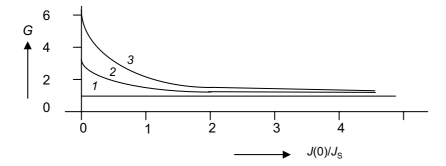


Fig. 1.1.7. Saturation of the gain factor G for a homogeneously and inhomogeneously broadened transition. 1: $G_0 = 1$, 2: $G_0 = 4$, 3: $G_0 = 6$.

1.1.5 Adiabatic equations

1.1.5.1 Rate equations

 T_2 is the time constant, which characterizes the transient behavior of the polarization. In most cases (see Table 1.1.6) T_2 is much smaller than T_1 , and the transient oscillations of the electrons can be neglected. In (1.1.48a) the polarization is replaced by its steady-state value (1.1.50)/(1.1.51) and the rate equations are obtained. They have to be completed by the time-dependent pump term, here labeled as Δn_0 . It depends on the specific pump scheme (see Sect. 1.1.5.3). The rate equations are widely used in laser design to evaluate output power, spiking behavior and Q-switching dynamics. The spontaneous emission contributes to the intensity of the interacting field, but only with a very

small amount and is neglected here. Nevertheless it is important, because the laser is started by spontaneous emission and in the lower limit it determines the laser band width (Chap. 5.1).

With these approximations the field equations (1.1.48a)/(1.1.48b)/(1.1.48c) for the interaction with a monochromatic field reduce to one equation for the inversion density and a transport equation for the intensity:

$$\frac{\partial \Delta n}{\partial t} = -\frac{J f(\omega)}{J_{\rm s} T_1} \Delta n - \frac{(\Delta n - \Delta n_0)}{T_1} , \qquad (1.1.67)$$

$$\left(\frac{\partial}{\partial z} + \frac{1}{c}\frac{\partial}{\partial t}\right)J = (\Delta n\,\sigma_0 f(\omega))J \tag{1.1.68}$$

(rate equations for a homogeneously broadened two-level system and a plane monochromatic wave)

J(z,t): local intensity,

 J_s : saturation intensity, depends on the level system (2,3, or 4 levels), see Sects. 1.1.4.1/1.1.5.3, $\Delta n(z,t)$: local inversion density.

1.1.5.2 Thermodynamic considerations

$$-1$$
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Einstein published in 1917 [17Ein] his famous work on the quantum theory of radiation, where for the first time induced emission was introduced, the cornerstone of laser physics. He discussed the two-level system in equilibrium with thermal radiation of spectral energy density ρ_{ω} (energy per volume and spectral range d ω). The density is given by Planck's law [61Mor]:

$$\rho_{\omega} = \frac{\hbar \,\omega^3}{\pi^2 c^3} \frac{1}{\exp\left[\hbar \,\omega/\kappa T\right] - 1} \qquad \left[\frac{\text{VAs}^2}{\text{m}^3}\right] \tag{1.1.69}$$

with

 $\kappa = 1.38 \times 10^{-23} \text{ VAs/K: Boltzmann's constant.}$

In thermal equilibrium the levels $|\varphi_1\rangle$, $|\varphi_2\rangle$ are populated according to Boltzmann's law [61Mor]:

$$\frac{n_2}{n_1} = \exp\left[-\hbar\,\omega_{\rm A}/\kappa T\right] \ . \tag{1.1.70}$$

These two fundamental laws can only be fulfilled, if induced emission is introduced, and Einstein postulated the following equation in steady state for the interaction of thermal radiation with a two-level system:

$$B_{12} \rho_{\omega} n_1 = B_{21} \rho_{\omega} n_2 + A_{21} n_2$$
 (1.1.71)
(absorption = induced emission + spontaneous emission)

with

 B_{12} , B_{21} , A_{21} : Einstein coefficients of induced and spontaneous emission.

The transition of atoms from the lower level to the upper level by absorption of radiation must be balanced by induced emission and spontaneous emission from the upper level. This equation was

derived by thermodynamical considerations. The quantum-mechanical equation (1.1.67) delivers in steady state, replacing Δn by $n_2 - n_1$ and n_0 by $n_1 + n_2$, and furthermore taking into account that for steady state without interaction holds $\Delta n_0 = -n_0$:

$$J\frac{\sigma}{\hbar\omega_{\mathcal{A}}}n_1 = J\frac{\sigma}{\hbar\omega_{\mathcal{A}}}n_2 + \frac{n_2}{T_1} \ . \tag{1.1.72}$$

This equation has the same structure as the Einstein equation. If the monochromatic intensity $J(\omega)$ is replaced by the spectral density ρ_{ω} and integration over the full spectral range is performed, a relation between the Einstein coefficients and the atomic parameters is obtained. These relations read in general for degenerated levels with weighting factors g_1 , g_2 (degeneracies) [92Koe, 81Ver, 00Dav]:

$$B_{12} = \frac{g_2}{12\pi} \frac{|\mu_{\rm A}|^2}{\hbar^2 \varepsilon \varepsilon_0} \,, \tag{1.1.73a}$$

$$B_{21} = \frac{g_1}{12\pi} \frac{|\mu_{\mathcal{A}}|^2}{\hbar^2 \varepsilon \varepsilon_0} \,, \tag{1.1.73b}$$

$$A_{21} = \frac{1}{T_1} = \frac{g_1}{3} \left| \mu_{\mathcal{A}} \right|^2 \frac{\omega_{\mathcal{A}}^3}{\pi \varepsilon \varepsilon_0 \hbar c^3} \,, \tag{1.1.74}$$

 $\mu_{\rm A} = \mu_{12} = \mu_{21}$,

$$\sigma_{21}(\omega) = \frac{\lambda^2}{4} A_{21} h(\omega) , \qquad (1.1.75)$$

$$\sigma_{12}(\omega) = \frac{g_2}{g_1} \sigma_{21}(\omega) , \qquad (1.1.76)$$

$$\sigma_{21}(\omega_{\rm A}) = \frac{\lambda^2}{4\pi} \frac{T_2}{T_1} \le \frac{\lambda^2}{4}$$
 (holds for Lorentzian line shape), (1.1.77)

$$B_{12}g_1 = B_{21}g_2 (1.1.78)$$

$$\frac{A_{21}}{B_{21}} = \frac{2\hbar\,\omega_{\rm A}^3}{\pi\,c^3} \ . \tag{1.1.79}$$

The above relations were derived for isotropic media. Anisotropic media are discussed in [86Sie]. Equation (1.1.80) holds for all dipole transitions, as long as the quantum system is coupled to a large number of modes (free space or a resonator with dimensions large compared with the wavelength). With these equations the gain coefficient can be related to the Einstein coefficient of spontaneous emission [92Koe]:

$$g(\omega) = \frac{\lambda^2}{4} h(\omega, \omega_{\rm A}) \left[n_2 - \frac{g_2}{g_1} n_1 \right] A_{21}$$
 (1.1.80)

with

 $h(\omega, \omega_{\rm A})$: the spectral line shape, depending on the type of broadening (see Sect. 1.1.6).

1.1.5.3 Pumping schemes and complete rate equations

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Till now a two-level system was discussed, assuming a steady-state inversion Δn_0 , which is always negative. To obtain positive inversion $\Delta n = n_2 - n_1 > 0$ and gain, additional levels are necessary.

 $\Delta n > 0$ is a state of non-equilibrium. To support this state, energy has to be pumped into the system. This pumping energy can be incoherent light, kinetic energy of electrons/ions, chemical energy or electric energy. The pumping schemes can become very complicated, and in most cases many energy levels are involved. To understand the principal process for the generation of inversion, two idealized pumping schemes will be discussed.

1.1.5.3.1 The three-level system

The simplified diagram of the three-level system is shown in Fig. 1.1.8. The level E_3 is excited by absorption of light or by electron collisions, depending on the specific system. The decay from E_3 to E_2 , the upper laser level, is very fast. Nearly all excited atoms are transferred into this level, which has a very long life time. If the pumping power is sufficiently high to overcome the decay of level E_2 , atoms will be accumulated and finally n_2 is larger than n_1 . The adiabatic rate equations give for the upper-level population without induced emission between the two levels (J=0):

$$\frac{\mathrm{d}\,n_2}{\mathrm{d}\,t} = W\left(n_0 - n_2\right) - \frac{n_2}{T_1} \ . \tag{1.1.81}$$

W is the pumping rate, the product of the cross-section σ_{13} and the specific pump parameters. T_1 is the upper laser-level lifetime. This holds under the assumption that the population of level E_3 is zero and that $n_1 + n_2 = n_0$. Equation (1.1.81) reads with the inversion density $\Delta n = n_2 - n_1$:

$$\frac{\mathrm{d}\Delta n}{\mathrm{d}t} = W\left(n_0 - \Delta n\right) - \frac{n_0 - \Delta n}{T_1} \tag{1.1.82}$$

and in steady state one obtains:

$$\frac{\Delta n_{\text{steady,3}}}{n_0} = \frac{WT_1 - 1}{WT_1 + 1} \,. \tag{1.1.83}$$

The relation between the inversion density and the pump rate is shown in Fig. 1.1.9. Inversion occurs for $WT_1 > 1$. With increasing pump rate the inversion increases also and approaches finally one, all atoms are in the upper level. To obtain $\Delta n_{\rm steady,3} > 0$ requires at least 50% of the active atoms to be pumped into the upper level, high pump rates are necessary and the efficiency is low. Equation (1.1.82) has to be completed by the coherent interaction term of (1.1.67). The complete rate equation for the three-level system with pump rate W, interacting with a monochromatic field of intensity J is given in (1.1.84). For the intensity (1.1.48c), (1.1.48d) hold, depending on the type of line-broadening (Sect. 1.1.6).

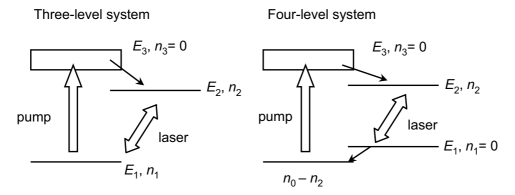


Fig. 1.1.8. The idealized three- and four-level system.

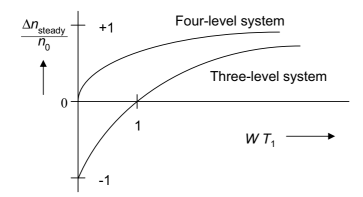


Fig. 1.1.9. Inversion density vs. pump rate for a three- and four-level system.

$$\frac{\partial \Delta n}{\partial t} = -\frac{J}{J_{\rm s}} \frac{f(\omega)}{T_1} \Delta n + W (n_0 - \Delta n) - \frac{n_0 + \Delta n}{T_1}$$
(1.1.84)

(rate equation of a three-level system).

1.1.5.3.2 The four-level system

The commonly used pump scheme, due to its high efficiency, is the four-level system as shown in Fig. 1.1.8. The two laser levels are E_2 and E_1 , where the lower level E_1 has a very short lifetime and its population n_1 is nearly zero. This requires that the energy $E_1 - E_0$ is much larger than the thermal energy κT . The pump level E_3 decays very rapidly to the upper laser level E_2 and its population is again nearly zero. The inversion density now is $\Delta n = n_2 - n_1 \approx n_2$. Then the rate equation for the pump process reads:

$$\frac{\partial \Delta n}{\partial t} = W \left(n_0 - \Delta n \right) - \frac{\Delta n}{T_1} \tag{1.1.85}$$

with the steady-state solution (without coherent interaction):

$$\frac{\Delta n_{\text{steady},4}}{n_0} = \frac{WT_1}{1 + WT_1} \,. \tag{1.1.86}$$

Inversion is reached now at very small pump-power levels as shown in Fig. 1.1.9. The efficiency of such systems is much higher than of three-level systems. The complete rate equation for pumping and interaction with a field of intensity J is obtained by taking into account the corresponding term of (1.1.67). It has to be considered that $n_1 = 0$, and therefore the saturation intensity is higher by a factor of 2.

$$\frac{\partial \Delta n}{\partial t} = -\frac{J}{J_{s,4}} \frac{f(\omega)}{T_1} \Delta n + W(n_0 - \Delta n) - \frac{\Delta n}{T_1}$$
(1.1.87)

(rate equation of a four-level system)

with

 $J_{\mathrm{s},4} = \frac{\hbar \omega_{\mathrm{A}}}{\sigma_0 T_1}$: saturation intensity of the four-level system.

1.1.5.4 Adiabatic pulse amplification

The pulse is adiabatic if its width τ is small compared with T_1 and large compared with T_2 . Then the variation of the upper-level population due to spontaneous emission and pump can be neglected and this term can be neglected. If such a pulse travels through an active medium of length ℓ , it depletes the upper level, is amplified and shaped as depicted in Fig. 1.1.10. The initial conditions at $t=-\infty$ are:

 $\begin{array}{lll} \text{Inversion density:} & \Delta \, n(z) = \Delta \, n_0 \;, & 0 \leq z \leq \ell \;. \\ \text{Input intensity:} & J_0 \;, & z = 0 \;. \\ \text{Input energy:} & E_{\text{in}} \;, & z = 0 \;. \end{array}$

The equations (1.1.67)/(1.1.68) can be solved for a loss-free-medium ($\alpha = 0$) with a four-level system and yield for the output intensity [63Fra]:

$$J_{\text{out}}(t) = J_{\text{in}}(t - \ell/c) \frac{G_0}{G_0 - (G_0 - 1) \exp\left[-\frac{1}{E_s} \int_{-\infty}^{t - \ell/c} J_{\text{in}}(t') dt'\right]}.$$
(1.1.88)

The total output energy density $E_{\rm out}$ of the pulse is

$$E_{\text{out}} = E_{\text{s}} \, \ell \, n \left[1 + G_0 \left(\exp \left(E_{\text{in}} / E_{\text{s}} \right) - 1 \right) \right] \tag{1.1.89}$$

with the two limiting cases

$$E_{\text{out}} = \begin{cases} G_0 E_{\text{in}} , & E_{\text{in}} \ll E_{\text{s}} , \\ E_{\text{in}} + E_{\text{s}} \ell n G_0 = E_{\text{in}} + \frac{\Delta n_0 \ell \hbar \omega_{\text{A}}}{2} , & E_{\text{in}} \gg E_{\text{s}} \end{cases}$$
(1.1.90)

with

 G_0 : small-signal gain factor, (1.1.63), $E_s = J_{s,4}T_1$: saturation energy density, $E_{\rm in,out}$: input/output energy density.

Equations (1.1.88)–(1.1.90) also hold for saturable absorbers with $G_0 < 1$. The pulse will be shaped in any case and the peak velocity will differ from the phase- and group velocities.

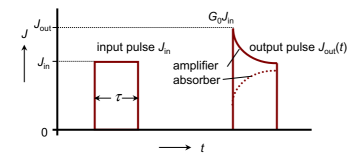


Fig. 1.1.10. Pulse amplification and shaping by a saturable amplifier/absorber.

1.1.5.5 Rate equations for steady-state laser oscillators

In the oscillator system, two counter-propagating traveling waves J^+ , J^- appear, see Fig. 1.1.11, which are amplified by an intensity- and z-dependent gain coefficient according to (1.1.58a), (1.1.58b):

$$\frac{\mathrm{d}J^{+}}{\mathrm{d}z} = [g(J) - \alpha] J^{+}, \qquad (1.1.91a)$$

$$\frac{\mathrm{d}J^{-}}{\mathrm{d}z} = -[g(J) - \alpha] J^{-}. \tag{1.1.91b}$$

For the two traveling waves the boundary conditions at the mirrors are:

$$J^+(z=0) = J^-(z=0)R_1$$
,

$$J^{-}(z=\ell) = J^{+}(z=\ell)R_2$$
.

The combination of (1.1.91a) and (1.1.91b) yields [81Ver]:

$$J^+(z)J^-(z) = \text{const.}$$
,

a useful relation for analytical solutions. The gain coefficient is saturated by both waves. In steady state (1.1.84)/(1.1.87) hold with $J=J^++J^-$, depending on the level system and on the type of broadening. For homogeneous broadening a solution is given in [81Ver]. In general, numerical calculations are necessary. For optimization a diagram is offered in [92Koe]. The intensity rate equations are very useful for laser design and optimization, but deliver no spectral effects such as line width [58Sch, 74Sar, 95Man], mode competition [86Sie, 00Dav], mode hopping [86Sie, 64Lam, 74Sar], or intensity-dependent frequency shifts (Lamb dip) [64Lam]. Multimode oscillation can be described by rate equations with restrictions [64Sta, 63Tan, 93Sve].

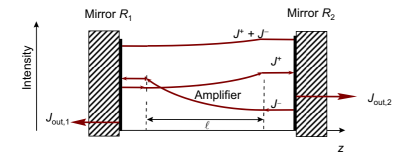


Fig. 1.1.11. The laser oscillator with two counter-propagating waves.

1.1.6 Line shape and line broadening

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1.1.6.1 Normalized shape functions

The line shape depends on the specific interaction process. Two standard line shapes, easy to handle, are the Lorentzian and the Gaussian profiles [92Koe], shown in Fig. 1.1.12. They can be normalized differently.

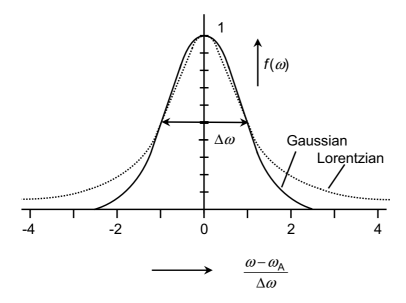


Fig. 1.1.12. Gaussian and Lorentzian line shape.

1.1.6.1.1 Lorentzian line shape

$$f_{\rm L}(\omega, \omega_{\rm A}) = \frac{\left(\Delta \,\omega_{\rm A}/2\right)^2}{\left(\omega - \omega_{\rm A}\right)^2 + \left(\Delta \,\omega_{\rm A}/2\right)^2} \,, \qquad h_{\rm L}(\omega, \omega_{\rm A}) = \frac{2}{\pi \Delta \,\omega_{\rm A}} f_{\rm L}(\omega, \omega_{\rm A}) \,. \tag{1.1.92}$$

1.1.6.1.2 Gaussian line shape

$$f_{\rm G}(\omega) = \exp\left[-\left(\frac{\omega - \omega_{\rm A}}{\Delta \omega_{\rm A}/2}\right)^2 \ln 2\right] , \qquad h_{\rm G}(\omega) = \sqrt{\frac{\ln 2}{\pi}} \frac{2}{\Delta \omega_{\rm A}} f_{\rm G}(\omega) .$$
 (1.1.93)

1.1.6.1.3 Normalization of line shapes

$$f_{\rm G,L}(\omega = \omega_{\rm A}) = 1 \; , \quad f_{\rm G,L}(\omega = \omega_{\rm A} \pm \Delta \, \omega_{\rm A}/2) = 0.5 \; , \quad \int_{-\infty}^{+\infty} h_{\rm G,L}(\omega, \omega_{\rm A}) d\, \omega = 1 \; .$$
 (1.1.94)

1.1.6.2 Mechanisms of line broadening

1.1.6.2.1 Spontaneous emission

The spontaneous emission decay time $T_{\rm sp}$ of quantum dot lasers can be influenced by the geometry [97Scu], but for all macroscopic laser systems it is equal to the free-atom decay and related to the dipole moment (see Sect. 1.1.5.2). The line width of the power spectrum is $\Delta \omega = 1/T_{\rm sp}$. The line shape is Lorentzian for undisturbed systems.

1.1.6.2.2 Doppler broadening

In thermal equilibrium the particles in a gas have a Maxwellian velocity distribution of the velocity v:

$$h(v) = \sqrt{\frac{m_{\rm A}}{2\pi\kappa T}} \exp\left[-\frac{m_{\rm A}v^2/2}{\kappa T}\right]$$
 (1.1.95)

with

 $m_{\rm A}$: atomic mass,

 κT : thermal energy of the particles.

The resonance frequency of a transition is shifted by the Doppler effect

$$\Delta \omega = \omega_{\rm A} v/c_0$$
.

Replacing the velocity in (1.1.76) by the frequency, delivers for the resulting spectral distribution a Gaussian line shape (1.1.74) with the width

$$\frac{\Delta \omega_{\rm D}}{\omega_{\rm A}} = \sqrt{\frac{8 \kappa T \ln 2}{m_{\rm A} c_0^2}} \ . \tag{1.1.96}$$

Some numbers are compiled in Table 1.1.5.

Table 1.1.5. Doppler and collision broadening for a thermal energy of $\kappa T = 1$ eV. The Doppler broadening refers to $\omega_{\rm A} = 10^{15} \; {\rm s}^{-1}$, the collision broadening holds for a pressure of $p = 133 \; {\rm Pa} \; (1 \; {\rm torr}) \; [81 {\rm Ver}, \; 01 {\rm Men}]$.

Gas	Doppler broadening $\Delta \omega_{\rm D} \ [10^{10} \ {\rm s}^{-1}]$	Collision broadening $\Delta \omega_{\rm C} \ [10^7 \ {\rm s}^{-1}]$	
$\overline{\mathrm{H}_2}$	5.6	2.8	
Не	4	1.3	
Ne	1.8	0.8	
CO_2	1.2	1.2	
$ CO_2 $ Ar	1.5	9	

1.1.6.2.3 Collision or pressure broadening

Elastic collisions between radiating atoms imply no energy loss, but a discontinuous jump in the phase of the emitted field. The average temporal length of the wave trains, in the undisturbed case

given by the spontaneous life time $T_{\rm sp}$, is reduced to the collision time τ . The Fourier transform of these shortened waves gives a Lorentzian line shape with the spectral width $\Delta \omega_{\rm C} = 2/\tau$ or

$$\Delta \omega_{\rm C} = \frac{32 \sigma_{\rm C} p}{\sqrt{\pi m_{\rm A} \kappa T}} \tag{1.1.97}$$

with

 $\sigma_{\rm C}$: collision cross section of the atom,

p: pressure of the gas.

The collision broadening is proportional to the gas pressure. Some numbers are given in Table 1.1.5.

1.1.6.2.4 Saturation broadening

A strong field of intensity J, comparable with the saturation intensity J_s , depletes the upper laser level. The gain is reduced according to (1.1.58a), (1.1.58b) and the gain profile becomes flatter and broader with the spectral width (see Fig. 1.1.13) [81Ver]:

$$\Delta\omega_{\rm S} = \Delta\omega_{\rm A}\sqrt{1+J/J_{\rm s}}$$
.

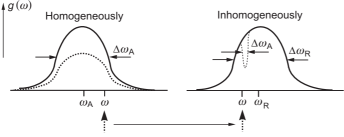
1.1.6.3 Types of broadening

The interaction of the field depends strongly on the type of broadening. Two idealized cases are the homogeneous and the inhomogeneous broadening [00Dav].

1.1.6.3.1 Homogeneous broadening

All transitions have the same resonance frequency ω_A . The gain is saturated for all atoms in the same way as given by (1.1.58a) and shown in Fig. 1.1.13. Examples for this type of broadening are:

- spontaneous emission,
- collision broadening,
- saturation broadening,
- thermal broadening in crystals by interaction with the lattice vibrations.



Frequency of the radiation field ω

Fig. 1.1.13. Saturation of homogeneously and inhomogeneously broadened systems by a radiation field of frequency ω .

1.1.6.3.2 Inhomogeneous broadening

Groups of atoms with spectral density $h(\omega_R, \omega_A)$ and different frequencies ω_A produce a resulting line profile with center frequency ω_R and width $\Delta \omega_R$ as shown in Fig. 1.1.14. A strong monochromatic field of frequency ω interacts mainly with the group $\omega_A = \omega$ and saturates this particular group. A dip appears in the profile, which is called spectral hole-burning. Examples of inhomogeneous broadening are:

- Doppler broadening,
- Stark broadening in crystals due to statistical local crystalline fields.

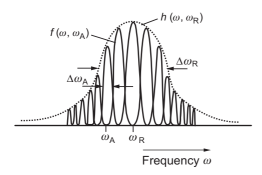


Fig. 1.1.14. An inhomogeneously broadened profile.

The resulting line profile is a convolution of the individual group profiles and the broadening process, which results in complicated integrals. The saturation process for inhomogeneously broadened lines is quite different, as will be shown by a simple example. In this case (1.1.58a) holds only for one group of atoms with the spectral density $h(\omega_A, \omega_R)$. Integration over all groups results in the total gain coefficient $g_{\rm inh}$:

$$g_{\rm inh}(\omega, \omega_{\rm R}) = \int_{-\infty}^{+\infty} f(\omega, \omega_{\rm A}) h(\omega_{\rm A}, \omega_{\rm R}) d\omega_{\rm A} . \qquad (1.1.98)$$

If the width $\Delta \omega_{\rm A}$ is much smaller than the total width $\Delta \omega_{\rm R}$, the function $h(\omega_{\rm A}, \omega_{\rm R})$ can be taken outside of the integral at $\omega_{\rm A} = \omega$. Assuming a Lorentzian profile for the single group, (1.1.98) becomes:

$$g_{\rm inh}(\omega) = \Delta \, n_0 \sigma_0 h(\omega, \omega_{\rm R}) \int \frac{f(\omega, \omega_{\rm A})}{1 + (J/J_{\rm s}) \, f(\omega, \omega_{\rm A})} {\rm d} \, \omega_{\rm A} \label{eq:ginh}$$

and can be integrated:

$$g_{\rm inh}(\omega) = \frac{\Delta n_0 \sigma_0}{\sqrt{1 + J/J_{\rm s}}} h(\omega, \omega_{\rm R}) \frac{\pi \Delta \omega_{\rm A}}{2} = \frac{\Delta n_0 \sigma_0}{\sqrt{1 + J/J_{\rm s}}} f(\omega) \frac{\Delta \omega_{\rm A}}{\Delta \omega_{\rm R}} . \tag{1.1.99}$$

The gain saturates slower than in the case of homogeneous broadening, but the maximum gain is lower by the ratio of the line widths. Inhomogeneous gain profiles can also be caused by spatial hole burning in solid-state laser systems. The standing waves between the mirrors produce an inversion grating and holes in the spectral gain profile [86Sie].

The spectral characteristics of lasers depend strongly on the type of broadening, see Fig. 1.1.15. In steady state the gain compensates losses and the gain profile saturates to fulfill the condition GRV=1. A homogeneously broadened gain profile saturates till the steady-state condition is fulfilled for the central frequency. The bandwidth $\Delta \omega_{L,h}$ is very small and depends on the thermal and

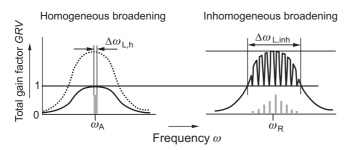


Fig. 1.1.15. Spectrum of an inhomogeneously and homogeneously broadened laser transition in steady state. Total gain factor GRV vs. frequency of the radiation field.

mechanical fluctuations [00Dav]. In the case of solid-state lasers spatial hole burning will influence the spectral behavior and can produce even for homogeneous transitions multi-mode oscillation [63Tan, 66Men]. In the case of inhomogeneous broadening each spectral group of atoms saturates separately and many modes will oscillate, which produces a large lasing bandwidth $\Delta \omega_{L,inh}$. If single-mode operation is enforced by suitable frequency selecting elements, the left \rightarrow right and the right \rightarrow left traveling waves produce two symmetric holes, due to the Doppler effect. This effect can be used for frequency stabilization (Lamb dip [64Lam]).

1.1.6.4 Time constants

The line profile of a real laser transition is in most cases a mixture of homogeneous and inhomogeneous profiles, depending on the temperature and the pressure. The following time constants are used in literature:

 $T_{\rm sp}$: spontaneous life time,

 T_1 : upper-laser-level life time (energy relaxation time, longitudinal relaxation time),

 T_2' : Stochastic processes broaden the line homogeneously. The inverse of the line width is the dephasing time T_2' .

 T_2^* : The line is broadened inhomogeneously. The inverse of this line width $\Delta \omega_R$ is the dephasing time T_2^* .

 T_2 : For the resulting dephasing time (transverse relaxation time, entropy time constant), approximately holds (depends on the line profiles):

$$\frac{1}{T_2^2} \approx \frac{1}{{T_2^*}^2} + \frac{1}{{T_2'}^2} \; . \label{eq:T22}$$

Some examples of decay times are given in Table 1.1.6.

1.1.7 Coherent interaction

If the interaction time of the radiation field with the two-level system is small compared with all relaxation times, including the pump term, the stochastic processes can be neglected and

Table 1.1.6. Spontaneous life time $T_{\rm sp}$, upper-laser-level life time T_1 , transverse relaxation time T_2 , homogeneous relaxation time T_2' and inhomogeneous relaxation time T_2^* [01Iff, 92Koe, 86Sie], [01Men, Chap. 6].

	$T_{\rm sp} \ [{\rm s}]$	T_1 [s]	T_2 [s]	T_2' [s]	T_2^* [s]
Neon-atom (He/Ne-laser), $\lambda_0=632.8$ nm, He ($p=130$ Pa), Ne ($p=25$ Pa)	10^8	10 ⁻⁸	3×10^{-9}	10^{-8}	4×10^{-9}
Chromion-ion, $\lambda_0 = 694.3$ nm, R_1 -transition in ruby					
T = 300 K T = 4 K		3×10^{-3} 4×10^{-3}	$10^{-12} \\ 2 \times 10^{-7}$	$10^{-12} \\ 3 \times 10^{-3}$	$2 \times 10^{-7} \\ 2 \times 10^{-7}$
SF ₆ -molecule, $\lambda_0=10.5~\mu\mathrm{m},$ $p=0.4~\mathrm{Pa}$	10^{-3}	10^{-3}	6×10^{-9}	7×10^{-6}	6×10^{-9}
Rhodamin-molecule in ethanol, singlet-transition, $\lambda_0 = 570.0$ nm	5×10^{-9}	5×10^{-9}	10^{-12}	10^{-12}	-
Neodymium-ion in YAG-crystal, $\lambda_0 = 1060 \text{ nm}, T = 300 \text{ K}$	5×10^{-4}	2.3×10^{-4}	7×10^{-12}	_	_
· · · · · · · · · · · · · · · · · · ·	·	·	·		

(1.1.45a)/(1.1.45b) hold. This kind of coherent interaction is of strong interest in nonlinear spectroscopy [84She, 86Sie, 71Lam, 72Cou], [01Men, Chap. 7] and confirmed by many experiments. Examples of nonlinear coherent interaction are transient response of atoms, optical nutation, photon echoes, $n\pi$ -pulses and quantum beats. Here only some very simple examples will be presented. A more detailed treatment is given in [95Man].

1.1.7.1 The Feynman representation of interaction

A very compact description of the two-level interaction was given by Feynman [57Fey]. The real electric field is

$$\boldsymbol{E}_{\mathrm{real}} = \frac{1}{2} \left\{ \boldsymbol{E}_0 \exp \left[\mathrm{i} \left(\omega t - kz \right) \right] + \boldsymbol{E}_0^* \exp \left[-\mathrm{i} \left(\omega t - kz \right) \right] \right\} \; .$$

It generates a real polarization, (1.1.23), shifted in phase against the field:

$$\mathbf{P}_{A,\text{real}} = \frac{1}{2} \left\{ \mathbf{P}_{A0} \exp\left[i(\omega t - kz)\right] + \mathbf{P}_{A0}^* \exp\left[-i(\omega t - kz)\right] \right\}$$
$$= \mathbf{C} \cos\left(\omega t - kz\right) + \mathbf{S} \sin\left(\omega t - kz\right)$$
(1.1.100)

with C, S real vectors:

$$m{C} = rac{1}{2} \left(m{P}_{
m A0} + m{P}_{
m A0}^*
ight) \; , \quad m{S} = rac{1}{2} {
m i} \, \left(m{P}_{
m A0} - m{P}_{
m A0}^*
ight) \; .$$

In the following an isotropic medium is assumed. Then μ_A , P_A and E are parallel and can be treated as scalars. With these new real quantities the equations of interaction (1.1.45a), (1.1.45b) become:

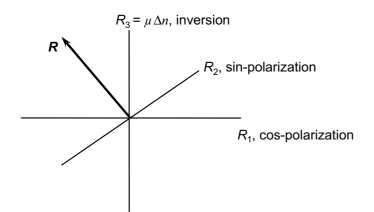


Fig. 1.1.16. In the case of coherent interaction, the system is characterized by its R-vector which rotates in the polarization/inversion space with constant length.

$$\frac{\partial C}{\partial t} = -\delta S + i \,\mu_{A} \cdot \Delta \, n \, \left(\frac{\Lambda - \Lambda^{*}}{2}\right) \,, \tag{1.1.101a}$$

$$\frac{\partial S}{\partial t} = \delta C - \mu_{A} \cdot \Delta n \left(\frac{\Lambda + \Lambda^{*}}{2} \right) , \qquad (1.1.101b)$$

$$\mu_{\mathcal{A}} \frac{\partial \Delta n}{\partial t} = -i C \left(\frac{\Lambda - \Lambda^*}{2} \right) + S \left(\frac{\Lambda + \Lambda^*}{2} \right) , \qquad (1.1.101c)$$

where Λ is a complex quantity. Its modulus is called the Rabi frequency:

$$\Lambda(z,t) = \frac{\mu_{\rm A} E_0}{\hbar} , \quad |\Lambda| : \text{ Rabi frequency }. \tag{1.1.102}$$

Two vectors $\boldsymbol{R},\,\boldsymbol{F}$ are introduced:

$$\mathbf{R} = (C, S, \mu_{\rm A} \, \Delta \, n) = (R_1, R_2, R_3) \,, \quad \mathbf{F} = \left(\frac{\Lambda + \Lambda^*}{2}, \, \mathrm{i} \, \frac{\Lambda - \Lambda^*}{2}, \, \delta\right) = (F_1, F_2, F_3) \,.$$

The R-vector characterizes the state of the two-level system and can be depicted in an inversion/polarization space, as shown in Fig. 1.1.16. R corresponds to the Bloch vector of the spin-1/2 system [46Blo]. The equations (1.1.101a), (1.1.101b) of interaction can be condensed to:

$$\frac{\partial \mathbf{R}}{\partial t} = [\mathbf{F} \times \mathbf{R}]$$
 (coherent interaction). (1.1.103)

Scalar multiplication of this equation with R results in:

$$\left\langle \boldsymbol{R} \frac{\partial \boldsymbol{R}}{\partial t} \right\rangle = \left\langle \boldsymbol{R} \left[\boldsymbol{F} \times \boldsymbol{R} \right] \right\rangle = 0$$

which means that the length of the vector is constant during interaction:

$$|C|^2 + |S|^2 + |\mu_A \Delta n|^2 = |\mathbf{R}_0|^2. \tag{1.1.104}$$

The tip of the vector moves on a sphere in the inversion/polarization space with complicated trajectories [69McC, 74Sar, 69Ics]. The incoherent relaxation and pumping of the system can be included in (1.1.103) by an additional relaxation term [72Cou].

1.1.7.2 Constant local electric field

$$E_0$$
 , E_0 , E_0

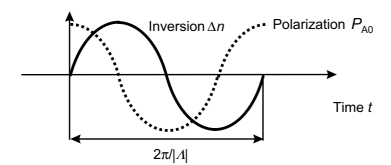


Fig. 1.1.17. Oscillation of inversion density Δn and polarization amplitude $P_{\rm A0}$ in resonance for a constant local electric field.

For a constant electric field at a fixed position z the rotating-wave approximation has a periodic solution. Inversion and polarization with the initial condition t = 0, $\Delta n = n_0$, $P_{A0} = 0$ are:

$$\frac{\Delta n}{n_0} = \frac{\delta^2 + |\Lambda|^2 \cos \beta t}{\beta^2} , \quad \Lambda = \frac{\mu_A E_0}{\hbar} , \qquad (1.1.105)$$

$$P_{A0} = n_0 \frac{\mu_A \Lambda}{\beta} \left[\frac{\delta}{\beta} \left(1 - \cos \alpha t \right) + i \sin \alpha t \right], \quad \beta = \sqrt{\delta^2 + |\Lambda|^2}.$$
 (1.1.106)

In resonance $\omega = \omega_{\rm A}$, $\delta = 0$, the inversion density Δn and the amplitude $\boldsymbol{P}_{\rm A0}$ of the polarization oscillate with this frequency, see Fig. 1.1.17. The real polarization $\boldsymbol{P}_{\rm A,real}$ of (1.1.100) contains the frequencies $\omega_{\rm A} \pm |\Lambda|$. Some values of dipole moments are given in Table 1.1.2 to estimate $|\Lambda|$. Off resonance the temporal behavior of inversion and polarization is more complicated (optical nutation) [72Cou]. If at t=0 all atoms are in the lower level ($\Delta n_0 = -n_0$) a complete inversion is produced at $t=\pi/|\Lambda|$ by a coherent field. It is called pulse inversion [60Vuy]. At $t=2/|\Lambda|$, all atoms are again in the lower level, no energy transfer has taken place.

1.1.7.3 Propagation of resonant coherent pulses

$$=_{i_{1}, i_{2}, i_{3}, i_{4}, i_{5}} au < T_{2}, \ldots, i_{i_{1}, i_{2}, i_{4}, i_{5}, i_{5},$$

The propagation of pulses in a two-level system is described by the rotating-wave approximation, (1.1.45a)/(1.1.45b), and by the wave equation in the SVE approximation (1.1.28). The set of these three non-linear equations is difficult to solve, only special cases will be discussed here. At t=0 the electric field E_0 is assumed to be real, $\Lambda = \Lambda^*$. In case of resonance, $\delta = 0$, (1.1.101a) delivers C = 0, $R_1 = 0$. The interaction equations (1.1.101b), (1.1.101c) reduce to

$$R_1 = 0 ,$$

$$\frac{\partial R_2}{\partial t} = -\Lambda R_3 ,$$

$$\frac{\partial R_2}{\partial t} = -\Lambda R_3 ,$$

$$\frac{\partial R_3}{\partial t} = \Lambda R_2 \ .$$

The R-vector moves in the R_2 - R_3 -plane, see Fig. 1.1.18. If the angle θ with the R_3 -axis is introduced, one solution of the above equations is:

$$R_2 = R_0 \sin \theta$$
,

$$R_3 = -R_0 \cos \theta$$

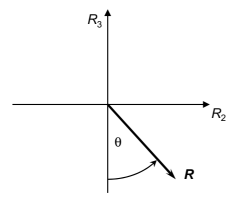


Fig. 1.1.18. In resonance, $\delta = 0$, the **R**-vector of the two-level system rotates in the R_2 - R_3 -plane.

with

$$\Lambda = \frac{\partial \theta}{\partial t} = \frac{\mu_{\rm A} E_0}{\hbar} \ . \tag{1.1.107}$$

 R_0 is given by the initial conditions at t=0. The SVE-approximation of (1.1.28) then becomes:

$$\left(\frac{\partial^2}{\partial t \,\partial z} + \frac{1}{c}\frac{\partial^2}{\partial t^2}\right)\theta = -\frac{\alpha}{2}\frac{\partial\theta}{\partial t} + \frac{\gamma}{2}R_0\sin\theta, \quad \gamma = \frac{\mu_A \,k_0}{n_T \,\varepsilon_0 \,\hbar}.$$
(1.1.108)

From θ the amplitude E_0 of the electric field can be calculated with (1.1.107)/(1.1.105).

1.1.7.3.1 Steady-state propagation of $n\pi$ -pulses

Steady state means that a pulse is propagating with velocity v and constant pulse envelope $E_0(t,z) = E_0(t-z/v)$. The amplitude depends on one parameter w only:

$$w = t - z/v$$

and (1.1.108) becomes:

$$\left(1 - \frac{c}{v}\right) \frac{\mathrm{d}^2 \theta}{\mathrm{d} w^2} + \frac{\alpha c}{2} \frac{\mathrm{d} \theta}{\mathrm{d} w} = c \frac{\gamma}{2} R_0 \sin \theta .$$
(1.1.109)

This equation is equivalent to the equation of the pendulum with friction in a gravitational field. In the following examples two different initial conditions are assumed:

$$R_0 = \mu_{\rm A} \, \Delta \, n_0 \left\{ > 0 \quad \text{(amplifier)}, \\ < 0 \quad \text{(absorber)}, \right.$$

which corresponds to the pendulum up or down at t = 0.

1.1.7.3.1.1 2π -pulse in a loss-free medium

A medium without losses ($\alpha=0$) interacts with a coherent pulse in resonance ($\delta=0$). The initial condition is $\Delta n_0(t=-\infty)=+\Delta n_0$ ($\Delta n_0<0$, absorber). One steady-state solution is the 2π -pulse, see Fig. 1.1.19, which corresponds to a local field of duration $\tau=2\pi/\Lambda$. The leading edge of the pulse produces an inversion and energy is transferred to the atomic system, the amplitude is reduced. The trailing part of the pulse is then amplified by this inversion. In total the pulse

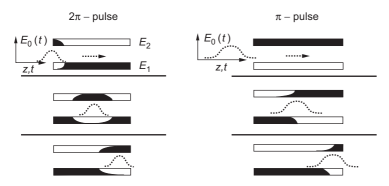


Fig. 1.1.19. Propagation of 2π - and π -pulses in a two-level system.

has lost no energy, but is delayed in time. Such a pulse is only stable, if the broadband losses are negligible and if the initial inversion is negative. The steady-state solution is:

$$E = E_{\text{peak}} \frac{\exp\left[i\omega \left(t - z/c\right)\right]}{\cosh\left[\left(t - z/v\right)/\tau\right]} \quad \text{(field)} , \tag{1.1.110}$$

$$E_{\text{peak}} = 2\sqrt{\hbar \omega} \sqrt{\frac{\Delta n_0}{\varepsilon_0 (1 - c/v)}}$$
 (peak amplitude), (1.1.111)

$$J_{\text{peak}} = \frac{2 \hbar \omega \Delta n_0 c}{1 - c/v} \quad \text{(peak intensity)} , \qquad (1.1.112)$$

$$T_{2\pi} = 2\tau = 2\sqrt{\frac{(1-c/v)T_2}{g_0c}}$$
 (pulse duration), (1.1.113)

$$v = \frac{c}{1 - g_0 c \tau^2 / T_2} \quad \text{(pulse peak velocity)} \tag{1.1.114}$$

with

 $g_0 = \Delta n_0 \sigma < 0$: small-signal absorption coefficient,

c: phase velocity in the medium,

v: pulse peak velocity.

This two-level system is the most simple model of a saturable absorber, which in the case of incoherent interaction absorbs the radiation. But the coherent 2π -pulse transmits the absorber without losing energy. Therefore this effect is called self-induced transparency [75Kri]. The pulse is characterized by three parameters: peak velocity v, peak amplitude E_{peak} and the width $T_{2\pi}$. One of these parameters can be chosen arbitrarily, the other two result from (1.1.112)/(1.1.113)/(1.1.114). But the interaction is coherent only as long as $T_{2\pi} \ll T_2$.

1.1.7.3.1.2 π -pulse in an amplifying medium

A steady-state solution in an amplifying medium, initial condition $\Delta n(t = -\infty) = \Delta n_0 > 0$, with broadband losses ($\alpha \neq 0$) is the π -pulse [74Loy], see Fig. 1.1.19:

$$E = E_{\text{peak}} \frac{\exp\left[i\omega\left(t - z/c\right)\right]}{\cosh\left[(t - z/c)/\tau\right]} \quad \text{(field)} , \tag{1.1.115}$$

$$E_{\text{peak}} = \frac{\hbar}{\tau \,\mu} \quad \text{(peak amplitude)} \,, \tag{1.1.116}$$

$$J_{\text{peak}} = \frac{\hbar \,\omega}{2\sigma_0 T_2} \left[\frac{g_0}{\alpha} \right]^2 \quad \text{(peak intensity)} \,, \tag{1.1.117}$$

$$T_{\pi} = 2\tau = 2 T_2 \frac{\alpha}{g_0} \quad \text{(pulse duration)}. \tag{1.1.118}$$

The pulse propagates approximately with c, depletes at each position the upper level, and converts this energy via the broadband losses α into heat. The saturated gain just compensates the losses. The pulse is only stable for $\alpha > 0$ and $g_0 > 0$.

So far solutions of the steady-state SVE-equation were presented, assuming resonance and a homogeneously broadened two-level system. Off-resonance interaction and inhomogeneously broadened systems are much more complicated and are discussed in detail in the literature [74Sar, 69Ics, 72Cou]. Moreover, the stability of the pulses with respect to small perturbations was not yet mentioned. It is controlled by the area theorem [67McC, 74Sar].

1.1.7.3.2 Superradiance

The spontaneous emission was neglected in the coherent interaction. An initial state, $\mathbf{R} = (0, 0, \mu \Delta n)$, complete inversion, without external field \mathbf{F} would be stable according to the interaction equations (1.1.103). But due to spontaneous emission and amplified spontaneous emission, the \mathbf{R} -vector will be pushed a bit out of equilibrium and decay into the stable position $\mathbf{R} = (0, 0, -\mu \Delta n)$. This phenomenon is called superradiance and discussed in detail in Chap. 6.2.

1.1.8 Notations

Symbol	Unit	Meaning
\underline{A}_{21}	s^{-1}	Einstein coefficient of spontaneous emission
B	$ m Vs/m^2$	magnetic induction
B_{12}, B_{21}	$\mathrm{m^3/VAs^3}$	Einstein coefficient of induced emission
\boldsymbol{C}	$\mathrm{As/m^2}$	component of the Feynman vector \boldsymbol{R}
c_0	m/s	vacuum velocity of a plane wave
c	m/s	phase velocity of light in a medium
$c_{1,2}$	_	coefficients of the eigenvector
D	$\mathrm{As/m^2}$	electric displacement
$oldsymbol{E}$	V/m	electric field
$oldsymbol{E}_0$	m V/m	electric-field amplitude
$E_{1,2}$	VAs	energy eigenstates of the two-level system
$E_{ m in}$	VAs	amplifier input energy
$E_{ m out}$	VAs	amplifier output energy
$E_{ m S}$	$ m VAs/m^2$	amplifier saturation energy density
$f(\omega,\omega_{ m A})$	_	line shape factor
G	_	gain factor
G_0	_	small-signal gain factor
g	m^{-1}	gain coefficient
g_0	m^{-1}	small-signal gain coefficient
$g_{1,2}$	_	degeneracies of lower/upper laser level

 $1.1.8 \; \mathrm{Notations} \qquad \qquad [\mathrm{Ref.} \; \mathrm{p.} \; 40$

$g_{ m h}$	m^{-1}	gain coefficient of a homogeneously broadened
		transition
$g_{ m inh}$	m^{-1}	gain coefficient of an inhomogeneously broadened
-		transition
H	$\mathrm{A/m}$	magnetic field
$oldsymbol{H}_0$	A/m	magnetic-field amplitude
H_0	VAs	Hamilton operator of the undisturbed transition
$H_{ m int}$	VAs	Hamilton operator of interaction
$egin{aligned} h(\omega, \omega_{ ext{A}}) \ oldsymbol{j} \end{aligned}$	S	line shape factor
$oldsymbol{j}$	$\mathrm{A/m^2}$	current density
J	Vs/m^2	magnetic polarization
J	$ m VA/m^2$	intensity
J^{+}, J^{-}	VA/m^2	intensity inside the resonator
$J_{ m s},\ J_{ m s4}$	VA/m^2	saturation intensity of 2-, 3- and 4-level system
k	m^{-1}	wave number
\boldsymbol{k}	m^{-1}	wave vector inside the medium
$oldsymbol{k}_0$	m^{-1}	wave vector in vacuum
ℓ	m	geometrical length of the active medium
n	_	complex refractive index
$n_{ m r}$	_	real refractive index
n_0	m^{-3}	density of active atoms
$n_{1,2}$	m^{-3}	density of lower/upper population
$oldsymbol{P}_{ ext{A,real}}^{ ext{real}}$	$\mathrm{As/m^2}$	real polarization of the active atoms
$oldsymbol{P}_{ m A}^{ m A,rear}$	$\mathrm{As/m^2}$	complex polarization of the active atoms
$oldsymbol{P}_{ m A0}^{\Lambda}$	$\mathrm{As/m^2}$	amplitude of the complex polarization
$oldsymbol{P}_{ m H}$	$\mathrm{As/m^2}$	complex polarization of the host material
R	As/m^2	Feynman vector
R	_	$=\sqrt{R_1 R_2}$, average mirror reflectivity
$R_{1,2}$	_	reflectivity of mirror 1, 2
r	m	position vector
$oldsymbol{S}$	$ m VA/m^2$	Poynting vector
T_1	S	upper-laser-level life time
T_2'	S	dephasing time due to homogeneous broadening
T_2^*	S	dephasing time due to inhomogeneous broadening
T_2	S	resulting dephasing time
$T_{ m sp}$	S	spontaneous decay time
T_{π} , $T_{2\pi}$		pulse duration of π -, 2π -pulses
$V^{\pi, 12\pi}$	s _	resonator loss factor per transit
•	$\mathrm{m/s}$	pulse peak velocity
$\stackrel{v}{Z}$	V/A	impedance
	•	
Z_0	V/A	vacuum impedance
O/	m^{-1}	absorption coefficient
α		susceptibility of the active atoms
$\chi_{\rm A}$	_	- *
χe	_	electric susceptibility
XH	_	susceptibility of the host material
χ _m	$^{-}$ s ⁻¹	magnetic susceptibility
δ		detuning
$\frac{\Delta}{\Lambda}n$	m^{-3}	inversion density
$\Delta_{ m tr}$	m^{-2}	transverse delta-operator
$\Delta \omega_{\mathrm{A}}$	s^{-1}	line width of homogeneous broadening
$\Delta\omega_{ m C}$	s^{-1}	line width of collision broadening

$\Delta\omega_{ m R}$	s^{-1}	line width of inhomogeneous broadening
$\Delta\omega_{ m S}$	s^{-1}	line width of saturation broadening
$\Delta \omega_{ m L,inh}, \Delta \omega_{ m L,h}$	s^{-1}	lasing bandwidth of inhomogeneous/homogeneous
, . ,		transitions
arepsilon	_	permittivity
$arepsilon_0$	$8.8542 \times 10^{-12} \text{ As/Vm}$	electric constant
arphi angle	_	state vector of the two-level system
$ \varphi_{1,2}\rangle$	_	eigenfunctions of the two-level system
κ	$1.38 \times 10^{-23} \text{ VAs}^2/\text{K}$	Boltzmann's constant
λ_0	m	vacuum wavelength
Λ	s^{-1}	Rabi frequency
μ	_	permeability
μ_0	$4\pi \times 10^{-7} \mathrm{Vs/Am}$	magnetic constant
$oldsymbol{\mu}_{12},oldsymbol{\mu}_{21}$	Asm	$=\mu_{\rm A}$, dipole moment of the two-level transition
$oldsymbol{\mu}_{ ext{A}}$	Asm	dipole moment of the two-level transition
θ	_	beam divergence, slope of the Feynman vector
$ ho_\omega$	$ m VAs^2/m^3$	spectral energy density (per $d\omega$)
$\sigma(\omega)$	m^2	cross section of the two-level system
$\sigma_{ m e}$	A/Vm	electric conductivity
σ_0	m^2	cross section of the two-level system in resonance
au	s	pulse width
ω	s^{-1}	frequency of the radiation field
$\omega_{ m A}$	s^{-1}	resonance frequency of the homogeneously
		broadened transition
$\omega_{ m R}$	s^{-1}	resonance frequency of the inhomogeneously
		broadened transition

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2.1 Definition and measurement of radiometric quantities

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2.1.1 Introduction

Radiometry is the science and technology of the measurement of electromagnetic energy. Here we confine ourselves on the subfield of optical radiometry which covers the measurement of electromagnetic radiation in the wavelength range from about $0.01~\mu m$ to $1000~\mu m$. Radiometric quantities are derived from the quantity energy. The corresponding photometric quantities on the other hand involve the additional evaluation of the radiant energy in terms of a defined weighting function, usually the standard photometric observer. In the following only the definitions of the radiometric quantities are explained in detail. Starting from the radiant energy the other fundamental radiometric quantities radiant power, radiant excitance, irradiance, radiant intensity, and radiance are derived by considering the additional physical quantities time, area, and solid angle.

The radiometric quantities defined in abstract terms are practically embodied by radiometric standards. Radiometry is based on primary detector standards and primary source standards. Primary detector standards are mostly electrical-substitution thermal detectors whereas for primary source standards the emitted radiant power is accurately calculable. For the radiometric measurement of cw laser emission radiation detectors or radiometers calibrated against primary detector standards are the preferred secondary standards. The detection principle of the radiometers could be thermal (thermopiles, bolometers, and pyroelectric detectors) or photoelectric (semiconductors). As secondary standards for pulsed laser radiation mostly thermally absorbing glass-disk calorimeters are used. These standards are derived from the cw standards using accurately measured shuttering of the laser radiation to produce pulses of known radiant energy.

2.1.2 Definition of radiometric quantities

Radiometric and photometric quantities are represented by the same principal symbol and may be distinguished by their subscripts. While radiometric quantities either have the subscript "e" or no subscript (as in the whole Chap. 2.1), photometric quantities have the subscript "v", where "e" stands for "energetic" and "v" for "visible". The most frequently used radiometric quantities are listed in Table 2.1.1 together with their symbols, defining equations, and units. The additional physical quantities applied in Table 2.1.1 are the time t, the element of solid angle $d\omega$, and the angle θ between the line of sight and the normal of the radiating or receiving surface with the area element dA, see Fig.2.1.1.

In the case that the quantities are functions of wavelength their designations must be preceded by the adjective "spectral". For example, the symbol for spectral radiance is $L(\lambda)$. This has to be well distinguished from the convention for the spectral concentration of a quantity, which is also preceded by the adjective "spectral". In that case, however, the symbol has the subscript λ , i.e. $\mathrm{d}L/\mathrm{d}\lambda = L_{\lambda}$.

Quantity	Symbol	Defining equation	Unit
Radiant energy	Q		J
Radiant power	Φ	$\Phi = \mathrm{d}Q/\mathrm{d}t$	W
Radiant excitance	M	$M = \mathrm{d}\Phi/\mathrm{d}A$	${ m W~m^{-2}}$
Irradiance	E	$E = d\Phi/dA$	$\mathrm{W}~\mathrm{m}^{-2}$
Radiant intensity	I	$I = \mathrm{d}\Phi/\mathrm{d}\omega$	${ m W~sr}^{-1}$
Radiance	L	$L = d^2 \Phi / (\cos \theta dA d\omega)$	${ m W} { m m}^{-2} { m sr}^{-1}$

Table 2.1.1. Radiometric quantities, their defining equations and units.

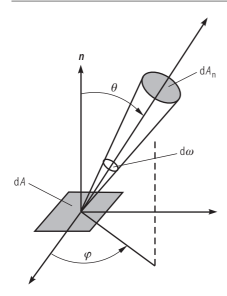


Fig. 2.1.1. Geometry for definition of the radiance.

To explain the defining equations given in Table 2.1.1 a radiation source of finite extent is considered. If we surround the radiation source with a closed surface and calculate the radiant energy Q penetrating the surface per unit time we get the total radiant power Φ emitted by the source. For clarity, the above mentioned symbols for the spectral properties of the radiation are omitted in this chapter. The radiant power per unit area of the radiation source associated with the emission into the hemispheric space above dA is defined as the radiant excitance M. At this point it is appropriate to introduce the radiation incident from all directions in the hemispheric space above the surface of a detector. The irradiance E is defined as the radiant power incident on a surface per unit area of the surface. The irradiance represents also the energy which propagates per unit time through the unit area perpendicular to the direction of energy transport. This is known as the density of energy flow identical to the magnitude of the Poynting vector averaged over time.

Coming back to the source-based radiometric quantities we consider now the radiant power proceeding from a point source per unit solid angle $d\omega$ in a specified direction. The corresponding quantity appropriate especially for nearly point-shaped sources is denoted as radiant intensity I. If we generalize and consider again a source of finite extent the directional nature of radiation has to be taken into account accurately. From Fig. 2.1.1 we formally define as radiance L the radiant power emitted in the (θ, φ) direction, per unit area of the surface normal to this direction and per unit solid angle. Note that the area dA_n used to define the radiance is the component of dA perpendicular to the direction of the radiation. This projected area is equal to $\cos\theta dA$ and in effect, this is how dA would appear to an observer situated on the surface in the (θ, φ) direction.

Although the directional distribution of surface emission varies according to the nature of the surface, there is a special case which provides a reasonable approximation for many surfaces. For

an isotropically diffuse emitter the radiance is independent of direction:

$$L(\theta, \varphi) = L . \tag{2.1.1}$$

Such an emitter is denoted as a lambertian radiator which emits in accordance with Lambert's cosine law:

$$I(\theta) = I(0)\cos\theta. \tag{2.1.2}$$

The radiant intensity of a perfectly diffuse surface element in any direction varies as the cosine of the angle between that direction and the normal to the surface element. It is noted that this law is consistent with the definitions of radiance and radiant intensity given in Table 2.1.1. It may be helpful to derive the relationship between radiance and radiant excitance for a lambertian radiator. The radiant excitance into the hemispheric space above dA is calculated from the radiance by integration over the solid angle $d\omega = \sin\theta \ d\theta \ d\varphi$:

$$M = \frac{\mathrm{d}\,\Phi}{\mathrm{d}A} = \int_{0}^{2\pi} \int_{0}^{\pi/2} L(\theta,\varphi) \,\cos\theta \,\sin\theta \,\mathrm{d}\theta \,\mathrm{d}\varphi \,. \tag{2.1.3}$$

By removing $L(\theta, \varphi)$ from the integrand according to (2.1.1) and performing the integration we get

$$M = \pi L . (2.1.4)$$

Note that the constant appearing in the above expression is π , not 2π , and has the unit steradian (sr).

2.1.3 Radiometric standards

2.1.3.1 Primary standards

Depending on the application primary source and primary detector standards are used to establish radiometric scales. Black-body radiators of known temperature with calculable spectral radiance are operated as primary source standards at temperatures up to about $3200~\rm K$ [96Sap]. Due to the steep decrease of their Planckian radiation spectrum in the UV spectral range radiometry with black-body radiators is limited to wavelengths above $200~\rm nm$. In comparison with a black-body radiator, the maximum of the synchrotron radiation spectrum emitted by an electron storage ring is shifted to shorter wavelengths by several orders of magnitude [96Wen]. In a storage ring electrons move with nearly the velocity of light along a circular trajectory and emit a calculable radiant power through an aperture stop situated near the orbital plane. Radiometry can thus be extended into the X-ray region up to photon energies of $100~\rm k\,eV$.

Electrical-substitution thermal detectors operated at ambient temperature have been the most frequently used primary detector standards. However, their performance is limited by the thermal properties of materials at room temperature resulting in complicated corrections that have to be applied. Hence, their uncertainties remain near 0.1~% to 0.3~% [89Fro, 79Wil]. Cryogenic radiometers have been developed to satisfy the increasing demands for more accurate detector standards from users especially in new and expanding fields of optical fibers, laser technology, and space science. Today, these instruments with absorption cavities at nearly the temperature of liquid helium

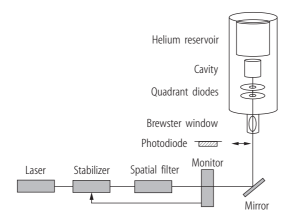


Fig. 2.1.2. Cryogenic radiometer for the calibration of photodiodes. The stabilized laser beam enters the cryostat via a Brewster window and is aligned by quadrant photodiodes. In the cavity the laser radiation is absorbed and electrically substituted.

are the most accurate among all primary standards, with relative uncertainties of less than 0.01 % [85Qui, 96Fox]. The principle of operation of both the cryogenic radiometers and the instruments at ambient temperature is that a thermometer measures the temperature rise of an absorption cavity, relative to a constant-temperature heat sink, during radiant and electrical heating cycles. By adjusting the electrical power so that the absorption cavity temperature rise is the same for both types of heating, the radiant power can be equated to the easily measured quantity of electrical power. For cryogenic radiometers the corrections due to the limited absorptance of the cavity, the lead heating of electrical connections, the radiative heat loss, and the background radiation can be made sufficiently small to reach very accurate equivalence of optical and electrical heating [96Fox]. Today, high-precision calibrations of laser radiometry secondary standards are mostly traceable to cryogenic radiometers. In Fig. 2.1.2 a typical experimental arrangement for the calibration of transfer photodiodes is shown [93Fu].

2.1.3.2 Secondary standards

Secondary standards serve to disseminate a metrologic scale or quantity to the user in science and industry. In this section, first, the common detectors used in the secondary standards for laser radiometry are shortly described, and second, some examples for laser radiometers and calorimeters are given. The detection principle of the secondary standards is usually thermal or photoelectric. The thermal detectors have the remarkable advantage of a flat spectral responsivity function which makes the calibration for different laser wavelengths not necessary or at least easier compared to that of photoelectric detectors. Among the thermal detectors we distinguish between thermopile detectors, bolometric and pyroelectric detectors.

A thermopile consists of a number of thermocouples in series to provide a thermoelectric voltage proportional to the temperature difference between the receiver and its thermal environment. Its optimization in detector applications has received considerable attention [68Smi, 58Sch, 70Ste]. At this point the term responsivity s is introduced which is the ratio of the detector output to the detector input. Whereas the detector input is a radiometric quantity, the detector output is usually an electrical quantity, for example current, voltage, or change in resistance. In order to optimize the responsivity of a thermopile one has to maximize the Seebeck coefficient of the two materials used for each thermocouple, the thermal resistance between the receiver and the environment, and the absorptance of the surface. The materials used for thermocouples are either metals, alloys, or semiconductors, for examples see [89Hen].

A bolometer is a temperature transducer based on the change of electrical resistance with temperature. The important quantity is the temperature difference between the receiver and its

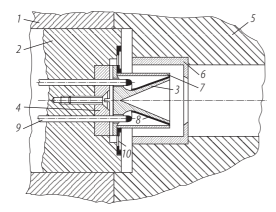


Fig. 2.1.3. Cross section of a cone-shaped laser radiometer. 3: blackened cone, 6: aperture, 7: heat protection tube, 8: electrical heater, 9: electrical connections, 10: thermopile; 1, 2, 4, 5: parts of the heat sink.

thermal environment. Therefore one resistance element is needed to measure the temperature of the receiver and one to measure that of the thermal environment. AC and DC bridge techniques are applied for the comparison, the most common employing Wheatstone bridge configurations. The second resistance element should be physically close to the radiation-measuring element to compensate for convective disturbances, pressure fluctuations, changes in temperature of the housing, and instabilities in the bridge supply. The resistors are preferably made of metal wires or films of nickel, platinum, or gold [65Ble]. Thermistors are also used which have a larger temperature coefficient of the resistance. At lower operation temperatures the signal-to-noise ratio of bolometers can be increased considerably [82Mat, 87McD].

Pyroelectric detectors produce a current proportional to the rate of temperature change. The detection mechanism is based on the temperature dependence of the electrical polarization in ferroelectric crystals. Since pyroelectric detectors respond to modulated radiant power only, their use in laser radiometers for measuring cw radiation requires chopping of the incident beam. This can provide considerable drift immunity and allows for the use of drift-free AC amplification techniques [70Put, 75Tif].

Beside the thermal detectors also photoelectric devices or quantum detectors are used in laser radiometry. Photoelectric detectors for laser radiometric applications are either photoconductors or photodiodes. In a photoconductor made of a thin film of a semiconductor material the incident radiation generates additional carriers. These intrinsic band-to-band transitions or extrinsic transitions involving forbidden-gap energy levels result in an increase of conductivity [81Sze]. For sensitive infrared detection, the photoconductor must be cooled in order to reduce thermal ionization of the energy levels. In photodiodes the carriers are mainly generated in the depletion layer of the diode junction. The electron-hole pairs separated by an internal or external electric field recombine by driving an external current. Photodiodes are operated in two different modes: In the photovoltaic mode no bias voltage is applied and the photodiode can be considered as current source. In contrast, in the reverse-bias mode the photocurrent generates a voltage drop at an external load resistance which is used as measuring quantity. The reverse-bias mode is preferred for the detection of pulsed laser radiation.

A practical example of a radiometer for which is shown in Fig. 2.1.3. It measures radiant power in the range from 1 mW to 10 W, whereas the lower limit is set by detector and amplifier noise and the upper by the load limit of the electrical heater [89Moe]. The radiation absorber is a polished hollow cone electro-plated with a nearly specular reflecting black nickel layer. The temperature difference between the absorber cone and the heat sink is measured by a thermopile. The electric heater for moderate-accuracy in-situ calibrations of the instrument is wound around the cone. Another design of a thermopile-type radiometer with an integral alignment module can be found in [88Ino]. Further similar systems are described in [77Gun, 91Rad]. A commercial version of a laser radiometer based on a pyroelectric lithium tantalate crystal is described in [89Hen]. For higher radiant power levels of up to 1 kW cavity absorbers cooled by a surrounding jacket of flowing

water are employed. The difference in temperature between the outflowing and inflowing water is measured and serves as quantity for the absorbed laser radiant power [96Bra]. A special design of the surface geometry of the cavity reduces the irradiance of the laser beam, thus improving the protection from damaging the surface.

The preferred instruments for , , , are thermally absorbing devices such as calorimeters. The receiver element is often a glass-disk, where the radiation is absorbed in the volume instead of on the surface. The absorptance exhibits an excellent stability under chemical and mechanical stress. This type of calorimeter is described in [70Edw, 74Gun]. The radiative load can be reduced by using glass with a low absorption coefficient which increases the length of the absorption path. On the other hand the heat capacity increases linearly with the thickness of the glass-disk which, in conjunction with the poor thermal conductivity of glass, results in long response and cooling times of these detectors. The radiometric scale for laser radiant energy is usually derived from the scale for cw laser radiant power. In [91Moe] a fast electromechanical shutter is used to produce pulses of known laser radiant energy of up to 5 J. The influence of the pulse duration has to be corrected in the calibration procedure. A laser energy meter not depending on a cw laser radiant power scale is described in [90Yua]. In this instrument the light pressure of the laser beam sensed by two mirrors is converted by a moving coil to an electrical signal. The main advantages of this system are fast response and no interruption of the laser beam. The device has been investigated for single laser pulses of radiant energies between 10 mJ and 6 J. Another method not interrupting the laser beam is the photoacoustic calorimetry [86Kim]. There, the radiant energy incident upon a mirror is absorbed at the mirror surface. The absorbed energy generates elastic strain waves which propagate through the mirror substrate. The strain waves eventually pass through a piezoelectric transducer attached to the back of the mirror substrate. The voltage of the piezoelectric crystal gives a direct indication of the amount of energy absorbed at the mirror surface. Since a priori the absorptance of the mirror is not known the instrument has to be calibrated against a standard energy meter.

2.1.4 Outlook – State of the art and trends

Although optical radiometry has been developed for 100 years, measurements of the various radiometric quantities only recently have achieved the required small uncertainties. Today the most accurate detector-based primary radiometric standard is the electrically calibrated cryogenic radiometer. In this instrument the radiant power of – preferably – a laser beam is measured by substituting the absorbed optical power of the laser beam by the electrical power of a heating system. Cryogenic radiometers operate at liquid helium temperatures and have a measurement uncertainty of a few parts in 10^4 , a significant improvement over earlier room-temperature radiometers.

Accurate characterization of laser sources is crucial to the effective development and use of industrial technologies such as light-wave telecommunications, laser-based medical instrumentation, materials processing, photolithography, data storage, and laser safety equipment. Traceable measurement standards are essential both for users to have confidence in their measurements and to support quality assurance in the manufacture of lasers and laser systems. Because lasers present a potential safety hazard, it is also important to have measurement standards to satisfy nationally and internationally agreed safety limits. The traceability for laser radiometric measurements in Germany is maintained by the Physikalisch-Technische Bundesanstalt. It meets the requirements for calibration and testing laboratories, certification and accreditation bodies defined in the ISO/IEC Guide 17025 and the DIN/EN 45000 and DIN/EN/ISO 9000 series of standards, see http://www.ptb.de/en/org/q/q3/q33/_index.htm.

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2.2 Beam characterization

В. Еррісн

2.2.1 Introduction

The success of almost any laser application depends mainly on the power density distributions in a certain area of the laser beam, usually the focal region. It is the aim of laser beam characterization to describe and predict the profiles a beam takes on under free-space propagation or behind optical systems.

A complete characterization of laser beams would allow the prediction of power density distributions, including size and shape, behind arbitrary optical systems as far as they are sufficiently known. Admittedly for such detailed characterization a huge amount of data and sophisticated measurement procedures are necessary. But for many applications the knowledge and prediction of the transverse extent of the laser beam profile might be sufficient. Restriction to nearly aberration-free optical systems then enables beam characterization by only ten or less parameters.

In the following the validity of the paraxial approximation will be presumed. In practical this means that the full divergence angle of the beam should not exceed 30 degrees. Furthermore, any polarization effects are neglected. Beam characterization methods based on the considerations presented in this chapter have recently become an international standard, published as ISO 11146 [99ISO].

2.2.2 The Wigner distribution

A complete description of partially coherent radiation fields (within the restrictions stated above) can be given by a two-point-correlation integral of the field in a transverse plane at location z [99Bor]:

$$\tilde{\Gamma}(\boldsymbol{r}_1, \boldsymbol{r}_2, z, \tau) = \frac{1}{T} \int_{t_0}^{t_0+T} E^*(\boldsymbol{r}_1, z, t) E(\boldsymbol{r}_2, z, t + \tau) dt, \qquad (2.2.1)$$

where $E(\mathbf{r}, z, t)$ is the electrical field, z the coordinate along the direction of propagation, $\mathbf{r} = (x, y)^{\mathsf{T}}$ a transverse spatial vector (see Fig. 2.2.1), and T the integration time which shall be large enough to ensure that the integration results are independent of the starting time t_0 . The temporal Fourier transform of this correlation integral is known as the cross-spectral density or the (mutual) power spectrum:

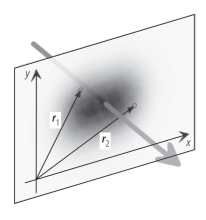


Fig. 2.2.1. Spatial coordinates r_1 and r_2 of a pair of points in a plane transverse to the direction of propagation.

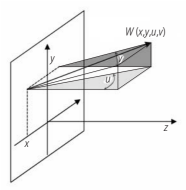


Fig. 2.2.2. The phase space coordinates of the Wigner distribution. x and y are spatial transverse coordinates, u and v are the corresponding angular coordinates.

$$\Gamma(\mathbf{r}_1, \mathbf{r}_2, z, \omega) = \int \tilde{\Gamma}(\mathbf{r}_1, \mathbf{r}_2, z, \tau) e^{i\omega\tau} d\tau.$$
(2.2.2)

Since laser beams in general can be considered as quasi-monochromatic, the frequency dependency will be dropped in the following:

$$\Gamma(\mathbf{r}_1, \mathbf{r}_2, z, \omega_0) \to \Gamma(\mathbf{r}_1, \mathbf{r}_2, z)$$
 (2.2.3)

From the cross-spectral density in a transverse plane at location z the power density in that plane can easily be obtained by

$$I(\mathbf{r},z) = \Gamma(\mathbf{r},\mathbf{r},z) \ . \tag{2.2.4}$$

Given the cross-spectral density at an entry plane the further propagation through arbitrary, but well-defined optical systems can be calculated by several methods and hence the power density distribution in the output plane of the systems predicted [99Bor].

The Wigner distribution W(r, q, z) of partially coherent beams is defined as the Fourier transform of the cross spectral density with respect to the separation vector s [78Bas]:

$$W(\mathbf{r}, \mathbf{q}, z) = \int \Gamma\left(\mathbf{r} + \frac{1}{2}\mathbf{s}, \mathbf{r} - \frac{1}{2}\mathbf{s}, z\right) e^{-ik\mathbf{q}\cdot\mathbf{s}} d\mathbf{s}.$$
 (2.2.5)

The Wigner distribution contains the same information as the cross-spectral density, but in a different, more descriptive manner. Considering $\mathbf{q} = (u, v)^{\mathsf{T}}$ as an angular vector with respect to the z-axis (Fig. 2.2.2), the Wigner distribution gives the part (amount) of the radiation power which passes the plane at z through the point \mathbf{r} in the direction given by \mathbf{q} . Within this picture the Wigner distribution might be considered as a generalization of the geometric optical radiance, although this analogy is limited. E.g. the Wigner distribution may take on negative values.

The power density distribution in a transverse plane is obtained by integration over the angles of direction,

$$I(\mathbf{r}, z) = \int W(\mathbf{r}, \mathbf{q}, z) d\mathbf{q}, \qquad (2.2.6)$$

and the far-field power density distribution by integration over the spatial coordinates,

$$I_{\mathrm{F}}(\boldsymbol{q}) = \int W(\boldsymbol{r}, \boldsymbol{q}, z) \, d\boldsymbol{r} .$$
 (2.2.7)

The Wigner distribution represents the beam in a transverse plane at location z. As the beam propagates in free space or through an optical system the Wigner distribution changes. This is reflected in the z-dependency of the Wigner distribution in the equations above. In the following equations this z-dependency will be dropped wherever appropriate.

The propagation of the Wigner distribution through aberration-free first-order optical systems (combinations of parabolic elements and free-space propagation) is very similar to that of geometric-optical rays. Such rays are specified by their position r and direction q. After propagation through an aberration-free optical system position and direction will change according to

$$\begin{pmatrix} r_{\text{out}} \\ q_{\text{out}} \end{pmatrix} = S \cdot \begin{pmatrix} r_{\text{in}} \\ q_{\text{in}} \end{pmatrix} , \qquad (2.2.8)$$

where S is a 4×4 -matrix representing the optical system, the system matrix (see Chap. 3.1). Considering the Wigner distribution as a density distribution of geometric optical rays, its propagation law is given by ray tracing [78Bas]:

$$W_{\text{out}}(\boldsymbol{r}_{\text{out}}, \boldsymbol{q}_{\text{out}}) = W_{\text{in}}(\boldsymbol{r}_{\text{in}}, \boldsymbol{q}_{\text{in}}) \quad \text{with} \quad \begin{pmatrix} \boldsymbol{r}_{\text{in}} \\ \boldsymbol{q}_{\text{in}} \end{pmatrix} = \mathsf{S}^{-1} \cdot \begin{pmatrix} \boldsymbol{r}_{\text{out}} \\ \boldsymbol{q}_{\text{out}} \end{pmatrix} .$$
 (2.2.9)

2.2.3 The second-order moments of the Wigner distribution

From the Wigner distribution smaller sets of data can be derived, which can be associated to certain physical properties of the beams. These sets of data are the so-called moments of the Wigner distribution [86Bas]:

$$\langle x^k y^{\ell} u^m v^n \rangle = \frac{\int W(x, y, u, v) \ x^k y^{\ell} u^m v^n \, \mathrm{d}x \, \mathrm{d}y \, \mathrm{d}u \, \mathrm{d}v}{\int W(x, y, u, v) \, \mathrm{d}x \, \mathrm{d}y \, \mathrm{d}u \, \mathrm{d}v} \quad \text{with} \quad k, \ell, m, n \ge 0 , \qquad (2.2.10)$$

where

$$W(x, y, u, v) = W(\mathbf{r}, \mathbf{q})$$
 with $\mathbf{r} = (x, y)^{\mathsf{T}}$, $\mathbf{q} = (u, v)^{\mathsf{T}}$. (2.2.11)

The order of the moments is defined by the sum of the exponents, $k + \ell + m + n$. There are four first-order moments, $\langle x \rangle$, $\langle y \rangle$, $\langle u \rangle$, and $\langle v \rangle$, which together specify position and direction of propagation of the beam profile's centroids within the given coordinate system.

The moments of the Wigner distribution are defined to be independent of the coordinate system:

$$\langle x^{k} y^{\ell} u^{m} v^{n} \rangle_{c} = \frac{\int W(x, y, u, v) (x - \langle x \rangle)^{k} (y - \langle y \rangle)^{\ell} (u - \langle u \rangle)^{m} (v - \langle v \rangle)^{n} dx dy du dv}{\int W(x, y, u, v) dx dy du dv}.$$
(2.2.12)

There are ten centered second-order moments, specified by $k+\ell+m+n=2$. Three pure spatial moments, $\langle x^2 \rangle_{\rm c}$, $\langle y^2 \rangle_{\rm c}$, $\langle xy \rangle_{\rm c}$, three pure angular moments, $\langle u^2 \rangle_{\rm c}$, $\langle v^2 \rangle_{\rm c}$, $\langle uv \rangle_{\rm c}$, and four mixed moments, $\langle xu \rangle_{\rm c}$, $\langle yv \rangle_{\rm c}$, $\langle xv \rangle_{\rm c}$, and $\langle yu \rangle_{\rm c}$. The centered second-order moments are associated to the beam extents in the near and far field and to the propagation of beam widths as will be discussed in the next section.

Only the three pure spatial moments can directly be measured since they can be obtained from the power density distribution in the observation plane by

$$\langle x^k y^\ell \rangle_{c} = \frac{1}{P} \int I(x, y) (x - \langle x \rangle)^k (y - \langle y \rangle)^\ell dx dy$$
 (2.2.13)

with

$$\langle x \rangle = \frac{1}{P} \int I(x, y) \ x \, \mathrm{d}x \, \mathrm{d}y \ , \tag{2.2.14}$$

$$\langle y \rangle = \frac{1}{P} \int I(x, y) \ y \, \mathrm{d}x \, \mathrm{d}y \ , \tag{2.2.15}$$

and

$$P = \int I(x,y) \, \mathrm{d}x \, \mathrm{d}y \,. \tag{2.2.16}$$

As the beam propagates through optical systems the Wigner distribution changes and consequently the moments change, too. A simple propagation law for the centered second-order moments through aberration-free optical systems can be derived from the propagation law of the Wigner distribution (2.2.9). Combining the ten moments in a symmetric 4×4 -matrix, the variance matrix

$$\mathsf{P} = \begin{pmatrix} \langle x^2 \rangle_{\mathsf{c}} & \langle xy \rangle_{\mathsf{c}} & \langle xu \rangle_{\mathsf{c}} & \langle xv \rangle_{\mathsf{c}} \\ \langle xy \rangle_{\mathsf{c}} & \langle y^2 \rangle_{\mathsf{c}} & \langle yu \rangle_{\mathsf{c}} & \langle yv \rangle_{\mathsf{c}} \\ \langle xu \rangle_{\mathsf{c}} & \langle yu \rangle_{\mathsf{c}} & \langle u^2 \rangle_{\mathsf{c}} & \langle uv \rangle_{\mathsf{c}} \\ \langle xv \rangle_{\mathsf{c}} & \langle yv \rangle_{\mathsf{c}} & \langle uv \rangle_{\mathsf{c}} & \langle v^2 \rangle_{\mathsf{c}} \end{pmatrix} , \tag{2.2.17}$$

delivers the propagation law

$$\mathsf{P}_{\mathrm{out}} = \mathsf{S} \cdot \mathsf{P}_{\mathrm{in}} \cdot \mathsf{S}^{\mathsf{T}} \,, \tag{2.2.18}$$

where $P_{\rm in}$ and $P_{\rm out}$ are the variance matrices in the input and output planes of the optical system, respectively, and S is the system matrix.

2.2.4 The second-order moments and related physical properties

In this section the relations between the centered second-order moments and some more physical properties are discussed.

2.2.4.1 Near field

The three spatial-centered second-order moments are related to the spatial extent of the power density in the reference plane as can be derived from (2.2.13). For example, the centered second-order moments $\langle x^2 \rangle_c$, defined by

$$\langle x^2 \rangle_{c} = \frac{1}{P} \int I(x,y) (x - \langle x \rangle)^2 dx dy,$$
 (2.2.19)

can be considered as the intensity-weighted average of the squared distances in x-direction of all points in the plane from the beam-profile center. Obviously, this quantity increases with increasing beam extent in x-direction. A beam width in x-direction can be defined as

$$d_x = 4\sqrt{\langle x^2 \rangle_{\rm c}} \ . \tag{2.2.20}$$

The factor of 4 in this equation has been chosen by convention to adapt this beam-width definition to the former $1/e^2$ -definition for the beam radius of Gaussian beams. For an aligned elliptical Gaussian beam profile,

$$I(x,y) \propto e^{-2\frac{x^2}{w_x^2}} \cdot e^{-2\frac{y^2}{w_y^2}},$$
 (2.2.21)

where w_x and w_y are the $1/e^2$ -beam radii in x- and y-direction, respectively, the relation

$$d_x = 2 w_x$$

holds. Similar, a beam width in y-direction can be defined as

$$d_y = 4\sqrt{\langle y^2 \rangle_{\rm c}} \ . \tag{2.2.22}$$

The beam width along an arbitrary azimuthal direction enclosing an angle of α with the x-axis can be derived from a rotation of the coordinate system delivering

$$d_{\alpha} = 4\sqrt{\langle x^2 \rangle_{\rm c} \cos^2 \alpha + 2 \langle xy \rangle_{\rm c} \sin \alpha \cos \alpha + \langle y^2 \rangle_{\rm c} \sin^2 \alpha} . \tag{2.2.23}$$

In general, the beam width considered as a function of the azimuthal direction α has unique maximum and minimum. The related directions are orthogonal to each other and define the principal axes of the beam. The signed angle between the x-axis and that principal axis which is closer to the x-axis is given by

$$\varphi = \frac{1}{2} \operatorname{atan} \left(\frac{2 \langle xy \rangle_{c}}{\langle x^{2} \rangle_{c} - \langle y^{2} \rangle_{c}} \right) . \tag{2.2.24}$$

The beam width along that principal axis which is closer to the x-axis is determined by

$$d_x' = 2\sqrt{2} \left\{ \left(\left\langle x^2 \right\rangle_c + \left\langle y^2 \right\rangle_c \right) + \varepsilon \left[\left(\left\langle x^2 \right\rangle_c - \left\langle y^2 \right\rangle_c \right)^2 + 4 \left\langle xy \right\rangle_c^2 \right]^{\frac{1}{2}} \right\}^{\frac{1}{2}}$$

$$(2.2.25)$$

with

$$\varepsilon = \operatorname{sgn}\left(\left\langle x^2\right\rangle_{c} - \left\langle y^2\right\rangle_{c}\right) . \tag{2.2.26}$$

Correspondingly, the beam width along the principal axis closer to the y-axis is given by

$$d_y' = 2\sqrt{2} \left\{ \left(\left\langle x^2 \right\rangle_c + \left\langle y^2 \right\rangle_c \right) - \varepsilon \left[\left(\left\langle x^2 \right\rangle_c - \left\langle y^2 \right\rangle_c \right)^2 + 4 \left\langle xy \right\rangle_c^2 \right]^{\frac{1}{2}} \right\}^{\frac{1}{2}} . \tag{2.2.27}$$

Hence, the three spatial-centered second-order moments define the size and orientation of the so-called variance ellipse as the representation of a beam profile's extent (Fig. 2.2.3).

Beam profiles having approximately equal beam widths in both principal planes, $d'_x \approx d'_y$, may be considered as circular and a beam diameter may be defined by

$$d = 2\sqrt{2}\sqrt{\langle x^2 \rangle + \langle y^2 \rangle} . {(2.2.28)}$$

Sometimes this is an useful definition even for non-circular beam profiles, denoted then as "generalized beam diameter".

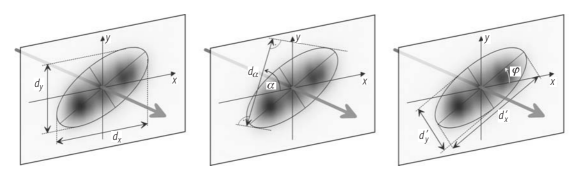


Fig. 2.2.3. Widths and variance ellipse of a power density profile. Left: widths d_x and d_y along the coordinate axes, middle: width d_α along an arbitrary direction, right: widths d'_x and d'_y along the principal axes

2.2.4.2 Far field

The three angular-centered second-order moments are related to the beam-profile extent in the far field, far away from the reference plane, or in the focal plane of a focusing lens. From the propagation law of the second-order moments, (2.2.18), the dependency of the spatial moments on the propagation distance z from the reference plane can be derived:

$$\langle x^{2} \rangle_{c}(z) = \langle x^{2} \rangle_{c,0} + 2 z \langle xu \rangle_{c,0} + z^{2} \langle u^{2} \rangle_{c,0} ,$$

$$\langle xy \rangle_{c}(z) = \langle xy \rangle_{c,0} + z \left(\langle xv \rangle_{c,0} + \langle yu \rangle_{c,0} \right) + z^{2} \langle uv \rangle_{c,0} ,$$

$$\langle y^{2} \rangle_{c}(z) = \langle y^{2} \rangle_{c,0} + 2 z \langle yv \rangle_{c,0} + z^{2} \langle v^{2} \rangle_{c,0} .$$

$$(2.2.29)$$

For large distances z the spatial moments depend only on the angular moments in the reference plane:

$$\langle x^{2} \rangle_{c}(z) \approx z^{2} \langle u^{2} \rangle ,$$

$$\langle xy \rangle_{c}(z) \approx z^{2} \langle uv \rangle ,$$

$$\langle y^{2} \rangle_{c}(z) \approx z^{2} \langle v^{2} \rangle .$$

$$(2.2.30)$$

The azimuthal angle φ_F of that principal axis in the far field, which is closer to the x-axis is then obtained by

$$\varphi_{\rm F} = \lim_{z \to \infty} \frac{1}{2} \operatorname{atan} \left(\frac{2 \langle xy \rangle_{\rm c}(z)}{\langle x^2 \rangle_{\rm c}(z) - \langle y^2 \rangle_{\rm c}(z)} \right) = \frac{1}{2} \operatorname{atan} \left(\frac{2 \langle uv \rangle_{\rm c}}{\langle u^2 \rangle_{\rm c} - \langle v^2 \rangle_{\rm c}} \right) , \qquad (2.2.31)$$

and the (full) divergence angles along the principal axes of the far field might be defined as

$$\theta_x' = \lim_{z \to \infty} \frac{d_x'(z)}{z} = 2\sqrt{2} \left\{ \left(\left\langle u^2 \right\rangle_{\text{c}} + \left\langle v^2 \right\rangle_{\text{c}} \right) + \eta \left[\left(\left\langle u^2 \right\rangle_{\text{c}} - \left\langle v^2 \right\rangle_{\text{c}} \right)^2 + 4 \left\langle uv \right\rangle_{\text{c}}^2 \right]^{\frac{1}{2}} \right\}^{\frac{1}{2}}, \quad (2.2.32)$$

$$\theta_y' = \lim_{z \to \infty} \frac{d_y'(z)}{z} = 2\sqrt{2} \left\{ \left(\left\langle u^2 \right\rangle_c + \left\langle v^2 \right\rangle_c \right) - \eta \left[\left(\left\langle u^2 \right\rangle_c - \left\langle v^2 \right\rangle_c \right)^2 + 4 \left\langle uv \right\rangle_c^2 \right]^{\frac{1}{2}} \right\}^{\frac{1}{2}}$$
(2.2.33)

with

$$\eta = \operatorname{sgn}\left(\left\langle x^2\right\rangle_{c} - \left\langle y^2\right\rangle_{c}\right) . \tag{2.2.34}$$

The generalized beam divergence angle might be defined as

$$\theta = 2\sqrt{2}\sqrt{\langle u^2\rangle_c + \langle v^2\rangle_c} \ . \tag{2.2.35}$$

The azimuthal orientation of the far field may differ from the orientation of the near field.

2.2.4.3 Phase paraboloid and twist

The four mixed moments $\langle xu\rangle_{\rm c}$, $\langle xv\rangle_{\rm c}$, $\langle yu\rangle_{\rm c}$, and $\langle yv\rangle_{\rm c}$ are closely related to the phase properties of the beam in the reference plane. Together with the three spatial moments they determine the radii of curvature and azimuthal orientation of the best-fitting phase paraboloid. Although the phase properties of partially coherent beams might be quite complicated, it is always possible to find a best-fitting phase function being quadratic (bilinear) in x and y:

$$\Phi(x,y) = k \left(a x^2 + 2 b x y + c y^2 \right) . \tag{2.2.36}$$

The best-fitting parameters a, b, c are defined by minimizing the generalized divergence angle, (2.2.35), if a phase function according to (2.2.36) would be subtracted from the actual phase distribution in the reference plane (e.g. by introducing a cylindrical lens) resulting in

$$a = \frac{\left\langle y^{2} \right\rangle \left\langle xu \right\rangle \left(\left\langle x^{2} \right\rangle + \left\langle y^{2} \right\rangle\right) - \left\langle xy \right\rangle^{2} \left(\left\langle xu \right\rangle - \left\langle yv \right\rangle\right) - \left\langle xy \right\rangle \left\langle y^{2} \right\rangle \left(\left\langle xv \right\rangle + \left\langle yu \right\rangle\right)}{\left(\left\langle x^{2} \right\rangle + \left\langle y^{2} \right\rangle\right) \left(\left\langle x^{2} \right\rangle \left\langle y^{2} \right\rangle - \left\langle xy \right\rangle^{2}\right)},$$

$$(2.2.37)$$

$$b = \frac{\langle x^2 \rangle \langle y^2 \rangle (\langle xv \rangle + \langle yu \rangle) - \langle xy \rangle (\langle x^2 \rangle \langle yv \rangle + \langle y^2 \rangle \langle xu \rangle)}{(\langle x^2 \rangle + \langle y^2 \rangle) (\langle x^2 \rangle \langle y^2 \rangle - \langle xy \rangle^2)}, \qquad (2.2.38)$$

$$c = \frac{\left\langle x^2 \right\rangle \left\langle yv \right\rangle \left(\left\langle x^2 \right\rangle + \left\langle y^2 \right\rangle \right) + \left\langle xy \right\rangle^2 \left(\left\langle xu \right\rangle - \left\langle yv \right\rangle \right) - \left\langle xy \right\rangle \left\langle x^2 \right\rangle \left(\left\langle xv \right\rangle + \left\langle yu \right\rangle \right)}{\left(\left\langle x^2 \right\rangle + \left\langle y^2 \right\rangle \right) \left(\left\langle x^2 \right\rangle \left\langle y^2 \right\rangle - \left\langle xy \right\rangle^2 \right)} \ . \tag{2.2.39}$$

A phase distribution as given in (2.2.36) can be considered as a rotated phase paraboloid, with

$$\varphi_{\rm P} = \frac{1}{2} \arctan\left(\frac{2b}{a-c}\right) \tag{2.2.40}$$

as the signed angle between the x-axis and that principal axis of the phase paraboloid, which is closer to the x-axis, and with

$$R_x' = \frac{2}{(a+c) + \mu\sqrt{(a-c)^2 + 4b^2}}$$
 (2.2.41)

and

$$R'_{y} = \frac{2}{(a+c) - \mu\sqrt{(a-c)^{2} + 4b^{2}}}$$
(2.2.42)

with

$$\mu = \operatorname{sgn}(a - c) \tag{2.2.43}$$

as the radii of curvature along that principal axis of the phase paraboloid, which is closer to the x- and y-axis, respectively. The radii of curvature R'_x and R'_x independently may be positive or negative or infinite, the later indicating a plane phase front along that azimuthal direction. The azimuthal orientation of the phase paraboloid's principal axes may differ from the orientation of the near field and/or far field.

If the radii of phase curvature along both principal axes are approximately equal, $R'_x \approx R'_y$, a generalized phase curvature of the best-fitting rotational symmetric phase paraboloid is defined by

$$R = \frac{\langle x^2 \rangle_c + \langle y^2 \rangle_c}{\langle xu \rangle_c + \langle yv \rangle_c} \,. \tag{2.2.44}$$

Another phase-related parameter is the so-called twist, defined as

$$t_{\rm w} = \langle xv \rangle - \langle yu \rangle$$
 (2.2.45)

The twist parameter is proportional to the orbital angular momentum transferred by the beam [93Sim].

2.2.4.4 Invariants

From the ten centered second-order moments two basic quantities can be derived, that are invariant under propagation through aberration-free first-order optics [03Nem].

The effective beam propagation ratio is defined as

$$M_{\text{eff}}^2 = \frac{4\pi}{\lambda} \left(\det(\mathsf{P}) \right)^{\frac{1}{4}} \ge 1$$
 (2.2.46)

and can be considered as a measure of the focusability of a beam. The lower limit holds only for coherent Gaussian beams.

The intrinsic astigmatism a, given by

$$a = \frac{8\pi^2}{\lambda^2} \left[\left(\left\langle x^2 \right\rangle_{\rm c} \left\langle u^2 \right\rangle_{\rm c} - \left\langle xu \right\rangle_{\rm c}^2 \right) + \left(\left\langle y^2 \right\rangle_{\rm c} \left\langle v^2 \right\rangle_{\rm c} - \left\langle yv \right\rangle_{\rm c}^2 \right) + 2 \left(\left\langle xy \right\rangle_{\rm c} \left\langle uv \right\rangle_{\rm c} - \left\langle xv \right\rangle_{\rm c} \left\langle yu \right\rangle_{\rm c} \right) \right] - \left(M_{\rm eff}^2 \right)^2 \ge 0 , \qquad (2.2.47)$$

is related to the visible and hidden astigmatism of the beam (see below).

2.2.4.5 Propagation of beam widths and beam propagation ratios

Under free-space propagation any directional beam width d_{α} , as well as the generalized beam diameter d, obeys an hyperbolic propagation law:

$$d_{\alpha}(z) = d_{0,\alpha} \sqrt{1 + \left(\frac{z - z_{0,\alpha}}{z_{R,\alpha}}\right)^2} = \sqrt{d_{0,\alpha}^2 + \theta_{\alpha}^2 (z - z_{0,\alpha})^2}, \qquad (2.2.48)$$

where $z_{0,\alpha}$ is the z-position of the smallest width, the waist position, $d_{0,\alpha}$ is the waist width, θ_{α} the divergence angle, and $z_{R,\alpha}$ the Rayleigh length, i.e. the distance from the waist position, where the width has grown by factor of $\sqrt{2}$. For the width along the x-direction, $\alpha=0$, see Fig. 2.2.4, the parameters can be obtained by

$$z_0 = -\frac{\langle xu\rangle_c}{\langle u^2\rangle_c} \,, \tag{2.2.49}$$

$$d_0 = 4\sqrt{\langle x^2 \rangle_{\rm c} - \frac{\langle xu \rangle_{\rm c}^2}{\langle u^2 \rangle_{\rm c}}}, \qquad (2.2.50)$$

and

$$z_{\rm R} = \sqrt{\frac{\langle x^2 \rangle_{\rm c}}{\langle u^2 \rangle_{\rm c}} - \frac{\langle x u \rangle_{\rm c}^2}{\langle u^2 \rangle_{\rm c}^2}} \,. \tag{2.2.51}$$

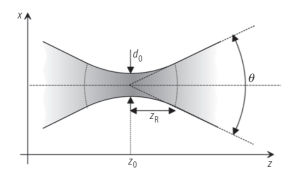


Fig. 2.2.4. Free-space propagation of beam widths with the beam waist position z_0 , the beam waist width d_0 , the Rayleigh length z_R , and the full divergence angle θ .

For other azimuthal directions α the same equations apply with the following substitutions:

$$\langle x^{2} \rangle_{c} \rightarrow \langle x^{2} \rangle_{c} \cos^{2} \alpha + 2 \langle xy \rangle_{c} \cos \alpha \sin \alpha + \langle y^{2} \rangle_{c} \sin^{2} \alpha ,$$

$$\langle xu \rangle_{c} \rightarrow \langle xu \rangle_{c} \cos^{2} \alpha + 2 (\langle xv \rangle_{c} + \langle yu \rangle_{c}) \cos \alpha \sin \alpha + \langle yv \rangle_{c} \sin^{2} \alpha ,$$

$$\langle u^{2} \rangle_{c} \rightarrow \langle u^{2} \rangle_{c} \cos^{2} \alpha + 2 \langle uv \rangle_{c} \cos \alpha \sin \alpha + \langle v^{2} \rangle_{c} \sin^{2} \alpha .$$

$$(2.2.52)$$

For the generalized diameter d the propagation parameters are obtained by

$$z_0 = -\frac{\langle xu\rangle_c + \langle yv\rangle_c}{\langle u^2\rangle_c + \langle v^2\rangle_c} , \qquad (2.2.53)$$

$$d_0 = 2\sqrt{2}\sqrt{\left(\langle x^2\rangle_c + \langle y^2\rangle_c\right) - \frac{\left(\langle xu\rangle_c + \langle yv\rangle_c\right)^2}{\langle u^2\rangle_c + \langle v^2\rangle_c}},$$
(2.2.54)

and

$$z_{\rm R} = \sqrt{\frac{\langle x^2 \rangle_{\rm c} + \langle y^2 \rangle_{\rm c}}{\langle u^2 \rangle_{\rm c} + \langle v^2 \rangle_{\rm c}} - \left(\frac{\langle xu \rangle_{\rm c} + \langle yv \rangle_{\rm c}}{\langle u^2 \rangle_{\rm c} + \langle v^2 \rangle_{\rm c}}\right)^2} \ . \tag{2.2.55}$$

It should be noted that beam widths along the principal axes, d'_x and d'_y , do , , obey the hyperbolic propagation law in the case of a general astigmatic beam with rotating variance ellipse (see next section).

The product of the (directional) beam waist diameter d, d_{α} and the corresponding far-field divergence angle θ , θ_{α} is called the beam parameter product. Due to diffraction the beam parameter product has a lower limit given by

$$d_0 \cdot \theta = \frac{d_0^2}{z_{\rm R}} \ge 4\frac{\lambda}{\pi} \,, \quad d_{0,\alpha} \cdot \theta_{\alpha} = \frac{d_{0,\alpha}^2}{z_{\rm R,\alpha}} \ge 4\frac{\lambda}{\pi} \,.$$
 (2.2.56)

Normalization to this lower limit delivers the so-called beam parameter ratios

$$M^2 = \frac{\pi}{\lambda} \frac{d_0 \cdot \theta}{4} , \quad M_\alpha^2 = \frac{\pi}{\lambda} \frac{d_{0,\alpha} \cdot \theta_\alpha}{4} . \tag{2.2.57}$$

The beam parameter ratios M^2 and M_{α}^2 are invariant in stigmatic aberration-free first-order optical systems (combinations of perfect spherical lenses). In contrast to the effective beam parameter ratio $M_{\rm eff}^2$, they may change under propagation through cylindrical lenses.

2.2.5 Beam classification

Lasers beams can be classified according to their propagation behavior. The classification is based on the discrimination between circular and non-circular power density profiles and the azimuthal orientation of the non-circular profiles. A beam profile is considered circular if the beam widths along both principal axes are approximately equal, or, in practice, if

$$\frac{\min(d_x', d_y')}{\max(d_x', d_y')} > 0.87. \tag{2.2.58}$$

In this sense a homogeneous profile with square footprint is regarded circular, see Fig. 2.2.5.

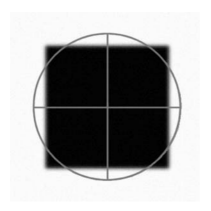


Fig. 2.2.5. Within the concept of second-order-moment beam characterization a square top-hat profile is considered circular: Its width is independent of the azimuthal direction.

2.2.5.1 Stigmatic beams

A laser beam is considered stigmatic if all its profiles under free-space propagation are circular and if all non-circular profiles behind an arbitrary cylindrical lens, inserted somewhere in the beam, have the same azimuthal orientation as the lens. The system matrix $P_{\rm st}$ of a perfectly stigmatic beam has only three independent parameters:

$$\mathsf{P}_{\mathrm{st}} = \begin{pmatrix} \langle x^2 \rangle_{\mathrm{c}} & 0 & \langle xu \rangle_{\mathrm{c}} & 0 \\ 0 & \langle x^2 \rangle_{\mathrm{c}} & 0 & \langle xu \rangle_{\mathrm{c}} \\ \langle xu \rangle_{\mathrm{c}} & 0 & \langle u^2 \rangle_{\mathrm{c}} & 0 \\ 0 & \langle xu \rangle_{\mathrm{c}} & 0 & \langle u^2 \rangle_{\mathrm{c}} \end{pmatrix} . \tag{2.2.59}$$

Physical parameters of a stigmatic beam are the beam diameter in the reference plane

$$d = 4\sqrt{\langle x^2 \rangle_{\rm c}} \tag{2.2.60}$$

and the full divergence angle

$$\theta = 4\sqrt{\langle u^2 \rangle} \ . \tag{2.2.61}$$

Since the properties of a stigmatic beam are independent of the azimuthal direction, it has a unique waist position

$$z_0 = -\frac{\langle xu\rangle_c}{\langle u^2\rangle_c} \tag{2.2.62}$$

with a waist diameter of

$$d_0 = 4\sqrt{\langle x^2 \rangle_c - \frac{\langle xu \rangle_c^2}{\langle u^2 \rangle_c}} \,. \tag{2.2.63}$$

The Rayleigh length $z_{\rm R}$ is the distance from the waist position where the diameter has grown by a factor of $\sqrt{2}$, given by

$$z_{\rm R} = \sqrt{\frac{\langle x^2 \rangle_{\rm c}}{\langle u^2 \rangle_{\rm c}} - \frac{\langle xu \rangle_{\rm c}^2}{\langle u^2 \rangle_{\rm c}^2}} \,. \tag{2.2.64}$$

Finally, the phase paraboloid is of rotational symmetry with the radius of curvature being

$$R = \frac{\langle x^2 \rangle_c}{\langle xu \rangle_c} \,. \tag{2.2.65}$$

2.2.5.2 Simple astigmatic beams

A laser beam is classified as simple astigmatic if at least some of the power density profiles the beam takes on under free-space propagation are non-circular, but all non-circular profiles have the same azimuthal orientation. In practice, the orientations of two non-circular beam profiles are regarded as equal, if the azimuthal angles differ by less than 10 degrees. A simple astigmatic beam whose principal axes are parallel to the x- and y-axis is called aligned simple astigmatic. The variance matrix $\mathsf{P}_{\mathsf{asa}}$ of a perfect aligned simple astigmatic beam has six independent parameters:

$$\mathsf{P}_{\mathrm{asa}} = \begin{pmatrix} \left\langle x^2 \right\rangle_{\mathrm{c}} & 0 & \left\langle xu \right\rangle_{\mathrm{c}} & 0\\ 0 & \left\langle y^2 \right\rangle_{\mathrm{c}} & 0 & \left\langle yv \right\rangle_{\mathrm{c}}\\ \left\langle xu \right\rangle_{\mathrm{c}} & 0 & \left\langle u^2 \right\rangle_{\mathrm{c}} & 0\\ 0 & \left\langle yv \right\rangle_{\mathrm{c}} & 0 & \left\langle v^2 \right\rangle_{\mathrm{c}} \end{pmatrix} . \tag{2.2.66}$$

All the physical parameters given for stigmatic beams can be assigned separately for each principal axis of a simple astigmatic beam. The diameters in x- and y-direction are

$$d_x = 4\sqrt{\langle x^2 \rangle_c}, \quad d_y = 4\sqrt{\langle y^2 \rangle_c}$$
 (2.2.67)

and the according full divergence angle

$$\theta_x = 4\sqrt{\langle u^2 \rangle} , \quad \theta_y = 4\sqrt{\langle v^2 \rangle} .$$
 (2.2.68)

Aligned simple astigmatic beams have in general two different waist positions for each principal axis:

$$z_{0,x} = -\frac{\langle xu\rangle_{c}}{\langle u^{2}\rangle_{c}}, \quad z_{0,y} = -\frac{\langle yv\rangle_{c}}{\langle v^{2}\rangle_{c}}$$
(2.2.69)

with the associated waist diameters

$$d_{0,x} = 4\sqrt{\langle x^2 \rangle_{\rm c} - \frac{\langle xu \rangle_{\rm c}^2}{\langle u^2 \rangle_{\rm c}}}, \quad d_{0,y} = 4\sqrt{\langle y^2 \rangle_{\rm c} - \frac{\langle yv \rangle_{\rm c}^2}{\langle v^2 \rangle_{\rm c}}}.$$
 (2.2.70)

Similarly, two Rayleigh lengths are defined by

$$z_{\mathrm{R},x} = \sqrt{\frac{\langle x^2 \rangle_{\mathrm{c}}}{\langle u^2 \rangle_{\mathrm{c}}} - \frac{\langle xu \rangle_{\mathrm{c}}^2}{\langle u^2 \rangle_{\mathrm{c}}^2}}, \quad z_{\mathrm{R},y} = \sqrt{\frac{\langle y^2 \rangle_{\mathrm{c}}}{\langle v^2 \rangle_{\mathrm{c}}} - \frac{\langle yv \rangle_{\mathrm{c}}^2}{\langle v^2 \rangle_{\mathrm{c}}^2}}, \tag{2.2.71}$$

Landolt-Börnstein New Series VIII/1A1 and the radii of phase curvature are

$$R_x = \frac{\langle x^2 \rangle_c}{\langle xu \rangle_c}, \quad R_y = \frac{\langle y^2 \rangle_c}{\langle yv \rangle_c}.$$
 (2.2.72)

The propagation laws for the beam diameters along both principal axes are:

$$d_x(z) = d_{0,x} \sqrt{1 + \left(\frac{z - z_{0,x}}{z_{\text{R.}x}}\right)^2} = \sqrt{d_{0,x}^2 + \theta_x^2 (z - z_{0,x})^2}$$
(2.2.73)

and

$$d_y(z) = d_{0,y} \sqrt{1 + \left(\frac{z - z_{0,y}}{z_{R,y}}\right)^2} = \sqrt{d_{0,y}^2 + \theta_y^2 (z - z_{0,y})^2}.$$
 (2.2.74)

For non-aligned simple astigmatic beams similar relations hold.

2.2.5.3 General astigmatic beams

All other beams are classified as general astigmatic. Usually all ten second-order moments are necessary to describe a general astigmatic beam.

2.2.5.4 Pseudo-symmetric beams

Pseudo-symmetric beams are general astigmatic but "look like" stigmatic or simple astigmatic under free-space propagation. They possess an inner astigmatism which is hidden under free propagation and propagation through stigmatic (isotropic) optical systems (i.e. combinations of spherical lenses). Pseudo-symmetric beams differ from real stigmatic or simple astigmatic beams by a non-vanishing twist parameter, $t_{\rm w} \neq 0$.

The variance matrix $P_{\rm pst}$ of pseudo-stigmatic beams is therefore

$$\mathsf{P}_{\mathrm{pst}} = \begin{pmatrix} \left\langle x^{2} \right\rangle_{\mathrm{c}} & 0 & \left\langle xu \right\rangle_{\mathrm{c}} & \frac{t}{2} \\ 0 & \left\langle x^{2} \right\rangle_{\mathrm{c}} & -\frac{t}{2} & \left\langle xu \right\rangle_{\mathrm{c}} \\ \left\langle xu \right\rangle_{\mathrm{c}} & -\frac{t}{2} & \left\langle u^{2} \right\rangle_{\mathrm{c}} & 0 \\ \frac{t}{2} & \left\langle xu \right\rangle_{\mathrm{c}} & 0 & \left\langle u^{2} \right\rangle_{\mathrm{c}} \end{pmatrix} . \tag{2.2.75}$$

Under free-space propagation there is no difference between a real stigmatic beam, $t_{\rm w}=0$, and the corresponding pseudo-stigmatic one, $t_{\rm w}\neq 0$, (2.2.29). The difference can be uncovered by inserting an arbitrary cylindrical lens somewhere in the beam path. The stigmatic beam is converted into a simple astigmatic beam with non-rotating variance ellipse while the pseudo-stigmatic one is turned into a general astigmatic beam with rotating variance ellipse. Figure 2.2.6 illustrates the different behaviors.

The variance matrix P_{psa} of aligned pseudo-simple astigmatic beams is given by

$$\mathsf{P}_{\mathrm{psa}} = \begin{pmatrix} \langle x^2 \rangle_{\mathsf{c}} & 0 & \langle xu \rangle_{\mathsf{c}} & \frac{t}{2} \\ 0 & \langle y^2 \rangle_{\mathsf{c}} & -\frac{t}{2} & \langle yv \rangle_{\mathsf{c}} \\ \langle xu \rangle_{\mathsf{c}} & -\frac{t}{2} & \langle u^2 \rangle_{\mathsf{c}} & 0 \\ \frac{t}{2} & \langle yv \rangle_{\mathsf{c}} & 0 & \langle v^2 \rangle_{\mathsf{c}} \end{pmatrix} . \tag{2.2.76}$$

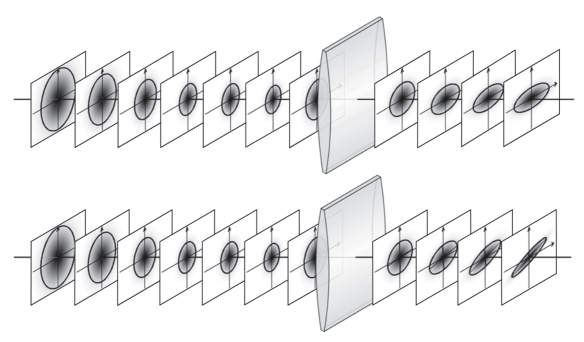


Fig. 2.2.6. Propagation of a stigmatic (top) and pseudo-stigmatic (bottom) laser beam. In free-space propagation both beams are indistinguishable. But a cylindrical lens transforms the stigmatic beam into a simple astigmatic one, whereas the pseudo-stigmatic beam becomes general astigmatic with rotating variance ellipse.

Again, under free-space propagation there is no difference between a real simple astigmatic beam, $t_{\rm w}=0$, and the corresponding pseudo-simple astigmatic one, $t_{\rm w}\neq 0$, (2.2.29). Inserting an aligned cylindrical lens somewhere in the beam pass unveils the difference. The simple astigmatic beam keeps being simple astigmatic while the pseudo-simple astigmatic one is turned into a general astigmatic beam with rotating variance ellipse. Figure 2.2.7 illustrates the different behaviors.

2.2.5.5 Intrinsic astigmatism and beam conversion

Applying astigmatic (anisotropic) optical systems (including cylindrical lenses) may convert beams from one class to another. But only beams with vanishing intrinsic astigmatism a, (2.2.47), can be converted into stigmatic ones [94Mor]. In practice, beams with

$$\frac{a}{\left(M_{\rm eff}^2\right)^2} < 0.039\tag{2.2.77}$$

are considered intrinsic stigmatic, all others intrinsic astigmatic (the limit of 0.039 is a consequence of (2.2.58)). Intrinsic astigmatic beams can always be converted into pseudo-stigmatic or simple astigmatic ones.

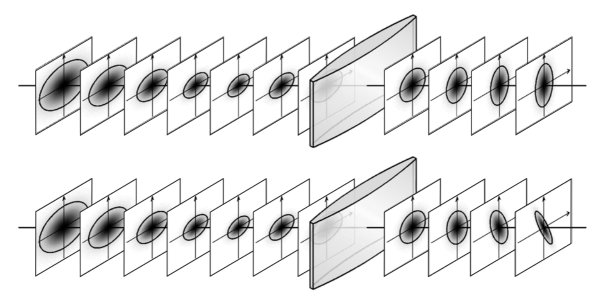


Fig. 2.2.7. Propagation of a simple astigmatic (top) and a pseudo-simple astigmatic (bottom) laser beam. In free-space propagation both beams are indistinguishable. But an aligned cylindrical lens transforms the simple astigmatic beam into a simple astigmatic one, whereas the pseudo-simple astigmatic beam becomes general astigmatic with rotating variance ellipse.

2.2.6 Measurement procedures

Only the three pure spatial moments out of the ten second-order moments are accessible for direct measurement. The other seven moments are retrieved indirectly based on the propagation law of the spatial moments (2.2.29).

The measurement method is based on the acquisition of a couple of power density profiles at different z-locations near the generalized beam waist, (2.2.53), e.g. by means of CCD cameras or similar devices (Fig. 2.2.8, left). From the measured profiles the spatial moments at each measurement plane are calculated. Fitting parabolas with three free parameters to the curve of each spatial moment delivers nine independent quantities: the moments $\langle x^2 \rangle_{\rm c,0}$, $\langle xy \rangle_{\rm c,0}$, $\langle y^2 \rangle_{\rm c,0}$, $\langle xu \rangle_{\rm c,0}$, $\langle xu \rangle_{\rm c,0}$, $\langle xu \rangle_{\rm c,0}$, and the sum of the crossed mixed moments $\langle xv \rangle_{\rm c,0} + \langle yu \rangle_{\rm c,0}$. If the waist of the beam is not accessible, an artificial waist has to be created by inserting an almost aberration-free focusing lens into the beam path. Approximately half of the profiles should be acquired close to the waist within one generalized Rayleigh length, the rest outside two Rayleigh lengths. This ensures balanced accuracy for all parameters of the fitting process.

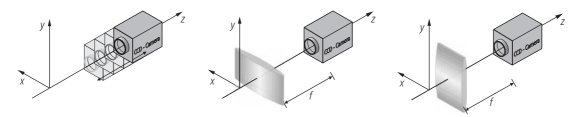


Fig. 2.2.8. Determination of the ten second-order moments in three steps. First step is a z-scan measurement (left), in the second step the CCD camera is placed in the focal plane behind a horizontally oriented cylindrical lens (middle), in the third step the lens is rotated by 90 degrees (right).

At least one cylindrical lens is needed for the measurement of the missing difference of the crossed mixed moments $\langle xv\rangle_{c,0} - \langle yu\rangle_{c,0}$. To retrieve it, a cylindrical lens with focal length f is inserted into the beam path at an arbitrary position in the beam waist region. Firstly, this cylindrical lens shall be aligned with the x-axis and the spatial moment $\langle xy\rangle_1$ is measured in the focal distance behind the lens (Fig. 2.2.8, middle). Next, the lens is rotated by 90 degrees and the spatial moment $\langle xy\rangle_2$ is again measured in the focal distance from the lens (Fig. 2.2.8, right). The missing difference of the crossed mixed moments of the reference plane is then given by

$$\langle xv\rangle_{c,0} - \langle yu\rangle_{c,0} = \frac{\langle xy\rangle_2 - \langle xy\rangle_1}{f} .$$
 (2.2.78)

2.2.7 Beam positional stability

2.2.7.1 Absolute fluctuations

For various reasons a laser beam may fluctuate in position and/or direction. The positional fluctuations in a transverse plane may be measured by the variance of the first-order spatial moments of the beam profile:

$$\left\langle x^{2}\right\rangle_{s} = \frac{1}{N} \sum_{i=1}^{N} \left\langle x\right\rangle_{i}^{2} - \left(\frac{1}{N} \sum_{i=1}^{N} \left\langle x\right\rangle_{i}\right)^{2} , \qquad (2.2.79)$$

$$\langle xy \rangle_{\rm s} = \frac{1}{N} \sum_{i=1}^{N} \langle x \rangle_{i} \langle y \rangle_{i} - \frac{1}{N} \sum_{i=1}^{N} \langle x \rangle_{i} \frac{1}{N} \sum_{i=1}^{N} \langle y \rangle_{i} , \qquad (2.2.80)$$

$$\left\langle y^{2}\right\rangle_{s} = \frac{1}{N} \sum_{i=1}^{N} \left\langle y\right\rangle_{i} - \left(\frac{1}{N} \sum_{i=1}^{N} \left\langle y\right\rangle_{i}\right)^{2} , \qquad (2.2.81)$$

where $\langle x \rangle_i$ and $\langle y \rangle_i$ are the first-order moments determined in N individual measurements and $\bar{x} = \frac{1}{N} \sum_{i=1}^{N} \langle x \rangle_i$, $\bar{y} = \frac{1}{N} \sum_{i=1}^{N} \langle y \rangle_i$ define the long-term average beam position. Obviously, the positional fluctuations are different from plane to plane. It can be shown that, under some reasonable assumptions, the positional fluctuations can be characterized closely analogous to the characterization of the beam extent based on the second-order moments of the Wigner distribution [94Mor, 96Mor]. Within this concept, the fluctuation properties of a laser beam are completely determined by ten different parameters, arranged in a symmetric 4×4 matrix

$$\mathsf{P}_{\mathsf{s}} = \begin{pmatrix} \left\langle x^{2} \right\rangle_{\mathsf{s}} & \left\langle xy \right\rangle_{\mathsf{s}} & \left\langle xu \right\rangle_{\mathsf{s}} & \left\langle xv \right\rangle_{\mathsf{s}} \\ \left\langle xy \right\rangle_{\mathsf{s}} & \left\langle y^{2} \right\rangle_{\mathsf{s}} & \left\langle yu \right\rangle_{\mathsf{s}} & \left\langle yv \right\rangle_{\mathsf{s}} \\ \left\langle xu \right\rangle_{\mathsf{s}} & \left\langle yu \right\rangle_{\mathsf{s}} & \left\langle u^{2} \right\rangle_{\mathsf{s}} & \left\langle uv \right\rangle_{\mathsf{s}} \\ \left\langle xv \right\rangle_{\mathsf{s}} & \left\langle yv \right\rangle_{\mathsf{s}} & \left\langle uv \right\rangle_{\mathsf{s}} & \left\langle v^{2} \right\rangle_{\mathsf{s}} \end{pmatrix},$$

$$(2.2.82)$$

obeying the same simple propagation law as the centered second-order moments:

$$P_{s,out} = S \cdot P_{s,in} \cdot S^{T}. \tag{2.2.83}$$

The elements of the beam fluctuation matrix may be considered as the centered second-order moments of a probability distribution p(x, y, u, v) giving the probability that the fluctuation beam

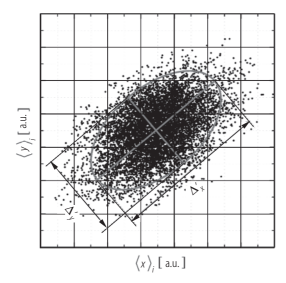


Fig. 2.2.9. Centroid coordinates of fluctuating beam and corresponding variance ellipse characterizing the fluctuations.

has a position (x,y) and direction (u,v) at a random measurement. Similar to the second-order moments of the Wigner distribution, only the three spatial moments are directly measurable. The complete set can be obtained from a z-scan measurement as described in the section above, by acquiring a couple of power density distributions in any measurement plane, calculating the first-order spatial moments from each profile, derive the three variances according to (2.2.79)–(2.2.81), and obtaining the second-order fluctuation moments in the reference plane from a fitting process. Again, measurements behind a cylindrical lens are necessary to achieve all ten parameters.

Fluctuation widths can be derived from the second-order fluctuation moments. In analogy to the beam width definitions, the fluctuation widths are

$$\Delta_x' = 2\sqrt{2} \left\{ \left(\left\langle x^2 \right\rangle_{\rm s} + \left\langle y^2 \right\rangle_{\rm s} \right) + \tau \left[\left(\left\langle x^2 \right\rangle_{\rm s} - \left\langle y^2 \right\rangle_{\rm s} \right)^2 + 4 \left\langle xy \right\rangle_{\rm s}^2 \right]^{\frac{1}{2}} \right\}^{\frac{1}{2}} , \tag{2.2.84}$$

$$\Delta_y' = 2\sqrt{2} \left\{ \left(\left\langle x^2 \right\rangle_{\mathrm{s}} + \left\langle y^2 \right\rangle_{\mathrm{s}} \right) - \tau \left[\left(\left\langle x^2 \right\rangle_{\mathrm{s}} - \left\langle y^2 \right\rangle_{\mathrm{s}} \right)^2 + 4 \left\langle xy \right\rangle_{\mathrm{s}}^2 \right]^{\frac{1}{2}} \right\}^{\frac{1}{2}}$$

$$(2.2.85)$$

with

$$\tau = \operatorname{sgn}\left(\left\langle x^{2}\right\rangle_{s} - \left\langle y^{2}\right\rangle_{s}\right) , \qquad (2.2.86)$$

where Δ'_x and Δ'_y are the beam fluctuation widths along the principal axes of the beam positional fluctuations and where

$$\beta = \frac{1}{2} \operatorname{atan} \left(\frac{2 \langle xy \rangle_{s}}{\langle x^{2} \rangle_{s} - \langle y^{2} \rangle_{s}} \right)$$
 (2.2.87)

is the signed angle between the x-axis and that principal axis of the beam fluctuation which is closer to the x-axis (Fig. 2.2.9). The principal axes of the beam positional fluctuations may not coincide with the principal axes of the power density distribution.

The width of the positional fluctuations along an arbitrary direction, given by the azimuthal angle α , is given by

$$\Delta_{\alpha} = 4\sqrt{\langle x^2 \rangle_{\rm s} \cos^2 \alpha + 2\langle xy \rangle_{\rm s} \sin \alpha \cos \alpha + \langle y^2 \rangle_{\rm s} \sin^2 \alpha} . \tag{2.2.88}$$

2.2.7.2 Relative fluctuations

For many applications the widths of the positional fluctuations compared to the momentary beam profile width might be more relevant than the absolute fluctuation widths. The relative fluctuation along an arbitrary direction, given by the azimuthal angle α , is defined by

$$\Delta_{\mathrm{rel},\alpha} = \sqrt{\frac{\langle x^2 \rangle_{\mathrm{s}} \cos^2 \alpha + 2 \langle xy \rangle_{\mathrm{s}} \sin \alpha \cos \alpha + \langle y^2 \rangle_{\mathrm{s}} \sin^2 \alpha}{\langle x^2 \rangle_{\mathrm{c}} \cos^2 \alpha + 2 \langle xy \rangle_{\mathrm{c}} \sin \alpha \cos \alpha + \langle y^2 \rangle_{\mathrm{c}} \sin^2 \alpha}}.$$
(2.2.89)

The effective relative fluctuation may by specified by

$$\Delta_{\rm rel} = \sqrt{\frac{\langle x^2 \rangle_{\rm s} + \langle y^2 \rangle_{\rm s}}{\langle x^2 \rangle_{\rm c} + \langle y^2 \rangle_{\rm c}}} \,. \tag{2.2.90}$$

2.2.7.3 Effective long-term beam widths

For applications with response times much longer than the typical fluctuation durations the timeaveraged intensity distribution rather than the momentary beam profile determines the process results:

$$\bar{I}(x,y) = \frac{1}{T} \int_{t_0}^{t_0+T} I(x,y,t) dt.$$
 (2.2.91)

The effective width of the time-averaged power density profile along an azimuthal direction enclosing an angle of α with the x-axis can be obtained from the widths of the momentary beam profile and the fluctuation width by

$$d_{\text{eff},\alpha} = \sqrt{d_{\alpha}^2 + \Delta_{\alpha}^2} \ . \tag{2.2.92}$$

References for 2.2

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3.1 Linear optics

R. GÜTHER

The propagation of light and its interaction with matter is completely described by , . , \bullet_{111} (1.1.4)–(1.1.7) and the material equations (1.1.8) and (1.1.9), see Chap. 1.1.

In this chapter the propagation of light in dielectric homogeneous and nonmagnetic media is discussed. Furthermore, monochromatic waves are assumed and linear interaction. The implications thereof for the medium are:

 $\varepsilon_{\rm r}$ ($\varepsilon(E, H)$ in (1.1.8)) is a complex tensor, which in most cases depends on the frequency only, but in special cases also on the spatial coordinate.

- , , , , $\mu_{\mathrm{r}}=1$ $(\mu(\boldsymbol{E},\boldsymbol{H})$ in (1.1.9)).
- $\dots, \qquad \rho = 0.$
- , j=0.

3.1.1 Wave equations

Maxwell's equations together with the material equations and the above assumptions result in the time-dependent wave equation for the electric field

$$\Delta \mathbf{E}(\mathbf{r},t) - \frac{\varepsilon_{r}}{c_{0}^{2}} \frac{\partial^{2}}{\partial t^{2}} \mathbf{E}(\mathbf{r},t) = 0$$
(3.1.1)

with

 $c_0 = 2.99792458 \times 10^8$ m/s: vacuum velocity of light,

$$\Delta = \frac{\partial^2}{\partial \, x^2} + \frac{\partial^2}{\partial \, y^2} + \frac{\partial^2}{\partial \, z^2} \ : \quad \text{delta operator}.$$

An identical equation holds for the magnetic field $\boldsymbol{H}(\boldsymbol{r},t)$.

For the following discussion we assume monochromatic fields, so that

$$E(r,t) = E(r) e^{i\omega t}$$
(3.1.2)

with

 ω : angular temporal frequency.

The magnetic field is related to E by the corresponding Maxwell equation (1.1.7)

$$\operatorname{curl} \boldsymbol{E}(\boldsymbol{r}) = -\mathrm{i}\,\omega\,\mu_0 \boldsymbol{H}(\boldsymbol{r}) \ . \tag{3.1.3}$$

Together with the ansatz (3.1.2), for isotropic media (ε_r is a complex scalar) (3.1.1) results in

$$\Delta E(r) + k_0^2 \,\hat{n}^2 E(r) = 0 \quad (\qquad (3.1.4)$$

with

 $k_0=2\pi/\lambda_0: \qquad ,$

 λ_0 : wavelength in vacuum,

 \hat{n} : complex refractive index, see (1.1.20).

For isotropic media and fields with uniform polarization the vector property of the field can be neglected. This results in

$$\Delta E(\mathbf{r}) + k_0^2 \,\hat{n}^2 E(\mathbf{r}) = 0 \quad (\quad (3.1.5)$$

In most cases the field can be approximated by a quasiplane wave, propagating in z-direction

$$E = E_0(r) e^{i(\omega t - k_0 \hat{n} z)}. \tag{3.1.6}$$

Remark: There are different conventions for writing the complex wave (3.1.6):

1. Electrical engineering and most books on quantum electronics:

$$\mathbf{E} \propto \exp(\mathrm{i}\,\omega\,t - \mathrm{i}\,k_0\,\hat{n}\,z)$$
,

for example [96Yar, 86Sie, 66Kog2, 84Hau, 91Sal, 98Sve, 96Die] and this chapter, Chap. 3.1.

2. Physical optics:

$$\mathbf{E} \propto \exp(\mathrm{i} k_0 \,\hat{n} \, z - \mathrm{i} \, \omega \, t)$$
,

for example [99Bor, 92Lan, 75Jac, 05Hod, 98Hec, 70Col].

[94Fel] discusses both cases.

Consequences of the convention: shape of results on phases of wave propagation, diffraction, interferences, Jones matrix, Collins integral, Gaussian beam propagation, absorption, and gain.

With

$$\left| \frac{\partial \boldsymbol{E}_0}{\partial z} \right| \ll |k_0 \, \hat{n} \, \boldsymbol{E}_0|$$

(3.1.4) can be reduced to

$$\Delta_{\mathbf{t}} \boldsymbol{E}_0 + 2 \mathrm{i} \, k_0 \hat{n} \, \frac{\partial \, \boldsymbol{E}_0}{\partial \, z} = 0 \quad (\boldsymbol{\rho}_0 \, \boldsymbol{\rho}_1 \, \boldsymbol{\rho}_2 \, \boldsymbol{\rho}_3 \, \boldsymbol{\rho}_4 \, \boldsymbol{\rho}_4$$

with

$$\varDelta_{\rm t} = \frac{\partial^2}{\partial \, x^2} + \frac{\partial^2}{\partial \, y^2} \; : \quad {\rm transverse \; delta \; operator \; (rectangular \; symmetry)},$$

see Chap. 1.1, (1.1.24a). , are: paraxial wave equation [86Sie], paraxial Helmholtz equation [96Ped, 78Gra].

The analogue approximation with respect to time t instead of the spatial coordinate z is used in ultrashort laser pulse physics [96Die, 86Sie].

3.1.2 Polarization

Restriction of (3.1.2) to a plane wave along the z-axis, see Fig. 3.1.1, results in

$$\begin{bmatrix} E_x \\ E_y \end{bmatrix} = \begin{bmatrix} E_{0x} \cos(\omega t - kz + \delta_x) \\ E_{0y} \cos(\omega t - kz + \delta_y) \end{bmatrix} \Rightarrow$$

$$\begin{bmatrix} E_{0x} \exp(i \delta_x) \\ E_{0y} \exp(i \delta_y) \end{bmatrix} \exp[i(\omega t - kz)] \equiv E_0 \mathbf{J} \exp[i(\omega t - kz)] . \tag{3.1.8}$$

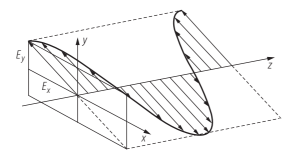


Fig. 3.1.1. Electric field of a linear polarized wave with propagation along the z-axis.

 $E_0 = \sqrt{E_{0x}^2 + E_{0y}^2} ,$ $J = \frac{1}{E_0} \begin{bmatrix} E_{0x} \exp(\mathrm{i}\,\delta_x) \\ E_{0y} \exp(\mathrm{i}\,\delta_y) \end{bmatrix} : \quad (\qquad , \qquad) , \qquad ,$ $\delta_x \text{ and } \delta_y : \text{ phase angles },$ $\Rightarrow : \text{ transition to the complex representation },$ $\varepsilon_0 n c_0 E_0^2 J J^* / 2 : \text{ light intensity } [\mathrm{W/m}^2] .$

Different *conventions* for *right*-hand polarization:

- 1. Looking against the direction of light propagation the light vector moves clockwise in the x-y-plane of Fig. 3.1.1 ([99Bor, 91Sal, 96Ped, 98Hec, 88Kle, 87Nau]).
- 2. The clockwise case occurs looking with the propagation direction (right-hand screw, elementary particle physics) ([84Yar, 88Yeh, 05Hod] and in this chapter).

Remark: J without normalization is also called Jones vector in [84Yar, 88Yeh, 90Roe, 77Azz, 86Sol], [95Bas, Vol. II, Chap. 27].

[41Jon, 97Hua, 88Yeh, 90Roe, 75Ger]:

$$\mathsf{J}_2 = \mathsf{M}\,\mathsf{J}_1 \tag{3.1.9}$$

with

J₁: Jones matrix for the initial polarization state,

M: Jones matrix describing an optical element or system,

 J_2 : Jones matrix of the polarization state after light has passed the element or system.

In Table 3.1.1 the characterization of the polarization states of light with the polarization of optical elements with the polarization of light with light with the polarization of light with the polarization of light with light with light with light with light with light with l

Table 3.1.1. Characterization of the polarization states of light with the *Jones vector*.

Jones vector J	$\begin{bmatrix} \cos \psi \\ \sin \psi \end{bmatrix}$	$\frac{1}{\sqrt{2}} \begin{bmatrix} 1 \\ \mathrm{i} \end{bmatrix}$	$\frac{1}{\sqrt{2}}\begin{bmatrix} i\\1 \end{bmatrix}$	$\begin{bmatrix} a \cdot \mathbf{i} \\ b \end{bmatrix} \text{ and } a^2 + b^2 = 1$
State of polarization	Linear polarization	Left circular polarization	Right circular polarization	Right elliptical polarization
Projection of the vector E onto the x - y -plane viewed along the propagation direction z	E \psi \psi \frac{\psi}{x}	<i>y x</i>	y x	b x

Table 3.1.2. Characterization of optical elements with the *Jones matrix*.

Opt. Element	Polarizer along the x -direction	0	Quarter-wave plate	Half-wave plate	Brewster-angle- tilted plate: index n	Faraday rotator (angle β)	Coordinate rotation by an angle α : $M(\alpha)$
Jones Matrix	$\begin{bmatrix} 1 & 0 \\ 0 & 0 \end{bmatrix}$	$\begin{bmatrix} 0 & 0 \\ 0 & 1 \end{bmatrix}$	$\begin{bmatrix} 1 & 0 \\ 0 & \pm i \end{bmatrix}$	$\begin{bmatrix} 1 & 0 \\ 0 & -1 \end{bmatrix}$	$ \begin{bmatrix} \left[\frac{2n}{n^2 + 1} \right]^2 0 \\ 0 & 1 \end{bmatrix} $	$\begin{bmatrix} \cos \beta - \sin \beta \\ \sin \beta & \cos \beta \end{bmatrix}$	$\begin{bmatrix} \cos \alpha & \sin \alpha \\ -\sin \alpha & \cos \alpha \end{bmatrix}$ Rotated element: $M(\alpha) M M(-\alpha)$

- 1 - , , , $M = M_3 \cdot M_2 \cdot M_1$, (3.1.10)

M: Jones matrix of the system which consists of elements with the matrices M1, M2, M3. Light passes first the element with M_1 and last the element with M_3 .

$$J_1 = \frac{1}{\sqrt{2}} \begin{bmatrix} 1 \\ 1 \end{bmatrix}$$
 (linear 45°-polarization), $M = \begin{bmatrix} 1 & 0 \\ 0 \pm i \end{bmatrix}$ ($\begin{cases} \text{right} \\ \text{left} \end{cases}$) quarter-wave plate), $J_2 = M \cdot J_1 = \frac{1}{\sqrt{2}} \begin{bmatrix} 1 \\ \pm i \end{bmatrix}$ ($\begin{cases} \text{right} \\ \text{left} \end{cases}$) circular polarization).

$$J = a_1 J_1 + a_2 J_2$$
 (3.1.11)
with $J_1 J_2^* = 0$.

linearly polarized light = left polarized light + right polarized light .

mixing of polarized and polarized light. It is described by the four components of the $\{s_0, s_1, s_2, s_3\}$, using $\langle \dots \rangle$ to signify averaging by detection:

$$s_0 = \langle E_x^2 \rangle + \langle E_y^2 \rangle \qquad \Rightarrow \qquad E_{0x}^2 + E_{0y}^2 , \qquad (3.1.12)$$

$$s_1 = \langle E_x^2 \rangle - \langle E_y^2 \rangle \qquad \Rightarrow \qquad E_{0x}^2 - E_{0y}^2 , \qquad (3.1.13)$$

$$s_1 = \langle E_x^2 \rangle - \langle E_y^2 \rangle \qquad \Rightarrow \qquad E_{0x}^2 - E_{0y}^2 \,, \tag{3.1.13}$$

$$s_2 = 2 \langle E_x E_y \cos [\delta_y - \delta_x] \rangle \quad \Rightarrow \quad 2E_{0x} E_{0y} \cos(\delta_y - \delta_x) , \qquad (3.1.14)$$

$$s_3 = 2 \langle E_x E_y \sin [\delta_y - \delta_x] \rangle \quad \Rightarrow \quad 2E_{0x} E_{0y} \sin(\delta_y - \delta_x) \tag{3.1.15}$$

with

$$s_0^2 > s_1^2 + s_2^2 + s_3^2$$
 $\Rightarrow s_0^2 = s_1^2 + s_2^2 + s_3^2$, (3.1.16)

where \Rightarrow means the transition from partially polarized light to completely polarized light, shown with the terms of Fig. 3.1.1.

Meaning of the s_i :

 s_0 : power flux, $\sqrt{s_1^2 + s_2^2 + s_3^2}/s_0$: degree of polarization, $\sqrt{s_1^2 + s_2^2}/s_0$: degree of linear polarization, s_3/s_0 : degree of circular polarization.

.,, ([75Ger, 77Azz, 90Roe, 95Bas]): extension of the , ,, where the four dimensional Stokes vector replaces the Jones vector and the real 4×4 Mueller matrices the complex 2×2 Jones matrices. The Jones calculus is usually sufficient to describe coherent laser radiation.

of the polarization state:

- [95Bas, Vol. 2, Chap. 22.15], [75Ger]. Result: Stokes vector.
- : see [05Hod]. There are commercial systems for this task.

 $\overline{}$ of polarized light are those two polarization states (Jones vectors) which reproduce themselves, multiplied with a complex factor (eigenvalue), if monochromatic light passes an optical element or system.

Calculation: see [97Hua, 77Azz], application: decoupling of the polarization mixing during round trips in resonators [74Jun].

3.1.3 Solutions of the wave equation in free space

Following (3.1.2), each of the wave solutions given in this section must be multiplied with the factor $e^{i\omega t}$ to obtain the propagating wave of (3.1.1).

3.1.3.1 Wave equation

The solutions of the wave equation (3.1.4) are vector fields.

3.1.3.1.1 Monochromatic plane wave

$$\boldsymbol{E} = \boldsymbol{E}_0 \exp \left\{ -i \, k_0 \, \hat{n} \, \boldsymbol{er} + i \, \varphi \right\}, \tag{3.1.17}$$

$$\boldsymbol{H} = \frac{\hat{n}}{c_0 \mu_0} \left(\boldsymbol{e} \times \boldsymbol{E}_0 \right) \exp \left\{ -i \, k_0 \, \hat{n} \, \boldsymbol{e} \boldsymbol{r} + i \, \varphi \right\} \tag{3.1.18}$$

with

r: position vector,

e: unit vector normal to the wave fronts,

 $k_0 = 2\pi/\lambda_0$: wave number,

 \hat{n} : complex refractive index,

 φ : phase.

For the phase velocity and the wave group velocity see Sect. 3.1.5.3.

3.1.3.1.2 Cylindrical vector wave

$$\mathbf{E} = E_0 \, \mathbf{e}_z \, H_0^{(2)}(k_0 \rho) \,, \tag{3.1.19}$$

$$\boldsymbol{H} = i \frac{E_0}{c_0 \mu_0} \left(\boldsymbol{e}_z \times \frac{\boldsymbol{\rho}}{\rho} \right) H_1^{(2)}(k_0 \rho) \quad (\rho > \lambda)$$
(3.1.20)

for time-harmonic electric source current density on the z-axis of a cylindrical coordinate system with the coordinates (ρ, φ, z) : (radial distance, azimuthal angle, z-axis) [94Fel, Chap. 5].

 $H_m^{(2)}$: m^{th} order Hankel function of the second kind [70Abr]; the change of convention in Sect. 3.1.1 includes: $H_m^{(2)} \Rightarrow H_m^{(1)}$ [94Fel, p. 487]; ρ : radial position vector,

 e_z : unit vector along the z-axis.

3.1.3.1.3 Spherical vector wave

$$\boldsymbol{E} = E_0 \cdot (\boldsymbol{n} \times \boldsymbol{p}) \times \boldsymbol{n} \cdot \frac{\exp(-i \, k_0 \, \hat{\boldsymbol{n}} \, r)}{r} \,, \tag{3.1.21}$$

$$\boldsymbol{E} = E_0 \cdot (\boldsymbol{n} \times \boldsymbol{p}) \times \boldsymbol{n} \cdot \frac{\exp(-i k_0 \,\hat{\boldsymbol{n}} \, r)}{r} , \qquad (3.1.21)$$

$$\boldsymbol{H} = \frac{E_0}{c_0 \mu_0} \cdot (\boldsymbol{n} \times \boldsymbol{p}) \cdot \frac{\exp(-i k_0 \,\hat{\boldsymbol{n}} \, r)}{r} \quad (r \gg \lambda_0)$$

is the , , , , $(1/r^2$ and higher inverse power terms $\ll 1/r$ -term) of an oscillating electric dipole ([99Bor, 94Leh, 75Jac]) with

 E_0 : amplitude [V],

p: unit vector of the dipole moment,

n: unit vector pointing from dipole to spatial position,

r: radial distance.

3.1.3.2 Helmholtz equation

The approximative transition from the $E_{i,j}$, wave equation (3.1.4) to the Helmholtz equation (3.1.5) ([99Bor]) results in $E_{i,j}$, solutions. $E_{i,j}$ is called: "field" [72Mar], "complex displacement" or "scalar wave function" [99Bor], "disturbance" [95Bas, Vol. I].

3.1.3.2.1 Plane wave

$$E = E_0 \exp \left\{ -i k_0 \,\hat{n} \, \boldsymbol{er} + i \,\varphi \,. \right\} \tag{3.1.23}$$

For the parameters see (3.1.18).

3.1.3.2.2 Cylindrical wave

$$E = E_0 H_0^{(2)}(k_0 \,\hat{n} \,\rho) \quad (\rho > \lambda_0) \tag{3.1.24}$$

is the diverging field of a homogeneous line source [41Str, Chap. IV], [94Fel, Chap. 5]. For the parameters see (3.1.19).

3.1.3.2.3 Spherical wave

$$E = E_0 \cdot \frac{\exp(-i k_0 \hat{n} r)}{r} \quad (r > \lambda_0) , \qquad (3.1.25)$$

parameters see (3.1.21).

3.1.3.2.4 Diffraction-free beams

3.1.3.2.4.1 Diffraction-free Bessel beams

Diffraction-free Bessel beams without transversal limitation are discussed in [05Hod, 91Nie, 88Mil].

$$E(x, y, z) = E_0 \cdot J_0(a \rho) \cdot \exp\{-i \cos(\theta_B) k_0 z\}$$
(3.1.26)

with

 E_0 : amplitude vector [V/m],

 J_0 : zero-order Bessel function of the first kind [70Abr]; higher-order Bessel beams see [96Hal]; $\rho = \sqrt{x^2 + y^2}$: radial distance from the z-axis,

 $a = k_0 \sin \Theta_{\rm B} \,[{\rm m}^{-1}],$

 $\Theta_{\rm B}$: convergence angle of the conus of the plane wave normal to the z-axis, see Fig. 3.1.2.

3.1.3.2.4.2 Real Bessel beams

Real Bessel beams are limited by a finite aperture D of the optical elements needed or Gaussian beam illumination (Gaussian Bessel beams [87Gor]).

Methods of generation: axicons [85Bic] (Fig. 3.1.2), annular aperture in the focus of a lens [87Dur, 91Nie], holographic [91Lee] or diffractive [96Don] elements. Because of finite aperture diffraction the latter display approximately the shape of (3.1.26) with cutoff at a geometric determined radius r_N , which includes N maxima (Fig. 3.1.3) and different amplitude patterns in dependence on z.

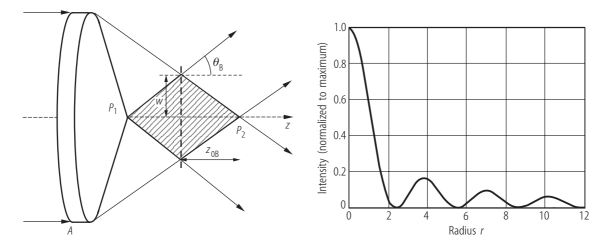


Fig. 3.1.2. Generation of a Bessel beam with help of an axicon A by a conus of plane-waves propagation directions.

Fig. 3.1.3. Transversal intensity structure of a Bessel beam ($\propto J_0^2(r)$).

of Bessel beams: Large depth of focus $2z_{0B}$ between P1 and P2 in Fig. 3.1.2 (thin "needle of light") for measurement purposes.

• Every maximum in Fig. 3.1.3 contains in the corresponding circular ring nearly the same power as the central peak. High power loss occurs if the central part is used only [05Hod].

3.1.3.2.4.3 Vectorial Bessel beams

Vectorial Bessel beams are discussed in [96Hal].

3.1.3.3 Solutions of the slowly varying envelope equation

are solutions of the (3.1.7) [91Sal, 96Ped, 86Sie, 78Gra], which is equivalent to paraxial approximation or Fresnel's approximation, see Sect. 3.1.4.

The, ___, •, • from SVE-approximated Gaussian beams towards an exact solution of the wave equation in the non-paraxial range is given in a Lax-Wünsche series [75Lax, 79Agr, 92Wue]. For contour plots of the relative errors in the Gaussian beam volume see [97For, 97Zen].

The of Gaussian beams is discussed in [79Dav, 95Gou], containing a Lax-Wünsche series; Gaussian beam in elliptical cylinder coordinates are given in [94Soi, 00Gou].

3.1.3.3.1 Gauss-Hermite beams (rectangular symmetry)

- 11/11/2-1 21 - 1 - - 1/1 . - 1

$$E_{mn}(x, y, z) = E_0 U_m(x, z) U_n(y, z) \exp\{-i k_0 z\}, \qquad (3.1.27)$$

$$U_m(x,z) = \sqrt{\frac{w_{0x}}{w_x(z)}} H_m\left(\frac{\sqrt{2}x}{w_x(z)}\right) \exp\left\{-\frac{x^2}{w_x^2(z)} - i\frac{k_0 x^2}{2R_x(z)}\right\} \exp\left\{i\varphi_m(z)\right\} , \qquad (3.1.28)$$

$$U_n(y,z) = U_{m \Rightarrow n}(x \Rightarrow y,z) \tag{3.1.29}$$

with

 w_{0x} : the $1/e^2$ -intensity waist radius,

 $z_{0x} = \frac{\pi w_{0x}^2}{\lambda}$: the Rayleigh distance (half depth of focus),

$$w_x(z) = w_{0x} \sqrt{1 + \frac{z^2}{z_0^2}}$$
 : the E_{00} -beam $1/\mathrm{e}^2$ -intensity radius,

$$R_x(z) = z \sqrt{1 + \frac{z^2}{z_0^2}}$$
 : the radius of curvature of the wavefront at position z ,

 $\varphi_m(z) = \left(\frac{1}{2} + m\right)$ arctan $\left(\frac{z}{z_0}\right)$: Gouy's phase, changing sign for the transition through z = 0,

$$H_m\left(\frac{\sqrt{2}}{w_x(z)}\right)$$
: the Hermite polynomial of order m [70Abr],

$$H_0(\xi) = 1$$
, $H_1(\xi) = 2\xi$, $H_2(\xi) = 4\xi^2 - 2$, $H_3(\xi) = 8\xi^3 - 12\xi$, $H_4(\xi) = 16\xi^4 - 48\xi^2 + 12$, ...

$$\int_{-\infty}^{\infty} d\xi \left\{ \frac{\exp\left(-\xi^2/2\right)}{\sqrt{\sqrt{\pi} m! \, 2^m}} H_m(\xi) \right\} \left\{ \frac{\exp\left(-\xi^2/2\right)}{\sqrt{\sqrt{\pi} n! \, 2^n}} H_n(\xi) \right\} = \delta_{mn} ,$$

$$\delta_{mn} = \begin{cases} 1 & \text{for } m = n \\ 0 & \text{for } m \neq n \end{cases} \quad \left(\frac{1}{2^{n-1} + 2^{n-1}} + \frac{1}{2^{n-1}} \right) . \tag{3.1.30}$$

Specialization of (3.1.27): $m=n=0\,,\,w_{0x}=w_{0y}=w_0\,,\,r=\sqrt{x^2+y^2}\,.$

$$E_{00}(r,z) = E_0 \frac{w_0}{w(z)} \exp\left\{-\frac{r^2}{w^2(z)} - i\frac{kr^2}{2R(z)}\right\} \exp\left\{i\frac{1}{2}\arctan\frac{z}{z_0}\right\} \exp\left\{-ikz\right\} , \qquad (3.1.31)$$

$$w(z) = w_0 \sqrt{1 + \frac{z^2}{z_0^2}}, \quad R(z) = z \sqrt{1 + \frac{z_0^2}{z^2}}.$$

Fig. 3.1.4. , , of E_{00} (fundamental mode): The shape of the Gaussian E_{00} -beam is depicted in Fig. 3.1.4. , , , of E_{00} in Fig. 3.1.4 are:

C : curves with constant amplitude decrease as $E(r,z)=E(0,z)/{\rm e}$ or constant intensity decrease as $I(r,z)=I(0,z)/{\rm e}^2$,

P: phase fronts with radius of curvature R(z),

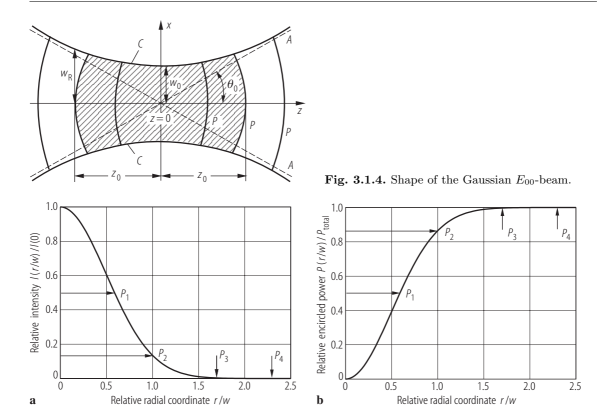


Fig. 3.1.5. (a) Cross section of a Gaussian beam perpendicular to the z-axis. (b) Power transmitted by a circular aperture with the relative radius r/w in a cross section.

Table 3.1.3. Characteristic points in Fig. 3.1.5.

Point in Fig. 3.1.5a, b	Relative abscissa r/w	Relative intensity, Fig. 3.1.5a	Relative transmission, Fig. 3.1.5b	Characterization
$\overline{P_1}$	0.588	0.5	0.5	FWHM ^a
P_2	1	0.135	0.865	$1/e^2$ -int. ^b
P_3	1.57	0.01	0.99	trunc. c
P_4	2.3	0.001	0.999	trunc. ^d

^a Full width half maximum/2.

 w_0 : ..., v_1 , v_2 , v_3 , v_4 , v_4 , v_5 , v_6 , v_6 , v_6 , v_7 , v_8 , v_9 , $v_$

 $\Theta_0 = \lambda/(\pi w_0)$: 1/e²-intensity divergence angle toward the asymptotes A.

In Fig. 3.1.5a the cross section of a Gaussian beam perpendicular to the z-axis is given, in Fig. 3.1.5b the power transmitted by a circular aperture with the relative radius r/w in a cross section. Characteristic points in Fig. 3.1.5 are listed in Table 3.1.3.

of the elliptical Gaussian beam: see Sect. 3.1.7.

 $^{^{\}rm b}$ 1/e²-intensity or 1/e-amplitude.

^c Diffraction of E_{00} -beam by circular aperture \Rightarrow 17 % intensity ripple [86Sie, p. 667].

d Diffraction of E_{00} -beam by circular aperture $\Rightarrow 1$ % intensity ripple [86Sie, p. 667] (no essential effect of truncation).

3.1.3.3.2 Gauss-Laguerre beams (circular symmetry)

$$E_{lp}(r, \psi, z) = E_0 \exp \left\{ -i \left[kz - \varphi_{lp}(z) \right] \right\} \frac{w_0}{w(z)} \left(\frac{\sqrt{2} \, r}{w(z)} \right)^l L_p^l \left(\frac{2 \, r^2}{w^2(z)} \right)$$

$$\times \exp \left\{ -\frac{r^2}{w^2(z)} - i \frac{k \, x^2}{2 \, R(z)} \right\} \begin{cases} \cos(l\psi) \\ \sin(l\psi) \end{cases}$$
(3.1.32)

with

z: propagation direction,

 r, φ : polar coordinates in the plane $\perp z$ -axis,

$$z_0 = \frac{\pi w_0^2}{\lambda}$$
: the Rayleigh distance (half depth of focus),

$$w(z) = w_0 \sqrt{1 + \left(\frac{z}{z_0}\right)^2}$$
: the E_{00} -beam $1/e^2$ -intensity radius,

$$R(z) = z \left\{ 1 + \left(\frac{z_0}{z}\right)^2 \right\}$$
: the radius of curvature of the wavefront at position z ,

$$\varphi_{lp} = (2p + l + 1) \arctan\left(\frac{z}{z_0}\right)$$
: Gouy's phase,

 L_p^l : Laguerre polynomial of degree p and order l [70Abr]:

p!: the factorial p.

- are formed by the cos- and sin-terms in (3.1.32).
- l = p = 0 means the rotational symmetrical Gaussian beam E_{00} .
- The , , , , determines what system of Gauss-Laguerre polynomials or Gauss-Hermite polynomials is more appropriate for a wave field development.

3.1.3.3.3 Cross-sectional shapes of the Gaussian modes

In Fig. 3.1.6 intensity distributions of Gauss-Hermite modes E_{mn} are given (rectangular symmetry), in Fig. 3.1.7 intensity distributions of Gauss-Laguerre modes E_{pl} (circular symmetry).

84 3.1.4 Diffraction [Ref. p. 131

Rectangular symmetry (Gauss-Hermite modes)

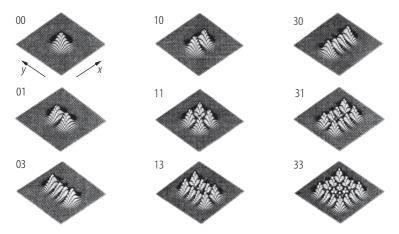


Fig. 3.1.6. Intensity distributions of Gauss-Hermite modes E_{mn} . The two digits at each distribution are m and n.

Circular symmetry (Gauss-Laguerre modes)

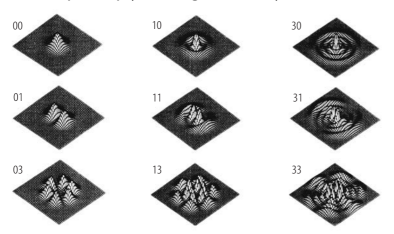


Fig. 3.1.7. Intensity distributions of Gauss-Laguerre modes E_{pl} . The two digits at each distribution are p and l.

3.1.4 Diffraction

Diffraction of light by aperture rims or amplitude and phase modifications inside the aperture:

- Solutions of Maxwell's equations taking into account the material properties of the aperture:
 - special cases: exact solutions [99Bor, 86Sta],
 - mostly: numerical solutions.
- Starting with a field near the aperture with reasonable assumptions for this field or its measurement: large variety of methods for different ranges of validity [99Bor, 86Sta, 61Hoe].

3.1.4.1 Vector theory of diffraction

- E and E are all E and E and E are all E are all E and E are all E

and meridional incidence of light [61Hoe]: Separation of the polarizations E parallel and E perpendicular to the plane of incidence for half plane [99Bor], slit [99Bor], gratings [80Pet], and volume gratings [69Kog, 81Sol, 81Rus].

3.1.4.2 Scalar diffraction theory

Two sources of scalar diffraction theory are:

- Transition from . . . , theory to theory: [99Bor, 86Sol]. The information about the polarization is lost.
- Mathematical formulation and generalization of, , , , , , : Each point on a wavefront may be regarded as a source of secondary waves, and the position of the wavefront at a later time is determined by the envelope of these secondary waves.

In Table 3.1.4 • with fields given near the diffraction aperture are listed. Figures 3.1.8 and 3.1.9 are related to Table 3.1.4.

(3.1.37): Approximation of (3.1.34): Huygens' principle with an additional directional factor (Fresnel).

(3.1.38): Approximation of (3.1.36): Huygens' principle with a modified directional factor.

(3.1.39):______ (= paraxial approximation). The approximation conditions from (3.1.34) to (3.1.39) resp. (3.1.40) are explained in [96For, 86Sta, 87Ree].

The condition $N_{\rm F}(a/d)^2/4 \ll 1$ [91Sal] is valid for sharp-edged apertures A, but it is weakened for the transmission of Gaussian-beam-like fields [86Sie, p. 635] or Gaussian-like soft apertures. Fresnel's approximation describes the propagation of the field from plane z=0 to plane z=z. This transformation can be cascaded to describe complex systems and is an often used $\frac{1}{12} \frac{1}{12} \frac{1}$

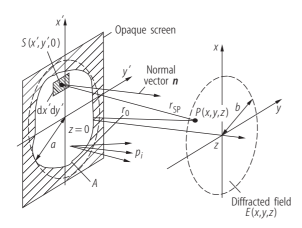


Fig. 3.1.8. Diffraction at an aperture A with source terms E(x', y', 0) and/or $\frac{\partial}{\partial z} E(x', y', z)|_{z=0}$, respectively, and a or b the maximum radial distances of source S or image point P, respectively. p_i symbolizes different plane waves for (3.1.41)–(3.1.43).

Table 3.1.4. Diffraction formulae with fields given near the diffraction aperture (r_{SP}) : see Fig. 3.1.8.

Integrals	Formula		Restrictions	Ref.
Rayleigh- Sommerfeld of 1 st kind	$E_{\rm RS1}(x,y,z) = -\frac{1}{4\pi} \iint_A E(x',y',0) \frac{\partial}{\partial z} \left(\frac{\exp(-ikr_{\rm SP})}{r_{\rm SP}} \right) dx' dy' $	(3.1.34)	$r_{\rm SP} > \lambda_0 ,$ plane aperture	[99Bor] [86Sta]
Rayleigh- Sommerfeld of 2 nd kind	$E_{\text{RS2}}(x, y, z) = -\frac{1}{2\pi} \iint_{A} \left[\frac{\partial E(x', y', z')}{\partial z} \right]_{z'=0} \frac{\exp(-ikr_{\text{SP}})}{r_{\text{SP}}} dx' dy' $	(3.1.35)	$r_{\mathrm{SP}} > \lambda_0 ,$ plane aperture	
Fresnel-Kirchhoff	$E_{\text{FK}}(x, y, z) = \frac{1}{2} \left[E_{\text{RS1}}(x, y, z) + E_{\text{RS2}}(x, y, z) \right]$ (6)	(3.1.36)	$r_{\rm SP} > \lambda_0 ,$ curved aperture	
Rayleigh- Sommerfeld 1 st kind approx.	$E_{\text{RS1a}}(x, y, z) = \frac{1}{\mathrm{i} \lambda} \iint_{A} E(x', y', 0) \frac{\exp(-\mathrm{i} k r_{\text{SP}})}{r_{\text{SP}}} \cos(n, r_{\text{SP}}) \mathrm{d} x' \mathrm{d} y'$	(3.1.37)	$r_{ m SP}\gg\lambda_0$	
Fresnel-Kirchhoff approximation, refers to Fig. 3.1.8	$E_{\text{FKa}}(x, y, z) = \frac{1}{\mathrm{i} \lambda} \iint_{A} E(x', y', 0) \frac{\exp(-\mathrm{i} k r_{\text{SP}})}{r_{\text{SP}}} \cdot \frac{1 + \cos(n, r_{\text{SP}})}{2} \mathrm{d} x' \mathrm{d} y'$	(3.1.38)	$r_{ m SP}\gg\lambda_0$	
Fresnel's approximation, refers to Fig. 3.1.8	$E_{\text{Fre}}(x,y,z) = \frac{\mathrm{i}\exp\left(-ikz\right)}{\lambda d} \iint_A E(x',y',0) \exp\left\{-\mathrm{i}\pi\frac{(x-x')^2 + (y-y')^2}{\lambdaz}\right\} \mathrm{d}x'\mathrm{d}y' \tag{9}$	(3.1.39)	$z\gg\lambda_0$	[99Bor] [96For] [97For] [87Ree] [86Sta]

Ref. p. 131]

Table 3.1.4 continued.

Integrals	Formula		Restrictions	Ref.
Fraunhofer far-field approximation, refers to Fig. 3.1.8	$E_{\rm Fra}(x,y,z) = \frac{{\rm i} \exp\left(-{\rm i} kz\right)p}{\lambda z} \iint_A E(x',y',0) \exp\left\{{\rm i} 2\pi \frac{xx'+yy'}{\lambdaz}\right\} {\rm d} x' {\rm d} y'$ with the additional phase term $p = \left\{ \begin{array}{cc} 1 & {\rm for} & \frac{b^2}{\lambda z} \ll 1 \\ \exp\left\{-{\rm i} \pi \frac{x^2+y^2}{\lambdaz}\right\} & {\rm otherwise} \end{array} \right.$	(3.1.40)	$\frac{a^2}{\lambda d} \ll 1$	[99Bor] [68Goo] [96For] [97For] [86Sta]
Plane-wave representation (also: angular-spectrum representation), refers to	2-D Fourier transform (see remark on (3.1.40)) of the source distribution E_s in plane $z=0$: $A_0(f_x, f_y) = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} E_s(x', y', 0) \exp\left\{i 2\pi (f_x x' + f_y y')\right\} dx' dy', \qquad (3.1.4)$ propagation of plane waves with the spatial frequencies f_x and f_y along the z-direction by distance z		$r_{ m SP}>\lambda_0$	[91Sal] [78Loh] [86Sta] [97For] [99Bor]
Figs. 3.1.8 and 3.1.9	$\exp\left\{-i 2\pi (f_x x + f_y y)\right\} \Rightarrow \exp\left\{-i 2\pi (f_x x + f_y y + \sqrt{1/\lambda^2 - f_x^2 - f_y^2} z)\right\},$	(3.1.42)		
	addition of plane waves at distance z : $E(x,y,z) = \int \int \int A_0(f_x, f_y) \exp\left\{-i 2\pi \left(f_x x + f_y y + \sqrt{1/\lambda^2 - f_x^2 - f_y^2} z\right)\right\} df_x df_y,$	(3.1.43)		
	equivalent to (3.1.34) [97For]			
Far field in the focal plane of a lens,	$E_{P}(x,y) = \frac{\mathrm{i}p}{\lambda f} \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} E_{S}(x',y') \exp\left\{\mathrm{i}2\pi \left(\frac{x}{\lambda f}x' + \frac{y}{\lambda f}y'\right)\right\} \mathrm{d}x' \mathrm{d}y' ,$	(3.1.44)	$d, f \gg \lambda$	[91Sal]
refers to Fig. 3.1.9	$p = \exp\left(i\pi \frac{(x^2 + y^2)(d - f)}{\lambda f^2}\right)$	(3.1.45)		

88 3.1.4 Diffraction [Ref. p. 131

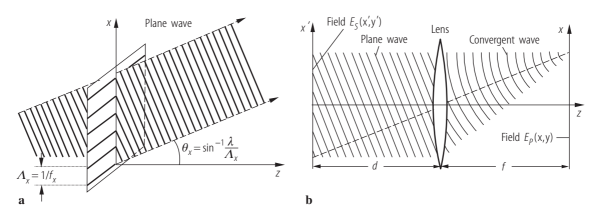


Fig. 3.1.9. (a) Spatial frequencies of a plane wave with propagation direction Θ_x with respect to the plane x=0 (and Θ_y analogously) are f_x and f_y with $\Theta_x=\sin^{-1}(\lambda f_x)\approx \lambda f_x$ and $\Theta_y=\sin^{-1}(\lambda f_y)\approx \lambda f_y$ (\approx : paraxial approximation). (b) Generation of the far field in the focal plane of a lens: The Fourier transformation (d=f) is changed by an additional phase term for $d\neq f$ with d: distance, f: focal length.

$$(3.1.40): \underline{\qquad} \quad (3.1.40): \underline{\qquad} \quad (3.1.46)$$

$$-\underline{\qquad} \quad (3.1.46)$$

- Validity of Fraunhofer's approximation: $N_{\rm F} \ll 1$. $p \neq 1$ (parabolic phase): the \bullet , , , of diffracted light is the square of the Fourier transform of E(x,y,0) only.
- - The convention of the plane-wave structure $\exp(i kx i \omega t)$ is connected with the determination of $F(f_x)$ by

$$F(f_x) = \int_{-\infty}^{\infty} dx f(x) e^{-i 2\pi f_x x}$$

[68Goo, 68Pap, 78Loh, 78Gas, 93Sto, 05Hod].

- The plane-wave structure $\exp(i\omega t - ikx)$ can be combined with

$$F(f_x) = \int_{-\infty}^{\infty} dx f(x) e^{i 2\pi f_x x}$$

[71Col, 73Men, 92Lug], but

$$F(f_x) = \int_{-\infty}^{\infty} dx f(x) e^{-i 2\pi f_x x}$$

is defined also in [88Kle, 91Sal, 95Wil, 96Ped].

- , \cdot , \cdot , \cdot , \cdot , \cdot , \cdot in (3.1.37) and (3.1.38):

$$r_{\rm SP} \approx r_0 + \frac{2x\xi - \xi^2 + 2y\eta - \eta^2}{2r_0}$$

[99Bor, 68Pap, 78Gra] with r_0 from Fig. 3.1.8 versus

$$r_{\rm SP} \approx z + \frac{2xx' - x'^2 + 2yy' - y'^2}{2z}$$

(references on lasers: [86Sie, 05Hod], optoelectronics: [68Goo, 72Mar, 91Sal]) for x_1, x_2, \dots . The sine of the diffraction angle $\sin \Theta_x = x/r_0$ is derived from principle and not by a postpositive reasoning of the paraxial range $x/z = \tan \Theta_x \approx \sin \Theta_x$. x/z should be "translated" into $\sin \Theta_x$ for better approximation.

(3.1.44), (3.1.45): $d \neq f$ (object is outside the object-side focal plane) \Rightarrow additional phase term p to the pure (inverse) Fourier transform (d = f), similarly to (3.1.40).

Applications: generation of the spectrum of a function, possibility of mathematical operations in the Fourier-space with complex filtering masks, correlation and convolution.

Another important diffraction theory

Diffraction theory after $A_{1,1,1}$, $A_{1,1,2}$, A_{1

3.1.4.3 Time-dependent diffraction theory

Two formulations of the time-dependent treatment of diffraction are possible:

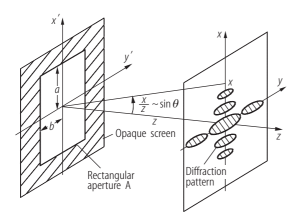
- 1. A general Fresnel-Kirchhoff's integral formula exists for \bullet , aperture A, see [99Bor, 99Pau].

3.1.4.4 Fraunhofer diffraction patterns

3.1.4.4.1 Rectangular aperture with dimensions $2a \times 2b$

In Fig. 3.1.10 the geometry of the diffraction from a rectangular aperture $2a \times 2b$ is shown. The x-part of the diffraction pattern in Fig. 3.1.10 is given in Fig. 3.1.11. In Table 3.1.5 the zeros and maxima of the intensity distribution are listed.

90 3.1.4 Diffraction [Ref. p. 131



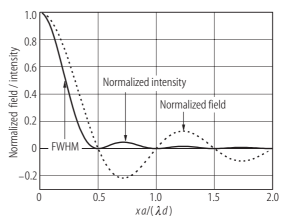


Fig. 3.1.10. Geometry of the diffraction from a rectangular aperture $2a \times 2b$.

Fig. 3.1.11. x-part of the diffraction pattern in Fig. 3.1.10. This is the diffraction pattern of a slit. For more exact electromagnetic solutions of a slit see [61Hoe, p. 266].

Table 3.1.5. Zeros and maxima of the intensity distribution.

Number n	$xa/\lambda z$	I_n/I_0
0	0	1
FWHM	2×0.221	0.5
1	0.5	0
1	0.715	0.0472
2	1	0
2	1.230	0.0168
3	1.5	0
3	1.735	0.0083
4	2	0
4	2.239	0.0050

Field distribution:

$$E(x,y,z) = \frac{4 a b}{\mathrm{i} \lambda z} E_0 \exp\left\{-\mathrm{i} k \left(z + \frac{x^2 + y^2}{2 z}\right)\right\} \operatorname{sinc}\left\{\frac{2 \pi a x}{\lambda z}\right\} \operatorname{sinc}\left\{\frac{2 \pi b y}{\lambda z}\right\}$$
(3.1.47)

with $\operatorname{sinc}(x) = \frac{\sin x}{x}$ and E_0 the electric-field amplitude.

Intensity:

$$I(x,y,z) = I(0,0,z)\operatorname{sinc}^{2}\left\{\frac{2\pi a x}{\lambda z}\right\}\operatorname{sinc}^{2}\left\{\frac{2\pi b y}{\lambda z}\right\}.$$
(3.1.48)

If the Fraunhofer diffraction is observed in the focal plane, z has to be replaced by f.

3.1.4.4.2 Circular aperture with radius a

The circular aperture with radius a is discussed in [61Hoe, p. 453]. In Fig. 3.1.12 diffraction by a circular aperture is shown. In Fig. 3.1.13a the diffracted field and intensity and in Fig. 3.1.13b the encircled energy in the diffraction plane with a circular screen are given. The zeros and maxima of intensity for diffraction by a circular aperture are listed in Table 3.1.6.

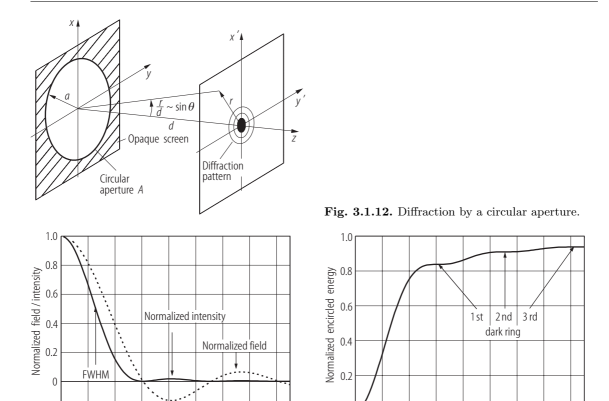


Fig. 3.1.13. (a) Diffracted field and intensity. (b) Encircled energy in the diffraction plane with a circular screen.

b

0.2 0.4

0.6

0.8 1.0

 $ra/(\lambda d)$

Table 3.1.6. Zeros and maxima of intensity for diffraction by a circular aperture.

Number n	$r_n a/(\lambda d)$	I_n/I_0
0	0	1
FWHM	2×0.257	0.5
1	0.610	0
1	0.817	0.0175
2	1.117	0
2	1.340	0.00415
3	1.619	0
3	1.849	0.00160
4	2.121	0
4	2.355	0.00078

Field distribution:

0.4

0.8

 $ra/(\lambda d)$

1.0

$$E(r,z) = \frac{\pi a^2}{i \lambda z} E_0 \exp\left\{-i k \left(z + \frac{kr^2}{2z}\right)\right\} \left\{2 \frac{J_1 \left[2 \pi a r/(\lambda z)\right]}{2 \pi a r/(\lambda z)}\right\}$$
(3.1.49)

with E_0 the electric-field amplitude and r the radius in the far-field plane.

Intensity:

$$I(r) = I(0, z) \left\{ 2 \frac{J_1[2 \pi a r/(\lambda z)]}{2 \pi a r/(\lambda z)} \right\}^2 . \tag{3.1.50}$$

Landolt-Börnstein New Series VIII/1A1 92 3.1.4 Diffraction [Ref. p. 131

3.1.4.4.2.1 Applications

•
$$r_{1 \text{ Airy}} = 0.610 \,\lambda/\sin \sigma$$
, (3.1.51)

1st-minimum radius of the intensity distribution in the focal plane of an aberration-free lens (Lommel 1885, Debye 1909, [86Sta, 99Bor]): Substitute in (3.1.50) $a/z \Rightarrow \sin \sigma$ (numerical aperture = sinus of the intersection angle σ with optical axis in the focal point, r_1, r_2, \dots, r_n , and $r = r_1$ airy as above.

: obscuration of the central part in the circular aperture A of Fig. 3.1.12:

- Reduction of the central diffraction maximum width by $\approx 20 \%$.
- Increase of secondary maximum by factor ≈ 7 .
- See Bessel beams, Sect. 3.1.3.2.4, [05Hod].

3.1.4.4.3 Gratings

$$\sin \alpha + \sin \beta = m \frac{\lambda}{q} \tag{3.1.52}$$

with

 α : angle of incidence (see Fig. 3.1.14),

 β : diffraction angle,

g: grating constant (grating period, groove distance),

m: order of diffraction. [82Hut, p. 25] often used: If the diffraction order is on the same side with the zero order (m=0) as the grating normal: m>0, otherwise m<0. In Fig. 3.1.14, the directions of the +1st transmitted order and the grating normal (dashed and dotted lines) are on the same side of the 0th transmitted order. Therefore m=1>0.

distribution between the different orders m [82Hut, 99Bor]. For the real phase and reflection gratings, it is substituted by the \bullet . \bullet \bullet \bullet curves in dependence on α or λ . There is an extreme diversity of cases. Catalogs of such curves: see [80Pet, 97Loe].

$$R_{\text{theor}} = \lambda/(\Delta \lambda) = m N = W \left(\sin \alpha + \sin \beta\right)/\lambda \tag{3.1.53}$$

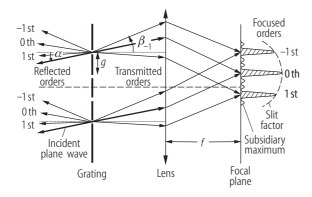


Fig. 3.1.14. Reflected and transmitted orders of a grating, here with N=4 slits. The far-field distribution is *visualized* after focusing by an *ideal* lens. Between the main maxima occur N-2 subsidiary maxima. The dashed envelope is the *slit factor*.

with

N: number of grooves of the grating,

W: width of the grating,

 α, β : see (3.1.52).

contains theoretical resolution and the aberrations of the optical elements for collimation and focusing of the grating-diffracted plane waves or by the aberrations of the collimation and focusing properties. [87Chr, 82Hut].

plication: external laser resonators). [82Hut] show lower disturbations than mechanically produced gratings (application: external laser resonators).

wavelengths [80Pet, 82Hut, 97Loe].

(, , , , , , , , ; [81Sol, 81Rus].

of spectral devices: [82Hut].

3.1.4.5 Fresnel's diffraction figures

Fresnel's approximation is given in (3.1.39) in Table 3.1.4.

3.1.4.5.1 Fresnel's diffraction on a slit

In Fig. 3.1.15 Fresnel's diffraction pattern of a slit with width 2a is shown.

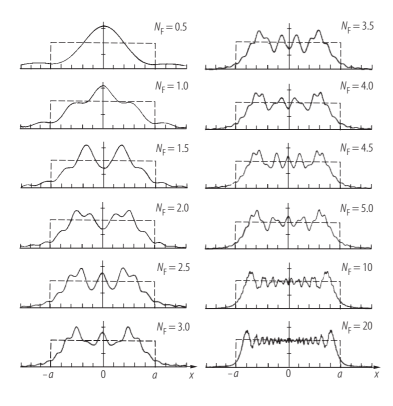


Fig. 3.1.15. Fresnel's diffraction pattern of a slit with width 2a(see Fig. 3.1.10 with $b \Rightarrow \infty$). Fresnel's number $N_{\rm F} = a^2/(\lambda z)$ is the essential parameter to characterize the transition from farfield (Fraunhofer) approximation $(N_{\rm F} < 0.2...0.5)$ to near-field (Fresnel) approximation $(N_{\rm F})$ 0.5). $N_{\rm F} = 0.5$: one central maximum only, $N_{\rm F}=3$: three maxima, $N_{\rm F} = N: N$ maxima. Hard-edgediffraction results in a ripple in the near field, which can be avoided by soft apertures, for instance Gaussian-like [86Sie] (apodization in optics [99Bor]). Figure after [86Sie, p. 721].

94 3.1.4 Diffraction [Ref. p. 131

3.1.4.5.2 Fresnel's diffraction through lens systems (paraxial diffraction)

Given: a system of lenses and the field distribution E(x, y) to be propagated.

The easily taken is:

- Given: E(x, y) in the plane z = 0. Required: the field in the plane z = z. Solution: (3.1.39).
- Given: E(x, y) in the plane z = 0 and near to this plane a lens. Required: the field in the plane z = z. Solution: modification of (3.1.39) by an additional factor L(x', y') to:

$$E_{\text{Fre}}(x, y, z) = \frac{i \exp\{-i kz\}}{\lambda d} \iint_{A} E(x', y', 0) L(x', y')$$

$$\times \exp\left\{-i \pi \frac{(x - x')^{2} + (y - y')^{2}}{\lambda z}\right\} dx' dy', \qquad (3.1.54)$$

$$L(x',y') = p(x',y') \exp\{-i kn t_L\} \exp\left\{\frac{i k(x'^2 + y'^2)}{2f}\right\}$$
(3.1.55)

with

n: refractive index of the lens,

 $t_{\rm L}$: thickness of the lens,

f: focal length of the lens,

p(x', y'): amplitude part, which can describe a marginal aperture or a Gaussian apodization.

A general complex function L(x',y') can model diffractive optical elements.

Cases of integration:

- No transversal limitations (without stops) and quadratic arguments of the exponential functions due to analytical results. The property of the closed form of such a calculation (see Sect. 3.1.7.4).
- One stop (finite integration limits): The result includes the error function [70Abr].
- Two and more finite integration limits are not useful. Then, (commercial) numerical field propagation programs through systems should be consulted.

Examples: [68Goo, 91Sal, 71Col, 85Iiz, 92Lug, 68Pap].

3.1.4.6 Fourier optics and diffractive optics

engineering to spatial frequency methods in optics, see Figs. 3.1.9, 3.1.10 and (3.1.41), (3.1.43), (3.1.44).

References: principles of Fourier optics: [68Goo, 78Loh, 83Ste, 85Iiz, 89Ars, 93Sto, 98Hec, 99Lau], filtering: [92Lug], filtering in connection with holography: [96Har, 71Col], noise suppression: [91Wyr].

In Fig. 3.1.16 low-pass filtering of a laser beam with a four-f-setup is shown.

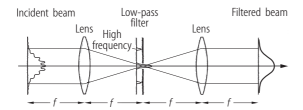


Fig. 3.1.16. Low-pass filtering of a laser beam with a four-f-setup [92Lug]. The mask is a low-pass filter, which transmits a zero mode only and suppresses the higher modes. The incident beam can also be modified by a transmission element which changes amplitude and phase.

changing microstructures whose dimensions are of the order of the wavelength mostly. They extend the classical means of optical design. References: [67Loh, 84Sch, 97Tur, 00Tur, 00Mey, 01Jah].

- 1 - , , , . . .

- ____, ____ generated by mechanical or interference ruling [69Str, 67Rud] on either plane or concave substrates for the combination of dispersive properties with imaging [82Hut, 87Chr].
- - acting as microoptic lenses of [97Her].
- ("modane") for transformation and filtering of modes of a laser [94Soi].
- Generation of theoretical ideal wavefronts for with interferometrical methods [95Bas, Vol. II, Chap. 31].
- Mode-discriminating and emission-forming elements in resonators [94Leg, 97Leg, 99Zei].

For pure imaging applications, refracting surfaces are still preferred, even in the micro-range [97Her]. Tasks with special dispersion requirements and special optical field transformations are the main application of the diffractive elements with increasing share.

The technology of dispersion compensation and weight reduction in large optical systems by special diffractive elements is partially solved, now.

3.1.5 Optical materials

$$\hat{\varepsilon}_{\mathbf{r}} = \hat{n}^2 \tag{3.1.56}$$

with

 $\hat{\varepsilon}_{r}$: complex relative dielectric constant (or tensor), \hat{n} : complex refractive index,

$$\alpha \ll k_0 : \hat{n} = n - i k_e = n - i n \kappa = n - i \frac{\alpha}{2 k_0} , \qquad (3.1.57)$$

(unity field amplitude):

$$\exp\left\{-\mathrm{i} kz\right\} = \exp\left\{-\mathrm{i} k_0(n-\mathrm{i} \kappa) z\right\} \exp\left\{-\mathrm{i} k_0 \left(n-\mathrm{i} \frac{\alpha}{2 k_0}\right) z\right\}$$
$$= \exp\left\{-\mathrm{i} k_0 z - \frac{\alpha}{2} z\right\} , \tag{3.1.58}$$

$$I(z) = I(0) \exp\{-\alpha z\} \quad (3.1.59)$$

in pumped media:

$$I(z) = I(0) \exp\{g z\}$$
 (3.1.60)

with

Different convention after (3.1.6): α , g, k_e and κ are defined with other signs, for example $\hat{n} = n (1 + i \kappa)$ if the other time separation (1st convention) is used [99Bor, Chap. 13], [95Bas, Vol. I, Chap. 9].

of α : see [85Koh, 04Ber, 82Bru], [90Roe, p. 34], [95Bas, Vol. II, Chap. 35].

3.1.5.1 Dielectric media

In Fig. 3.1.17 the real- and imaginary part of the refractive index in the vicinity of a resonance in the UV are shown.

$$\hat{n} = n - i k_{e} = 1 + \frac{N e^{2}}{2 \varepsilon_{0} m (\omega_{0}^{2} - \omega^{2} + i \gamma \omega)}$$

$$= \left\{ 1 + \frac{N e^{2} \gamma (\omega_{0}^{2} - \omega^{2})}{2 \varepsilon_{0} m [(\omega_{0}^{2} - \omega^{2})^{2} + \gamma^{2} \omega^{2}]} \right\} - i \left\{ \frac{N e^{2} \gamma \omega}{2 \varepsilon_{0} m [(\omega_{0}^{2} - \omega^{2})^{2} + \gamma^{2} \omega^{2}]} \right\}$$
(3.1.61)

with

 $e=-1.602\times 10^{-19}$ C: elementary charge, $m=9.109\times 10^{-31}$ kg: mass of the electron, $\omega=2\,\pi\nu$ [s $^{-1}$]: circular frequency of the light, ω_0 [s $^{-1}$]: circular resonant frequency of the electron,

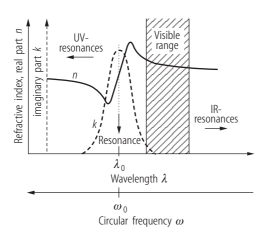


Fig. 3.1.17. Real- and imaginary part of the refractive index in the vicinity of a resonance in the UV. The principal shape is explained by the classical oscillator model after J.J. Thomson, P. Drude, and H.A. Lorentz [99Bor, 88Yeh].

```
\gamma [s<sup>-1</sup>]: damping coefficient,
    N [m^{-3}]: density of molecules,
    \varepsilon_0 = 8.8542 \times 10^{-12} \text{ As/Vm}: electric permittivity of vacuum.
see [96Ped, 88Kle], generalization to dense media see [96Ped, 88Kle, 99Bor].
The connects n(\omega) with k(\omega) [88Yeh].
```

3.1.5.2 Optical glasses

Dispersion formula [95Bac]:

$$n^{2}(\lambda) = 1 + \frac{B_{1}\lambda^{2}}{\lambda^{2} - C_{1}} + \frac{B_{2}\lambda^{2}}{\lambda^{2} - C_{2}} + \frac{B_{3}\lambda^{2}}{\lambda^{2} - C_{3}} \quad (, , , ,) .$$

$$(3.1.62)$$

The dimensions of the constants are given in example 3.1.7. The available wavelength range is given by the transmission limits, usually.

[96Sch]: Glass N-BK7: λ [μ m], $B_1 = 1.03961212$, $B_2 = 2.31792344 \times 10^{-1}$, $B_3 = 1.01046945$, $C_1 = 6.00069867 \times 10^{-3}$ [μ m²], $C_2 = 2.00179144 \times 10^{-2}$ [μ m²], $C_3 = 1.03560653 \times 10^2$ $[\mu m^2]$, $n(0.6328 \ \mu m) = 1.51509$, $n(1.06 \ \mu m) = 1.50669$.

Other interpolation formulae for $n(\lambda)$ are given in [95Bac], [95Bas, Vol. II, Chap. 32], [05Gro1, p. 121].

_____ is available from glass catalogs (see Sect. 3.1.5.10) and from subroutines in commercial optical design programs:

- $u_{\rm d} = \frac{n_{\rm d}-1}{n_{\rm F}-n_{\rm C}}$ with $n_{\rm d}(587.56~{\rm nm}={\rm yellow~He}$ line), $n_{\rm F}(486.13 \text{ nm} = \text{blue H-line}), n_{\rm C}(656.27 \text{ nm} = \text{red H-line})$ [95Bac, 80Sch]; achromatic correction of systems [84Haf], - , , , of n and $\nu_{\rm d}$, 1-11 1-112-121 -27 1113 (Verdet's constant),
- chemical resistance, thermal conductivity, micro hardness etc.

Sellmeier-like formulae for crystals are available in [95Bas, Vol. II, Chap. 32]. Information in connection with laser irradiation damage is presented in [82Hac]. Specific values of given in tables in [01Iff].

3.1.5.3 Dispersion characteristics for short-pulse propagation

The parameters can be calculated from the dispersion interpolation (3.1.62) [91Sal, 96Die]:

$$c_{\rm ph} = \frac{c_0}{n(\nu)} \quad (\dots \quad [\text{m s}^{-1}]) ,$$
 (3.1.64)

$$v = \frac{2\pi}{\frac{\mathrm{d}\beta}{\mathrm{d}\nu}} = \frac{1}{\frac{\mathrm{d}\beta}{\mathrm{d}\omega}} \quad (\text{m s}^{-1}]), \qquad (3.1.65)$$

$$D_{v} = \frac{1}{2\pi} \frac{\mathrm{d}^{2} \beta}{\mathrm{d} \nu^{2}} = 2\pi \frac{\mathrm{d}^{2} \beta}{\mathrm{d} \omega^{2}} = \frac{\mathrm{d}}{\mathrm{d} \nu} \left(\frac{1}{v} \right) \quad (3.1.66)$$

with

 ν : frequency of light,

 c_0 : velocity of light in vacuum.

Application: Temporal pulse forming by the GVD of dispersive optical elements [96Die, 01Ben].

3.1.5.4 Optics of metals and semiconductors

The refractive index of α_0 is characterized by free-electron contributions ($\omega_0 = 0$ in (3.1.61)). One obtains from [67Sok, 72Woo], [95Bas, Vol. II, Chap. 35] with a plasma resonance (here collision-free: $\gamma = 0$):

$$n^2(\omega) = 1 - \left(\frac{\omega_{\rm p}}{\omega}\right)^2 \tag{3.1.67}$$

with

 $\omega_{\rm p}$ [s⁻¹]: plasma frequency, depending on free-electron density [88Kle].

From (3.1.67) follows

- $-n(\omega) < 1$ for $\omega > \omega_p$, which means $\lambda < \lambda_p$ (example: $\lambda_p = 209$ nm for Na): transparency,
- pure imaginary $n(\omega)$ for $\omega < \omega_{\rm p}$, $\lambda_{\rm p} < \lambda$.

Other effects change the ideal case (3.1.67) [88Kle].

The complex refractive index of [1, • [1] is determined by transitions of electrons between or within the energy bands and by photon interaction with the crystal lattice (reststrahlen wavelength region). It depends strongly on the wavelength and is modified by heterostructures and dopands [71Pan, 95Kli], [95Bas, Vol. II, Chap. 36].

3.1.5.5 Fresnel's formulae

Fresnel's formulae describe the transmission and reflection of plane light waves at a plane interface between

- homogeneous isotropic media: [99Bor, 88Kle] and other textbooks on optics,
- homogeneous isotropic medium and anisotropic medium: special cases [99Bor, 86Haf] and other textbooks on optics,
- general case of anisotropic media: [58Fed],
- modification by photonic crystals: [95Joa, 01Sak].

Fresnel's formulae for the ..., ... (field) ..., ..., ..., ..., are listed in Table 3.1.7.

: plane, containing the wave number vector \boldsymbol{k} of the light and the normal vector \boldsymbol{n} on the interface.

Table 3.1.7. Fresnel's formulae for the amplitude (field) reflection and transmission coefficients.

Case	The four values Θ , Θ' , \hat{n} , and \hat{n}' are considered	Using the angles Θ and Θ' only	$\sin\Theta'$ is eliminated $\bar{n} = \frac{\hat{n}'}{\hat{n}}$	
Reflection				
$r_{ m s}=rac{{E_{ m s}}^{\prime\prime}}{E_{ m s}}=$	$\frac{\hat{n} \cos \Theta - \hat{n}' \cos \Theta'}{\hat{n} \cos \Theta + \hat{n}' \cos \Theta'}$	$-\frac{\sin(\Theta-\Theta')}{\sin(\Theta+\Theta')}$	$\frac{\cos\Theta - \sqrt{\bar{n}^2 - \sin^2\Theta}}{\cos\Theta + \sqrt{\bar{n}^2 - \sin^2\Theta}}$	(3.1.68)
Reflection				
$r_{\rm p} = \frac{E_{\rm p}^{\prime\prime}}{E_{\rm p}} =$	$\frac{\hat{n}' \cos \Theta - \hat{n} \cos \Theta'}{\hat{n}' \cos \Theta + \hat{n} \cos \Theta'}$	$\frac{\tan(\Theta-\Theta')}{\tan(\Theta+\Theta')}$	$\frac{\bar{n}^2 \cos \Theta - \sqrt{\bar{n}^2 - \sin^2 \Theta}}{\bar{n}^2 \cos \Theta + \sqrt{\bar{n}^2 - \sin^2 \Theta}}$	(3.1.69)
Transmission				
$t_{ m s} = rac{E_{ m s}'}{E_{ m s}} =$	$\frac{2\hat{n} \cos \Theta}{\hat{n}' \cos \Theta + \hat{n} \cos \Theta'}$	$\frac{2 \sin \Theta' \cos \Theta}{\sin(\Theta + \Theta')}$	$\frac{2 \cos \Theta}{\cos \Theta + \sqrt{\bar{n}^2 - \sin^2 \Theta}}$	(3.1.70)
Transmission				
$t_{\rm p} = \frac{E_{\rm p}^{\prime}}{E_{\rm p}} =$	$\frac{2\hat{n} \cos \Theta}{\hat{n}' \cos \Theta + \hat{n} \cos \Theta'}$	$\frac{2 \sin \Theta' \cos \Theta}{\sin(\Theta + \Theta') \cos(\Theta - \Theta')}$	$\frac{2\bar{n} \cos \Theta}{\cos \Theta + \bar{n}^2 \sqrt{\bar{n}^2 - \sin^2 \Theta}}$	(3.1.71)
Application of cases	Mostly used for pure dielectric media.	In a stack of films, the angles to the axis were calculated previously.	See remark in Sect. 3.1.5.5.	

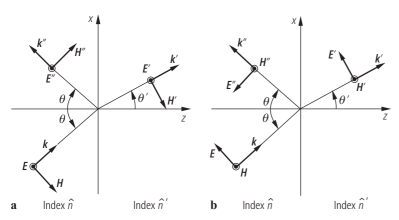


Fig. 3.1.18. Refraction at an interface, represented in the plane of incidence: (a) $E_{\rm s}$ -case, (b) $E_{\rm p}$ -case. The commonly used *convention* is shown for the orientation of the relevant vectors (k: the wave number vector, E: the electrical field, and H: the magnetic field) ensuring that k, E, and H are a right-handed system in every case. The E-field is important for the action on a nonmagnetic material.

/ | /- 16-19₁₁:

- E , , , to the plane of incidence: s-polarization (TE-case or σ -case [88Kle]), the corresponding E-component is called E_{\perp} [99Bor] or $E_{\rm s}$ (s: "senkrecht" (German) which means "perpendicular") [88Yeh] or index E [97Hua] or index x [90Roe, 77Azz, 91Sal].
- E_{μ} , to the plane of incidence: p-polarization (TM-case or π -case [88Kle]), the corresponding E-component is called E_{\parallel} [99Bor] or $E_{\rm p}$ [88Yeh] or index M [97Hua] or index y[90Roe, 77Azz, 91Sal].

$$\hat{n} \sin \Theta = \hat{n}' \sin \Theta' \tag{3.1.72}$$

with

 \hat{n}, \hat{n}' : refractive indices of both media, respectively, Θ , Θ' : see Fig. 3.1.18.

Other convention than Fig. 3.1.18b [58Mac, 89Gha, 91Ish] (electrical engineering) on the orientation of the **E**-vectors: **E** and **E**" point into the same direction for $\Theta \to 0$, **H** changes sign; application: **E**interferences.

- \hat{n} is real and \hat{n}' is complex (_______,,•_____ [76Fed, 77Azz] or ______ [88Boi]). \hat{n} and \hat{n}' are real and $\bar{n} = \frac{\hat{n}'}{\hat{n}} < 1$ and $(\bar{n}^2 \sin^2 \Theta) < 0$ (________). Then $\sqrt{\bar{n}^2 \sin^2 \Theta} = \frac{\hat{n}'}{\hat{n}}$ i $\sqrt{\sin^2\Theta - \bar{n}^2}$ yields for (3.1.68) and (3.1.69) $r_{\rm s} = \exp{({\rm i}\,\delta_{\rm s})}$ and $r_{\rm p} = \exp{({\rm i}\,\delta_{\rm p})}$ (modulus = 1, all energy reflected) and $\tan{\frac{\delta_{\rm s}}{2}} = -\frac{\sqrt{\sin^2\Theta - \bar{n}^2}}{\cos\Theta}$ and $\tan{\frac{\delta_{\rm p}}{2}} = -\frac{\sqrt{\sin^2\Theta - \bar{n}^2}}{\bar{n}^2\cos\Theta}$.

The intensities in the media are calculated with help of the z-component of Poynting's vector [88Kle, 90Roe, 76Fed].

(reflected part of intensity):

$$R_{\rm s,p} = |r_{\rm s,p}|^2$$
 (3.1.73)

(transmitted part of intensity):

$$T_{s,p} = \frac{\operatorname{Re}(\hat{n}' \cos \Theta')}{\operatorname{Re}(\hat{n} \cos \Theta)} |t_{s,p}|^2$$
(3.1.74)

with

Re: real part.

- 1 1/2111 .- 111:

$$T_{s,p} + R_{s,p} = 1$$
.

3.1.5.6 Special cases of refraction

3.1.5.6.1 Two dielectric isotropic homogeneous media (\hat{n} and \hat{n}' are real)

$$r_{\rm s} = \frac{n - n'}{n + n'} = -r_{\rm p} \ .$$
 (3.1.75)

(The negative sign of r_p results from the convention of Fig. 3.1.18 that E_p is diffracted into $-E_p''$).

$$R_{\rm s} = R_{\rm p} = \left(\frac{n - n'}{n + n'}\right)^2$$
 and $T_{\rm s} = T_{\rm p} = 1 - R_{\rm s}$. (3.1.76)

$$n = 1, n' = 1.5 \text{ (glass)}$$
: $R_s = 0.04$.

3.1.5.6.2 Variation of the angle of incidence

3.1.5.6.2.1 External reflection (n < n')

(angle of polarization) $\Theta_{\rm B}$:

$$\Theta_{\rm B} + \Theta_{\rm B}' = 90^{\circ} \,, \quad R_{\rm p} = 0 \,, \quad \tan \Theta_{\rm B} = \frac{n'}{n} \,.$$
 (3.1.77)

 $n = 1, n' = 1.5, \Theta_{\rm B} = 56.3^{\circ}$. See Fig. 3.1.19.

3.1.5.6.2.2 Internal reflection (n > n')

•,•,-, of total reflection:

$$\sin \Theta_{\rm C} = \frac{n'}{n} \ . \tag{3.1.78}$$

and $r_{\rm p}=\exp\left(\mathrm{i}\,\Phi_{\rm p}\right)$.

$$\tan \Theta_{\rm B} = \frac{n'}{n} \,. \tag{3.1.79}$$

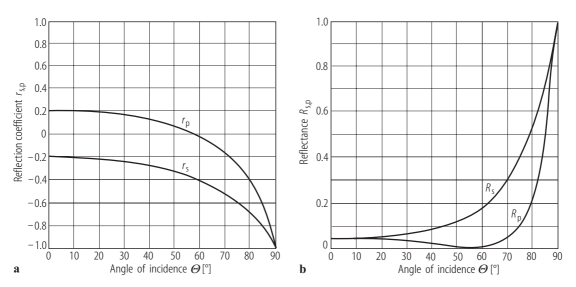


Fig. 3.1.19. (a) Reflection coefficients r_p and r_s and (b) reflectances R_p and R_s for n=1 and n'=1.5.

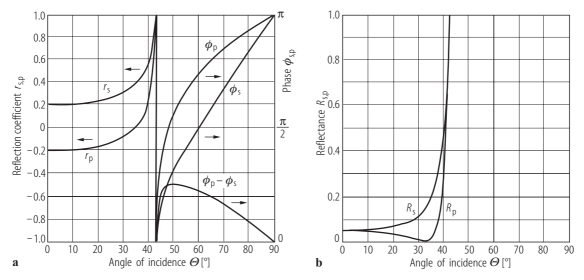


Fig. 3.1.20. Internal reflection (n = 1.5, n' = 1). (a) Reflection coefficients $r_{\rm p}$ and $r_{\rm s}$ for $\Theta < \Theta_{\rm C}$ and phases $\Phi_{\rm p}$ and $\Phi_{\rm s}$ for $\Theta > \Theta_{\rm C}$. (b) Reflectances $R_{\rm p}$ and $R_{\rm s}$ (= 1 for $\Theta > \Theta_{\rm C}$).

$$n = 1.5, n' = 1, \Theta_{\rm C} = 41.8^{\circ}, \Theta_{\rm B} = 33.7^{\circ}$$
. See Fig. 3.1.20.

in Fig. 3.1.21 [88Kle, p. 67]:

$$d_{\rm pen} = \frac{\lambda_0}{2\pi\sqrt{n^2\sin^2\Theta - n'^2}} \ . \tag{3.1.80}$$

[88Yeh, p. 74], see Fig. 3.1.22:

$$d_{G.-H.,s,p} = \frac{\mathrm{d}\,\Phi_{s,p}}{\mathrm{d}\,\Theta} \tag{3.1.81}$$

with $\Phi_{\rm p}$ and $\Phi_{\rm s}$ from Fig. 3.1.20. For a more precise treatment of the Goos–Hänchen shift for Gaussian beams see [05Gro1, p. 100].

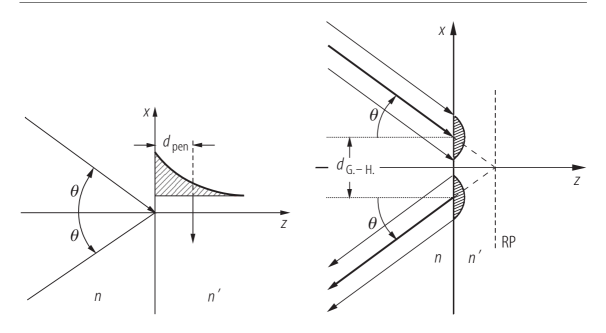
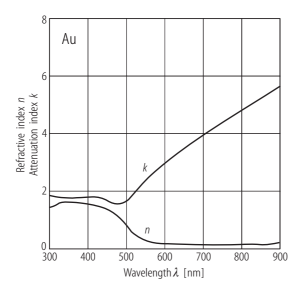


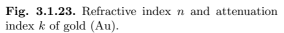
Fig. 3.1.21. Total reflection of plane waves with an inhomogeneous wave in the medium with the refractive index n' (d_{pen} : amplitude $\Rightarrow 1/e$).

Fig. 3.1.22. Goos–Hänchen shift of a total reflected beam with finite (exaggerated small) cross section (RP: effective reflection plane).

3.1.5.6.3 Reflection at media with complex refractive index (Case $\hat{n}=1$ and $\hat{n}'=n'+\mathrm{i}\,k'$)

In Fig. 3.1.23 the refractive index n and the attenuation index k of gold (Au) is shown, in Fig. 3.1.24 the reflectance for both polarization cases of gold is given.





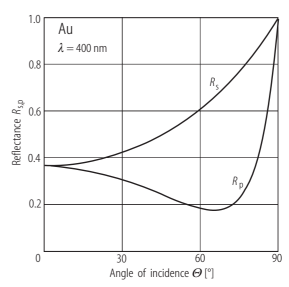


Fig. 3.1.24. Reflectance for both polarization cases of gold (Au). There is a minimum of $R_{\rm p}$ which is connected with a *pseudo Brewster angle*.

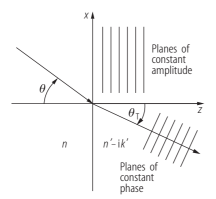


Fig. 3.1.25. Refraction at a medium with absorption: generation of an inhomogeneous wave.

(Fig. 3.1.25):
$$m_T = \frac{n}{n_T} \sin \Theta$$
 is modified:
$$(3.1.82)$$

with

$$2\,n_{\rm T}^2 = n^{\,\prime\,2} - k^{\,\prime\,2} + n^2\sin^2\Theta + \sqrt{\left(n^{\,\prime\,2} - k^{\,\prime\,2} - n^2\sin^2\Theta\right)^2 + 4\,n^{\,\prime\,2}\,k^{\,\prime\,2}} \quad ({\rm Ketteler's\ formula}) \; .$$

The effective refractive index $n_{\rm T}$ determines the direction angle $\Theta_{\rm T}$ of planes of constant phase in Fig. 3.1.25 via (3.1.82) [88Kle, p. 78], [41Str, p. 503], [99Bor, p. 740]. The full inhomogeneous wave can be calculated using [99Bor, p. 740].

$$\Theta = 45^{\circ}$$
, Au: $\lambda = 800$ nm, $n' = 0.19$, $k' = 4.9$, $n_{\rm T} = 0.73$, $\Theta_{\rm T} = 75.1^{\circ}$ (see [28Koe, p. 209]).

in the case
$$\Theta = 0$$
°:
$$I = I_0 \exp\left\{-2 \left(\omega/c\right) k' z\right\} . \tag{3.1.83}$$

$$\Theta=0^{\circ}$$
, Au: $\lambda=800$ nm, $n'=0.19,\ k'=4.9,\ I=I_0\exp\left(-7.7\times10^4\,z\text{[mm]}\right)$, $1/e-\text{depth}=13$ nm.

can be measured. The complex refractive index of a material results [77Azz, 90Roe]. Application: Measurements for the optical constants of metals, semiconductors, and thin-film systems.

3.1.5.7 Crystal optics

3.1.5.7.1 Classification

The dielectric tensor $\varepsilon_{\rm r}=\varepsilon_{ij}$ in (1.1.8) is symmetrical and real in the case of a nonabsorbing medium

In Fig. 3.1.26 vectors connected with wave propagation in crystal optics are depicted. In Table 3.1.8 optical crystals are listed. In Table 3.1.9 three of the eight surfaces for visualization of wave propagation in crystals are presented.

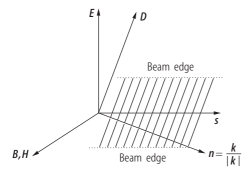


Fig. 3.1.26. Vectors connected with wave propagation in crystal optics [99Bor]: s: ray direction unit vector \parallel Poynting vector $E \times H$, n: unit vector in the normal direction $\parallel k$ and \perp phase planes, orthogonalities: $B, H \perp E, D, n, s$; $E \perp s$; $D \perp n$.

Table 3.1.8. Optical crystals.

Classification: system (syngony) of crystal	Refractive index in the main axis system	Optical type of crystal	Example	Values of the refractive index for $\lambda = 589.3 \text{ nm}$
triclinic, monoclinic, orthorhombic	$n_x \neq n_y \neq n_z \neq n_x$	biaxial crystal, no ordinary waves	$ m NaNO_3$	$n_x = 1.344$, $n_y = 1.411$, $n_z = 1.651$
trigonal, tetragonal, hexagonal	$n_x = n_y = n_o$ (ordinary index)	positive uniaxial crystal: $n_{\rm o} < n_{\rm e}$	SiO_2 (quartz)	$n_{\rm o} = 1.544,$ $n_{\rm e} = 1.553$
	$n_x \neq n_z = n_e$ (extraordinary index)	negative uniaxial crystal: $n_{\rm o} > n_{\rm e}$	CaCO ₃ (calcite)	$n_{\rm o} = 1.658,$ $n_{\rm e} = 1.486$
cubic	$n_x = n_y = n_z = n$	isotropic crystal	NaCl	n = 1.544

Table 3.1.9. Three of the eight surfaces for visualization of wave propagation in crystals.

Surface	Given	Found by construction are the
Index ellipsoid (indicatrix) (one-shell surface)	normal direction n	D -vectors for the two polarization cases and the two refractive indices for phase propagation
Index surface, wave vector surface (two-shell surface)	normal direction \boldsymbol{n}	ray directions s , which are perpendicular to the surface for both polarization cases
Ray surface, wave surface, representing Huygens' elementary wave for both polarization cases (two-shell surface)	ray direction \boldsymbol{s}	normal direction $\boldsymbol{n},$ which is perpendicular to the surface

Main feature of $s_1, \ldots, s_n : s_1, \ldots, s_n$ for wave propagation, mostly.

- s is essential for description of the energy propagation (edges of bundles, rays),
- n is essential for description of the interferences of infinite broad waves.

References: [28Szi, 54Bel, 58Shu, 61Ram, 76Fed, 79Wah, 84Yar, 04Ber, 99Pau, 99Bor]. A detailed comparison between that surfaces is given in [28Szi].

3.1.5.7.2 Birefringence (example: uniaxial crystals)

11-11-12 1/1-1/1/10 1/1-1 1/1/1/1/2 1/2:

- n of wavefronts: The wavevector surface is shown in Fig. 3.1.27.

$$\sin \Theta_{\rm o} = \frac{n}{n_{\rm o}} \sin \Theta \quad (\quad \bullet_{\rm o} \quad (\mathbf{k}_{\rm o})) \tag{3.1.84}$$

 $(n_0, \dots, n_1, \dots, n_n)$ on the angle of incidence),

 $(n_{\rm e}$, on the angle of incidence).

Poynting vector): \mathbf{s}_{e} and \mathbf{s}_{o} are given by tangent construction in Fig. 3.1.28. for the calculation of \mathbf{k}_{o} ($\|\mathbf{s}_{o}\|$), \mathbf{k}_{e} , \mathbf{s}_{e} of Fig. 3.1.28 with n, n_{o} , n_{e} , η , θ of Fig. 3.1.29 [86Haf]:

$$n_{\theta e}^{2} = \frac{A}{B} + \frac{n^{2}(n_{o}^{2} - n_{e}^{2})^{2} \sin^{2}\Theta \sin^{2}(2\eta)}{2B^{2}}$$

$$\pm \frac{n(n_{o}^{2} - n_{e}^{2})^{2} \sin\Theta \sin(2\eta)}{B} \times \sqrt{n^{2} \sin^{2}\Theta \left[\frac{(n_{o}^{2} - n_{e}^{2})^{2} \sin^{2}(2\eta)}{4B^{2}} - 1\right] + \frac{A}{B}}$$
(3.1.86)

with

$$A = (n_{\rm e}^2 - n_{\rm o}^2) \, n^2 \, \sin^2 \Theta \, \cos \left(2 \eta \right) - n_{\rm o}^2 \, n_{\rm e}^2 \; , \label{eq:A}$$

$$B = n_0^2 + (n_e^2 - n_0^2) \sin^2 \eta$$
,

where the decision on the \pm sign in (3.1.86) can be made by controlling the satisfaction of

$$n_{\theta e}^{2} \left[n_{o}^{2} + (n_{e}^{2} - n_{o}^{2}) \sin^{2}(\eta + \Theta_{e}) \right] = n_{e}^{2} n_{o}^{2}.$$

The resulting angles are:

$$\Theta_{o} = \arcsin(n \sin \Theta/n_{o}), \qquad (3.1.87)$$

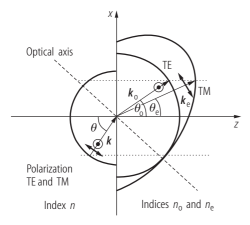


Fig. 3.1.27. Construction of wavefront birefringence with the wavevector surface: The wavefronts show no transversal limitation.

Ref. p. 131] 3.1 Linear optics 107

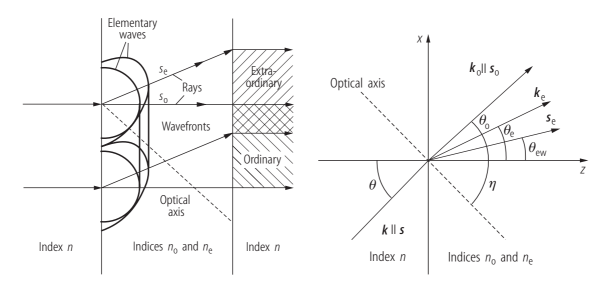


Fig. 3.1.28. Huygens' tangent construction of birefringence in a crystal slab for transversal-limited beams.

Fig. 3.1.29. Refraction for normal and ray directions. η : angle between z-axis and optical axis.

$$\Theta_{\rm e} = \arcsin\left(n\sin\Theta/n_{\Theta\,\rm e}\right) ,$$
 (3.1.88)

$$\Theta_{\text{ew}} = \arctan \frac{\tan \eta - C}{1 + C \tan \eta} \tag{3.1.89}$$

with

$$C = \frac{n_{\rm o}^2}{n_{\rm e}^2} \times \frac{\sqrt{n_{\Theta\,\mathrm{e}}^2 - n^2 \sin^2 \Theta} \; \tan \eta + n \, \sin \Theta}{\sqrt{n_{\Theta\,\mathrm{e}}^2 - n^2 \sin^2 \Theta} \; - n \, \sin \Theta \, \tan \eta} \; . \label{eq:constraint}$$

 $\Theta_{\rm o}$, $\Theta_{\rm e}$, $\Theta_{\rm e}$, $\Theta_{\rm o}$, $\Theta_{\rm o}$, $\Theta_{\rm o}$, and $\Theta_{\rm e}$ \Rightarrow phase differences (interferences) and reflection coefficient, $\Theta_{\rm o}$ and $\Theta_{\rm ew}$ \Rightarrow ray separation in a crystal.

Calcite:
$$n_{\rm o}=1.658,\,n_{\rm e}=1.486,\,\eta=45^\circ$$
: #1: $\Theta=0^\circ$: $C=1.244822,\,n_{\Theta\,{\rm e}}=1.565,\,\Theta_{\rm o}=\Theta_{\rm e}=0^\circ,\,\Theta_{\rm ew}=-6.224^\circ$; #2: $\Theta=45^\circ$: $n_{\Theta\,{\rm e}}=1.636,\,C=0.438329,\,\Theta_{\rm o}=25.23^\circ,\,\Theta_{\rm e}=25.6^\circ,\,\Theta_{\rm ew}=21.33^\circ$.

General formulation of (3.1.85)–(3.1.89): see [76Fed, Table 9.1] for more detailed discussions.

3.1.5.8 Photonic crystals

Starting with the forbidden (stop) bands in case of multi-layer Bragg reflection [88Yeh, p. 123] a material class is under development which stops light propagation along as many directions and for as many wavelengths as possible. This suppresses the spontaneous emission for laser applications and opens new possibilities in the micro- and nano-optics [95Joa, 01Sak, 04Bus]. Photonic crystal fibers [04Bus] can be designed for special light propagation properties and high-power fiber lasers [03Wad].

3.1.5.9 Negative-refractive-index materials

The common excitation of electrical dipoles and magnetical dipoles by light in a medium can result in a negative dielectric permittivity $\text{Re}(\varepsilon) < 0$ in combination with a negative magnetic permeability $\text{Re}(\mu) < 0$. Then, in Snell's law (3.1.72) an effective index $\hat{n}' < 0$ is possible [68Ves] which results in imaging by a slab of this material without curved surfaces [00Pen] and other interesting effects [05Ram]. Such metamaterials can be generated by microtechnology, now for mm- and terahertz-waves, but with the trend towards visible radiation [05Ele].

3.1.5.10 References to data of linear optics

[62Lan] contains optical constants, only. In later editions, the optical constants are listed together with other properties of substances. An overview is given in the content volume [96Lan].

Optical glass: [62Lan, Chap. 283], [97Nik], [95Bas, Vol. 2, Chap. 33], cat-

alogs of producers: [96Sch, 98Hoy, 96Oha, 92Cor], and com-

mercial optical design programs.

Infrared materials: [98Pal, 91Klo], [96Sch, infrared glasses], commercial optical

design programs.

Crystals: [62Lan, Chap. 282], [95Bas, Vol. 2, Chap. 33], [97Nik, 91Dmi,

81Kam].

Photonic crystals: [95Joa, 01Sak, 04Bus].

Negative-refractive-index materials: [05Ram].

Polymeric materials: [62Lan, Chap. 283], [95Bas, Vol. 2, Chap. 34], [97Nik]. Metals: [62Lan, Chap. 281], [98Pal], [95Bas, Vol. 2, Chap. 35]. Semiconductors: [96Lan, 98Pal, 87EMI], [95Bas, Vol. 2, Chap. 36].

Solid state laser materials: [01Iff, 97Nik, 81Kam].

Liquids: [62Lan, Chaps. 284, 285], [97Nik].

Gases: [62Lan, Chap. 286].

3.1.6 Geometrical optics

Geometrical optics represents the limit of the wave optics for $\lambda \Rightarrow 0$.

The development $\sin \sigma = \sigma - \frac{1}{3!}\sigma^3 + \frac{1}{5!}\sigma^5 - \ldots$ with σ the angle in Snell's law characterizes the different approaches of geometrical optics. Table 3.1.10 gives an overview of different approximations of geometrical optics.

3.1.6.1 Gaussian imaging (paraxial range)

The signs of the parameters determined in [03DIN, 96Ped] are applied in Sect. 3.1.6.1.1, later on f = f' is used.

Table 3.1.10. Different approximations of geometrical optics.

Problem to be treated	Algorithm for solving
Given: object point O in the paraxial range, asked: image point O' in the paraxial range approximation: $\sin \sigma \approx \sigma$.	 Gaussian collineation and Listing's construction: see Sect. 3.1.6.1. Gaussian matrix formalism (ABCD-matrix): see Sect. 3.1.6.2, ref.: [04Ber, 99Bor].
Imaging in Seidel's range, asked: imaging quality approximation: $\sin \sigma \approx \sigma - \frac{1}{3!}\sigma^3$.	Formulae for Seidels aberrations: see Sect. 3.1.6.3, ref.: [70Ber, 80Hof, 84Haf, 84Rus, 86Haf, 91Mah].
General image formation.	(Commercial) ray tracing programs with geometric and wave optical merit functions and tolerancing, ref.: [84Haf, 86Haf].

3.1.6.1.1 Single spherical interface

Figure 3.1.30 shows the imaging by a spherical interface in the paraxial range (small x, x', h).

$$n\left(\frac{1}{r} - \frac{1}{s}\right) = n'\left(\frac{1}{r} - \frac{1}{s'}\right) \quad \text{or} \quad \frac{n'}{s'} = \frac{n}{s} + \frac{n' - n}{r} . \tag{3.1.90}$$

 $n\left(\frac{1}{r}-\frac{1}{s}\right)$ is a constant on both sides of the interface.

$$f = -\frac{nr}{n'-n} \,. \tag{3.1.91}$$

$$f' = \frac{n'r}{n'-n} \ . \tag{3.1.92}$$

Remark: The symbol f means outside this section, Sect. 3.1.6.1, the positive focal length for a positive (converging) lens.

$$zz' = ff'. (3.1.93)$$

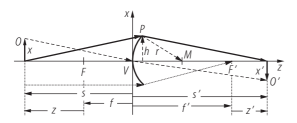


Fig. 3.1.30. Imaging by a spherical interface in the paraxial range (small x [object height], x' [image height], h [zonal height]). Full line: axial imaging, dashed line: off-axis imaging, dotted line: focusing to image side F'. Sign conventions: s, s' > 0, if they point to the right-hand side of the vertex V, r > 0, if the center of curvature of the interface is on the right-hand side in comparison with V. Here: s < 0, s' > 0, r > 0. M: center of curvature of the sphere. The left-hand-side-directed arrows symbolize negative values for the corresponding parameters here.

$$x' n' s' = x n s \tag{3.1.94}$$

with

n: object-space refractive index,

n': image-space refractive index,

s: object distance,

s': image distance,

r: radius of curvature of the interface,

x: height of the object point,

x': height of the image point,

z: focus-related object distance,

z': focus-related image distance.

cession via (3.1.90) by using $s_{\text{following surface}} = s'_{\text{prior surface}} - d$, d: the distance between the surfaces, and (3.1.94) for an object height $x \neq 0$.

3.1.6.1.2 Imaging with a thick lens

Figure 3.1.31 shows the axial imaging with a thick lens, Fig. 3.1.32 depicts $\bullet_{i_1}\bullet_{i_2}\bullet_{i_3}\bullet_{i_4}$

$$-\frac{1}{a} + \frac{1}{a'} = \frac{1}{f'} = (n'-1)\left(\frac{1}{r_1} - \frac{1}{r_2}\right) + \frac{t(n'-1)^2}{n'r_1r_2}.$$
(3.1.95)

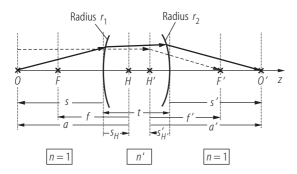


Fig. 3.1.31. Axial imaging with a thick lens. Cardinal planes and points are: object-space principal plane with object principal point H on axis, image-space principal plane with image principal point H' on axis, object-space focal point F, image-space focal point F'. Nodal points [98Mah, 96Ped] are equal to the principal points if O and O' are embedded in media with equal refractive index as here. Then f = -f'. The sign convention used here means: Parameters characterized by an arrow pointing to the left (right) hand side show a negative (positive) sign [80Hof, 86Haf]. The dashed line shows the use of H' for simplifying the plot for a ray focusing.

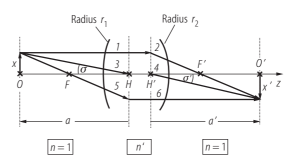


Fig. 3.1.32. Listing's construction for thick-lens imaging of a finite-height object point O to image point O'. Scheme of construction: Ray 1 (parallel with axis) is sharply bent at plane H' towards F'. Ray 3 towards H is continued at H' with the angle $\sigma' = \sigma$. Ray 5 through F is bent sharply parallel with axis at H-plane. The magnification x'/x = a'/a can be calculated by elimination of a' from $(3.1.95) \Rightarrow x'$.

 $\mathcal{L}_{1}, \mathcal{L}_{1}, \mathcal{L}_{1}, \mathcal{L}_{2}, \mathcal{L}_{3}, \mathcal{L}_{4}, \mathcal{L}_{5}, \mathcal{L}$

$$s_H = -\frac{n'-1}{n'r_2}f't \ . \tag{3.1.96}$$

 $\mathcal{L}_{i_1}, \mathcal{L}_{i_1}, \mathcal{L}_{i_2}, \mathcal{L}_{i_3}, \mathcal{L}_{i_4}, \mathcal{L}_{i_5}, \mathcal{L}$

$$s_{H'} = -\frac{n'-1}{n'r_1}f't. (3.1.97)$$

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$$\overline{H}H' = t \left(1 - f' \frac{n' - 1}{n'} \left(\frac{1}{r_1} - \frac{1}{r_2} \right) \right) . \tag{3.1.98}$$

 \bullet_{t} , $t \Rightarrow 0: (3.1.95) \Rightarrow$ "Lens maker's formula".

3.1.6.2 Gaussian matrix (ABCD-matrix, ray-transfer matrix) formalism for paraxial optics

. , , can be treated with the help of the ray-transfer matrix:

- 1. full description of , ..., (this section, Sect. 3.1.6.2),

The optical system can be the separating distance in an optical medium, a single spherical optical surface or a true, more complicated optical system.

There are •. , or the ABCD-matrices:

: The slope components of the input and output rays are the real angles without any relation to the refractive indices at input and output spaces of Fig. 3.1.33 [66Kog1, 66Kog2, 84Hau, 91Sal, 95Bas, 96Ped, 96Yar, 98Hec, 98Sve, 01Iff, 05Gro1, 05Hod]. Then, the determinant of the matrix $\mathsf{M} \colon \|\mathsf{M}\| = n'/n$ with n the index of the medium of the input plane and n' the index of the medium of the output plane.

[75Ger, 86Sie, 88Kle, 04Ber] use:

slope parameter = (angle) \times (related refractive index). Then the equation $\|M\| = 1$ applies.

In Fig. 3.1.34 the concatenation of different ray-transfer matrices for different types of subsystems is shown.

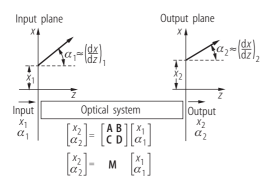


Fig. 3.1.33. Transfer of the input height x_1 and slope α_1 into the output height x_2 and slope α_2 with the ray-transfer matrix M. The sign of slope α_1 is positive in this figure. The German standard DIN 1335 uses a different sign with change of some signs in the ABCD matrices [96Ped].

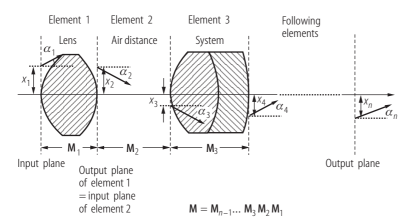


Fig. 3.1.34. Concatenation of different ray-transfer matrices for different types of sub-systems. Matrices known for systems before can be used to construct the matrix for a larger system containing the known systems. The sequence of the matrices is shown at the bottom of the figure.

3.1.6.2.1 Simple interfaces and optical elements with rotational symmetry

In Table 3.1.11 ABCD-matrices for simple interfaces and optical elements with rotational symmetry are listed.

3.1.6.2.2 Non-symmetrical optical systems

Rotational symmetry lacks and the axis is tilted due to the system, the central ray of imaging is called the basic ray. The optics in a narrow region around the basic ray is called heaving a system, vol. [95Bas, Vol. 1, p. 1.47] as analogon to paraxial optics. For treatment of astigmatic pencils see [72Sta].

A special case of the non-symmetrical optical system is a $_{i_1,i_2}$, $_{i_1,i_2}$, $_{i_1,i_2}$: Two orthogonal cases do not mix during propagation. Examples are different setups of spectroscopy and laser physics (ring resonators).

In Table 3.1.12 ABCD-matrices for non-symmetrical optical elements without torsion are listed.

3.1.6.2.3 Properties of a system

In Table 3.1.13 distances between __ • _ _ , _ _ of an optical system are listed, in Table 3.1.14 the meaning of the vanishing of different elements of the ABCD-matrix is depicted.

3.1.6.2.4 General parabolic systems without rotational symmetry

The generalization of the two-dimensional ray transfer after Fig. 3.1.33 to three dimensions [69Arn] is shown in Fig. 3.1.35. The ray in the input plane is characterized by two coordinates x_1 and y_1 of the piercing point P and two small (paraxial range) angles α_1 and β_1 .

The matrix S relates these parameters to the corresponding parameters in the output plane like in Fig. 3.1.33:

 $\textbf{Table 3.1.11.} \ \mathsf{ABCD}\text{-}matrices \ \mathsf{for \ simple \ interfaces \ and \ optical \ elements \ with \ rotational \ symmetry.}$

Effect	Figure	ABCD-matrix	Remark
Propagation	- d - ► 1 0	$\begin{bmatrix} 1 & d \\ 0 & 1 \end{bmatrix}$	The rays propagate from I to O within the same medium.
Spherical surface	n_1 n_2	$\left[\frac{1}{n_1 - n_2} \frac{0}{n_1} \right]$	Sign: $r > 0$ for convex surface seen by the propagating light.
Plane	n_1 n_2	$\left[\begin{matrix} 1 & 0 \\ 0 & \frac{n_1}{n_2} \end{matrix}\right]$	Corresponds to a spherical surface with $r \Rightarrow \infty$.
Planar plate	$\begin{bmatrix} n_1 \\ n_2 \end{bmatrix}$ $\begin{bmatrix} n_1 \\ n_2 \end{bmatrix}$	$\begin{bmatrix} 1 & \frac{n_1}{n_2} d \\ 0 & 1 \end{bmatrix}$	Contains two refractions.
Thin lens	r_1 r_2 r_1 r_2 r_1	$\begin{bmatrix} 1 & 0 \\ -\frac{1}{f} & 1 \end{bmatrix}$	$ \frac{1}{f} = \frac{n_2 - n_1}{n_1} \left[\frac{1}{r_1} - \frac{1}{r_2} \right] , $ air: $n_1 = 1$.
Thick lens in air	$(r_1>0)/r_2<0$ $(r_1>0)/r_2<$	$\begin{bmatrix} 1 - \frac{s_{H'}}{f} & \frac{d}{n} \\ -\frac{1}{f} & 1 + \frac{s_H}{f} \end{bmatrix}$	$\begin{split} &\frac{1}{f} = (n-1) \left[\frac{1}{r_1} - \frac{1}{r_2} \right] + \frac{(n-1)^2 t}{n r_1 r_2} , \\ &s_H = -\frac{(n-1) f t}{n r_2} , \text{ see } (3.1.96) , \\ &s_{H'} = -\frac{(n-1) f t}{n r_1} , \text{ see } (3.1.97) , \\ &H, H' \colon \text{principal planes.} \end{split}$
Spherical mirror	substituted by	$\begin{bmatrix} 1 & 0 \\ -\frac{2}{r} & 1 \end{bmatrix}$	Unfolding of the mirror; $ {\rm sign}(r)>0 , {\rm if \ the \ incident \ light \ sees} $ a concave mirror surface.
Gradient- index lens or thermal lens	$\begin{bmatrix} n_1 \\ \\ \\ \end{bmatrix}$ $\begin{bmatrix} n_1 \\ \\ \\ \end{bmatrix}$ $\begin{bmatrix} n_1 \\ \\ \\ \end{bmatrix}$	$\begin{bmatrix} A & B \\ C & D \end{bmatrix}$ $n = n_0 (1 - \gamma x^2);$ $\gamma > 0: \text{ higher}$ index on axis	$A = \cos\left(\sqrt{2\gamma} t\right);$ $B = n_1 \sin\left(\sqrt{2\gamma} t\right) / \left(n_0 \sqrt{2\gamma}\right);$ $C = -\left(\sqrt{2\gamma} n_0 / n_1\right) \sin\left(\sqrt{2\gamma} t\right);$ $D = \cos\left(\sqrt{2\gamma} t\right);$ development of the trigonometric functions for $\sqrt{2\gamma} t \ll 1 \Rightarrow \text{simplifications}$
			Gradient optics: see [02Gom, 05Gro1].

Table 3.1.11 continued.

Effect	Figure	ABCD-matrix	Remark
Gaussian apodization, usable for q-parameter transfer (Table 3.1.18)	1 0	$\begin{bmatrix} 1 & 0 \\ -\frac{i\lambda a}{2\pi} & 1 \end{bmatrix}$ $\lambda : \text{ wavelength of light}$	The amplitude transmission function between I and O is $\exp\left(-ax^2/2\right)$, x : transverse coordinate [86Sie, p. 787]

Remark: Other treatments of the mirror see [86Sie, 98Sve, 75Ger].

 ${\bf Table~3.1.12.~ABCD-} {\bf matrices~for~non-symmetrical~optical~elements~without~torsion.}$

Effect	Figure	ABCD-matrix	Remark
Refraction at a sphere Tangential (meridional) plane	θ_1 n_1 n_2	$\begin{bmatrix} \frac{\cos(\theta_1)}{\cos(\theta_2)} & 0\\ \frac{\Delta n_t}{r n_2} & \frac{n_1 \cos(\theta_2)}{n_2 \cos(\theta_1)} \end{bmatrix}$	$n_1 \sin(\theta_1) = n_2 \sin(\theta_2)$ (Snell's law) $\Delta n_t = \frac{n_2 \cos(\theta_2) - n_1 \cos(\theta_1)}{\cos(\theta_1) \cos(\theta_2)}$
Sagittal plane	n_1	$\left[\begin{array}{cc} 1 & 0 \\ \frac{\Delta n_{\mathrm{s}}}{r n_2} & \frac{n_1}{n_2} \end{array}\right]$	$\Delta n_{\rm s} = n_2 \cos(\theta_2) - n_1 \cos(\theta_1)$
Rowland concave grating (unfolded) Tangential (meridional) plane	θ_1 θ_2 Radius of curvature r	$\begin{bmatrix} A & B \\ C & D \end{bmatrix},$ $A = \frac{\cos(\theta_1)}{\cos(\theta_2)};$ $B = 0;$ $C = -\frac{2\cos(\theta_2)}{r_t\cos(\theta_1)};$ $D = A.$	Grating equation (3.1.52): $\sin(\theta_1) + \sin(\theta_2) = m \frac{\lambda}{g},$ $r_t = \frac{2r \cos^2(\theta_2)}{\cos(\theta_1) + \cos(\theta_2)}$
Sagittal plane	-	$\begin{bmatrix} 1 & 0 \\ -\frac{2}{r_{\rm s}} & 1 \end{bmatrix}$	$r_{\rm s} = rac{2r}{\cos{(heta_1)} + \cos{(heta_2)}} \; ,$ general corrected holographical gratings: see [81Gue]
Spherical concave mirror		Specialization of the Rowland grating to $g \Rightarrow \infty$, $\theta_1 = \theta_2$.	

Table 3.1.13. Distances between cardinal elements of an optical system: F, F': object- and image-space focal points, respectively; H, H': object- and image-space principal points, respectively; I, O: input and output plane, respectively. The order of points determines the signs.

Distance between two points	$A, B, C, \text{ and } D$ for $n_1 = n_2$
\overline{IF}	$\frac{D}{C}$
\overline{FH}	$-\frac{1}{C}$
$\overline{OF'}$	$-rac{A}{C}$
$\overline{HF^{\prime}}$	$-\frac{1}{C}$

Table 3.1.14. The meaning of the vanishing of different elements of the ABCD-matrix.

Element	Figure	Remark
A = 0	O B O	$x_2 = B \alpha_1$ Focusing of collimated light into the image-side focal plane.
B = 0	A 0 O	$x_2 = A x_1$ The input plane is <i>imaged</i> to the output plane (conjugated planes). A: magnification of imaging; appl.: calculation of image plane.
C = 0	AB OD	$\alpha_2 = D \alpha_1$ Transformation of collimated light into collimated light. D : angular magnification; telescope (afocal system).
D = 0	$\begin{bmatrix} \mathbf{A} \ \mathbf{B} \\ \mathbf{C} \ 0 \end{bmatrix}$	$\alpha_2 = C x_1$ Collimation of divergent pencil of rays. C: power of the element or system.

$$\begin{bmatrix} x_2 \\ y_2 \\ \alpha_2 \\ \beta_2 \end{bmatrix} = \begin{bmatrix} A_{xx} & A_{xy} & B_{xx} & B_{xy} \\ A_{yx} & A_{yy} & B_{yx} & B_{yy} \\ C_{xx} & C_{xy} & D_{xx} & D_{xy} \\ C_{yx} & C_{yy} & D_{yx} & D_{yy} \end{bmatrix} \begin{bmatrix} x_1 \\ y_1 \\ \alpha_1 \\ \beta_1 \end{bmatrix} \quad \text{or} \quad \begin{bmatrix} \mathbf{r}_2 \\ \mathbf{\gamma}_2 \end{bmatrix} = \begin{bmatrix} \mathsf{A} & \mathsf{B} \\ \mathsf{C} & \mathsf{D} \end{bmatrix} \begin{bmatrix} \mathbf{r}_1 \\ \mathbf{\gamma}_1 \end{bmatrix} = \mathsf{S} \begin{bmatrix} \mathbf{r}_1 \\ \mathbf{\gamma}_1 \end{bmatrix}$$
(3.1.99)

with the matrices A, B, C, D, and S given by comparison with the more detailed representations.

Identities between the matrices, characteristic for the symplectic geometry (see Sect. 3.1.6.2.6), are:
$$AD^{\mathsf{T}} - BC^{\mathsf{T}} = I$$
; $AB^{\mathsf{T}} = BA^{\mathsf{T}}$; $CD^{\mathsf{T}} = DC^{\mathsf{T}}$, and $\det \begin{vmatrix} A & B \\ C & D \end{vmatrix} = \frac{n'}{n}$, where T means the

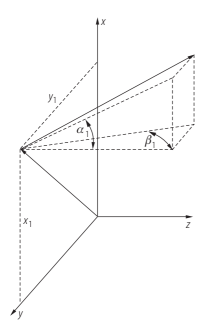


Fig. 3.1.35. Three-dimensional ray in the input plane I.

transposition of the matrix and I the identity matrix [86Sie, 05Hod]. The matrix S contains at most 10 independent parameters [76Arn, 86Sie, 05Hod].

In Table 3.1.15 general ray-transfer matrices are given.

3.1.6.2.5 General astigmatic system

can be generated by two cylindrical lenses with their axes non-parallel and non-orthogonal, separated by a distance $L: S_{GA} = R^{-1} S_{cyl,1} R S_L S_{cyl,2}$.

3.1.6.2.6 Symplectic optical system

geometry [03Wal]. They can be generated by a finite number of cylindrical and spherical lenses separated by free spaces. The mathematical formulation is connected with the matrix properties given in Sect. 3.1.6.2.4. For theoretical foundation and practical calculations see [64Lun, p. 216], [83Mac, 85Sud, 86Sie, 99Gao, 05Gro1, 05Hod].

3.1.6.2.7 Misalignments

The geometric optical calculations of [0.5], with matrix techniques require, generally, higher dimensional matrices [05Gro1, p. 51], for example 3×3 -matrices [86Sie] or 4×4 -matrices [85Wan] for two-dimensional problems or 6×6 -matrices for three-dimensional problems [76Arn].

 ${\bf Table~3.1.15.~General~ray\text{-}transfer~matrices~[99Gao,~05Hod]}.$

Effect of the matrix	Matrix
Free propagation, index n_0 , length z	$S_{\mathrm{L}} = \begin{bmatrix} 1 & 0 & \frac{z}{n_0} & 0 \\ 0 & 1 & 0 & \frac{z}{n_0} \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 0 \end{bmatrix}$
Aligned spherical thin lens, focal length f	$S_{\mathrm{sph}} = \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ \frac{-1}{f} & 0 & 1 & 0 \\ 0 & \frac{-1}{f} & 0 & 1 \end{bmatrix}$
Aligned cylindrical thin lens	$S_{ ext{cyl}} = \left[egin{array}{ccc} 1 & 0 & 0 & 0 \ 0 & 1 & 0 & 0 \ -1 \ f_x & 0 & 1 & 0 \ 0 & 0 & 0 & 1 \end{array} ight]$
Cylindrical telescope, m and n are the magnifications along x - and y -axis, respectively	$S_{\mathrm{M}} = \begin{bmatrix} m & 0 & 0 & 0 \\ 0 & n & 0 & 0 \\ 0 & 0 & m^{-1} & 0 \\ 0 & 0 & 0 & n^{-1} \end{bmatrix}$
Rotation of the <i>x-y</i> -plane by the angle θ : given a system matrix S , then the rotated system matrix $S_{\rm rot} = R^{-1}\left(\theta\right) S\left(\theta = 0\right) R\left(\theta\right)$	$R = \begin{bmatrix} \cos\theta & \sin\theta & 0 & 0 \\ -\sin\theta & \cos\theta & 0 & 0 \\ 0 & 0 & \cos\theta & \sin\theta \\ 0 & 0 & -\sin\theta & \cos\theta \end{bmatrix}$

3.1.6.3 Lens aberrations

with $R^{-1}(\theta) = R(-\theta) = R^{T}(\theta)$

are required by large object-space aperture light sources like (large vertical far-field angles) or large image-space aperture laser focusing optics like

$$q = \frac{r_2 + r_1}{r_2 - r_1} \,. \tag{3.1.100}$$

Shape factor and spherical aberration for focusing of light:

$$\frac{r_1}{r_2} = \frac{n(2n-1)-4}{n(2n+1)} \ .$$

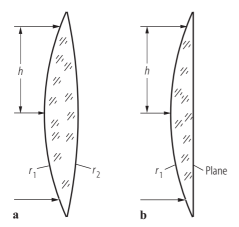


Fig. 3.1.36. Focusing of incident collimated light by (a) a general lens with curvature radii r_1 and r_2 , (b) a plano-convex lens with shape factor q = 1.

- Refractive index $n=1.5 \Rightarrow \frac{r_1}{r_2} = -\frac{1}{6} \Rightarrow q=0.7$.
- $-\left|\frac{r_1}{r_2}\right|\Rightarrow \frac{1}{\infty}\Rightarrow q=1$ (plano-convex lens), spherical aberration near to minimum.

In Fig. 3.1.36 the focusing of incident collimated light by (a) a general lens with curvature radii r_1 and r_2 and (b) a plano-convex lens with shape factor q = 1 is shown.

In Table 3.1.16 the third-order spherical aberration and coma for a thin plano-convex lens is given in comparison with the diffraction-limited resolution for a plane wave or Gaussian illumination.

Remark 1: Third-order formulae for finite object distance: see [88Kle, 76Jen], more general: [80Hof, 86Haf, 96Ped, 99Bor].

Remark 2: About further third-order aberrations as a stigmatism, field curvature, image distortion: see [76Jen, 78Dri, 80Hof, 86Haf, 88Kle, 96Ped, 99Bor].

Remark 3: The third-order aberrations are not exactly valid for higher apertures. Example: The third-order spherical aberration deviates for 2h/f=1/5 by $\approx 2\,\%$ from the ray-tracing values (the limit, recommended in [74Sle] for estimations), h/f=3/10: $\approx 15\,\%$ deviation [76Jen]. Therefore, the ray tracing should be preferred for larger deviations from the paraxial case. It is the base of modern commercial optical design programs.

Given: a plano-convex lens after Fig. 3.1.36b with the radius of the spherical surface $r_1=5$ mm, n=1.5, collimated light with wavelength $\lambda=1$ µm, stop with a height h=1.5 mm, and a fiber with core diameter $2\,r=100$ µm and numerical aperture N.A.=0.2. Required: a geometric-optical estimation on the hits of the core of the fiber by the rays in the paraxial focal point and in the point of least confusion (Fig. 3.1.37). From (3.1.101)–(3.1.105): f=10 mm, $\Delta s_1'=-262$ µm, $|\Delta s_1'|=39$ µm, $\Delta s_{1c}'=-210$ µm, $|\Delta s_{tc}'|=16$ µm, $\Delta s_{tb}=4$ µm, and $\Delta s_{tg}'=2.1$ µm. In the paraxial focal plane as well as in the plane of least confusion, the hits of the fiber core by rays are closer than 50 µm to the optical axis and the angles of the rays with the optical axis are ≤ 0.15 within the fiber aperture. Therefore, all rays are accepted by a step-index fiber. About the analog task for Gaussian beams see references in Sect. 3.1.7.5.4 and commercial optical design programs, which show in this case, that a large part of radiation is coupled in higher-order modes.

Table 3.1.16. Third-order spherical aberration and coma for a thin plano-convex lens [76Jen, p. 152], [88Kle, p. 185], [87Nau, p. 109] in comparison with the diffraction-limited resolution for a plane wave or Gaussian illumination.

Figures

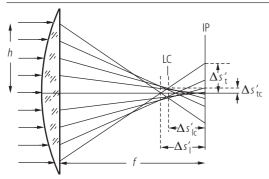


Fig. 3.1.37. Spherical aberration at a plano-convex lens. IP: paraxial image plane, LC: least confusion.

Formulae

Lens equation (3.1.95) with t=0, $a\Rightarrow -\infty$, $r_2\Rightarrow -\infty$, f=f', which is modified outside Sect. 3.1.6.1:

$$\frac{1}{f} = (n-1)\frac{1}{r_1} \,, \tag{3.1.101}$$

$$\frac{\Delta s_1'}{f} = -\frac{n^3 - 2n^2 + 2}{2n(n-1)^2} \left(\frac{h}{f}\right)^2 , \qquad (3.1.102)$$

$$\Delta s_{\rm t}' = \Delta s_{\rm l}' \, \frac{h}{f} \,, \tag{3.1.103}$$

plane of least confusion [87Nau, 99Pau, 99Bor], [01Iff, p. 214]:

$$\Delta s_{\rm lc}' \approx 0.8 \,\Delta s_{\rm l}' \,, \tag{3.1.104}$$

$$\Delta s_{\rm tc}^{\prime} \approx 0.4 \, \Delta s_{\rm t}^{\prime} \,.$$
 (3.1.105)

Gaussian weights of the illumination change the geometric optical position of least confusion [01Mah].

$$x' = \theta f , \qquad (3.1.106)$$

$$\frac{a}{f} = -\frac{n^2 - n - 1}{2n(n-1)} \left(\frac{h}{f}\right)^2 \theta \tag{3.1.107}$$

with

 θ : angle of incidence.

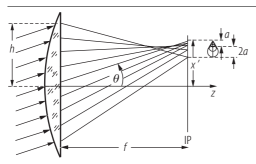


Fig. 3.1.38. Coma at a plano-convex lens.

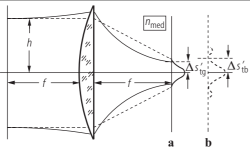


Fig. 3.1.39. Diffraction-limited resolution for (a) a Gaussian beam with waist h ($1/e^2$ -intensity level) in the object-side focal plane, (b) a plane wave at circular stop with radius h.

$$\Delta s_{\rm tg}' = \frac{\lambda}{\pi n_{\rm med}(h/f)} , \qquad (3.1.108)$$

$$\Delta s'_{\rm tb} = 0.61 \frac{\lambda}{n_{\rm med} \sin \sigma'}$$

$$\approx 0.61 \frac{\lambda}{n_{\rm med} (h/f)}$$
(3.1.109)

with

 λ : wavelength [m], h: zonal height [m],

f: focal length [m],

 $n_{\rm med}$: refractive index of the image space.

3.1.7 Beam propagation in optical systems

Paraxial propagation of light in a system given by its ABCD-matrix can be calculated

- Chap. 2.2 (beam characterization).

3.1.7.1 Beam classification

In Table 3.1.17 various types of beams are listed.

3.1.7.2 Gaussian beam: complex q-parameter and its **ABCD-transformation**

3.1.7.2.1 Stigmatic and simple astigmatic beams

3.1.7.2.1.1 Fundamental Mode

- and rotational-symmetric system:
 - ⇒ both longitudinal cross sections are treated equally,
- and elements with a symmetry plane:
 - \Rightarrow two different sets of ABCD-matrices for the tangential and sagittal cut (see Table 3.1.12).

The q_1, \dots, q_n of the q_1, \dots, q_n [66Kog1, 66Kog2]

$$\frac{1}{q_x(z)} = \frac{1}{R_x(z)} - \frac{\mathrm{i}\,\lambda}{\pi w_x(z)^2} \tag{3.1.110}$$

formalizes the x-part of the fundamental-mode equation (3.1.31)

$$U_0(x,z) = \sqrt{\frac{w_{0x}}{w_x(z)}} \exp\left\{-\frac{x^2}{w_x(z)^2} - i\frac{kx^2}{2R_x(z)}\right\}$$
(3.1.111)

to the simple complex shape

$$U_0(x,z) = \frac{1}{\sqrt{1 + i\frac{z}{z_0}}} \exp\left\{-i\frac{kx^2}{q_x(z)}\right\}.$$
(3.1.112)

In Fig. 3.1.40 the transfer of a field distribution by an optical system given by its ABCD-matrix is shown. In Table 3.1.18 the q-parameter transfer for stigmatic and simple astigmatic beams is given.

Table 3.1.17. Types of beams.

Beam [69Arn, 05Hod]	Generated by	Beam type is characterized by the shape of the matrix S (3.1.99)	Examples	References with practical example calculations
Stigmatic θ_x θ_y θ_y θ_y θ_y	Fundamental-mode laser	$A_{xx} = A_{yy}$; $A_{xy} = A_{yx} = 0$, and the same for B, C, D	TE00-mode handling in laser applications	[75Ger, 86Sie, 91Sal], [96Yar, 01Iff, 05Hod], see Sect. 3.1.7.2.1
Simple astigmatic	Semiconductor lasers or: Anamorphic optical system (f.e. cylindrical lens) in combination with a stigmatic beam	$A_{xx} \neq A_{yy}$; $A_{yx} = A_{xy} = 0$ (no mixing of both orthogonal planes), and the analog for B, C, D	 ring lasers, lasers, including dispersive elements (dye-lasers), tolerance calculations for resonators and beamguiding optics 	[05Hod, 99Gao, 86Sie], see Sect. 3.1.7.2.1
General astigmatic	General rotation of an anamorphic optics in relation with a simple astigmatic beam	General case	Transformation of higher- order radiation modes	[05Hod, 99Gao], see Sect. 3.1.7.2.2

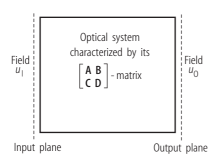


Fig. 3.1.40. Transfer of a field distribution by an optical system given by its ABCD-matrix.

Table 3.1.18. q-parameter transfer for stigmatic and simple astigmatic beams.

Œ	łi	V€	en

Propagated field

- Gaussian beam in the *input* plane:

$$u_{\rm I}(x,z) = \exp\left\{-i\frac{kx^2}{2q_{\rm I}x}\right\}$$
 . (3.1.113)

- ABCD-matrix of the optical system (see Tables 3.1.11 and 3.1.12).
- Starting point:

$$R_{\rm \,I}=1/{\rm Re}\left(1/q_{\rm \,I}\right)\,,$$

$$w_{\rm I} = 1/\sqrt{-\pi \operatorname{Im}\left(1/q_{\rm I}\right)/\lambda} \; .$$

- Transformation of the q-parameter:

$$q_{Ox} = \frac{Aq_{Ix} + B}{Cq_{Ix} + D} . (3.1.114)$$

- Field in the output plane:

$$u_{O}(x,z) = \exp\left\{-i\frac{kx^2}{2q_{Ox}}\right\}$$
 (3.1.115)

with the real parameters of the output beam [96Gro]:

- beam radius:

$$w_{Ox} = w_{Ix} \sqrt{\left(\frac{\lambda B}{\pi w_{Ix}^2}\right)^2 + \left(A + \frac{B}{R_{Ix}}\right)^2}, \quad (3.1.116)$$

- curvature radius of the wavefront:

$$R_{Ox} = \frac{\left(A + \frac{B}{R_{Ix}}\right)^2 + \left(\frac{\lambda B}{\pi w_{Ix}^2}\right)^2}{\left(A + \frac{B}{R_{Ix}}\right) \left(C + \frac{D}{R_{Ix}}\right) + D\left(\frac{\lambda B}{\pi w_{Ix}^2}\right)^2} \ . \tag{3.1.117}$$

Given: the waist of a Gaussian beam

$$u_{\mathrm{I}} = \exp\left\{-\frac{x^2}{w_0^2}\right\} = \exp\left\{-\mathrm{i}\,\frac{kx^2}{2q_1}\right\}$$

with $q_1 = i z_0$ in comparison with (3.1.110).

Asked: free-space propagation along the distance z with the ABCD-matrix $\begin{bmatrix} 1 & z \\ 0 & 1 \end{bmatrix}$.

$$q_2 = \frac{Aq_1 + B}{Cq_1 + D} = z + i z_0$$

and

$$U_{\rm O} = \left(A + \frac{B}{q_1}\right)^{-1/2} \exp\left\{-i\frac{kx^2}{2q_2}\right\} = \left(1 + \frac{z}{iz_0}\right)^{-1/2} \exp\left\{-i\frac{kx^2}{2(z+iz_0)}\right\}.$$

3.1.7.2.1.2 Higher-order Hermite-Gaussian beams in simple astigmatic beams

Treatment of the x- or y-component of Hermite-Gaussian-beams after (3.1.27): The complex q-parameter transformation is treated as above, the fundamental mode part is given as above, the new beam radius for the Hermite polynom of order m, $H_m(\sqrt{2} x/w_{Ix})$ is calculated from the new q-parameter and the phase is derived from it, too [70Col].

For complex Hermite-Gaussian beams: see [86Sie].

3.1.7.2.2 General astigmatic beam

In Table 3.1.19 the Q^{-1} -matrix transfer for general astigmatic beams is given. The matrix Q^{-1} is the matrix scheme of inverses of q-parameters and no inverted matrix [96Gro].

Table 3.1.19. Q^{-1} -matrix transfer for general astigmatic beams.

Given Propagated field

- General Gaussian beam in the input plane:

$$U_{\rm I}({m r}) = \exp\left\{ -{\rm i}\, rac{k}{2}\, {m r}\, {f Q}_{
m I}^{-1}\, {m r}
ight\} \; , \ \ (3.1.118)$$

 $r \sim (x,y)$ the transverse position vector perpendicular to the propagation axis z .

Q₁⁻¹-matrix:

$$Q_{I}^{-1} = \begin{bmatrix} \frac{1}{q_{xx}} & \frac{1}{q_{xy}} \\ \frac{1}{q_{xy}} & \frac{1}{q_{yy}} \end{bmatrix}$$
(3.1.119)

with q_{xx} , q_{xy} , q_{yy} complex terms describing the general amplitude- and phase-distribution of $U_{\rm I}$, and

$$\mathbf{r} \, \mathsf{Q}_{\mathrm{I}}^{-1} \, \mathbf{r} = \frac{x^2}{q_{xx}} + 2 \frac{xy}{q_{xy}} + \frac{y^2}{q_{yy}} \, . \quad (3.1.120)$$

 S-matrix of the optical system (see Table 3.1.15) with

$$S = \begin{bmatrix} A & B \\ C & D \end{bmatrix}$$

after (3.1.99).

- Transformation of the Q_I^{-1} -matrix to its output value:

$$Q_{O}^{-1} = (C + D Q_{I}^{-1}) (A + B Q_{I}^{-1})^{-1},$$
 (3.1.121)

see [88Sim, 96Gro, 05Hod].

- Field in the output plane:

$$U_{\mathcal{O}}(\mathbf{r}) = \exp\left\{-i\frac{k}{2}\mathbf{r}\,\mathsf{Q}_{\mathcal{O}}^{-1}\mathbf{r}\right\}. \tag{3.1.122}$$

Transformation of a simple astigmatic Gaussian beam (no mixing between x and y) with a θ -rotated cylindrical lens to a general astigmatic beam: We start with $\mathsf{Q}_{\mathrm{I}}^{-1} = \begin{bmatrix} \frac{1}{q_{xx}} & 0 \\ 0 & \frac{1}{q_{yy}} \end{bmatrix}$. The rotation of an x-aligned cylindrical lens, given as $\mathsf{S}_{\mathrm{cyl}}$ in Table 3.1.15, is performed by multiplication of the following property of the start with the

The rotation of an x-aligned cylindrical lens, given as $S_{\rm cyl}$ in Table 3.1.15, is performed by multiplying first $S_{\rm cyl}$ with the rotation matrix R of Table 3.1.15, and then the product with the inverse of R is:

$$\mathsf{S}_{\mathrm{rotated\ cyl.}} = \mathsf{R}^{-1}\,\mathsf{S}_{\mathrm{cyl}}\,\mathsf{R} = \left[egin{matrix}\mathsf{A}\ \mathsf{B}\\\mathsf{C}\ \mathsf{D}\end{smallmatrix}
ight]$$

with

$$\mathsf{A} = \begin{bmatrix} 1 \ 0 \\ 0 \ 1 \end{bmatrix} \ , \quad \mathsf{B} = \begin{bmatrix} 0 \ 0 \\ 0 \ 0 \end{bmatrix} \ , \quad \mathsf{C} = \begin{bmatrix} -\cos^2\theta/f_x & -\sin\theta\,\cos\theta/f_x \\ -\sin\theta\,\cos\theta/f_x & -\sin^2\theta/f_x \end{bmatrix} \ , \quad \mathsf{D} = \begin{bmatrix} 1 \ 0 \\ 0 \ 1 \end{bmatrix}$$

and

$$\mathsf{Q}_{\mathrm{O}}^{-1} = \begin{bmatrix} -\cos^2\theta/f_x + 1/q_{xx} & -\sin\theta\,\cos\theta/f_x \\ -\sin\theta\,\cos\theta/f_x & -\sin^2\theta/f_x + 1/q_{yy} \end{bmatrix} \,.$$

Therefore, the output field

$$u_{O}(\mathbf{r}) = \exp\left\{-i\frac{k}{2}\left[\left(-\frac{\cos^{2}\theta}{f_{x}} + \frac{1}{q_{xx}}\right)x^{2} - 2\frac{\sin\theta\cos\theta}{f_{x}}xy + \left(-\frac{\sin^{2}\theta}{f_{x}} + \frac{1}{q_{yy}}\right)y^{2}\right]\right\}$$

is a general astigmatic Gaussian beam with a mixing term between the coordinates x and y.

3.1.7.3 Waist transformation

Often, the transfer of the beam waist is required for instance for focusing of laser light. Then, the following algorithms are , than the q-parameter algorithm.

3.1.7.3.1 General system (fundamental mode)

In Table 3.1.20 the waist transformation for a general system is given.

3.1.7.3.2 Thin lens (fundamental mode)

The formulae (3.1.123)–(3.1.126) are further simplified using the focal length f for the thin lens only, see Table 3.1.21.

Remark: Discussion of equation (3.1.127):

The right-hand-side term of (3.1.127) containing z_0 represents the modification introduced by the Gaussian beam optics to the thin-lens equation ((3.1.95), $t \Rightarrow 0$) shown in Fig. 3.1.42.

In Fig. 3.1.43 the relation of the Gaussian waist transfer to the thin-lens equation of geometrical optics for different influences of diffraction is shown.

 ${\it Main\ modifications}$ of the geometrical optics:

- No "image distance" is at infinity.
- For z = f (point P) the image is at z' = f (transfer of the object-side focal plane to the image-side focal plane after (3.1.130), not $\Rightarrow \infty$).
- If a target z'-position is given, then two starting z-positions are possible.

Example 3.1.17. Given for Fig. 3.1.42: z=1179 mm, $w_0=0.22$ mm, $\lambda=1.06$ µm; it follows z'=109 mm, $w'_0=0.02$ mm, $\theta'=0.96^{\circ}$, and $z'_0=1.21$ mm. The second right-hand term of (3.1.127) translates the Gaussian waist image by 0.16 mm in comparison with the geometrical optical image towards the lens.

Table 3.1.20. Waist transformation for a general system.

- ABCD-matrix of the system,
- waist w_0 ,
- wavelength λ , including $z_0 = \pi w_0^2 / \lambda$,
- distance z to the input plane of the system.

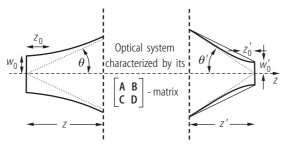


Fig. 3.1.41. Waist transformation by an optical system.

Asked: Waist w'_0 and distance z' to the output plane of the system including z'_0 .

 $\begin{cases} \frac{-(Az+B)(Cz+D) - ACz_0^2}{C^2 z_0^2 + (Cz+D)^2} & \text{for } C \neq 0, \\ -\frac{Az+B}{D} & \text{for } C = 0, \end{cases}$ for C=0, (3.1.123)

$$z_0' = z_0 \left[\frac{Cz' + A}{Cz + D} \right] = \frac{z_0}{C^2 z_0^2 + (Cz + D)^2} ,$$
(3.1.124)

$$w_0' = \sqrt{\frac{\lambda z_0'}{\pi}} \,, \tag{3.1.125}$$

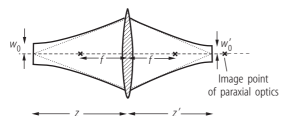
$$\Theta_0' = \sqrt{\frac{\lambda}{\pi z_0'}} \,. \tag{3.1.126}$$

The beam parameter product is invariant:

$$w_0' \Theta_0' = w_0 \Theta_0 = \lambda/\pi$$
.

Table 3.1.21. Waist transformation by a thin lens.

Given	Solution	
- Focal length f of the lens, - wavelength λ ,	$\frac{1}{z} + \frac{1}{z'} = \frac{1}{f} + \frac{z_0^2}{z \left[z^2 + z_0^2 - zf\right]} \; ,$	(3.1.127)
- waist w_0 , including $z_0 = \pi w_0^2/\lambda$,	see Fig. 3.1.42,	



distance z to the input plane of the system.

Fig. 3.1.42. Waist transformation by a thin lens.

 $w_0' = w_0 \frac{f}{\sqrt{z_0^2 + (z - f)^2}}$ (3.1.128)

$$z_0' = \frac{\pi w_0'^2}{\lambda} \ . \tag{3.1.129}$$

If z = f, then

$$z' = f$$
 and $w'_0 = \frac{w_0 f}{z_0}$. (3.1.130)

Asked: Waist w'_0 and distance z' to the output plane of the system and z_0' .

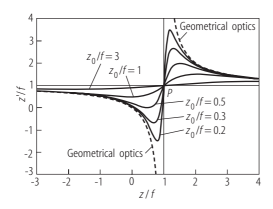


Fig. 3.1.43. Relation of the Gaussian waist transfer (full lines) to the thin-lens equation (dashed) of geometrical optics for different influences of diffraction (wavelength λ respectively z_0).

3.1.7.4 Collins integral

For Fresnel's approximation of diffraction in paraxial systems see [68Goo, 71Col, 78Loh, 94Roe]. It was generalized to the propagation of field distributions in ABCD-described systems by [70Col, 76Arn, 05Gro2, 05Hod].

3.1.7.4.1 Two-dimensional propagation

In Table 3.1.22 the propagation in rotational symmetric systems and simple astigmatic systems is given.

Table 3.1.22. Propagation in rotational symmetric systems and simple astigmatic systems.

Given	Solution	
 ABCD-matrix of the optical system (see Tables 3.1.11 and 3.1.12), field distribution in the input plane U_I(x), path length along the optical axis L. 	Field $U_{\rm O}(x_2)$ in the output plane (Collins integral): $U_{\rm O}(x_2) = \sqrt{\frac{\mathrm{i}}{\lambda B}} \ \mathrm{e}^{-\mathrm{i}kL} $ $\times \int\limits_{-\infty}^{\infty} \mathrm{d}x_1 U_{\rm I}(x_1) \exp\left\{-\mathrm{i}\frac{k}{2B} \left[Ax_1^2 - 2x_1x_2 + Dx_2^2\right]\right\} .$	(3.1.131)

The waist of a Gaussian beam is given with $U_{\rm I}(x_1)=\exp{(-x_1^2/w_1^2)}$ in the input plane. The system consists of a thin lens with the focal length f followed by a free-space propagation by distance z. The ABCD-matrix is calculated from Fig. 3.1.34 and Table 3.1.11:

$$\begin{bmatrix} A & B \\ C & D \end{bmatrix} = \begin{bmatrix} 1 - z/f & z \\ -1/f & 1 \end{bmatrix} .$$

$$U_{\mathcal{O}}(x_2) = \sqrt{\frac{\mathrm{i}}{\lambda z}} \, \mathrm{e}^{-\mathrm{i}kL} \int_{-\infty}^{\infty} \mathrm{d}x_1 \, \exp\left(-\frac{x_1^2}{w_1^2}\right) \exp\left\{-\mathrm{i}\frac{k}{2z} \left[\left(1 - \frac{z}{f}\right)x_1^2 - 2x_1x_2 + x_2^2\right]\right\} .$$

The result is an output Gaussian intensity distribution with the waist radius

$$w_{\rm O} = \frac{w_{\rm I}}{\sqrt{1 + \left(\frac{\pi w_{\rm I}^2}{\lambda f}\right)^2}} \; , \label{eq:wo}$$

the waist position $z=z_{\mathrm{waist}}=f\,\left(\frac{\pi\,w_{\,\mathrm{O}}\,w_{\,\mathrm{I}}}{\lambda\,f}\right)^2$, and $z_{\,\mathrm{O}}=\pi\,w_{\,\mathrm{O}}^2/\lambda$.

For inclusion of $\bullet_{i,j}$, $\bullet_{i,j}$, $\bullet_{i,j}$, $\bullet_{i,j}$, $\bullet_{i,j}$ in Collins Integral see [96Tov].

3.1.7.4.2 Three-dimensional propagation

In Table 3.1.23 the propagation in in general astigmatic systems is given.

Table 3.1.23. Propagation in general astigmatic systems.

Given	Solution
- S: matrix of the optical system, see Table 3.1.15 and (3.1.99) with $S = \begin{bmatrix} A & B \\ C & D \end{bmatrix},$	Field $U_{\mathrm{O}}(\boldsymbol{r}_{2})$ in the output plane (Collins integral): $U_{\mathrm{O}}(\boldsymbol{r}_{2}) = \frac{-\mathrm{i}\exp(-\mathrm{i}kL)}{\lambda\sqrt{\det\mathbf{B}}}\int\int\mathrm{d}\boldsymbol{r}_{1}U_{1}\left(\boldsymbol{r}_{1}\right)$ $\times\exp\left\{-\mathrm{i}\frac{k}{2}\left[\boldsymbol{r}_{1}B^{-1}A\boldsymbol{r}_{1}-2\boldsymbol{r}_{1}B^{-1}\boldsymbol{r}_{2}+\boldsymbol{r}_{2}DB^{-1}\boldsymbol{r}_{2}\right]\right\}(3.1.132)$
- field distribution in the input plane: $U_{\rm I}(\boldsymbol{r}_1)$, where \boldsymbol{r}_1 is the position vector in the input plane.	with det B the determinant and B^{-1} the inverse of the matrix B. Examples in [70Col, 05Gro2, 05Hod].

3.1.7.5 Gaussian beams in optical systems with stops, aberrations, and waveguide coupling

3.1.7.5.1 Field distributions in the waist region of Gaussian beams including stops and wave aberrations by optical system

- focused Gaussian beams with aberrations and stops: see [69Cam, 71Sch],
- obscuration of a rotationally symmetrical Gaussian beam including longitudinal focal shift: see [82Car, 86Sta],
- extended systematic discussion of diffraction with stops, obscuration, and aberrations: see [86Mah, 01Mah],
- spherical aberration: see [98Pu].

3.1.7.5.2 Mode matching for beam coupling into waveguides

The calculation of the excitation coefficient of an eigenmode in a waveguide (output mode) by the incident mode (input mode) at the surface of the waveguide is described in Table 3.1.24.

This task occurs

waveguide).

- if a laser beam is formed by an optical system and coupled afterwards into an optical fiber,
- if a laser beam of a master oscillator is to be coupled into a power amplifier,
- in the case of waveguide-waveguide coupling especially fiber-fiber coupling or coupling between semiconductor lasers.

Solutions are available in commercial optical design programs.

Table 3.1.24. Definitions for waveguide coupling.

Given	Solution	
- Incident beam (emitted by a laser	Coupling coefficient (power relation):	
(and) transformed by an optical system): $E_{\text{input}}(x, y)$. - Waveguide with an eigenmode field the	$\eta = \frac{O_{\mathrm{IO}} \ O_{\mathrm{IO}}^*}{N_{\mathrm{I}} N_{\mathrm{O}}} \ .$	(3.1.133)
coupling to which is asked: $E_{\text{output}}(x, y)$.	Overlap integral:	
Plane of mode matching	$O_{\rm IO} = \int_{-\infty}^{\infty} dx \int_{-\infty}^{\infty} dy E_{\rm I}(x,y) E_{\rm O}^*(x,y) .$	(3.1.134)
$E_{\text{input}}(x)$ $E_{\text{output}}(x)$ $E_{\text{output}}(x)$	Normalization:	
Waveguide	$N_{\rm I} = \int_{-\infty}^{\infty} \mathrm{d}x \int_{-\infty}^{\infty} \mathrm{d}y E_{\rm I}(x,y) E_{\rm I}^*(x,y) .$	(3.1.135)
Fig. 3.1.44. Mode matching.	Normalization:	
Asked: Part of power transmitted into the waveguide (fiber, laser, integrated optical	$N_{\rm O} = \int_{-\infty}^{\infty} dx \int_{-\infty}^{\infty} dy E_{\rm O}(x, y) E_{\rm O}^*(x, y) .$	(3.1.136)

3.1.7.5.3 Free-space coupling of Gaussian modes

For the case that a Gaussian output waist of a source waveguide and a Gaussian input waist of a receiver waveguide are separated by air, the coupling of both waveguides is generally treated in [64Kog]. Higher-order modes are also included. The approximation of small misalignments (offset and tilt) is given in Table 3.1.25, large offsets and tilts are treated in [64Kog, 91Wu].

waveguide.

Effective antireflection layers are assumed to be on the

Table 3.1.25. Coupling of waveguides .

Civen

- Source WG1 (laser, waveguide) which emits a Hermite-Gaussian beam,
- receiver WG2 (laser, waveguide) which can accept Hermite-Gaussian eigenmodes:

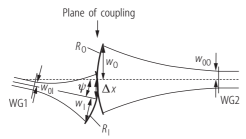


Fig. 3.1.45. Coupling of Gaussian beams. w_{01} and w_{00} : beam waist radii for WG1 and WG2, respectively; w_{1} and w_{0} : beam radii in the coupling plane; R_{1} and R_{0} : curvature radii of the beam wavefronts in the coupling plane; $k = 2\pi/\lambda$; λ : the wavelength of light; Δx : the lateral offset between the waveguides; ψ : the tilt of the axis.

Asked: The efficiency of the excitation of the modes in WG2, here the fundamental mode 00.

Solution

$$\begin{split} \eta_{00-00} &= \frac{4}{\left(\frac{w_{\rm I}}{w_{\rm O}} + \frac{w_{\rm O}}{w_{\rm I}}\right)^2 + \left(\frac{\pi w_{\rm I} w_{\rm O}}{\lambda}\right)^2 \left(\frac{1}{R_{\rm I}} - \frac{1}{R_{\rm O}}\right)^2} \\ &- \frac{8 \left(w_{0\rm I} w_{0\rm O} \Delta x\right)^2}{\left(w_{0\rm I}^2 + w_{0\rm O}^2\right)^3} - \frac{k^2 \, \psi^2}{2} \left(w_{\rm I}^2 + w_{\rm O}^2\right) \; . \end{split} \tag{3.1.137}$$

 $\eta_{\,00-00}=1$ for $\Delta\,x=\psi=0$ and the exact beam radii and curvature fitting $w_{\,\rm I}=w_{\,\rm O}$ and $R_{\,\rm I}=R_{\,\rm O}$, otherwise $\eta_{\,00-00}<1$.

Equation (3.1.137) contains the approximations:

- Gaussian beams (paraxial optics).
- Right-hand side of (3.1.137): $2^{\rm nd}$ and $3^{\rm rd}$ term $\ll 1^{\rm st}$ term.

About coupling coefficients for higher-order modes and without the approximation: see [64Kog]; on couplings with Hermite-Gaussian modes and Laguerre-Gaussian modes: see [94Kri, 80Gra].

3.1.7.5.4 Laser fiber coupling

1-1 1 1 . 1 - 1 . 1 . 1 :

- Launching of ____ of a single-mode fiber:
 - Calculation of the overlap integral (3.1.134) for a Gaussian mode and the mode field for different fiber cross sections: see [88Neu, p. 179], [80Gra].
 - Approximation of the exact fiber fundamental modes by a Gaussian field distribution (see [88Neu, pp. 68]) and the application of the waist transformation from laser via an optical system with the methods of Sects. 3.1.7.2–3.1.7.4 and calculation of the overlap integral equation (3.1.134) or mode-coupling equation (3.1.137) [91Wu].
- Launching of fundamental-mode laser radiation or . . , or incoherent light sources into . . , :
 - Overlap integral techniques in the framework of partial coherence theory: see [87Hil].
 - Geometric optical methods (ray tracing and phase space techniques): see [90Gec, 95Sny, 91Gra, 91Wu, 01Iff].

- Inclusion of the of the optical system used:
 - Monomodal and partial coherent case: calculation of the wave aberrations of the optical system by ray-tracing methods and inclusion of these aberrations into the overlap integral: see [82Wag, 95Gae, 89Hil, 99Gue].
 - Ray-tracing methods are adequate for stops and aberrations, but not reliable for a few mode waveguides: rough design [01Iff]: the spot diagram of the ray tracing in the fiber facet should be within the core area and the angles of incidence should be smaller then the aperture angle [88Neu] of the fiber.

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4.1 Frequency conversion in crystals

G.G. GURZADYAN

4.1.1 Introduction

4.1.1.1 Symbols and abbreviations

4.1.1.1.1 Symbols

η	conversion efficiency		
η (energy)	energy conversion efficiency		
η (power)	power conversion efficiency		
η (quantum)	quantum conversion efficiency		
$ au_{ m p}, au$	pulse duration		
α	angle between interacting beams		
$\Delta \lambda$	wavelength bandwidth		
Δu	frequency bandwidth		
$\Delta \theta$	angular bandwidth		
E	energy		
f	laser pulse repetition rate		
I_0	pump intensity		
$I_{ m thr}$	threshold intensity		
$arphi_{ m pm}$	phase-matching angle in the XY plane from X axis		
L	crystal length		
λ	wavelength		
n	refractive index		
n_{o}	ordinary refractive index		
$n_{ m e}$	extraordinary refractive index		
ν	wave number, frequency		
P	power		
$ heta_{ m pm}$	phase-matching angle from Z axis		
ρ	birefringence (walk-off) angle		
$T, T_{\rm pm}$	crystal temperature		
, F	•		
Type I	$o + o \rightarrow e$ or $e + e \rightarrow o$		
Type II	$o + e \rightarrow e$ or $o + e \rightarrow o$		
ooe	$o + o \rightarrow e$ or $e \rightarrow o + o$		
eeo	$e + e \rightarrow o \text{ or } o \rightarrow e + e$		
eoe	$e + o \rightarrow e$ or $e \rightarrow e + o$		
oeo	$o + e \rightarrow o \text{ or } o \rightarrow e + o$		
	•		

142 4.1.1 Introduction [Ref. p. 187]

4.1.1.1.2 Abbreviations

av average cw continuous wave

DFG difference frequency generation

DROPO doubly resonant OPO
ERR external ring resonator
FIHG fifth harmonic generation
FOHG fourth harmonic generation

ICDFG intracavity difference frequency generation ICSHG intracavity second harmonic generation

IR infrared mid IR middle infrared NC noncollinear

NCSHG noncollinear second harmonic generation

OPA optical parametric amplifier OPO optical parametric oscillator SFG sum frequency generation

SH second harmonic

SHG second harmonic generation
SIHG sixth harmonic generation
SP OPO synchronously pumped OPO

SROPO singly resonant OPO

SRS stimulated Raman scattering
THG third harmonic generation
TROPO triply resonant OPO
TWOPO traveling-wave OPO

UV ultraviolet

4.1.1.1.3 Crystals

Chemical formula	Symbol	Crystal name
A m A nC		Duovotito
Ag_3AsS_3		Proustite
${ m AgGaS_2}$		Silver Thiogallate
$AgGaSe_2$		Silver Gallium Selenide
Ag_3SbS_3		Pyrargyrite
$\mathrm{Ba_2NaNb_5O_{15}}$		Barium Sodium Niobate (Banana)
β -BaB ₂ O ₄	BBO	Beta-Barium Borate
$CdGeAs_2$		Cadmium Germanium Arsenide
CdSe		Cadmium Selenide
CsB_3O_5	CBO	Cesium Borate
CsH_2AsO_4	CDA	Cesium Dihydrogen Arsenate
$CsLiB_6O_{10}$	CLBO	Cesium Lithium Borate
$C_6H_6N_2O_3$	POM	3-Methyl-4-Nitro-Pyridine-1-Oxide
$C_8H_8O_3$	MHBA	4-Hydroxy-3-Methoxy-Benzaldehyde (Vanillin)
$C_{10}H_{11}N_3O_6$	MAP	Methyl N-(2,4-Dinitrophenyl)-L-Alaninate
$C_{10}H_{13}N_3O_3$	DAN	N-[2-(Dimethylamino)-5-Nitrophenyl]-Acetamide
$C_{11}H_{14}N_2O_3$	NPP	N-(4-Nitrophenyl)-(L)-Propinol
CsD_2AsO_4	DCDA	Cesium Dideuterium Arsenate
GaSe		Gallium Selenide

$HgGa_2S_4$		Mercury Thiogallate
α -HIO ₃		α -Iodic Acid
KB_5O_8 $4D_2O$	DKB5	Potassium Pentaborate Tetradeuterate
KB_5O_8 $4H_2O$	KB5	Potassium Pentaborate Tetrahydrate
$\mathrm{KD_2AsO_4}$	DKDA	Potassium Dideuterium Arsenate
$\mathrm{KD_2PO_4}$	DKDP	Potassium Dideuterium Phosphate
$\mathrm{KH_{2}PO_{4}}$	KDP	Potassium Dihydrogen Phosphate
KNbO_3		Potassium Niobate
$\mathrm{KTiOAsO_4}$	KTA	Potassium Titanyl Arsenate
$KTiOPO_4$	KTP	Potassium Titanyl Phosphate
LiB_3O_5	LBO	Lithium Triborate
$LiCOOH H_2O$	LFM	Lithium Fomate
$LiIO_3$		Lithium Iodate
$LiNbO_3$		Lithium Niobate
LiNbO ₃ :MgO		Mg:O-doped Lithium Niobate
$(NH_2)_2CO$		Urea
$\mathrm{NH_4H_2AsO_4}$	ADA	Ammonium Dihydrogen Arsenate
$NH_4H_2PO_4$	ADP	Ammonium Dihydrogen Phosphate
$NO_2C_6H_4NH_2$	mNA	meta-Nitroaniline
RbH_2AsO_4	RDA	Rubidium Dihydrogen Arsenate
RbH_2PO_4	RDP	Rubidium Dihydrogen Phosphate
$RbTiOAsO_4$	RTA	Rubidium Titanyl Arsenate
Te		Tellurium
Tl_3AsSe_3		Thallium Arsenic Selenide
ZnGeP_2		Zinc Germanium Phosphide

4.1.1.2 Historical layout

The pioneering work of Franken et al. [61Fra] on second harmonic generation of ruby laser radiation in quartz and invention of the phase-matching concept [62Gio, 62Mak] generated a new direction in the freshly born field of nonlinear optics: frequency conversion in crystals. Sum frequency generation by mixing the outputs of two ruby lasers in quartz was already realized in 1962 [62Mil, 62Bas]. Zernike and Berman [65Zer] were the first to demonstrate difference frequency mixing. Optical parametric oscillation was experimentally realized in 1965 by Giordmaine and Miller [65Gio]. First monographs on nonlinear optics by Akhmanov and Khokhlov [64Akh] and Bloembergen [65Blo] greatly stimulated development of the nonlinear frequency converters. At present the conversion of laser radiation in nonlinear crystals is a powerful method for generating widely tunable radiation in the ultraviolet, visible, near, mid, and far IR regions.

For theoretical and experimental details of nonlinear frequency conversions in crystals, see monographs by Zernike and Midwinter [73Zer], Danelyus, Piskarskas et al. [83Dan], Dmitriev and Tarasov [87Dmi], Shen [84She], Handbook of nonlinear optical crystals (by Dmitriev, Gurzadyan, Nikogosyan) [91Dmi, 99Dmi], Handbook of nonlinear optics (by Sutherland) [96Sut]. For frequency conversion of femtosecond laser pulses, see also [88Akh]. For linear and nonlinear optical properties of the crystals, see [77Nik, 79Kur, 84Jer, 87Nik, 87Che, 96Sut, 99Dmi, 00Cha, 00Sas]. For related nonlinear phenomena, see [96Sut]. For the historical perspective of the nonlinear frequency conversion over the first forty years, see [00Bye]. In the following section, Sect. 4.1.2, we present some basic equations which may be useful for simple calculations of frequency converters.

144 4.1.2 Fundamentals [Ref. p. 187

4.1.2 Fundamentals

4.1.2.1 Three-wave interactions

Dielectric polarization P (dipole moment of unit volume of the substance) is related to the field E by the material equation of the medium [64Akh, 65Blo] (Chap. 1.1):

$$P(E) = \varepsilon_0 \left(\chi^{(1)} E + \chi^{(2)} E^2 + \chi^{(3)} E^3 + \dots \right)$$
(4.1.1)

with

 $\varepsilon_0 = 8.854 \times 10^{-12} \ \mathrm{CV^{-1}m^{-1}}$: dielectric permittivity of free space, $\chi^{(1)} = n^2 - 1$: the linear, and $\chi^{(2)}$, $\chi^{(3)}$ etc.: the nonlinear dielectric susceptibilities.

In the present chapter, Chap. 4.1, we consider only three-wave interactions in crystals with square nonlinearity ($\chi^{(2)} \neq 0$). The following nonlinear frequency conversion processes are considered:

Second Harmonic Generation (SHG):

$$\omega + \omega = 2\omega \,, \tag{4.1.2}$$

Sum-Frequency Generation (SFG) or up-conversion:

$$\omega_1 + \omega_2 = \omega_3 \,, \tag{4.1.3}$$

Difference-Frequency Generation (DFG) or down-conversion:

$$\omega_3 - \omega_2 = \omega_1 \,, \tag{4.1.4}$$

Optical Parametric Oscillation (OPO):

$$\omega_3 = \omega_2 + \omega_1 \ . \tag{4.1.5}$$

For efficient frequency conversion phase matching should be fulfilled:

$$\mathbf{k}_1 + \mathbf{k}_2 = \mathbf{k}_3 \tag{4.1.6}$$

with

 \boldsymbol{k}_i : the wave vectors for ω_1 , ω_2 , ω_3 , respectively.

Two types of phase matching are introduced:

```
type I: o + o \rightarrow e or e + e \rightarrow o,
type II: o + e \rightarrow e or o + e \rightarrow o,
```

or with shortened notations:

```
ooe: 0+o\rightarrow e or e\rightarrow o+o,
eeo: e+e\rightarrow o or o\rightarrow e+e,
eoe: e+o\rightarrow e or e\rightarrow e+o,
oeo: o+e\rightarrow o or o\rightarrow e+o.
```

In the shortened notation (ooe, eoe, ...) applies: $\omega_1 < \omega_2 < \omega_3$, i.e. the first symbol refers to the longest-wavelength radiation, and the latter to the shortest-wavelength radiation. Here, *o-beam*, or ordinary beam, is the beam with polarization normal to the principal plane of the crystal, i.e.

the plane containing the wave vector \mathbf{k} and crystallophysical axis Z (or optical axis, for uniaxial crystals). The e-beam, or extraordinary beam, is the beam with polarization in the principal plane.

The methods of angular and temperature phase-matching tuning are used in frequency converters. Angular tuning is rather simple and more rapid than temperature tuning. Temperature tuning is generally used in the case of 90 $^{\circ}$ phase matching, i.e., when the birefringence angle is zero. This method is mainly used in crystals with a strong temperature dependence of phase matching: LiNbO₃, LBO, KNbO₃, and Ba₂NaNb₅O₁₅.

4.1.2.2 Uniaxial crystals

For uniaxial crystals the difference between the refractive indices of the ordinary and extraordinary beams, birefringence Δn , is zero along the optical axis (crystallophysical axis Z) and maximum in the normal direction. The refractive index of the ordinary beam does not depend on the direction of propagation, however, the refractive index of the extraordinary beam $n_e(\theta)$ is a function of the polar angle θ between the Z axis and the vector k (but not of the azimuthal angle φ) (Fig. 4.1.1):

$$n_{\rm e}(\theta) = n_{\rm o} \left[\frac{1 + \tan^2 \theta}{1 + \left(\frac{n_{\rm o}}{n_{\rm e}}\right)^2 \tan^2 \theta} \right]^{\frac{1}{2}},$$
 (4.1.7)

where $n_{\rm o}$ and $n_{\rm e}$ are the refractive indices of the ordinary and extraordinary beams in the plane normal to the Z axis and termed as corresponding principal values. Note that if $n_{\rm o} > n_{\rm e}$, the crystal is negative, and if $n_{\rm o} < n_{\rm e}$, it is positive. For an o-beam the indicatrix of the refractive indices is a sphere with radius $n_{\rm o}$, and an ellipsoid of rotation with semiaxes $n_{\rm o}$ and $n_{\rm e}$ for an e-beam (Fig. 4.1.2). In the crystal the beam, in general, is divided into two beams with orthogonal polarizations; the angle between these beams ρ is the birefringence (or walk-off) angle.

Equations for calculating phase-matching angles in uniaxial crystals are given in Table 4.1.1 [86Nik, 99Dmi].

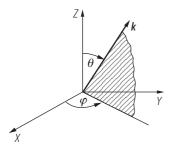


Fig. 4.1.1. Polar coordinate system for description of refraction properties of uniaxial crystals (k is the light propagation direction, Z is the optic axis, θ and φ are the coordinate angles).

4.1.2.3 Biaxial crystals

For biaxial crystals the optical indicatrix has a bilayer surface with four points of interlayer contact which correspond to the directions of two optical axis. In the simple case of light propagation in 146 4.1.2 Fundamentals [Ref. p. 187]

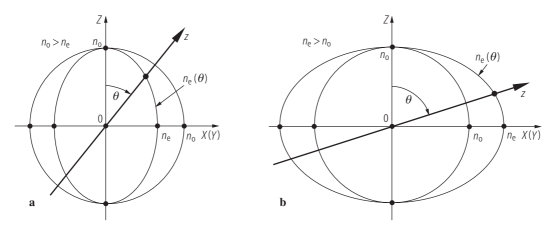


Fig. 4.1.2. Dependence of refractive index on light propagation direction and polarization (index surface) in uniaxial crystals: (a) negative: $n_o > n_e$ and (b) positive: $n_e > n_o$.

Table 4.1.1. Equations for calculating phase-matching angles in uniaxial crystals [86Nik, 99Dmi].

Negative uniaxial crystals	Positive uniaxial crystals
$\tan^2 \theta_{\rm pm}^{\rm ooe} = (1 - U)/(W - 1)$ $\tan^2 \theta_{\rm pm}^{\rm eoe} \approx (1 - U)/(W - R)$ $\tan^2 \theta_{\rm pm}^{\rm oee} \approx (1 - U)/(W - Q)$	$\tan^2 \theta_{\rm pm}^{\rm eeo} \approx (1 - U)/(U - S)$ $\tan^2 \theta_{\rm pm}^{\rm eeo} = (1 - V)/(V - Y)$ $\tan^2 \theta_{\rm pm}^{\rm eeo} = (1 - T)/(T - Z)$

Notations:

$$U = (A+B)^2/C^2 \; ; \; W = (A+B)^2/F^2 \; ; \; R = (A+B)^2/(D+B)^2 \; ;$$

$$Q = (A+B)^2/(A+E)^2 \; ; \; S = (A+B)^2/(D+E)^2 \; ; \; V = B^2/(C-A)^2 \; ;$$

$$Y = B^2/E^2 \; ; \; T = A^2/(C-B)^2 \; ; \; Z = A^2/D^2 \; ;$$

$$A = n_{\rm ol}/\lambda_1 \; ; \; B = n_{\rm o2}/\lambda_2 \; ; \; C = n_{\rm o3}/\lambda_3 \; ;$$

$$D = n_{\rm el}/\lambda_1 \; ; \; E = n_{\rm el}/\lambda_2 \; ; \; F = n_{\rm el}/\lambda_3 \; .$$

These expressions can be generalized to noncollinear phase matching. In this case, for example, the phase-matching angle θ_{pm}^{ooe} is determined from the above presented equation using the new coefficients U and W:

$$U = (A^2 + B^2 + 2AB\cos\gamma)/C^2$$
, $W = (A^2 + B^2 + 2AB\cos\gamma)/F^2$,

where γ is the angle between the wave vectors \mathbf{k}_1 and \mathbf{k}_2 .

the principal planes XY, YZ, and XZ the dependencies of refractive indices on the direction of light propagation represent a combination of an ellipse and a circle (Fig. 4.1.3). Thus in the principal planes a biaxial crystal can be considered as a uniaxial crystal, e.g. a biaxial crystal with $n_Z > n_Y > n_X$ in the XY plane is similar to a negative uniaxial crystal with $n_{\rm o} = n_Z$ and

$$n_{\rm e}(\varphi) = n_Y \left(\frac{1 + \tan^2 \varphi}{1 + (n_Y/n_X)^2 \tan^2 \varphi} \right)^{\frac{1}{2}}$$
 (4.1.8)

The angle V_Z between the optical axis and Z axis for the case $n_Z > n_Y > n_X$ can be found from:

$$\sin V_Z = \frac{n_Z}{n_Y} \left(\frac{n_Y^2 - n_X^2}{n_Z^2 - n_X^2} \right)^{\frac{1}{2}} \tag{4.1.9}$$

and for the case $n_X > n_Y > n_Z$:

$$\cos V_Z = \frac{n_X}{n_Y} \left(\frac{n_Y^2 - n_Z^2}{n_X^2 - n_Z^2} \right)^{\frac{1}{2}} . \tag{4.1.10}$$

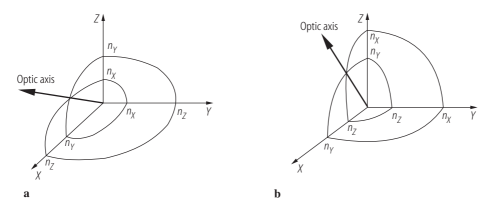


Fig. 4.1.3. Dependence of refractive index on light propagation direction and polarization (index surface) in biaxial crystals: (a) $n_X < n_Y < n_Z$, (b) $n_X > n_Y > n_Z$.

For a positive biaxial crystal the bisectrix of the acute angle between optical axes coincides with $n_{\rm max}$ and for a negative one the bisectrix coincides with $n_{\rm min}$.

Equations for calculating phase-matching angles upon propagation in principal planes of biaxial crystals are given in Table 4.1.2 [87Nik, 99Dmi].

4.1.2.4 Effective nonlinearity

Miller delta formulation [64Mil]:

$$\varepsilon_0 E_i(\omega_3) = \delta_{ijk} P_i(\omega_1) P_k(\omega_2) , \qquad (4.1.11)$$

where the Miller coefficient,

$$\delta_{ijk} = \frac{1}{2\varepsilon_0} \frac{\chi_{ijk}^{(2)}(\omega_3)}{\chi_{ii}^{(1)}(\omega_1) \chi_{jj}^{(1)}(\omega_2) \chi_{kk}^{(1)}(\omega_3)}, \qquad (4.1.12)$$

has small dispersion and is almost constant for a wide range of crystals. For anisotropic media the coefficients $\chi^{(1)}$ and $\chi^{(2)}$ are, in general, the second- and third-rank tensors, respectively. In practice, the tensor

$$d_{ijk} = \frac{1}{2}\chi_{ijk} \tag{4.1.13}$$

is used instead of χ_{ijk} . Usually, the "plane" representation of d_{ijk} in the form d_{il} is used, the relation between l and jk is:

\overline{jk}		l	
11	\leftrightarrow	1	
22	\leftrightarrow	2	
33	\leftrightarrow	3	
23 or 32	\leftrightarrow	4	
31 or 13	\leftrightarrow	5	
12 or 21	\leftrightarrow	6	

Table 4.1.2. Equations for calculating phase-matching angles in biaxial crystals upon light propagation in the principal planes [87Nik, 99Dmi]. (a) $n_X < n_Y < n_Z$

Principal plane	Type of interaction	Equations	Notations
	ooe	$\tan^2 \varphi = \frac{1 - U}{W - 1}$	$U = \left(\frac{A+B}{C}\right)^2; W = \left(\frac{A+B}{F}\right)^2; A = \frac{n_{Z1}}{\lambda_1}; B = \frac{n_{Z2}}{\lambda_2}; C = \frac{n_{Y3}}{\lambda_3}; F = \frac{n_{X3}}{\lambda_3}$
XY	eoe	$\tan^2 \varphi \approx \frac{1 - U}{W - R}$	$U = \left(\frac{A+B}{C}\right)^2; W = \left(\frac{A+B}{F}\right)^2; R = \left(\frac{A+B}{D+B}\right)^2; A = \frac{n_{Y1}}{\lambda_1}; B = \frac{n_{Z2}}{\lambda_2}; C = \frac{n_{Y3}}{\lambda_3}; D = \frac{n_{X1}}{\lambda_1}; F = \frac{n_{X3}}{\lambda_3}$
	oee	$\tan^2 \varphi \approx \frac{1 - U}{W - Q}$	$U = \left(\frac{A+B}{C}\right)^2; W = \left(\frac{A+B}{F}\right)^2; Q = \left(\frac{A+B}{A+E}\right)^2; A = \frac{n_{Z1}}{\lambda_1}; B = \frac{n_{Y2}}{\lambda_2}; C = \frac{n_{Y3}}{\lambda_3}; E = \frac{n_{X2}}{\lambda_2}; F = \frac{n_{X3}}{\lambda_3}$
	eeo	$\tan^2\theta \approx \frac{1-U}{U-S}$	$U = \left(\frac{A+B}{C}\right)^2; \ S = \left(\frac{A+B}{D+E}\right)^2; \ A = \frac{n_{Y1}}{\lambda_1}; \ B = \frac{n_{Y2}}{\lambda_2}; \ C = \frac{n_{X3}}{\lambda_3}; \ D = \frac{n_{Z1}}{\lambda_1}; \ E = \frac{n_{Z2}}{\lambda_2}$
YZ	oeo	$\tan^2 \theta = \frac{1 - V}{V - Y}$	$V = \left(\frac{B}{C - A}\right)^2; Y = \left(\frac{B}{E}\right)^2; A = \frac{n_{X1}}{\lambda_1}; B = \frac{n_{Y2}}{\lambda_2}; C = \frac{n_{X3}}{\lambda_3}; E = \frac{n_{Z2}}{\lambda_2}$
	eoo	$\tan^2 \theta = \frac{1 - T}{T - Z}$	$T = \left(\frac{A}{C - B}\right)^2; \ Z = \left(\frac{A}{D}\right)^2; \ A = \frac{n_{Y1}}{\lambda_1}; \ B = \frac{n_{X2}}{\lambda_2}; \ C = \frac{n_{X3}}{\lambda_3}; \ D = \frac{n_{Z1}}{\lambda_1}$
XZ	ooe	$\tan^2 \theta = \frac{1 - U}{W - 1}$	$U = \left(\frac{A+B}{C}\right)^2; W = \left(\frac{A+B}{F}\right)^2; A = \frac{n_{Y1}}{\lambda_1}; B = \frac{n_{Y2}}{\lambda_2}; C = \frac{n_{X3}}{\lambda_3}; F = \frac{n_{Z3}}{\lambda_3}$
$\theta < V_Z$	eoe	$\tan^2 \theta \approx \frac{1 - U}{W - R}$	$U = \left(\frac{A+B}{C}\right)^2; W = \left(\frac{A+B}{F}\right)^2; R = \left(\frac{A+B}{D+B}\right)^2; A = \frac{n_{X1}}{\lambda_1}; B = \frac{n_{Y2}}{\lambda_2}; C = \frac{n_{X3}}{\lambda_3}; D = \frac{n_{Z1}}{\lambda_1}; F = \frac{n_{Z3}}{\lambda_3}$
	oee	$\tan^2 \theta \approx \frac{1 - U}{W - Q}$	$U = \left(\frac{A+B}{C}\right)^2; W = \left(\frac{A+B}{F}\right)^2; Q = \left(\frac{A+B}{A+E}\right)^2; A = \frac{n_{Y1}}{\lambda_1}; B = \frac{n_{X2}}{\lambda_2}; C = \frac{n_{X3}}{\lambda_3}; E = \frac{n_{Z2}}{\lambda_2}; F = \frac{n_{Z3}}{\lambda_3}$
XZ	eeo	$\tan^2 \theta \approx \frac{1 - U}{U - S}$	$U = \left(\frac{A+B}{C}\right)^2; \ S = \left(\frac{A+B}{D+E}\right)^2; \ A = \frac{n_{X1}}{\lambda_1}; \ B = \frac{n_{X2}}{\lambda_2}; \ C = \frac{n_{Y3}}{\lambda_3}; \ D = \frac{n_{Z1}}{\lambda_3}; \ E = \frac{n_{Z2}}{\lambda_2}$
$\theta > V_Z$	oeo	$\tan^2 \theta = \frac{1 - V}{V - Y}$	$V = \left(\frac{B}{C - A}\right)^{2}; Y = \left(\frac{B}{E}\right)^{2}; A = \frac{n_{Y1}}{\lambda_{1}}; B = \frac{n_{X2}}{\lambda_{2}}; C = \frac{n_{Y3}}{\lambda_{3}}; E = \frac{n_{Z2}}{\lambda_{2}}$
	600	$\tan^2 \theta = \frac{1 - T}{T - Z}$	$T = \left(\frac{A}{C - B}\right)^2; \ Z = \left(\frac{A}{D}\right)^2; \ A = \frac{n_{X1}}{\lambda_1}; \ B = \frac{n_{Y2}}{\lambda_2}; \ C = \frac{n_{Y3}}{\lambda_3}; \ D = \frac{n_{Z1}}{\lambda_1}$

Ref. p. 187]

(b) $n_X > n_Y > n_Z$

Principal plane	Type of interaction	Equations	Notations
	eeo	$\tan^2 \varphi \approx \frac{1 - U}{U - S}$	$U = \left(\frac{A+B}{C}\right)^2; \ S = \left(\frac{A+B}{D+E}\right)^2; \ A = \frac{n_{Y1}}{\lambda_1}; \ B = \frac{n_{Y2}}{\lambda_2}; \ C = \frac{n_{Z3}}{\lambda_3}; \ D = \frac{n_{X1}}{\lambda_1}; \ E = \frac{n_{X2}}{\lambda_2}$
XY	oeo	$\tan^2 \varphi = \frac{1 - V}{V - Y}$	$V = \left(\frac{B}{C - A}\right)^2; \ Y = \left(\frac{B}{E}\right)^2; \ A = \frac{n_{Z1}}{\lambda_1}; \ B = \frac{n_{Y2}}{\lambda_2}; \ C = \frac{n_{Z3}}{\lambda_3}; \ E = \frac{n_{X2}}{\lambda_2}$
	eoo	$\tan^2 \varphi = \frac{1 - T}{T - Z}$	$T = \left(\frac{A}{C - B}\right)^2; \ Z = \left(\frac{A}{D}\right)^2; \ A = \frac{n_{Y1}}{\lambda_1}; \ B = \frac{n_{Z2}}{\lambda_2}; \ C = \frac{n_{Z3}}{\lambda_3}; \ D = \frac{n_{X1}}{\lambda_1}$
	ooe	$\tan^2\theta = \frac{1-U}{W-1}$	$U = \left(\frac{A+B}{C}\right)^2; W = \left(\frac{A+B}{F}\right)^2; A = \frac{n_{X1}}{\lambda_1}; B = \frac{n_{X2}}{\lambda_2}; C = \frac{n_{Y3}}{\lambda_3}; F = \frac{n_{Z3}}{\lambda_3}$
YZ	eoe	$\tan^2 \theta \approx \frac{1 - U}{W - R}$	$U = \left(\frac{A+B}{C}\right)^2; W = \left(\frac{A+B}{F}\right)^2; R = \left(\frac{A+B}{D+B}\right)^2; A = \frac{n_{Y1}}{\lambda_1}; B = \frac{n_{X2}}{\lambda_2}; C = \frac{n_{Y3}}{\lambda_3}; D = \frac{n_{Z1}}{\lambda_1}; F = \frac{n_{Z3}}{\lambda_3}$
	oee	$\tan^2 \theta \approx \frac{1 - U}{W - Q}$	$U = \left(\frac{A+B}{C}\right)^2; W = \left(\frac{A+B}{F}\right)^2; Q = \left(\frac{A+B}{A+E}\right)^2; A = \frac{n_{X1}}{\lambda_1}; B = \frac{n_{Y2}}{\lambda_2}; C = \frac{n_{Y3}}{\lambda_3}; E = \frac{n_{Z2}}{\lambda_2}; F = \frac{n_{Z3}}{\lambda_3}$
XZ	eeo	$\tan^2\theta \approx \frac{1-U}{U-S}$	$U = \left(\frac{A+B}{C}\right)^2; \ S = \left(\frac{A+B}{D+E}\right)^2; \ A = \frac{n_{X1}}{\lambda_1}; \ B = \frac{n_{X2}}{\lambda_2}; \ C = \frac{n_{Y3}}{\lambda_3}; \ D = \frac{n_{Z1}}{\lambda_1}; \ E = \frac{n_{Z2}}{\lambda_2}$
$\theta < V_Z$	oeo	$\tan^2 \theta = \frac{1 - V}{V - Y}$	$V = \left(\frac{B}{C - A}\right)^2; Y = \left(\frac{B}{E}\right)^2; A = \frac{n_{Y1}}{\lambda_1}; B = \frac{n_{X2}}{\lambda_2}; C = \frac{n_{Y3}}{\lambda_3}; E = \frac{n_{Z2}}{\lambda_2}$
	eoo	$\tan^2 \theta = \frac{1 - T}{T - Z}$	$T = \left(\frac{A}{C - B}\right)^2; \ Z = \left(\frac{A}{D}\right)^2; \ A = \frac{n_{X1}}{\lambda_1}; \ B = \frac{n_{Y2}}{\lambda_2}; \ C = \frac{n_{Y3}}{\lambda_3}; \ D = \frac{n_{Z1}}{\lambda_1}$
\overline{XZ}	ooe	$\tan^2 \theta = \frac{1 - U}{W - 1}$	$U = \left(\frac{A+B}{C}\right)^2; W = \left(\frac{A+B}{F}\right)^2; A = \frac{n_{Y1}}{\lambda_1}; B = \frac{n_{Y2}}{\lambda_2}; C = \frac{n_{X3}}{\lambda_3}; F = \frac{n_{Z3}}{\lambda_3}$
$\theta > V_Z$	eoe	$\tan^2 \theta \approx \frac{1 - U}{W - R}$	$U = \left(\frac{A+B}{C}\right)^2; W = \left(\frac{A+B}{F}\right)^2; R = \left(\frac{A+B}{D+B}\right)^2; A = \frac{n_{X1}}{\lambda_1}; B = \frac{n_{Y2}}{\lambda_2}; C = \frac{n_{X3}}{\lambda_3}; D = \frac{n_{Z1}}{\lambda_1}; F = \frac{n_{Z3}}{\lambda_3}$
	oee	$\tan^2 \theta \approx \frac{1 - U}{W - Q}$	$U = \left(\frac{A+B}{C}\right)^2; W = \left(\frac{A+B}{F}\right)^2; Q = \left(\frac{A+B}{A+E}\right)^2; A = \frac{n_{Y1}}{\lambda_1}; B = \frac{n_{X2}}{\lambda_2}; C = \frac{n_{X3}}{\lambda_3}; E = \frac{n_{Z2}}{\lambda_2}; F = \frac{n_{Z3}}{\lambda_3}$

Kleinman symmetry conditions [62Kle]: $d_{21} = d_{16}$, $d_{24} = d_{32}$, $d_{31} = d_{15}$, $d_{13} = d_{35}$, $d_{14} = d_{36}$, $d_{25} = d_{12} = d_{26}$, $d_{32} = d_{24}$ are valid in the case of non-dispersion of electron nonlinear polarizability. The equations for calculating the conversion efficiency include the effective nonlinear coefficients $d_{\rm eff}$, which comprise all summation operations along the polarization directions of the interacting waves and thus reduce the calculation to one dimension. Effective nonlinearities $d_{\rm eff}$ for different crystal point groups under valid Kleinman symmetry conditions are presented in Table 4.1.3.

The conversion factors for SI and CGS-esu systems are given in Table 4.1.4.

Table 4.1.3. Expressions for d_{eff} in nonlinear crystals when *Kleinman symmetry* relations are valid. (a) Uniaxial crystals

Point group	Type of interaction	
	00e, 0e0, e00	eeo, eoe, oee
$\overline{4,4mm}$	$d_{15} \sin \theta$	0
6, 6mm	$d_{15}\sin heta$	0
$\bar{6}m2$	$d_{22} \cos \theta \sin (3\varphi)$	$d_{22}\cos^2\theta\cos\varphi$
3m	$d_{15} \sin \theta - d_{22} \cos \theta \sin (3\varphi)$	$d_{22}\cos^2\theta\cos(3\varphi)$
$\bar{6}$	$(d_{11}\cos(3\varphi)-d_{22}\sin(3\varphi))\cos\theta$	$(d_{11}\sin(3\varphi) + d_{22}\cos(3\varphi))\cos^2\theta$
3	$(d_{11}\cos(3\varphi)-d_{22}\sin(3\varphi))\cos\theta+d_{15}\sin\theta$	$(d_{11}\sin(3\varphi) + d_{22}\cos(3\varphi))\cos^2\theta$
32	$d_{11} \cos \theta \cos (3\varphi)$	$d_{11} \cos^2 \theta \sin (3\varphi)$
$\bar{4}$	$(d_{14} \sin(2\varphi) + d_{15} \cos(2\varphi)) \sin\theta$	$(d_{14}\cos(2\varphi)-d_{15}\sin(2\varphi))\sin(2\theta)$
$\bar{4}2m$	$d_{36} \sin \theta \sin (2\varphi)$	$d_{36} \sin(2\theta) \cos(2\varphi)$

(b) Biaxial crystals (assignments of crystallophysical and crystallographic axes: for mm2 and 222 point groups: $X, Y, Z \to a, b, c$; for 2 and m point groups: $Y \to b$)

Point group	Principal plane	Type of interaction ooe, oeo, eoo	eeo, eoe, oee
2	XY	$d_{23}\cos\varphi$	$d_{36} \sin (2\varphi)$
	$egin{array}{c} YZ \ XZ \end{array}$	$d_{21} \cos \theta$ 0	$d_{36} \sin (2\theta) d_{21} \cos^2 \theta + d_{23} \sin^2 \theta - d_{36} \sin (2\theta)$
m	XY	$d_{13}\sinarphi$	$d_{31}\sin^2\varphi + d_{32}\cos^2\varphi$
	YZ	$d_{31} \sin \theta$	$d_{13} \sin^2 \theta + d_{12} \cos^2 \theta$
	XZ	$d_{12}\cos\theta - d_{32}\sin\theta$	0
mm2	XY	0	$d_{31}\sin^2\varphi + d_{32}\cos^2\varphi$
	YZ	$d_{31} \sin \theta$	0
	XZ	$d_{32} \sin \theta$	0
222	XY	0	$d_{36} \sin{(2\varphi)}$
	YZ	0	$d_{36} \sin{(2\theta)}$
	XZ	0	$d_{36} \sin (2\theta)$

Table 4.1.4. Units and conversion factors.

Nonlinear coefficient	MKS or SI units		CGS or electrostatic units
$\chi_{ij}^{(1)}$	1 (SI, dimensionless)	=	$\frac{1}{4\pi}$ (esu, dimensionless)
d_{ij} or $\chi^{(2)}_{ijk}$	$1 \text{ V}^{-1}\text{m}$	=	$\frac{3 \times 10^4}{4 \pi} (\mathrm{erg}^{-1} \mathrm{cm}^3)^{\frac{1}{2}}$
δ_{ij}	$1~\mathrm{C^{-1}m^2}$	=	$\frac{4\pi}{3\times 10^5} \ (\mathrm{erg^{-1}cm^3})^{\frac{1}{2}}$

Note that in SI units $\mathbf{P}^{(n)} = \overline{\varepsilon_0 \chi^{(n)} \mathbf{E}^n}$ (with $\mathbf{P}^{(n)}$ expressed in C m⁻²), whereas in CGS or esu units $P^{(n)} = \chi^{(n)} E^n$ (with $P^{(n)}$ expressed in esu).

4.1.2.5 Frequency conversion efficiency

4.1.2.5.1 General approach

The conversion efficiency of a three-wave interaction process for the case of square nonlinearity

$$\boldsymbol{P}_{\mathrm{nl}} = \varepsilon_0 \, \chi^{(2)} \boldsymbol{E}^2 \tag{4.1.14}$$

can be determined from the wave equation derived from Maxwell's equations [64Akh, 65Blo, 73Zer, 99Dmi], see also (1.1.4)–(1.1.7),

$$\nabla \times \nabla \times \mathbf{E} + \frac{(1 + \chi^{(1)})}{c^2} \frac{\partial^2 \mathbf{E}}{\partial t^2} = -\frac{1}{\varepsilon_0 c^2} \frac{\partial^2 \mathbf{P}_{\text{nl}}}{\partial t^2}$$
(4.1.15)

with the initial and boundary conditions for the electric field \boldsymbol{E} .

An exact calculation of the nonlinear conversion efficiency for SHG, SFG, and DFG generally requires a numerical calculation. In some simple cases analytical expressions are available. In order to choose the proper method, the contribution of different effects in the nonlinear mixing process should be determined. For this purpose the following approach is introduced [99Dmi]:

- Consider the *effective lengths* of the interaction process:
 - 1. Aperture length L_a :

$$L_{\rm a} = d_0 \,\rho^{-1} \,, \tag{4.1.16}$$

where d_0 is the beam diameter.

2. Quasistatic interaction length L_{qs} :

$$L_{\rm qs} = \tau \nu^{-1} \,, \tag{4.1.17}$$

where τ is the radiation pulse width and ν is the mismatch of reverse group velocities. For SHG

$$\nu = u_{\omega}^{-1} - u_{2\omega}^{-1} \,, \tag{4.1.18}$$

where u_{ω} and $u_{2\omega}$ are the group velocities of the corresponding waves ω and 2ω .

3. Diffraction length L_{dif} :

$$L_{\text{dif}} = k \, d_0^2 \,. \tag{4.1.19}$$

4. Dispersion-spreading length $L_{\rm ds}$:

$$L_{\rm ds} = \tau^2 g^{-1} \,, \tag{4.1.20}$$

152 4.1.2 Fundamentals [Ref. p. 187]

where g is the dispersion-spreading coefficient

$$g = \frac{1}{2} \left(\frac{\partial^2 k}{\partial \omega^2} \right) . \tag{4.1.21}$$

5. Nonlinear interaction length $L_{\rm nl}$:

$$L_{\rm nl} = (\sigma a_0)^{-1}$$
 (4.1.22)

Here σ is the nonlinear coupling coefficient:

$$\sigma_{1,2} = 4\pi k_{1,2} n_{1,2}^{-2} d_{\text{eff}} ,$$
(4.1.23)

$$\sigma_3 = 2\pi k_3 n_3^{-2} d_{\text{eff}} , \qquad (4.1.24)$$

and

$$a_0 = \left(a_1^2(0) + a_2^2(0) + a_3^2(0)\right)^{\frac{1}{2}}, \tag{4.1.25}$$

where $a_n(0)$ are the wave amplitudes of interacting waves λ_1 , λ_2 , and λ_3 at the input surface of the crystal.

- The length of the crystal L should be compared with L_{eff} from above equations. If $L < L_{\text{eff}}$ the respective effect can be neglected.

4.1.2.5.2 Plane-wave fixed-field approximation

When the conditions $L < L_{\rm nl}$ and $L < L_{\rm eff}$ are fulfilled, the so-called fixed-field approximation is realized. For SHG, $\omega + \omega = 2\omega$ and $\Delta k = 2k_{\omega} - k_{2\omega}$, the conversion efficiency η is determined by the equation:

$$\eta = P_{2\omega}/P_{\omega} = \frac{2\pi^2 d_{\text{eff}}^2 L^2 P_{\omega}}{\varepsilon_0 c n_{\omega}^2 n_{2\omega} \lambda_2^2 A} \operatorname{sinc}^2 \left(\frac{|\Delta k| L}{2}\right). \tag{4.1.26}$$

For SFG, $\omega_1 + \omega_2 = \omega_3$ and $\Delta k = k_1 + k_2 - k_3$, the conversion efficiency η is:

$$\eta = P_3/P_1 = \frac{8\pi^2 d_{\text{eff}}^2 L^2 P_2}{\varepsilon_0 c n_1 n_2 n_3 \lambda_3^2 A} \operatorname{sinc}^2 \left(\frac{|\Delta k| L}{2}\right) . \tag{4.1.27}$$

For DFG, $\omega_1=\omega_3-\omega_2$ and $\Delta k=k_1+k_2-k_3$, the conversion efficiency η is:

$$\eta = P_1/P_3 = \frac{8\pi^2 d_{\text{eff}}^2 L^2 P_2}{\varepsilon_0 c n_1 n_2 n_3 \lambda_1^2 A} \operatorname{sinc}^2 \left(\frac{|\Delta k| L}{2}\right) . \tag{4.1.28}$$

Note that all the above equations are for the SI system, i.e. $[d_{\rm eff}]={\rm m/V}$; $[P]={\rm W}$; $[L]={\rm m}$; $[\lambda]={\rm m}$; $[A]={\rm m}^2$; $\varepsilon_0=8.854\times 10^{-12}~{\rm A\,s/\,(V\,m)}$; $c=3\times 10^8~{\rm m/s}$.

When the powers of the mixing waves are almost equal, the conversion efficiency is for THG, $\omega + 2\omega = 3\omega$:

$$\eta = \frac{P_{3\omega}}{(P_{2\omega}P_{\omega})^{\frac{1}{2}}}; \tag{4.1.29}$$

for FOHG in the case of $\omega + 3\omega = 4\omega$:

$$\eta = \frac{P_{4\omega}}{(P_{3\omega}P_{\omega})^{\frac{1}{2}}} \,, \tag{4.1.30}$$

or for $2\omega + 2\omega = 4\omega$:

$$\eta = \frac{P_{4\omega}}{P_{2\omega}} \; ; \tag{4.1.31}$$

for SFG, $\omega_1 + \omega_2 = \omega_3$:

$$\eta = \frac{P_3}{(P_1 P_2)^{\frac{1}{2}}}; \tag{4.1.32}$$

for DFG, $\omega_1 = \omega_3 - \omega_2$:

$$\eta = \frac{P_1}{(P_2 P_3)^{\frac{1}{2}}} \,. \tag{4.1.33}$$

In some cases (mentioned additionally) the conversion efficiency is calculated from the power (energy) of fundamental radiation, e.g. for fifth harmonic generation, $\omega + 4\omega = 5\omega$:

$$\eta = \frac{P_{5\omega}}{P_{\omega}} \ . \tag{4.1.34}$$

Corresponding equations are valid for energy conversion efficiencies by substituting the pulse energy instead of power in the above equations.

The efficiency η in the case of OPO is calculated by the equation

$$\eta = \frac{E_{\text{OPO}}}{E_0} \,,$$
(4.1.35)

where E_{OPO} is the total OPO radiation energy (signal + idler) and E_0 is the energy of the pump radiation. Conversion efficiency can also be determined in terms of pump depletion:

$$\eta = 1 - \frac{E_{\text{unc}}}{E_{\text{pump}}} \,, \tag{4.1.36}$$

where $E_{\rm unc}$ is the energy of unconverted pumping beam after the OPO crystals. Pump depletions are usually significantly greater than the ordinary η values.

The quantum conversion efficiency (for the ratio of converted and mixing quanta) in the case of exact phase-matching ($\Delta k = 0$) for sum-frequency generation, $\omega_1 + \omega_2 = \omega_3$, is determined by the following equation (SI system):

$$\eta = \frac{P_3 \lambda_3}{P_1 \lambda_1} = \sin^2 \left(2 \pi d_{\text{eff}} L \sqrt{\frac{2P_2}{\varepsilon_0 c n_1 n_2 n_3 \lambda_1 \lambda_3 A}} \right) , \qquad (4.1.37)$$

and for difference-frequency generation, $\omega_1 = \omega_3 - \omega_2$:

$$\eta = \frac{P_1 \lambda_1}{P_3 \lambda_3} = \sin^2 \left(2 \pi d_{\text{eff}} L \sqrt{\frac{2P_2}{\varepsilon_0 c n_1 n_2 n_3 \lambda_1 \lambda_3 A}} \right) . \tag{4.1.38}$$

In the presence of linear absorption all the above equations for conversion efficiencies should be multiplied by the factor

$$\exp\left(-\alpha L\right) \approx 1 - \alpha L$$
, (4.1.39)

where α is the linear absorption coefficient of the crystal.

154 4.1.3 Selection of data [Ref. p. 187]

4.1.2.5.3 SHG in "nonlinear regime" (fundamental wave depletion)

Analytical equation for SHG power conversion efficiency for the case of fundamental power depletion in the plane-wave approximation and for exact phase matching ($\Delta k = 0$) is given below [99Dmi]:

$$\eta = \frac{P_{2\,\omega}}{P_{\omega}} = \tanh^2\left(\frac{L}{L_{\rm nl}}\right) \,. \tag{4.1.40}$$

In order to calculate

$$L_{\rm nl} = (\sigma \, a_0)^{-1} \tag{4.1.41}$$

one should determine a_0 [V cm⁻¹]:

$$a_0 = \left[\frac{752 \, P_\omega}{\pi \, \zeta^2 \, n} \right]^{\frac{1}{2}} \tag{4.1.42}$$

from input radiation power P_{ω} [W] and the characteristic radius of the beam ζ [cm], and the parameter σ [V⁻¹]

$$\sigma = \frac{8\pi^2 d_{\text{eff}}}{n \lambda_{c}} ; \qquad (4.1.43)$$

where λ_1 is in m, d_{eff} in mV⁻¹.

4.1.3 Selection of data

Literature up to the end of 1998 is compiled in this chapter. Attempts were made to select the most reliable and recent data.

Tables in Sect. 4.1.4–4.1.8 present data on second, third, fourth, fifth, and sixth harmonic generation of Nd:YAG laser (including intracavity and in external resonant cavities), harmonic generation of iodine, ruby, Ti:sapphire, semiconductor, dye, argon, He–Ne, NH₃, CO, and CO₂ lasers, sum-frequency mixing (including up-conversion of IR radiation into the visible), difference-frequency generation, optical parametric oscillation (cw, nanosecond, picosecond, and femtosecond in the UV, visible, near and mid IR regions) and picosecond continuum generation.

Second harmonic generation of Nd:YAG laser was realized with conversion efficiency of $\eta=80~\%$ in KDP and KTP, THG with $\eta=80~\%$ in KDP, FOHG with $\eta=80-90~\%$ (calculated from SH) in ADP and KDP, FIHG in KDP, ADP (upon cooling) and BBO and urea (at room temperature). Second harmonic generation of Ti:sapphire laser with $\eta=75~\%$ was achieved in LBO, minimum pulse durations for SH were as short as 10–16 fs (BBO, LBO). Third and fourth harmonics of Ti:sapphire laser were generated in BBO, thus covering the range of wavelengths 193–285 nm. Second harmonic of CO₂ laser with $\eta=50~\%$ was obtained in ZnGeP₂.

Sum-frequency generation (mixing) is used, in particular, for extending the range of generating radiation into the ultraviolet. By use of SFG the shortest wavelengths in VUV were achieved with KB5 crystal (166 nm), LBO (172.7 nm), CBO, CLBO (185 nm), BBO, KDP and ADP (189, 190, and 208 nm, respectively). At present, $\lambda=166$ nm is the minimum wavelength achieved by frequency conversion in crystals. Sum-frequency generation is also used for up-conversion of near IR (1–5 µm) and CO₂ laser radiation into the visible. Maximum conversion efficiencies up to 40–60 % were obtained for the latter case in AgGaS₂, CdSe, and HgGa₂S₄ crystals.

Difference-frequency generation makes it possible to produce IR radiation in the near IR (up to 7.7 μ m, in LiIO₃), mid IR (up to 18–23 μ m, in AgGaSe₂, GaSe, CdSe, Ag₃AsS₃) and far IR (0.05–30 mm, in LiNbO₃ and GaP).

Optical parametric oscillation is a powerful method for generating continuously tunable radiation in the UV (up to 314–330 nm, in LBO and urea), visible, and IR regions (up to 16–18 μ m, in CdSe and GaSe). Singly resonant OPO, or SROPO, uses resonant feedback at only the signal or idler frequency. Doubly resonant OPO, or DROPO, uses resonant feedback of both signal and idler frequencies. Exotic triply resonant OPO, or TROPO, with resonant feedback also at pump frequency, and quadruply resonant OPO, or QROPO, with SHG inside the OPO cavity and resonant feedback also at the second harmonic, are used very seldom.

Different OPO schemes and their energetic, temporal, spectral, and spatial characteristics are considered in detail in [73Zer, 78Dmi, 83Dan, 87Dmi] and in the three special issues of the Journal of the Optical Society of America B (Vol. 10, No. 9 and 11, 1993 and Vol. 12, No. 11, 1995) devoted to optical parametric oscillators. In Tables 4.1.30–4.1.33 we list only the main OPO parameters realized in practice: pump wavelengths, phase-matching angles, pump thresholds (peak intensity and/or average power), tuning ranges, OPO pulse durations, and conversion efficiencies for OPO experiments in the UV, visible, and near IR spectral ranges. The column headed *notes* gives data on the OPO type, pump intensities, crystal lengths, phase-matching temperatures, and output characteristics of OPO radiation (energy, power, bandwidth).

High conversion efficiencies were obtained with resonant schemes of cw OPO ($\eta = 40 - 80$ % with LiNbO₃:MgO crystal), nanosecond ($\eta = 60$ % with BBO), traveling-wave and synchronously pumped picosecond OPO ($\eta = 45 - 75$ % with KDP, KTP, KTA, BBO), and synchronously pumped femtosecond OPO ($\eta = 50$ % with BBO). Minimum pulse durations were 13 fs in SP OPO with BBO crystal, pumped by the second harmonic of a Ti:sapphire laser. Very low power thresholds (0.4 mW) were achieved with LiNbO₃:MgO containing quadruply resonant OPO. In general, in the case of OPO the total conversion efficiencies to both, idler and signal wavelengths, are presented. In most cases the conversion efficiency corresponds to the maximum for the range of wavelengths.

The picosecond continuum, first detected in media with cubic nonlinearity (D_2O , H_2O , etc.), was also observed in crystals with square nonlinearity (KDP, LiIO₃, LiNbO₃, etc.).

We don't pretend to comprehend all directions of frequency conversion in crystals. Some special aspects, e.g. second harmonic generation in layers and films, waveguides and fibers, periodically poled crystals, liquid crystals, as well as different design configurations of frequency converters have been beyond our consideration. For "justification" we refer to Artur L. Schawlow's famous saying: "To do successful research, you don't need to know everything. You just need to know of one thing that isn't known".

4.1.4 Harmonic generation (second, third, fourth, fifth, and sixth)

Table 4.1.5. Second harmonic generation of Nd:YAG laser radiation (1.064 \rightarrow 0.532 $\mu m).$

Crystal	Type of interaction	$\theta_{ m pm} \ [m deg]$	$[\mathrm{W} \ \mathrm{cm}^{-2}]$	$ au_{ m p} \ [m ns]$	L [mm]	Conversion efficiency [%]	Ref.	Notes
KDP	ooe	41	10 ⁹	0.15	25	32 (energy)	[75Att]	
	ooe	41	_	0.05	25	60	[76Att]	
	ooe	41	8×10^{9}	0.03	14	82 (energy)	[78Mat]	
	ooe	41	7×10^{9}	0.03	20	81 (energy)	[78Mat]	
	ooe	41.35	_	$0.1~\mathrm{ms}$	40	0.38 (energy)	[93Dim]	$\lambda = 946 \text{ nm}$
DKDP	eoe	53.5	10^{8}	18	30	50 (power)	[76Mac]	
	eoe	53.5	3×10^{9}	0.25	40	70 (power)	[76Mac]	
	eoe	53.5	8×10^7	20	30	50 (energy)	[78Kog]	$P_{2\omega} = 10 \text{ W}$
	ooe	36.6	3×10^{8}	8	20	40 (energy)	[91Bor1]	
	eoe	53.7	3×10^8	8	20	50 (energy)	[91Bor1]	
CDA	ooe	90	2×10^{8}	10	17.5	57 (power)	[74Kat2]	T = 48 °C
	ooe	90	4×10^9	0.007	13	25 (energy)	[72Rab]	
DCDA	ooe	90	8×10^7	20	21	40 (energy)	[78Kog]	$T = 90 \dots 100 ^{\circ}$ C
	ooe	90	3×10^8	20	16	40 (energy)	[78Kog]	$P_{2\omega} = 10 \text{ W}$
	ooe	90	2×10^{8}	10	13.5	45 (power)	[74Kat2]	$T = 112 ^{\circ}\text{C}$
	ooe	90	9×10^7	_	29	50 (power)	[74Amm]	1 11 2 0
	ooe	90	-	15	20	57	[76Hon]	$P_{2\omega} = 6 \text{ W}$
RDA	ooe	50	_	10	_	34 (power)	[75Kat2]	T = 25 °C
RDP	ooe	50.8	2×10^8	10	15.3	36 (power)	[74Kat5]	
LiIO ₃	ooe	30	7×10^{7}	_	18	44 (power)	[73Dmi]	
- 0	ooe	30	3×10^9	0.04	5	50	[84Van]	
LiNbO ₃	ooe	90	3.7×10^{7}	10	20	40	[81Bye]	$T = 120 {}^{\circ}\text{C}$
	ooe a	90			9 – 30	50		$P_{2\omega}=1~\mathrm{W}$
LFM	ooe	55.1	2×10^8	_	15	36	[75And]	
KTP	eoe	26 ^b	_	10	=	22	[78Har]	
	eoe	$26^{\ b}$	_	0.04	5	18	[83Joh]	
	eoe	$25.2^{\ { m b}}$	_	0.07	7.2	52	[85Ale]	
	eoe	25 $^{\rm b}$	2.5×10^8	15	4	60	[85Bel]	
	eoe	$30 ^{\mathrm{b}}$	2×10^7	35	9	40 (energy)	$[86 \mathrm{Dri}]$	multimode
	eoe	$30 ^{\mathrm{b}}$	9×10^{7}	35	4	45 (energy)	$[86 \mathrm{Dri}]$	multimode
	eoe	$30 ^{\mathrm{b}}$	10^{8}	30	5.1	60 (energy)	$[86 \mathrm{Dri}]$	two-pass
	eoe	$30^{\rm \ b}$	10^{8}	30	8	50 (energy)	[86Dri]	Gaussian
	eoe	$26^{\ b}$	_	0.2	5	55	[87Moo1]	$E_{2\omega} = 0.19 \text{ J}$
	eoe	$23^{\rm \ b}$	2.5×10^8	10	3	30	[86Lav]	
	eoe	$23^{\ b}$	3.2×10^{8}	8.5	4.5	55 (power)	[93Bol]	

(continued)

Table 4.1.5 continued.

Crystal	Type of interaction	$ heta_{ m pm} \ [m deg]$	I_0 [W cm ⁻²]	$ au_{ m p} \ [m ns]$	L [mm]	Conversion efficiency [%]	Ref.	Notes
KTP	eoe	_		8	7	80 (energy)	[92Bro]	$E_{2\omega} = 0.72 \text{ J},$ $T = 55 ^{\circ}\text{C}$
	eoe	_	9×10^7	17	10	97 (energy)	[97Coo]	Multistage system with 3 SHG crystals, $E_{2\omega}=0.2~\mathrm{J}$
KNbO ₃	ooe	19	4.7×10^7	11	4.8	40 (energy)	[92See]	Nd:YLF (1.047 μm)
BBO	ooe	_	1.9×10^{8}	14	6	47	[87Adh]	$P_{2\omega} = 4.5 \; {\rm W}$
	ooe	_	1.67×10^{8}	14	6	38	[87Adh]	$P_{2\omega} = 8.5 \text{ W}$
	ooe	_	2.53×10^{8}	14	6	37	[87Adh]	$P_{2\omega} = 36 \text{ W}$
	ooe	21	2×10^{9}	1	6.8	68 (energy)	[86Che]	
	ooe	21	2.5×10^{8}	8	6.8	58 (energy)	[86Che]	
	ooe	22.8	1.4×10^{8}	_	7	32 (power)	[90Bha2]	
	ooe	22.8	1.6×10^{8}	8	7.5	55-60 (energy)	[91Bor1]	
LBO	ooe	0 в	10 ⁹	0.035	15	65 (energy)	[91Hua]	T = 148.5 $\pm 0.5 ^{\circ}\text{C}$
	ooe	$0^{\rm b}$	5×10^8	10	12.5	60 (energy)	[90Lin2]	T = 149 °C
	ooe	12 ^b	$(5-8) \times 10^{8}$	9	14	70 (energy)	[91Xie]	
	ooe	$12^{\rm \ b}$	1.4×10^8	8	17	55-60 (energy)	[91Bor1]	
CLBO	ooe type II	29.4 41.9	10^{11} 3×10^{8}	0.0015 7	7 12	53 (energy) 55 (energy)	[96Sha] [98Yap]	$E_{2\omega} = 1.55 \text{ J}$

a LiNbO₃ grown from congruent melt. b φ_{pm} .

 ${\bf Table~4.1.6.~Second~harmonic~generation~of~Nd:} YAG~laser~radiation~in~organic~crystals.$

Crystal	Type of interaction	$d_{\rm eff}$ / $d_{36}({ m KDP})$	$ heta_{ m pm} \ [m deg]$	$arphi_{ m pm} \ [m deg]$	η [%]	Ref.	Notes
POM	eeo	13.6	18.1 (1.32 µm)	90	50	[88Jos]	$L=7$ mm, $\tau_{\rm p}=160$ ps
MAP	eoe	38.3	2.2	0	30	[77Oud]	L=1 mm
MAP	oeo	37.7	11	90	40	[77Oud]	L = 1.7 mm
mNA	ooe	37.7	90	55	15	[74Dav]	$L=2.5\mathrm{mm}, \Delta\theta=2.9\mathrm{mrad}$
mNA	ooe	6.8	90	8.5	85	[80Kat3]	NCSHG in the XY plane, L = 3 mm
DAN	_	_	40	0	20	[87Nor]	$L=2~\mathrm{mm}$
MHBA	_	30	_	_	59	[93Zha2]	$L=3~\mathrm{mm}$

Table 4.1.7. Intracavity SHG of Nd:YAG laser radiation (1.064 \rightarrow 0.532 $\mu m).$

Crystal	θ_{pm} [deg]	L [mm]	Mode of Nd:YAG laser operation	$P_{2\omega}$ [W]	η [%]	Ref.
LiIO ₃	29	_	Q-switched	0.3	100	[69Des]
	29	20	cw	4	$40 (0.12^{a})$	[81Dmi]
	29	_	Continuous pump, mode-locked, $\tau = 800 \text{ ps}$	5	40 (0.13 ^a)	[82Gol]
	29	15	$\tau = 180 \mu\text{s}, f = 50 \text{ Hz}$	100 (peak)	$0.06^{\rm \ a}$	[80 Koe]
	34	4	Diode-laser pumped cw Nd:YAG laser, $\lambda = 946$ nm	0.52	_	[97Kel]
LiNbO ₃	90	_	Continuous pump, Q-switched	0.31	100	[72Dmi]
	90	1	$\tau = 60 \text{ ns}, f = 400 \text{ Hz}$	100 (peak)	_	[68Smi]
Banana	90	3	cw	1.1	100	[68Geu]
	90	_	Continuous pump, Q-switched	0.016	100	$[70\mathrm{Che}]$
	90	5	_	$0.3 \dots 0.5$	_	[74Gul]
KTP	26	3.5	Q-switched	5.6	_	[84Liu]
-	_	4.6	Acoustooptic modulation, $f = 4 \dots 25 \text{ kHz}$	28	$54 (0.6^{\text{ a}})$	[87Per]
	_	_	Diode-laser pumped cw Nd:YAG laser	$0.03 \dots 0.1$	6 a	[92Ant]
	_	15	Diode-laser pumped mode-locked Nd:YAG laser, $\tau = 120$ ps, $f = 160$ MHz	3	56 (1.3 ^a)	[92Mar]
	_	15	Diode-laser pumped cw Nd:YAG laser	2.8	47 (0.94 a)	[92Mar]
	90	4.4	Q-switched Nd:YalO ₃ laser, $\lambda = 1.08 \ \mu m$	15	-	[86Gar]
	_	5	Diode-laser pumped Nd:YVO ₄ laser	0.07	$9.1^{\rm a}$	[94Tai]
KNbO ₃	90	5	cw	0.366	90	[77Fuk]
	60	3.7	Diode-laser pumped cw Nd:YAG laser, $\lambda = 946 \text{ nm}$	0.0031	$0.74^{\mathrm{\ a}}$	[89Ris]
	0	6.2	Diode-laser pumped cw Nd:YAG laser	0.002	1 ^a	[89Bia]
	90	1.3	Ti-sapphire laser pumped Nd:YalO ₃ laser, $\lambda = 946$ nm	0.015	_	[95Zar]
BBO	25	4	Diode-laser pumped cw Nd:YAG laser, $\lambda = 946$ nm	0.55	-	[97Kel]
LBO	$\varphi = 11.4$	9	Diode-laser pumped Q-switched Nd:YAG laser, $\tau=60~\mathrm{ns}$	4	10 ^a	[94Han]

 $^{^{}a}$ Conversion efficiency calculated with respect to the energy of pumping flash lamps or diode lasers.

Table 4.1.8. Second harmonic generation of Nd:YAG laser radiation (1.064 \rightarrow 0.532 μm) in external resonant cavities.

Crystal	$ heta_{ m pm} \ [m deg]$	T_{pm} [°C]	L [mm]	Mode of laser operation	$P_{2\omega}$ [W]	η [%]	Ref.
LiNbO ₃ :MgO	90	_	12.5	Diode-laser pumped, cw	0.03	56	[88Koz]
	90	110	12	Diode-laser pumped, cw (monolithic ring frequency doubler)	0.2	65	[91Ger]
	90	107	_	Diode-laser pumped, cw (monolithic ring frequency doubler)	0.005	50	[93Fie]
	90	110	7.5	Diode-laser pumped, cw (monolithic ring frequency doubler)	0.1	82	[94Pas]
LiNbO ₃	90	233.7	10	Injection-locked Nd:YAG laser	1.6	69	[91Jun]
KTP	90	63	10	cw YAlO ₃ :Nd laser ($\lambda = 1.08 \mu m$)	0.6	85	[92Ou]
LBO	$\begin{array}{c} 90 \ (\theta), \\ 0 \ (\varphi) \end{array}$	149.5	6	Injection-locked cw Nd:YAG laser	6.5	36	[91Yan]
	90	167	12	Diode-laser pumped mode-locked Nd:YLF laser ($\lambda=1.047~\mu\mathrm{m},$ $\tau=12~\mathrm{ps},f=225~\mathrm{MHz})$	0.75	54	[92Mal]

Table 4.1.9. Third harmonic generation of Nd:YAG laser radiation (1.064 \rightarrow 0.355 $\mu m).$

Crystal	Type of inter-action	$\theta_{ m pm} \ [m deg]$	$ au_{ m p} \ [m ns]$	L [mm]	η [%]	Ref.	Notes
KDP	eoe	58	0.15	12	32 (energy)	[75Att]	$I_0 = 1 \text{ GW cm}^{-2}$
KDP	eoe a	58	25	_	6 (energy)	[79And1]	P = 40 MW
KDP	eoe	58	0.05	_	10 (energy)	[72Kun]	
DKDP	eoe	59.5	8	20	17 (energy)	[91Bor1]	$I_0 = 0.25 \text{ GW cm}^{-2}$
RDA	ooe	66.2	8	14.8	12 (power)	[75Kat2]	$\Delta\thetaL=1.0$ mrad cm
RDP	ooe	61.2	_	15.3	44 (power)	[74Kat5]	$I_0 = 0.2 \text{ GW cm}^{-2}$
RDP	ooe	61.2	8	15.3	21 (power)	[74Kat1]	
$LiIO_3$	ooe	47	0.8	8	0.7 (power)	[85Bog]	$P_{\rm av} = 4.5 \ {\rm mW}$
$LiIO_3$	ooe	47.5	_	4	4 (power)	[71Oka]	
BBO	eoe	64	8	5.5	23 (energy)	[86Che]	$I_0 = 0.25 \text{ GW cm}^{-2}$
BBO	ooe	31.3	8	7.5	20 (energy)	[91Bor1]	$I_0 = 0.19 \text{ GW cm}^{-2}$
BBO	ooe	31.3	9	6	35 (quantum)	[93Wu]	Intracavity THG, $P = 0.2 \text{ W}$
CBO	type II	40.3 $^{\rm b}$	0.035	5	80 °	[97Wu]	$I_0 = 5 \text{ GW cm}^{-2}$
LBO	type I	$38.1^{\ b}$	8	12.2	22 (energy)	[91Bor1]	$I_0 = 0.19 \text{ GW cm}^{-2}$
LBO	type II	41	8	12.6	60 (energy)	[89Wu]	

^a Neodymium silicate glass laser.

 $^{^{\}rm b}$ $\varphi_{\rm pm}.$ $^{\rm c}$ Conversion efficiency from 0.532 $\mu m.$

Table 4.1.10. Fourth harmonic generation of Nd:YAG laser radiation (1.064 \rightarrow 0.266 μm).

Crystal	Type of interaction	θ_{pm} [deg]	I_0 [W cm ⁻²]	$ au_{ m p}$ [ns]	L [mm]	Conversion efficiency (from 532 nm) [%]	Ref.	Notes
KDP	ooe	78	=	7	_	30 35	[77Aba]	
DKDP	ooe	90	8×10^{9}	0.03	4	75	[77Rei]	
DKDP	ooe	90	5×10^7	25	20	40	[76Liu]	T = 60 °C,
							_	P = 2.5 MW
DKDP	ooe	90	_	600	50	3.4	[85Per]	$T = 49.8 ^{\circ}\text{C},$
								$P_{\rm av} = 0.5 \ {\rm W}$
ADP	ooe	90	8×10^9	0.03	4	85	[77Rei]	
ADP	ooe	90	_	8	30	15 ^a	[75Kat3]	$T = 51.2 ^{\circ}\text{C},$
								$P_{\rm av} = 5 \ {\rm W}$
BBO	ooe	48	_	5	_	16	[88Lag]	E = 80 mJ
BBO	ooe	48	1.6×10^{8}	1	5	52	[86Che]	
BBO	ooe	57.8	_	$80 \ \mu s$	6.6	0.17	[93Dim]	Nd:YAG laser cooled
							_	to 253 K, $\lambda = 946 \text{ nm}$
CBO ^b	ooe	52.3	4×10^{9}	0.035	10	60	[97Wu]	
CLBO	ooe	62	_	10	9	30	[95Mor1]	E = 110 mJ
CLBO	ooe	61.6	_	7	10	50	[96Yap]	E = 500 mJ
CLBO	ooe	62.5	10^{11}	1.5 ps	10	24	[96Sha]	
CLBO	_	_	_	0.014	6	38	[97Sri]	
CLBO	ooe	_	1.7×10^{8}	46	15	20	[01Koj]	$T = 140 ^{\circ}\text{C},$
							,	$P_{\rm av} = 20.5 \; {\rm W}$
$\rm Li_2B_4O_7$	ooe	66	_	10	35	20	[97 Kom]	E = 160 mJ

 $[^]a$ Efficiency of conversion from 1.064 $\mu m.$ b 1.064 + 0.355 \rightarrow 0.266 $\mu m;$ conversion efficiency from 0.355 $\mu m.$

Table 4.1.11. Fifth harmonic (1.064 \rightarrow 0.2128 $\mu m)$ and sixth harmonic generation (1.064 \rightarrow 0.1774 $\mu m)$ of Nd:YAG laser radiation.

Crystal	$ heta_{ m pm} \ [m deg]$	Type of interaction	Crystal temperature [°C]	Output parameters	$ au_{ m p}$ [ns]	Ref.
KDP	90	ooe ^a	-70	E = 0.1 mJ	_	[69Akm]
KDP	90	ooe	-35	$P_{\text{av}} = 2.6 \text{ mW},$ f = 120 kHz	30	[78Mas]
KDP	90	ooe	-40	$P_{\text{av}} = 2 \text{ mW},$ f = 6 kHz	30	[79Jon]
KDP	84	ooe ^b	20	E = 0.45 mJ	0.015	[88Gar, 89Aru]
ADP	90	ooe	-40	$P_{\text{av}} = 5 \dots 7 \text{ mW},$ f = 10 Hz	10	[76Mas1]
ADP	90	ooe ^c	-67.5	E = 20 J	0.5	[88Beg]
KB5	$53 \pm 1(\varphi)$	eeo	20	E = 0.7 mJ	6	[76Kat1]
KB5	$53 \pm 1(\varphi)$	eeo	20	E = 0.1 mJ	0.02	[82Tan]
KB5	$52.1(\varphi)$	eeo	20	E = 0.3 mJ	0.03	[80Aru]
Urea	72	eeo	20	E = 30 mJ	10	[80Kat1]
BBO	55	ooe	20	E = 20 mJ	5	[86Che, 88Lag]
CLBO	_	ooe	20	E = 35 mJ	10	[95Mor1]
CLBO	67.3	ooe	20	E = 230 mJ	7	[96Yap]
$\mathrm{Li_{2}B_{4}O_{7}}$	80	ooe	20	E = 70 mJ	10	[97 Kom]
${ m KB5}^{ m d}$	$90(\theta), 68.5(\varphi)$	eeo	20	$P_{\rm av} = 6~{\rm mW}$	6	$[96 \mathrm{Ume}]$
${ m KB5}^{ m d}$	$80(\theta), 90(\varphi)$	ooe	20		6	$[96 \mathrm{Ume}]$

^a Neodymium silicate glass laser.

Table 4.1.12. Generation of harmonics of Nd:YAG laser radiation with $\lambda=1.318~\mu m$.

Number of harmonic	λ [nm]	Crystal	$\theta_{ m ooe}$ [deg]	L [mm]	$ au_{ m p}$ [ns]	Output parameters	Energy conversion efficiency [%]	Ref.
2	659.4	LiNbO ₃	44.67	16	40	85 kW	10	[81Akm]
3	439.6	KDP	42.05	30	40	$3.4~\mathrm{kW}$	0.4	[81Akm]
4	329.7	KDP	53.47	30	40	$6~\mathrm{kW}$	0.6	[81Akm]
5 ^a	263.8	KDP	55.33	30	30	$0.2~\mathrm{kW}$	0.02	[81Akm]
6 ^b	219.3	KB5	78 (eeo)	15	45	3 kW	0.5	[87Aru]
2	659.4	DCDA	70.38	13.5	25	$1.4~\mathrm{MW}$	40	[76Kat2]
2^{c}	659.4	$LiIO_3$	22	10	30	1 W (av.)	100	[81Kaz]
2	659.4	${\rm LiNbO_3}$	$90 \ (T = 300 \ ^{\circ}\text{C})$	19	50	60 mJ	48	[83Kaz]
2	659.4	${\rm LiNbO_3}$	90	20	50	10 mJ	21	[83Kaz]
$2^{c,d}$	659	LBO	$\varphi = 3.7$	_	2	0.3 W (av.)	_	[94Lin]
2^{d}	659	LBO	along Z axis	16	76	$0.85~\mathrm{mJ}$	40	[95Mor2]
3	439.6	KDP	$42.05 (T = 300 ^{\circ}\text{C})$	40	50	$1.4~\mathrm{mJ}$	3	[83Kaz]
3	439.6	$LiIO_3$	_	8	50	$1.4~\mathrm{mJ}$	1.2	[83Kaz]

^b Nd:YAlO₃ laser.

^c Nd:YLF laser.

^d Sixth harmonic generation, $\omega + 5 \omega = 6 \omega$.

 $[\]label{eq:continuous} \begin{array}{l} ^{\rm a}\ \omega+4\,\omega=5\,\omega\,. \\ ^{\rm b}\ 3\,\omega+3\,\omega=6\,\omega\,. \\ ^{\rm c}\ \ {\rm Intracavity\ SHG}. \end{array}$

^d Nd:YLF laser.

Table 4.1.13. Generation of harmonics of high-power Nd:glass laser radiation in KDP crystals.

Fundam	ental radiatio	n	Second 1	narmonic				Third a	nd fourth har	monics			Ref.
λ [μm]	$I_0 [10^9 \text{ cm}^{-2}]$	$ au_{ m p}$ [ns]	λ [μm]	Type of interaction	η [%]	Crystal length [mm]	E [J]	λ [μ m]	Type of interaction	η [%]	Crystal length [mm]	E [J]	
1.054	2.5	0.14	0.53	eoe ^a	67	12	9	0.35	eoe	80	12	11	[80Sek]
1.054	3.5	0.7	0.53	eoe	67	12	25	0.35	eoe	80	12	30	[80Sek]
1.064	2.5	0.1	0.532	eoe	67	8	17	0.266	ooe	30	7	4	[80Lot]
1.064	9.5	0.7	0.532	ooe	83	10	346						[82Lin]
1.064	2.0	0.7	0.532	eoe	67	12	_	0.355	eoe	55	10	41	[82Lin]
1.064	1.2	0.7	0.532	ooe	_	10	_	0.266	ooe	51	10	50	[82Lin]
1.06	0.2	25	0.53	ooe	80	40	60						[82Ibr]
1.06	2.7	0.5	0.53	ooe	90	30	20						[83Gul]
1.06	2.7	0.5	0.53	eoe ^a	67	18	_	0.35	eoe	81	18	$10 \dots 20$	[83Gul]
1.053	1.5	0.6	0.53	eoe	70	16	80	0.26	ooe	46	7	53	[85Bru]
1.054	5	0.5	0.53	eoe	87	17.5	_	0.264	ooe	$92^{\rm \ b}$	10	_	[88Beg]

 $[^]a$ The angle between the polarization vector of the fundamental radiation and o-ray is 35 $^\circ$. Conversion efficiency from 0.527 μm

Table 4.1.14. Generation of harmonics of iodine laser radiation: $\lambda = 1.315 \ \mu m \ (\tau_p = 1 \ ns) \ [80Fil, 81Wit, 83Fil, 83Bre].$

	SHG $\omega + \omega = 2$	ω		THG $\omega + 2\omega =$	3ω	FOHG $2\omega + 2\omega = 4\omega$	FIHG $2\omega + 3\omega = 5\omega$	SIHG $3\omega + 3\omega = 6\omega$
Wavelength [nm] Crystal	657.6 DKDP	KDP	KDP	438.4 DKDP	KDP	328.8 KDP	263.0 KDP	219.2 KB5
Crystal length [mm] Type of interaction $\theta_{\rm pm}$ [deg]	19 eoe 51.3	20 eoe 61.4	10 ooe 44.3	20 eoe ^a 48	10 ooe 42.2	40 ooe 53.6	– ooe 74	10 eeo 80.5 (φ_{pm})
Conversion efficiency [%] at $I_0 = (11.5) \times 10^9 \text{ W cm}^{-2}$ $I_0 = 3 \times 10^9 \text{ W cm}^{-2}$	30 70	16 -	12 -	30 50	6 –	15 30	- 9	- 3

Table 4.1.15. Second harmonic generation of ruby laser radiation (694.3 \rightarrow 347.1 nm).

Crystal	Type of interaction	$ heta_{ m pm} \ [m deg]$	I_0 [W cm ⁻²]	L [cm]	Power conversion efficiency [%]	Ref.	Notes
RDA	ooe	80.3 (90)	1.5×10^{8}	1.45	58	[74Kat3]	$T = 20 ^{\circ}\text{C (90 °C)},$
RDP	ooe	67	1.8×10^8	1.0	37	[74Kat4]	$L \Delta \theta = 4.37 \text{ mrad cm}$ $T = 20 ^{\circ}\text{C},$ $L \Delta \theta = 2.4 \text{ mrad cm}$
${ m LiIO_3}$	ooe	52	1.3×10^8	1.1	40	[70Nat]	$L \Delta \theta = 0.2 \text{ mrad cm}$

Table 4.1.16. Harmonic generation of Ti:sapphire (Ti: Al_2O_3) laser radiation.

(a) Second harmonic generation.

Crystal	$\lambda_{2\omega}$ [nm]	τ	$ heta_{ m pm}$ [deg]	L [mm]	Output power $P_{2\omega}$ [mW]	η [%]	Ref.	Notes
KDP	390	150 fs	43	3 40	300	50	[95Kry]	
$LiIO_3$	$360 \dots 425$	1.5 ps	43	10	700	50	[91Neb]	
BBO	$360 \dots 425$	1.5 ps	30	8	450	27	[91Neb]	
BBO	430	54 fs	27.5	0.055	230	$75 (5.2^{a})$	[92Ell]	ICSHG
BBO	$383 \dots 407$	_	ooe	5	170	$7.4^{\rm \ a}$	[93Poi]	ICSHG
BBO	425	16 fs	28	$0.1 \dots 1$	40	_	[95Ash]	
BBO	438	10 fs	26.7; ooe	0.04	3.6	1	[98Ste]	
BBO	400	150 fs	ooe	0.5	150	38	[98Zha]	
LBO	400	150 fs	_	3	130	32	[98Zha]	
LBO	$350 \dots 450$	$12 \dots 25 \text{ ns}$	$90 (\theta), 22 \dots 40 (\varphi)$	5	$25~\mathrm{mJ}$	30	[91Skr]	
LBO	$360 \dots 425$	1.5 ps	$90 (\theta), 32 (\varphi)$	8	350	20	[91Neb]	
LBO	410	cw	90 (θ) , 31.8 (φ)	10.7	410	21.6	[93Bou]	ERR
LBO	416	14 fs	$90 (\theta), 29 (\varphi)$	0.1	30	_	[94Bac]	ICSHG
LBO	400	1.5 ps	type I	10	1280	75	[94Wat]	ERR
LBO	398	cw	90 (θ) , 31.7 (φ)	8	650	70	[95Zho, 96Zho]	ICSHG
KNbO_3	$430 \dots 470$	35 ns	along a axis	7.9	$7.8~\mathrm{kW}$	$45(2^{a})$	[90Wu]	ICSHG
KNbO_3	430	cw	_	6	650	48	[91Pol]	ERR

^a Total conversion efficiency from the pump source.

(b) Third harmonic generation: $\omega + 2 \omega = 3 \omega$.

Crystal	$\lambda_{3\omega}$ [nm]	τ	$ heta_{ m pm} \ [m deg]$	L [mm]	Output power $P_{3\omega}$ [mW]	η [%]	Ref.	Notes
BBO LBO BBO	240 285 266 283 252 267	1 ps	50, ooe 90 (θ) , 70 (φ) 58, eoe	6.5 12 7 0.3	150 35 18	30 10 6	[91Neb, 92Neb] [91Neb] [93Rin]	f = 82 MHz $f = 82 MHz$ $f = 1 kHz$

(c) Fourth harmonic generation.

Crystal	$\lambda_{4\omega}$ [nm]	τ	$ heta_{ m pm} \ [m deg]$	L [mm]	Output power $P_{4\omega}$ [mW]	η [%]	Ref.	Notes
BBO ^a	205 213	1 ps	ooe	8	10	4	[91Neb]	f = 82 MHz
BBO ^b	$193 \dots 210$	$1 \dots 2 \text{ ps}$	75, ooe	6.9	10	4	[92Neb]	f = 82 MHz
BBO ^b	$193 \dots 210$	165 fs	65, ooe	0.1	6	3	[98Rot]	f = 82 MHz
BBO ^b	$193 \dots 210$	340 fs	65, ooe	0.3	15	-	[98Rot]	f = 82 MHz
BBO ^b	$189 \dots 200$	180 fs	71, ooe	0.1	4	1	[93Rin]	NC, f = 1 kHz
BBO ^b	186	10 ns	81 (θ) , 30 (φ) , ooe	5	0.008	_	[99Kou]	$T=91~\mathrm{K}$

 $^{^{\}mathrm{a}}\,\,2\,\omega + 2\,\omega = 4\,\omega\,.$

Table 4.1.17. Second harmonic generation of semiconductor laser radiation in $KNbO_3$.

λ_{ω} [nm]	Phase-matching conditions	L [mm]	$P_{2\omega}$ [mW]	η [%]	Ref.	Notes
842 865 842	T = -23 °C along a axis along a axis	5 5 5	24 0.215 6.7	14 1.7 0.57	[89Gol] [89Dix] [90Hem]	external resonator External Ring Resonator (ERR) ERR, cw
856 972	along a axis, $T = 15$ °C along b axis	7 5	41 1.2	39 4.8	[90Koz] [92Zim]	external resonator distributed Bragg reflection semi- conductor laser
858 858		12.4 12.4	62 80	1.1	[93Gol] [95Gol]	THG in LBO, 90 (θ), 31.8 (φ); 15 mm, $\lambda = 286$ nm, 0.05 mW
972	along b axis	6.5	156	_	[95Zim]	ERR, FOHG in BBO (14 mm) in ERR, $\lambda = 243$ nm, 2.1 mW
860 858	along a axis along a axis	10 10	50 90	60 -	[97Lod] [98Mat]	FOHG in BBO ($\theta = 71^{\circ}$): $\lambda = 214.5 \text{ nm}, 0.1 \text{ mW}$

Table 4.1.18. Second harmonic generation of dye laser radiation.

Crystal	$\lambda_{2\omega} \ [\mathrm{nm}]$	Parameters of output radiation (energy, power, pulse duration); conversion efficiency	Ref.	Notes
KDP KDP KDP ADP a ADP a ADP a ADP a ADP a RDP	267.5–310 280–310 280 280–310 290–315 250–260 293 295 313.8–318.5	0.1 kW (peak), $\eta = 1 \%$ 50 mJ 90 mW, $\eta = 10 \%$ 50 mJ, $\eta = 8.4 \%$ up to 1 mW, $\eta = 0.03 \%$ 120 μ W 0.13 mW, $\eta = 0.08 \%$, $\tau = 3$ ps $\eta = 10^{-4}$, $\tau = 3 \dots 4$ ps 3.6 MW, $\tau = 8$ ns, $\eta = 52 \%$ in power 3.2 MW, $\tau = 10$ ns, $f = 10$ Hz, $\eta = 36 \%$	[80Yam] [80Wel]	L = 55 mm, external cavity $\theta_{\text{ooe}} = 90^{\circ}, T = 200280 \text{ K}$ L = 3 mm L = 13 mm $\theta = 90^{\circ}, T = 2098^{\circ}\text{C},$ $I_0 = 36 \text{ MW cm}^{-2}, L = 25 \text{ mm}$ $\theta = 90^{\circ}$

(continued)

 $^{^{\}mathrm{b}}\ \omega + 3\,\omega = 4\,\omega\,.$

Table 4.1.18 continued.

Crystal	$\lambda_{2\omega} \ [\mathrm{nm}]$	Parameters of output radiation (energy, power, pulse duration); conversion efficiency	Ref.	Notes
$\overline{\mathrm{ADA}}$	292-302	30 mW	[77Fer]	θ = 90°
ADA ^a	285–315	400 mW (single-mode regime), 50 mW (multimode regime)	[76Fro]	$\theta = 90^{\circ}$, temperature tuning, L = 30 mm
DKDA	310–355	$0.8 \dots 3.2 \text{ MW}, \ \tau = 10 \text{ ns}, $ $f = 10 \text{ Hz}, \ \eta = 9 \dots 36 \%$	[77Kat2]	$\theta = 90$ °, $L = 15$ mm
LiIO ₃ ^a	295	$\eta = 10^{-4}, \tau = 2.1 \text{ ps}$	[80 Wel]	$L=0.3~\mathrm{mm}$
$LiIO_3^a$	293 - 312	0.37 mW, cw regime	[86Bue]	L = 10 mm
$LiIO_3$	293 - 330	15 mW, cw regime	[83Maj]	L=1 mm
$LiIO_3$	293	$3 \text{ kW}, \eta = 30 \%$	[76Str]	L=6 mm
$LiIO_3$	293-310	4 mW, cw regime, $\eta = 0.4\%$	[75Bet]	$L=6$ mm, $\Delta \lambda = 0.03$ nm
$LiIO_3$	293-310	21 mW, cw regime, $\eta = 2\%$	[75Bet]	$L=6$ mm, $\Delta \nu=30$ MHz
BBO	204.8 - 215	100 kW, $\tau = 8 \text{ ns}, \eta = 4 \dots 17 \%$	[86Kat]	$\theta = 70^{\circ} \dots 90^{\circ}$
BBO	205–310	50 kW, $\tau = 9 \dots 22 \text{ ns},$ $\eta = 1 \dots 36 \%$	[86Miy]	L=6 and 8 mm
BBO ^a	315	$20 \text{ mW}, \tau = 43 \text{ fs}$	[88Ede]	$\theta=38^{\circ}, \varphi=90^{\circ}, L=55~\mu\mathrm{m}$
BBO	230-303	$0.02 \dots 0.18 \text{ mJ}, \tau = 17 \text{ ns}$	[90Mue]	$\theta_{\text{ooe}} = 40 \dots 60 ^{\circ}, L = 7 \text{mm}$
BBO ^a	243	30 mW, cw regime	[91Kal]	$\theta_{\text{ooe}} = 55$ °, $L = 8$ mm, $\Delta \nu = 200 \text{ Hz}$
KB5	217.3 - 234.5	0.3 kW, $\tau = 7 \text{ ns}, \eta = 1 \%$	[75Dew]	XY plane, eeo
KB5	217.1 - 240	$5 \dots 6 \mu J, \tau = 3 \dots 4 \text{ns}, \eta = 10 \%$	[76Dew]	XY plane, $\theta_{\text{ooe}} = 90 \dots 0$ °
KB5	217.1–315.0	$5\ldots 6~\mu J, 5~ns, 10\%$	[76Dew]	XY plane, $\varphi_{\text{eeo}} = 90 \dots 31$ °, $L = 10 \text{ mm}$
KB5	217 - 250	$0.1 \dots 5 \mu J, \eta = 0.2 \dots 5 \%$	[76Zac]	XY plane, $\varphi_{\rm eeo} = 90 \dots 65$ °
DKB5	216.15	$2 \mu J$, $\tau = 3 \text{ ns}$, $\eta = 5 \%$	[78Pai]	$\theta = 90^{\circ}, \varphi = 90^{\circ}$
LFM	230-300	$\eta=2\%$	[73Dun]	XZ plane, $\theta_{\text{ooe}} = 35 \dots 45$ °, $L = 10 \text{ mm}$
LFM $^{\rm a}$	290 - 315	$\eta = 10^{-4}$	[72Gab]	XZ plane, $\theta_{\text{ooe}} = 45$ ° (590 nm)
LFM $^{\rm a}$	238 - 249	70 μJ (244 nm), cw regime	[80Bas]	XZ plane, $\theta_{\text{ooe}} = 39^{\circ} (486 \text{ nm})$
LFM	237.5–260	20 W, nanosecond regime, $\eta = 0.7\%$	[76Str]	, ,
LFM $^{\rm a}$	243	1.4 mW, cw regime	[84Foo]	$\theta_{\rm ooe} = 36.8$ °, $L = 15$ mm
LFM	285 – 310	4 μJ, cw regime	[75Bet]	•
KNbO_3	425–468	400 kW, $\eta = 43\%$	[79Kat]	angular tuning in XY and YZ planes, temperature tuning $(20220 ^{\circ}\text{C})$ along the a axis
KNbO_3	419–475	12 µW, cw regime, $\eta=0.065\%$	[83Bau]	along the a axis, T from -36 °C to $+180$ °C, $L=9$ mm
$\mathrm{KNbO_3}^{\mathrm{a}}$	425–435	21 mW, cw regime, $\eta=1.1\%$	[85Bau]	along the a axis, $T = 050$ °C, $L = 9$ mm
Urea	238-300	_	[79Hal]	$\theta_{\rm eeo} = 90 \dots 45$ °, $L = 2$ mm
Urea	298-370	_	[79Hal]	$\theta_{\rm eoo} = 90 \dots 50$ °, $L = 2$ mm

^a Intracavity SHG.

 ${\bf Table\ 4.1.19.\ Second\ harmonic\ generation\ of\ gas\ laser\ radiation.}$

Type of laser	Crystal	λ	$ heta_{ m pm}$	T	Ref.
		$[\mu \mathrm{m}]$	$[\deg]$	$[^{\circ}C]$	
Argon laser	KDP ^a	0.5145	90	-13.7	[67Lab]
	ADP	0.4965	90	-93.2	[73Jai]
	ADP	0.5017	90	-68.4	[73Jai]
	ADP	0.5145	90	-10.2	[73Jai]
	ADP ^a	0.5145	90	-10	[82Ber]
	KB5	0.4579	$67.2~(\varphi_{\mathrm{pm}})$	20	[76Che]
	KB5	0.4765	$60.2~(\varphi_{\mathrm{pm}})$	20	[76Che]
	KB5	0.4880	$56.6~(\varphi_{\mathrm{pm}})$	20	[76Che]
	KB5	0.5145	$50.2~(\varphi_{\rm pm})$	20	[76Che]
	BBO	0.5145	49.5	20	[86Xin]
	BBO	0.4965	52.5	20	[86Xin]
	BBO	0.4880	54.5	20	[86Xin]
	BBO	0.4765	57.0	20	[86Xin]
	BBO ^a	0.4880	55	20	[89Zim]
	BBO ^a	0.5145	_	20	[92Tai]
He-Ne laser	LiIO ₃ ^a	$1.152 \dots 1.198$	25	20	[83Kac]
	${\rm LiNbO_3}$	1.152	90	169	$[74\mathrm{Ant}]$
	$LiNbO_3$	1.152	90	281	[75Kus]
	${ m AgGaS_2}$	3.39	33	20	[75Bad]
NH ₃ laser	Te	12.8	_	_	[80Sha]
	$CdGeAs_2$	11.7	35.7		[87And3]
CO laser	$\rm ZnGeP_2$	$5.2 \dots 6.3$	47.5	_	[87 And 2]

 $^{^{\}rm a}$ Intracavity SHG.

Table 4.1.20. Harmonic generation of CO_2 laser radiation.

Crystal	$\lambda \ [\mu \mathrm{m}]$	Nonli- near process	Type of interaction, $\theta_{\rm pm}$ [deg]	I_0 [W cm ⁻²]	L [mm]	η (power) [%]	Ref.
$\overline{\mathrm{Ag_3AsS_3}}$	10.6	SHG	ooe, 22.5	1.1×10^{7}	4.4	2.2	[75Nik2]
$AgGaSe_2$	10.6	SHG	ooe, 57.5	1.7×10^{6}	15.3	2.7	[74Bye]
$AgGaSe_2$	10.25	SHG	ooe, 52.7	$< 10^{7}$	21	35	[85Eck]
$AgGaSe_2$	10.6	SHG	ooe, 53	_	20	0.1 ^a	[97Sto]
$ZnGeP_2$	$9.19 \dots 9.7;$	SHG	eeo, 76	_	_	5	[84And]
	$10.15 \dots 10.8$						
$ZnGeP_2$	8.6	SHG	eeo, 55.8	_	_	10.1	[87And4]
$ZnGeP_2$	10.6	SHG	eeo, 76	10^{9}	3	49	[87And1]
$ZnGeP_2$	$10.26 \dots 10.61$	SHG	eeo	4.4×10^{7}	7.2	11.3	[93Bar2]
$ZnGeP_2$	9.6	SHG	eeo, 70	5.5×10^{7}	10	8.1	[94Mas]
$ZnGeP_2$	10.78	SHG	eeo, 90	_	10	_	[97Kat]
$CdGeAs_2$	10.6	SHG	oeo, 48.4	1.4×10^{7}	9	15	[74Kil]
$CdGeAs_2$	10.6	SHG	eeo, 32.5	_	13	21	[76Men]
$CdGeAs_2$	10.6	SHG	eeo, 32.5	_	13	$0.44^{\rm \ a}$	[76Men]

(continued)

Table 4.1.20 continued.

Crystal	$\lambda \ [\mu \mathrm{m}]$	Nonli- near process	Type of interaction, $\theta_{\rm pm}$ [deg]	I_0 [W cm ⁻²]	L [mm]	η (power) [%]	Ref.
Tl ₃ AsSe ₃	9.6	SHG	_	_	3.7	10.9	[87Pas]
Tl_3AsSe_3	9.6	SHG	ooe, 19	10^{7}	$5 \dots 6$	28	[89Auy]
Tl_3AsSe_3	10.6	SHG	ooe	6.3×10^{8}	4.57	57	[91Suh]
Tl_3AsSe_3	9.25	SHG	ooe, 19	2×10^7	46	20	[96Suh]
GaSe	$9.3 \dots 10.6$	SHG	ooe, 12.8 14.4	2×10^{7}	6.5	9	[89Abd]
GaSe	$9.2 \dots 11.0$	SHG	ooe, 13	_	2.5	_	[95Bha]
$CdGeAs_2$	_	THG	oeo, 45	_	4.5	1.5	[79Men]
Tl_3AsSe_3	9.6	THG	ooe, 21	10^{7}	$5 \dots 6$	_	[89Auy]
$ZnGeP_2$	10.6	FOHG	eeo, 47.5	_	10	$14^{ m \ b}$	[87And1]
$ZnGeP_2$	_	FOHG	eeo, 47.5	_	5	2	[85And]
$ZnGeP_2$	10.6	FOHG	eeo, 47.8	_	10	_	[97Sto] ^a
$ZnGeP_2$	9.55	FOHG	eeo, 49	_	10	10	[98Cho]
Tl_3AsSe_3	9.6	FOHG	ooe, 27	10^{7}	$5 \dots 6$	$27^{\mathrm{\ b}}$	[89Auy]
$Tl_{3}AsSe_{3} \\$	9.6	FIHG	ooe, 28	10^{7}	$5 \dots 6$	45 $^{\rm c}$	[89Auy]

4.1.5 Sum frequency generation

Table 4.1.21. Sum frequency generation of UV radiation in KDP.

$\lambda_{ m SF} \ [m nm]$	Sources of interacting radiation	$ au_{ m p}$ [ns]	Conversion efficiency, power, energy	Ref.
190–212	SRS of 1.064 μ m + sum frequency radiation (220–250 nm) [83Tak]	0.02	20–40 μJ	[85Tak]
215 – 223	2ω of dye laser + Nd:YAG laser	10	10 kW	[76 Mas 1]
215–245	SRS of 266 nm (4 ω of Nd:YAG laser) + OPO (0.9–1.4 μ m)	0.02	100 μJ	[83Tak]
217–275	2ω of dye laser + Nd:YAG laser (1.064 μ m)	25–30	50–55 %, 10 mW (average)	[83Kop]
217 – 226	OPO (1.1–1.5 μ m) + 4 ω of Nd:YAG laser (266 nm)	0.02	100 kW	[82Tan]
218 – 244	(269-315 nm) [79Ang] + Nd:YAG laser	0.03	$0.1 \mathrm{\ mJ}$	[79Ang]
239	Nd:YAG laser $(1.064 \mu m) + XeCl$ laser $(308 nm)$	0.7	50%	[81Lyu]
240 – 242	2ω of ruby laser (347 nm) + dye laser	30	1 MW	[78Sti3]
257 - 320	Dye laser + argon laser	cw regime	$0.2~\mathrm{mW}$	[77Bli]
269 – 315	SRS of 532 nm (2 ω of Nd:YAG laser) + 532 nm	0.03	1-3 mJ	[79Ang]
269 - 287	OPO (1.29–3.6 μ m) + 3 ω of Nd:YAG laser (355 nm)	0.02	100 kW	[82Tan]
271	Two copper vapor lasers (511 and 578 nm)	35	1.5%, 100 mW (average)	[89Cou]
288–393 $^{\rm a}$	OPO (0.63–1.5 μ m) + 2 ω of Nd:YAG laser (0.532 nm)	0.02	100 kW	[82Tan]
360 – 415	Dye laser $+$ Nd:YAG laser	25 - 30	60 – 70%	[79Dud]
362 – 432	Dye laser $+$ Nd:YAG laser	0.03	20%	[76Moo]

^a DKDP crystal was used.

 $^{{\}rm ^a~Continuous\text{-}wave~regime.}$ ${\rm ^b~Conversion~efficiency~from~2}\,\omega\,.$ ${\rm ^c~Conversion~efficiency~from~4}\,\omega\,.$

Table 4.1.22. Sum frequency generation of UV radiation in ADP.

$\overline{\lambda_{ m SF}}$ [nm]	Sources of interacting radiation	$ au_{ m p}$ [ns]	Conversion efficiency, power, energy	Ref.
208–214	2ω of dye laser + Nd:YAG laser, $\theta = 90^{\circ}, T = -120^{\circ}0^{\circ}C$	10	1.7 μJ	[76Mas1]
222 - 235	2ω of dye laser + Nd:YAG laser	10	10%	[76Mas1]
240-248	Dye laser $+ 2\omega$ of ruby laser, $\theta = 90^{\circ}$, $T = -20 + 80^{\circ}$ C	30	4%, 1 MW	[78Sti3]
243–247 $^{\rm a}$	Dye laser + argon laser (363.8 nm)	cw regime	$4~\mathrm{mW}$	[91Kal, 83Cou]
243^{a}	Dye laser + argon laser (351 nm), $\theta = 90^{\circ}$, $T = 8^{\circ}$ C	cw regime	$0.3~\mathrm{mW}$	[83Hem1]
247.5	Dye laser + krypton laser (413.1 nm), $\theta = 90$ °, $T = -103$ °C	cw regime	_	[79Mar]
246-259	Dye laser + 2ω of Nd:YAG laser, $\theta = 90$ °, T = -1200°C	10	1% , $3~\mu J$	[76Mas1]
$252 – 268 ^{\rm a} \\ 270 – 307$	Dye laser + argon laser (477, 488, 497 nm), $\theta_{\rm ooe} = 90$ ° Dye laser + 2ω of Nd:YAG laser, $\theta_{\rm ooe} = 81$ °	cw regime ps regime	8 mW -	[82Liu] [76Moo]

^a ADP crystal was placed in an external resonator.

Table 4.1.23. Sum frequency generation of UV radiation in BBO.

λ_{SF} [nm]	Sources of interacting radiation	$ au_{ m p}$ [ns]	Conversion efficiency, power, energy	Ref.
188.9–197	Dye laser (780–950 nm) + 2ω of another dye laser (248.5 nm)	10	up to 0.1 mJ	[88Mue]
190.8–196.1	Ti:sapphire laser (738–825 nm) + 2 ω of Ar laser (257 nm)	_	tens of nW	[91Wat]
193	Dye laser + KrF laser (248.5 nm)	9	$0.2\%, 2 \mu J$	[88Mue]
193	Dye laser (707 nm) + 4ω of Nd:YAG laser	90-250 fs	10 μJ (250 fs)	[92Hof]
193.3	Dye laser (724 nm, 5 ps) + 4ω of Nd:YLF laser (263 nm, 25 ps)	0.01	1.7 %, 4 μJ (2.5 mJ) ^a	[92Tom]
193.4	FOHG of dye laser radiation (774 nm, 300 fs), $\omega + 3 \omega = 4 \omega$	800 fs	0.5 μJ (1.5 mJ) ^a	[92Rin]
194	Ti:sapphire laser $+ 2\omega$ of Ar laser (257 nm), three crystal configuration with external cavity	_	0.016 mJ	[92Wat]
194	Diode laser (792 nm) + 2ω of Ar laser (257 nm)	cw	$2~\mathrm{mW}$	[97Ber]
195.3	THG of dye laser $(T [crystal] = 95 \text{ K})$	17	$5\%, 8 \mu J$	[88Lok]
196-205	Dye laser $+ 2\omega$ of another dye laser	5	$0.1 \mathrm{mJ}$	[92Hei]
197.7 - 202	THG of dye laser	0.008	1%, 1–4 mW	[88Gus]
198 – 204	THG of dye laser	5	20%, 1.7 mJ	[87Gla]
271	Two copper vapor lasers (511 and 578 nm)	35	0.9%, $64 mW$	[89Cou]
362.6-436.4	Dye laser + Nd:YAG laser, noncollinear SFG (NCSFG), $\alpha = 4.8 \dots 21.3$ °	-	1 %, 0.065 mJ	[90Bha1]
369	Diode laser $(1310 \text{ nm}) + \text{Ar laser } (515 \text{ nm})$	_	$1.3~\mu\mathrm{W}$	[91Sug]
370.6	Dye laser (568.6 nm) + Nd:YAG laser, NCSFG, $\alpha = 6.3$ $^{\circ}$	_	8–18%	[92Bha]

^a After amplification in an ArF excimer gain module.

Table 4.1.24. Sum frequency generation of UV radiation in LBO.

$\lambda_{ m SF} \ [m nm]$	Sources of interacting radiation	$ au_{ m p}$ [ns]	Conversion efficiency, power, energy	Ref.
170–185 ^a	OPO (1.6–2.5 μ m) + 4 ω of Ti-sapphire laser (189–210 nm), θ = 66–90 $^{\circ}$, ooe	100 fs	4	[98Pet3]
172.7 – 187	OPO (1.65–2.15 µm) + 4ω of Ti-sapphire laser (190–203.75 nm), $\theta = 90^{\circ}$, $\varphi = 73^{\circ}$, ooe	130 fs	$50 \mathrm{~nJ}$	[94Sei3]
185–187.5 $^{\rm b}$	OPO + 5 ω of Nd:YAG laser (212.8 nm), $\theta = 62-74$ °	-	_	[95Kat]
194 ^b	OPO + 5 ω of Nd:YAG laser (212.8 nm), θ = 51.2°, ω = 90°	5	2.2%	$[00\mathrm{Kag}]$
185 ^c	OPO + 5ω of Nd:YAG laser (212.8 nm), $\theta = 64$ °	_	_	[97 Ume]
194 ^c	OPO + 5ω of Nd:YAG laser (212.8 nm), $\theta = 53^{\circ}$, $\omega = 0^{\circ}$	5	1 %	[00Kag]
$195-210^{\text{ c}}$ 226-265	Nd:YAG laser $+ 2\omega$ of dye laser, 2ω or 3ω of dye laser	10	14%	[00Bha]
188–195	OPO (1.6–2.3 μ m) + 5 ω of Nd:YAG laser (212.8 nm), $\theta = 90^{\circ}$, $\varphi = 90$ –52 $^{\circ}$, ooe	6	$0.2 – 2 \%, \\ 2 – 40 \ \mu J$	[91Bor2]
187.7–195.2	· ·	8	3 kW (peak)	[92Wu]
191.4	SRS in H ₂ (1.908 µm) + 5 ω of Nd:YAG laser, $\theta = 90$ °, $\varphi = 88-50$ °, ooe	8	10 %, 67 kW (peak), 2 mW (average)	[92Wu]
218-242	OPO (1.2–2.6 µm) + 4 ω of Nd:YAG laser (266 nm), $\theta = 90^{\circ}$, $\varphi = 90$ –33°, ooe	6	0.2–2 %, 20–400 μJ	[91Bor2]
232.5 - 238	Nd:YAG laser $+ 2\omega$ of dye laser	10	_	[90Kat]
240 – 255	Nd:YAG laser + 2ω of dye laser, NCSFG	10	$8\%,0.12~\mathrm{mJ}$	[93Bha]

 $^{^{\}rm a}$ $\rm Li_2B_4O_7$ crystal was used. $^{\rm b}$ CBO crystal was used.

 ${\bf Table~4.1.25.~Sum~frequency~generation~of~UV~radiation~in~KB5.}$

$\lambda_{ m SF} \ [m nm]$	Sources of interacting radiation	$ au_{ m p} \ [m ns]$	Conversion efficiency, power, energy	Ref.
208-217	Two dye lasers, $\theta = 90^{\circ}$, $\varphi = 90^{\circ}$, eeo	10	0.025 %, 1 W	[76Dun]
196.6	Dye laser $+ 2\omega$ of Nd:YAG	8	$0.1\%, 0.5 \mathrm{mJ}$	[77Kat1]
207.3 – 217.4	Ruby laser (694.3 nm) + 2ω of dye laser	3	0.3%, 0.8 mJ	[77Kat2]
201 – 212	Nd:YAG + 2ω of dye laser	20	10% , $2-10 \mu J$	[77Sti]
185-200	Dye laser (740–910 nm) + 2 ω of dye laser (237 nm), $\theta = 90^{\circ}$, eeo	30	10 %, up to 10 μJ	[78Sti2]
211–216	Dye laser + Ar laser (351.1 nm)	cw regime	10 ⁻⁶ , 50–100 nW	[78Sti1]
196.7–226	OPO + 3ω and 4ω of Nd:YAG laser, $\theta = 90$ °, $\varphi = 65$ °, eeo	0.02	$20~\mathrm{kW}$	[82Tan]
194.1-194.3	Dye laser $+ 2\omega$ of Ar laser (257 nm)	cw regime	$2 \mu W$	[83Hem2]
200-222	OPO + 3ω and 4ω of Nd:YAG laser	0.045	$2 \times 10^{-5}, 1 \mu\text{J}$	[83Pet]
166–172	OPO (1.15–1.6 µm) + 4 ω of Ti-sapphire laser, $\theta=90$ °, $\varphi=90$ °, eeo	200 fs	0.05–0.4 MW	[98Pet2]

^c CLBO crystal was used.

 ${\bf Table~4.1.26.~Up\mbox{-}conversion~of~near~IR~radiation~into~the~visible.}$

Crystal	$\lambda_{ m IR} \ [\mu { m m}]$	Pump source	η [%]	Ref.
LiIO ₃	3.39	0.694 µm, mode-locked ruby laser	100	[73Gur]
	$3.2 \dots 5$	1.064 µm, Nd:YAG laser	0.001	[74Gur]
	2.38	0.488 μm, argon laser	4×10^{-8}	[75Mal2]
	1.98, 2.22, 2.67	0.694 µm, mode-locked ruby laser	$0.14 \dots 0.28$	[75Mal1]
	3.39	0.5145 μm, argon laser	2.4×10^{-2}	[80See]
	$1 \dots 2$	$0.694~\mu\mathrm{m},\mathrm{ruby~laser}$	18	[71Cam]
${\rm LiNbO_3}$	$1.69 \dots 1.71$	0.694 μm, Q-switched ruby laser	1	[67Mid]
	$1.6 \dots 3.0$	0.694 μm, Q-switched ruby laser	100	[75Aru]
	1.6	0.694 μm, ruby laser	10^{-5}	[68Mid]
	3.3913	$0.633~\mu m$, cw He-Ne laser	10^{-5}	[67Mil]
	3.3922	$0.633~\mu\mathrm{m},~\mathrm{cw}$ He-Ne laser	5×10^{-5}	[73Bai]
KTP	1.064	$0.809 \mu m$, diode laser	68	[93Kea]
	1.54	0.78 µm, diode laser	7×10^{-4}	[93Wan1]
	1.064	0.824 μm, dye laser (intracavity SFG)	0.26	[90Ben]
	1.064	0.809 μm, diode laser	55	[92Ris]
	1.064	$0.805 \mu m$, diode laser	24	[92Kea]
	1.319; 1.338	$0.532~\mu\mathrm{m},~2\omega$ of Q-switched Nd:YAG laser	10	[89Sto]

 $^{^{\}rm a}$ The angle between the polarization vector of the fundamental radiation and o-ray is 35 $^{\circ}.$

Table 4.1.27. Up-conversion of CO_2 laser radiation by sum-frequency generation.

Crystal	Pump source	$\lambda_{ m pump} \ [\mu { m m}]$	Type of interaction	$ heta_{ m pm} \ [m deg]$	${\rm [W~cm^{-2}]}$	L [mm]	η [%]	Ref.
$\overline{\mathrm{Ag_3AsS_3}}$	ns Nd:YAG laser, 740 W	1.064	eoe	20	_	6	0.84	[72Tse]
_	Ruby laser, 1 ms	0.694	_	_	10^{4}	10	0.14	[72Luc]
	ns Nd:YAG laser	1.064	eoe	20	400	6	0.5	[73Alc]
	Nd:YAG laser	1.064	eoe	20	_	14	1.5	[74Vor]
	Ruby laser, 25 ps	0.694	ooe	25.2	10^{8}	5	10.7	[75Nik1]
	ns Nd:YAG laser	1.064	eoe	20	=	_	30 a	[79Jaa]
	ns Nd:YAG laser	1.064	eoe	20	$(0.5\dots1.2)\times10^6$	_	8 ^b	[81And]
$AgGaS_2$	Nd:YAG laser	1.064	oee	40	6×10^5	3	40 a	[75Vor]
J	Dye laser, 3 ns	0.598	ooe	90	_	5	40	[77Jan]
	Ruby laser, 30 ns	0.694	eoe	55	_	3.3	9	[77And1]
	ns Nd:YAG laser	1.064	oee	40	_	_	30	[78Vor]
	ns Nd:YAG laser	1.064	oee	40	$(0.5\dots1.2)\times10^6$	-	14^{b}	[81And]
${\rm HgGa_2S_4}$	ns Nd:YAG laser	1.064	ooe	41.6	$(0.5\dots1.2)\times10^6$	3.6	$60 (20)^{\rm b}$	[80And, 81And]
$ZnGeP_2$	Nd:YAG laser	1.064	oeo	$82 \dots 89$	_	10	1.4	[71Boy]
	ns Nd:YAG laser	1.064	oeo	82.9	$(0.51.2) \times 10^6$	_	6 ^b	[81And]
	Nd:YAG laser, 30 ns	1.064	oeo	82.5	3×10^6	3	5	[79And2]
CdSe	Nd:YAG laser	1.833	oeo	77	2.4×10^7	10	35 a	[71Her]
	HF laser, 250 ns	2.72	oeo	70.5	6×10^6	30	40	[76Fer]

 ^a Power-conversion efficiency.
 ^b Power-conversion efficiency for two cascades:

 $^{10.6 + 1.064 \}rightarrow 0.967 \ \mu m$ $0.967 + 1.064 \rightarrow 0.507 \ \mu m.$

4.1.6 Difference frequency generation

Table 4.1.28. Generation of IR radiation by DFG.

(a) Crystal: LiIO₃

$\frac{\lambda}{\lambda}$ [μm]	Sources of interacting radiations, crystal parameters	Conversion efficiency, energy, power, $\tau_{\rm p}$	Ref.
4.1-5.2	Dye laser + ruby laser, ICDFG, $L = 12 \text{ mm}$	100 W (peak)	[72Mel]
1.25 - 1.60;	Dye laser $+$ Q-switched Nd:YAG laser	0.5-70 W (peak),	[75Gol]
3.40 – 5.65	(1.064 and 0.532 µm), ICDFG, $\theta_{\rm ooe} = 2128.5$ °	$\Delta \nu = 0.1 \text{ cm}^{-1}, 60 \text{ ns}$	
2.6 – 7.7	Dye laser + 2ω of Nd:YAG laser, $\theta_{\rm ooe} = 22$ °	$2~\mathrm{nJ}50~\mu\mathrm{J},10~\mathrm{ns}$	[95Cha2]
2.3 – 4.6	Dye laser + argon laser (514 and 488 nm)	$0.54~\mu\mathrm{W},~\mathrm{cw}$	[76Wel]
4.3 – 5.3	Dye laser + 2ω of Nd:YAG laser, $\theta_{\rm ooe} = 24.3$ °	_	[77Dob]
0.7 – 2.2	Dye laser + nitrogen laser, $\theta_{\rm ooe} = 51-31$ °	3 ns	[78Koe]
3.8 – 6.0	Dye laser + copper vapor laser (511 nm), $\theta_c = 21-24$ °	$10-100~\mu W,~20~ns$	[82Ata]
3.5 – 5.4	Dye laser + 2ω of Nd:YAG laser, $\theta_{\rm ooe} = 20$ $^{\circ}$	0.8 mJ, 10 ns	[83Man]
1.2 – 1.6	Two dye lasers, $\theta_{\rm ooe} = 29$ °	1.5-5 ps	[84Cot]
4.4 – 5.7	Dye laser + Nd:YAG laser, $\theta_{\rm ooe} = 2022^{\circ}$	550 kW, 8 ns	[85Kat]
~ 5	Two dye lasers, $\theta_{\rm ooe} = 20$ °, $L = 3$ mm	10%, $10 nJ$, $400 fs$	[91Els]
2.5 – 5.3	Signal and idler pulses of OPO, $\theta_{\rm ooe} = 21$ $^{\circ}$	0.2 mW, f = 82 MHz, 200 fs	[94Loh]
6.8 – 7.7	Dye laser + 2ω of Nd:YAG laser, $\theta = 2829~^\circ$	100 mW (peak)	[95Cha1]

(b) Crystal: $LiNbO_3$

λ [μm]	Sources of interacting radiations, crystal parameters	Conversion efficiency, energy, power, $\tau_{\rm p}$	Ref.
3–4	Dye laser + ruby laser	1%, 6 kW	[71Dew]
2.2 – 4.2	Dye laser $+$ argon laser	$1 \mu W, cw$	[74Pin]
2 - 4.5	Dye laser (1.2 ps) + argon laser (100 ps), $\theta = 90^{\circ}$,	25 μW (average),	[84Rud,
	$T = 200 \dots 400 ^{\circ} \text{C}$	1.2 ps, f = 138 MHz	85Ree]
2-4	Dye laser + Nd:YAG laser, $\theta_{\rm ooe} = 4657$ °	60%, 1.6 MW	[80Kat2]
2.04	Two dye lasers, $\theta_{\rm ooe} = 90^{\circ}$	$50\%, \Delta \lambda = 0.03 \mathrm{nm}$	[77Sey]
1.7 – 4.0	CPM dye laser + subpicosecond continuum,	10 kW (peak), 0.2 ps,	[87Moo2]
	$\theta_{\rm c} = 55$ °, $L = 1$ mm	$\Delta \nu = 100 \text{ cm}^{-1}$	
4.043	Two Nd:YAG lasers (1.064 and 1.444 μ m), $L=25$ mm	5.5%, $30 mJ$, $14 ns$	[94Won]
1.6 – 4.8	$Nd:glass\ laser\ +\ OPO$	$6\%,30~\mu\mathrm{J},13~\mathrm{ps}$	[95DiT]

(c) Crystal: BBO

$\frac{\lambda}{[\mu \mathrm{m}]}$	Sources of interacting radiations, crystal parameters	Conversion efficiency, energy, power, $\tau_{\rm p}$	Ref.
2.5	Dye laser (620 nm) + picosecond continuum (825 nm), $\theta_{\text{ooe}} = 20.3$ °, $L = 5$ mm	5% , 4 μ J, 0.5 ps	[91Pla]
0.9 – 1.5	Dye laser + Nd:YAG laser, $\theta_{\rm ooe} = 20.524.5^{\circ}$, $L = 10~\rm{mm}$	23%,4.5 mJ, 8 ns	[93Ash]
2.04–3.42 1.23–1.76	Two dye lasers, NCDFG, $\theta_{\rm ooe}=1217$ °, $L=6~\rm mm$ Dye laser + Ti:sapphire laser	$300{\text -}400 \text{ W (peak)}$ $10 \mu\text{W (average)},$ 150 fs, f = 80 MHz	[91Bha] [93Sei]

1.35–1.75 Dye laser + 2 ω of Ti:sapphire laser, ICDFG 2.8–3.6 Ti:sapphire laser + OPO, $\theta_{\rm coe} = 90^{\circ}$, $\varphi = 47^{\circ}$ 40–150 μW, 90–350 fs, [9] 1.2–2.2 Nd;YAG laser + dye laser, $\theta_{\rm coe} = 90^{\circ}$, $\varphi = 31^{\circ}$ 1.05–2.8 Two Ti:sapphire lasers, dye laser + Ti:sapphire laser 1.14–1.23 Dye laser (550–570 nm) + Nd;YAG laser, $\theta_{\rm eco} = 82-90^{\circ}$, $\varphi = 0^{\circ}$ (e) Crystal: KTA λ Sources of interacting radiations, crystal parameters 2.66–5.25 Ti:sapphire laser + Nd;YAG laser, $\theta_{\rm coe} = 40^{\circ}$, $\varphi = 0^{\circ}$ 60 % (quantum), 1–15 mJ, 2 ns (f) Crystal: Ag ₃ AsS ₃ λ Sources of interacting radiations, crystal parameters 2.75 Ti:sapphire laser + Nd;YAG laser, $\theta_{\rm coe} = 40^{\circ}$, $\varphi = 0^{\circ}$ 60 % (quantum), 1–15 mJ, 2 ns (f) Crystal: Ag ₃ AsS ₃ λ Sources of interacting radiations, crystal parameters 2.75 Towo dye lasers 3.7-10.2 OPO (1.06-1.67 μm) + 2ω of phosphate glass laser (527 nm) (g) Crystal: AgGaS ₂ λ Sources of interacting radiations, crystal parameters 2.5-18.3 Two dye lasers, $\theta_{\rm coe} = 38-52^{\circ}$ 3.9-9.4 Dye laser + Nd;YAG laser, $\theta_{\rm coe} = 38-52^{\circ}$ 3.9-9.4 Dye laser + Nd;YAG laser, $\theta_{\rm coe} = 38-52^{\circ}$ 3.9-9.4 Dye laser + Nd;YAG laser, $\theta_{\rm coe} = 38-52^{\circ}$ 3.9-9.4 Dye laser + Nd;YAG laser, $\theta_{\rm coe} = 38-52^{\circ}$ 3.9-9.4 Dye laser + Nd;YAG laser, $\theta_{\rm coe} = 38-52^{\circ}$ 3.9-9.4 Dye laser + Nd;YAG laser, $\theta_{\rm coe} = 38-52^{\circ}$ 3.9-9.4 Dye laser + Nd;YAG laser, $\theta_{\rm coe} = 38-52^{\circ}$ 3.0 mW, 10 ns 3.7-10.6 Two dye lasers, $\theta_{\rm coe} = 65-85^{\circ}$ 3.0 mW, 10 ns 3.7-10.6 Two dye lasers, $\theta_{\rm coe} = 65-85^{\circ}$ 3.0 mW, 10 ns 4.6-12 Two dye lasers, $\theta_{\rm coe} = 65-85^{\circ}$ 3.0 mW, 10 ns 4.6-12 Two dye laser + Ti:sapphire laser, $\theta_{\rm coe} = 90^{\circ}$ 4.7-6-6.45 Dye laser + Ti:sapphire laser, $\theta_{\rm coe} = 90^{\circ}$ 4.7-8-6.45 Dye laser + Ti:sapphire laser, $\theta_{\rm coe} = 90^{\circ}$ 4.7-8-7-8 Dye laser + Ti:sapphire laser, $\theta_{\rm coe} = 90^{\circ}$ 4.7-8-7-8 Dye laser + Ti:sapphire laser, $\theta_{\rm coe} = 90^{\circ}$ 4.7-8-7-8 Dye laser + Ti:sapphire laser, $\theta_{\rm coe} = 90^{\circ}$ 4.7-8-7-8 Dye laser +	Ref.	Conversion efficiency, energy, power, $\tau_{\rm p}$	Sources of interacting radiations, crystal parameters	λ [μm]					
1.35–1.75 Dye laser + 2 ω of Ti:sapphire laser, ICDFG 2.8–3.6 Ti:sapphire laser + OPO, $\theta_{\rm eoe} = 90^\circ$, $\varphi = 47^\circ$ for the sum of the sum	[75Bri]	, ,	Dye laser + Nd:YAG laser, $\theta_{\rm eoe} = 7678^{\circ}, \varphi = 0^{\circ}$	1.4–1.6					
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	[94Pet] [95Gal1]	$\begin{array}{c} 10~{\rm W~(peak),~1.6~ps} \\ 40{-}150~\mu{\rm W,~90}{-}350~{\rm fs}, \end{array}$							
Sources of interacting radiations, [µm] crystal parameters energy, power, τ_p [2.66–5.25 Ti:sapphire laser + Nd:YAG laser, $\theta_{\rm eoe} = 40^\circ$, $\varphi = 0^\circ$ 60% (quantum), 1 –15 mJ, 2 ns [5] (f) Crystal: Ag ₃ AsS ₃ $\lambda \qquad \text{Sources of interacting radiations,} \qquad \text{Conversion efficiency,} \qquad \text{Renergy, power, } \tau_p$ 11–23 Two dye lasers 3 W (peak), 30 ns [7] 3.7–10.2 OPO (1.06–1.67 µm) + 2ω of phosphate glass laser (527 nm) (g) Crystal: AgGaS ₂ $\lambda \qquad \text{Sources of interacting radiations,} \qquad \text{Conversion efficiency,} \qquad \text{Renergy, power, } \tau_p$ (g) Crystal: AgGaS ₂ $\lambda \qquad \text{Sources of interacting radiations,} \qquad \text{Conversion efficiency,} \qquad \text{Renergy, power, } \tau_p$ 5.5–18.3 Two dye lasers, $\theta = 90^\circ$ 4 W, 4 ns [7] 5.10 Dye laser + Nd:YAG laser, $\theta_{\rm eoe} = 38$ –52 $\theta_{\rm eoe} = 38$ –53 $\theta_{\rm eoe} = 38$ –54 $\theta_{\rm eoe} = 38$ –55 $\theta_{\rm eoe} = 38$ –57 $\theta_{\rm eoe} = 38$ –57 $\theta_{\rm eoe} = 38$ –58 $\theta_{\rm eoe} = 38$ –59 $\theta_{\rm eoe} = 38$ –59 $\theta_{\rm eoe} = 38$ –50 $\theta_{\rm eoe$	[95Cha3] [96Mom] [96Bha]	36 % (quantum), 1 mJ 20 μW, cw	1.05–2.8 Two Ti:sapphire lasers, dye laser + Ti:sapphire laser 1.14–1.23 Dye laser (550 – 570 nm) + Nd:YAG laser,						
$ [\mu m] \qquad \text{crystal parameters} \qquad \text{energy, power, τ_p} \\ \hline 2.66-5.25 \qquad \text{Ti:sapphire laser} + \text{Nd:YAG laser, $\theta_{eoe} = 40~^\circ$, $\varphi = 0~^\circ$} \qquad \begin{array}{c} 60\% \text{ (quantum), } \\ 1-15\text{ mJ, 2 ns} \end{array} \\ \hline \\ \textbf{(f) Crystal: Ag_3AsS_3} \\ \hline \\ \hline \lambda \qquad \qquad \text{Sources of interacting radiations, } \qquad \text{Conversion efficiency, energy, power, τ_p} \\ \hline 11-23 \qquad \text{Two dye lasers} \qquad 3\text{W (peak), 30 ns} \\ 3.7-10.2 \qquad \text{OPO (1.06-1.67 } \mu\text{m}) + 2\omega \text{ of phosphate glass laser} \\ \hline (527\text{nm}) \qquad \qquad \begin{array}{c} 25-50\mu\text{J, 10 ps} \end{array} \\ \hline \\ \textbf{(g) Crystal: AgGaS_2} \\ \hline \\ \hline \lambda \qquad \qquad \text{Sources of interacting radiations, } \qquad \text{Conversion efficiency, energy, power, τ_p} \\ \hline \textbf{(g) Crystal: AgGaS_2} \\ \hline \lambda \qquad \qquad \text{Sources of interacting radiations, energy, power, τ_p} \\ \hline \textbf{(5-11)} \qquad \qquad \text{Crystal parameters} \qquad \qquad \text{energy, power, τ_p} \\ \hline \textbf{(5-13.3)} \qquad \qquad \text{Two dye lasers, $\theta = 90~^\circ$} \qquad \qquad 4\text{W, 4 ns} \qquad \boxed{7} \\ \hline \textbf{(5-11)} \qquad \qquad \text{Dye laser} + \text{Nd:YAG laser, $\theta_{eoe} = 38-52~^\circ$} \qquad 180\text{kW, 12 ns} \qquad \boxed{8} \\ \hline \textbf{(8-11)} \qquad \qquad \text{OPO (2-4}\mu\text{m}) + \text{radiation at $\lambda = 1.4-2.13}\mu\text{m} \qquad 1\text{kW, 8 ns} \qquad \boxed{8} \\ \hline \textbf{(8-11)} \qquad \qquad \text{OPO (2-4}\mu\text{m}) + \text{radiation at $\lambda = 1.4-2.13}\mu\text{m} \qquad 1\text{kW, 8 ns} \qquad \boxed{8} \\ \hline \textbf{(8-11)} \qquad \qquad \text{OPO (2-4}\mu\text{m}) + \text{radiation at $\lambda = 1.4-2.13}\mu\text{m} \qquad 1\text{kW, 8 ns} \qquad \boxed{8} \\ \hline \textbf{(8-11)} \qquad \qquad \text{OPO (2-4}\mu\text{m}) + \text{radiation at $\lambda = 1.4-2.13}\mu\text{m} \qquad 1\text{kW, 8 ns} \qquad \boxed{8} \\ \hline \textbf{(8-11)} \qquad \qquad \text{OPO (2-4}\mu\text{m}) + \text{radiation at $\lambda = 1.4-2.13}\mu\text{m} \qquad 1\text{kW, 8 ns} \qquad \boxed{8} \\ \hline \textbf{(8-11)} \qquad \qquad \text{OPO (2-4}\mu\text{m}) + \text{radiation at $\lambda = 1.4-2.13}\mu\text{m} \qquad 1\text{kW, 8 ns} \qquad \boxed{8} \\ \hline \textbf{(8-11)} \qquad \qquad \text{OPO (2-4}\mu\text{m}) + \text{radiation at $\lambda = 1.4-2.13}\mu\text{m} \qquad 1\text{kW, 8 ns} \qquad \boxed{8} \\ \hline \textbf{(8-11)} \qquad \qquad \text{OPO (2-4}\mu\text{m}) + \text{radiation at $\lambda = 1.4-2.13$}\mu\text{m} \qquad 1\text{kW, 8 ns} \qquad \boxed{8} \\ \hline \textbf{(8-11)} \qquad \qquad \text{OPO (2-4}\mu\text{m}) + \text{radiation at $\lambda = 1.4-2.13$}\mu\text{m} \qquad 1\text{kW, 8 ns} \qquad \boxed{8} \\ \hline \textbf{(8-11)} \qquad \qquad \text{OPO (2-4}\mu\text{m}) + \text{radiation at $\lambda = 1.4-2.13$}\mu\text{m} \qquad 1\text{kW, 8 ns} \qquad \boxed{8} \\ \hline \textbf{(8-11)} \qquad \qquad \text{OPO (2-4}\mu\text{m}) + \text{radiation at $\lambda = 1.4-2.13$}\mu\text{m} \qquad 1\text{kW, 8 ns}$			l: KTA	(e) Crystal					
(f) Crystal: Ag ₃ AsS ₃ $ \lambda \text{Sources of interacting radiations,} \text{Conversion efficiency,} \text{R} \\ [\mu\text{m}] \text{crystal parameters} \text{energy, power, } \tau_{\text{p}} \\ 11-23 \text{Two dye lasers} 3 \text{ W (peak), } 30 \text{ ns} \\ 3.7-10.2 \text{OPO } (1.06-1.67 \text{ μm}) + 2 \omega \text{ of phosphate glass laser} \\ (527 \text{ nm}) \text{(5} \\ 25-50 \text{ μJ, } 10 \text{ ps} \text{[8} \\ 25-5$	Ref.		-						
Sources of interacting radiations, crystal parameters energy, power, $\tau_{\rm p}$ [11–23 Two dye lasers 3.7–10.2 OPO (1.06–1.67 μm) + 2ω of phosphate glass laser (527 nm) (527 nm) (527 nm) (25–50 μJ, 10 ps (527 nm) (527	[95Kun]	· - / /	2.66–5.25 Ti:sapphire laser + Nd:YAG laser, $\theta_{\rm eoe} = 40$ °, $\varphi = 0$ °						
[μm] crystal parameters energy, power, $\tau_{\rm p}$ 11–23 Two dye lasers 3 W (peak), 30 ns (5.7–10.2 OPO (1.06–1.67 μm) + 2ω of phosphate glass laser (5.27 nm)			l: Ag ₃ AsS ₃	(f) Crystal					
3.7–10.2 OPO (1.06–1.67 μm) + 2 ω of phosphate glass laser (527 nm) (g) Crystal: AgGaS ₂ λ Sources of interacting radiations, (pm] crystal parameters energy, power, τ_p 5.5–18.3 Two dye lasers, $\theta = 90^{\circ}$ 4 W, 4 ns [75–11 Dye laser + Nd:YAG laser, $\theta_{\rm eoe} = 38$ –52 ° 180 kW, 12 ns [83.9–9.4 Dye laser + Nd:YAG laser 1%, 8 ps [88.7–11.6 Two dye lasers, $\theta_{\rm eoe} = 65$ –85 ° 0.1 mW, 500 ns [74.6–12 Two dye lasers, $\theta_{\rm eoe} = 45$ –83 ° 300 mW, 10 ns [77–9 Dye laser + Ti:sapphire laser, $\theta_{\rm eoe} = 90^{\circ}$ 1 μW, cw, [98.7–11.6 Two dye lasers, $\theta_{\rm eoe} = 45$ –83 ° 300 mW, 10 ns [77–9 Dye laser + Ti:sapphire laser, $\theta_{\rm eoe} = 90^{\circ}$ 1 μW, cw, [98.7–11.6 Two dye lasers, $\theta_{\rm eoe} = 45$ –83 ° 300 mW, 10 ns [77–9 Dye laser + Ti:sapphire laser, $\theta_{\rm eoe} = 90^{\circ}$ 1 μW, cw, [98.7–11.6 Two dye laser + Ti:sapphire laser, $\theta_{\rm eoe} = 90^{\circ}$ 1 μW, cw, [98.7–11.6 Two dye laser + Ti:sapphire laser, $\theta_{\rm eoe} = 90^{\circ}$ 1 μW, cw, [98.7–11.6 Two dye laser + Ti:sapphire laser, $\theta_{\rm eoe} = 90^{\circ}$ 1 μW, cw, [98.7–11.6 Two dye laser + Ti:sapphire laser, $\theta_{\rm eoe} = 90^{\circ}$ 1 μW, cw, [98.7–11.6 Two dye laser + Ti:sapphire laser, $\theta_{\rm eoe} = 90^{\circ}$ 1 μW, cw, [98.7–11.6 Two dye laser + Ti:sapphire laser, $\theta_{\rm eoe} = 90^{\circ}$ 1 μW, cw, [98.7–11.6 Two dye laser + Ti:sapphire laser, $\theta_{\rm eoe} = 90^{\circ}$ 1 μW, cw, [98.7–11.6 Two dye laser + Ti:sapphire laser, $\theta_{\rm eoe} = 90^{\circ}$ 1 μW, cw, [98.7–11.6 Two dye laser + Ti:sapphire laser, $\theta_{\rm eoe} = 90^{\circ}$ 1 μW, cw, [98.7–11.6 Two dye laser + Ti:sapphire laser, $\theta_{\rm eoe} = 90^{\circ}$ 1 μW, cw, [98.7–11.6 Two dye laser + Ti:sapphire laser, $\theta_{\rm eoe} = 90^{\circ}$ 1 μW, cw, [98.7–11.6 Two dye laser + Ti:sapphire laser, $\theta_{\rm eoe} = 90^{\circ}$ 1 μW, cw, [98.7–11.6 Two dye laser + Ti:sapphire laser, $\theta_{\rm eoe} = 90^{\circ}$ 1 μW, cw	Ref.	0 /							
$\lambda \qquad \text{Sources of interacting radiations,} \qquad \text{Conversion efficiency,} \qquad \text{R} \\ [\mu\text{m}] \qquad \text{crystal parameters} \qquad \qquad \text{energy, power, } \tau_{\text{p}} \\ \hline 5.5-18.3 \qquad \text{Two dye lasers, } \theta = 90 ^{\circ} \qquad \qquad 4 \text{W, 4 ns} \qquad [75-11 \qquad \text{Dye laser + Nd:YAG laser, } \theta_{\text{eoe}} = 38-52 ^{\circ} \qquad 180 \text{kW, 12 ns} \qquad [83.9-9.4 \qquad \text{Dye laser + Nd:YAG laser} \qquad \qquad 1\%, 8 \text{ps} \qquad [84-11 \qquad \text{OPO } (2-4 \mu\text{m}) + \text{radiation at } \lambda = 1.4-2.13 \mu\text{m} \qquad 1 \text{kW, 8 ns} \qquad [88.7-11.6 \qquad \text{Two dye lasers, } \theta_{\text{ooe}} = 65-85 ^{\circ} \qquad \qquad 0.1 \text{mW, 500 ns} \qquad [77-9 \qquad \text{Dye laser + Ti:sapphire laser, } \theta_{\text{ooe}} = 90 ^{\circ} \qquad \qquad 1 \mu\text{W, cw,} \qquad [97-9] \qquad \text{Dye laser + Ti:sapphire laser, } \theta_{\text{ooe}} = 90 ^{\circ} \qquad 1 \mu\text{W, cw,} \qquad [97-9] \qquad 20 \mu\text{My, cw,} \qquad [97-9] \qquad 20 \muMy, c$	[76Hoc] [80Bar1]	* / /	10.2 OPO (1.06–1.67 μ m) + 2 ω of phosphate glass laser						
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$			l: AgGaS ₂	(g) Crystal					
5–11 Dye laser + Nd:YAG laser, $\theta_{\rm coe} = 38$ –52 ° 180 kW, 12 ns 3.9–9.4 Dye laser + Nd:YAG laser 1%, 8 ps [8 4–11 OPO (2–4 μm) + radiation at $\lambda = 1.4$ –2.13 μm 1 kW, 8 ns [8 8.7–11.6 Two dye lasers, $\theta_{\rm coe} = 65$ –85 ° 0.1 mW, 500 ns [7 4.6–12 Two dye lasers, $\theta_{\rm coe} = 45$ –83 ° 300 mW, 10 ns [7 7–9 Dye laser + Ti:sapphire laser, $\theta_{\rm coe} = 90$ ° 1 μW, cw, [9 4.76–6.45 Dye laser + Ti:sapphire laser, $\theta_{\rm coe} = 90$ °, $L = 45$ mm 20 μW, cw, [9 4.76–6.45 GaAlAs laser (858 nm) + Ti:sapphire laser (715 nm), 47 μW (cw), $\theta_{\rm coe} = 90$ ° 89 μW (50 μs) 4.73; 5.12 Diode laser + Ti:sapphire laser, $\theta_{\rm coe} = 90$ ° 1 μW, cw [9 4.73; 5.12 Diode laser + Ti:sapphire laser, $\theta_{\rm coe} = 90$ ° 1 μW, cw [9 4.73; 5.12 Diode laser + Ti:sapphire laser, $\theta_{\rm coe} = 90$ ° 1 μW, cw	Ref.	• ,							
$\begin{array}{llllllllllllllllllllllllllllllllllll$	[76Sey] [84Kat] [85Els] [86Bet] [74Han] [73Han] [92Can]	180 kW, 12 ns 1%, 8 ps 1 kW, 8 ns 0.1 mW, 500 ns 300 mW, 10 ns 1 μW, cw,	Dye laser + Nd:YAG laser, $\theta_{\rm eoe} = 3852^{\circ}$ Dye laser + Nd:YAG laser OPO (2–4 µm) + radiation at $\lambda = 1.42.13\mu\text{m}$ Two dye lasers, $\theta_{\rm ooe} = 6585^{\circ}$ Two dye lasers, $\theta_{\rm ooe} = 4583^{\circ}$	5-11 3.9-9.4 4-11 8.7-11.6 4.6-12					
~4.26 GaAlAs laser (858 nm) + Ti:sapphire laser (715 nm), 47 μW (cw), $\theta_{\rm ooe}=90^\circ$ 89 μW (50 $\mu s)$ 4.73; 5.12 Diode laser + Ti:sapphire laser, $\theta_{\rm ooe}=90^\circ$ 1 μW , cw [9]	[92Hie]	$20 \mu W, cw,$	Dye laser + Ti:sapphire laser, $\theta_{\rm ooe} = 90$ °, $L = 45~\rm mm$	4.76-6.45					
	[93Sim2]	$47 \mu W (cw),$		~ 4.26					
5.2-6.4 Nd·YAG laser + near IR (DFG in LiIO ₂) $35%$ 23 ps [8]	[93Sim1]	$1 \mu W, cw$		4.73; 5.12					
	[88Spe]	35%, 23 ps	Nd:YAG laser + near IR (DFG in LiIO ₃)	5.2 – 6.4					
2.16 ps, f = 76 MHz	[91Yod]	2.16 ps, f = 76 MHz							

(continued)

Table 4.1.2	8 (g) continued.			
$\lambda \ [\mu \mathrm{m}]$	Sources of interacting radiations, crystal parameters	Conversion efficiency, energy, power, $\tau_{\rm p}$	Ref.	
4.5–11.5	Dye laser (870–1000 nm) + Ti:sapphire laser (815 nm), $\theta_{\rm c}=45^{\circ},L=1$ mm	10 nJ, $f = 1$ kHz, 400 fs	[93Ham]	
9	Ti:sapphire laser with dual wavelength output (50–70 fs), $\theta_c = 44$ °, $L = 1$ mm	$0.03~\mathrm{pJ},f=85~\mathrm{MHz}$	[93Bar1]	
3.1-4.4 $2.5-5.5$	Ti:sapphire laser + Nd:YAG laser, ICDFG, $\theta_c = 74$ ° Signal and idler pulses of OPO, $\theta = 40$ °	0.3 mW, cw 0.5 mW, f = 82 MHz, 200 fs	[95Can] [94Loh]	
6.2–9.7 6.8–12.5 2.4–12 5–12	Two Ti:sapphire lasers (696–804 nm and 766–910 nm) Two diode lasers (766–786 nm and 830–868 nm) Signal and idler waves of BBO based OPA Signal and idler waves of LiNbO $_3$ based OPO (1.8–2.7 μ m)	3 μJ , 0.08 %, 13 ns 1 μW , cw 2.5 mW, 50 fs 0.1 mJ, 6 ns	[96Aka] [98Pet1] [98Gol] [99Hai]	
~ 5	Two diode lasers, $\theta = 90$ °, $L = 30$ mm	$0.2~\mu\mathrm{W,~cw}$	[96Sch]	
(h) Crysta	l: $AgGaSe_2$			
λ [μm]	Sources of interacting radiations, crystal parameters	Conversion efficiency, energy, power, $\tau_{\rm p}$	Ref.	
7–15	OPO (1.5–1.7 µm) + Nd:YAG laser (1.32 µm), $\theta_{\rm ooe} = 9057~^\circ$	1.2%	[74Bye]	
12.2–13 8–18 5–18	CO laser (5.67–5.85 µm) + CO ₂ laser, θ = 61 ° Idler and signal waves of OPO Idler and signal waves of OPO, $\theta_{\rm ooe}$ = 51 °	$0.2~\mu W,~cw$ $0.1~mJ,~3-6~ns$ $0.2~mJ,~8~ns$	[73Kil] [93Bos] [98Abe]	
(i) Crystal	: $CdGeAs_2$			
$\lambda \ [\mu \mathrm{m}]$	Sources of interacting radiations, crystal parameters	Conversion efficiency, energy, power, $\tau_{\rm p}$	Ref.	
11.4–16.8	CO laser + CO ₂ laser	4 μW, cw	[74Kil]	
(j) Crystal	: GaSe			
$\frac{\lambda}{[\mu \mathrm{m}]}$	Sources of interacting radiations, crystal parameters	Conversion efficiency, energy, power, $\tau_{\rm p}$	Ref.	
9.5–18 4–12 7–16	Dye laser + ruby laser Idler and signal waves of OPO Nd:YAG laser + laser on F_{2-} colour centers, $\theta_{\rm ooe} = 13-15$ °, $\theta_{\rm eoe} = 12-16$ °	300 W, 20 ns 60 W 0.1–1 kW, 10 ns	[76Abd] [78Bia] [80Gus]	
6–18 5.2–18	Dye laser (1.1–1.4 μ m) + Nd:glass laser (1.053 μ m) Idler and signal waves of OPO, $L=1$ mm	10 nJ 1 μ J, 1 ps 2 mW, 3.3 %, f = 76 MHz, 120 fs	[93Dah] [98Ehr]	
(k) Crysta	l: CdSe			
λ [μm]	Sources of interacting radiations, crystal parameters	Conversion efficiency, energy, power, $\tau_{\rm p}$	Ref.	
16	OPO signal wave (1.995 μ m) + OPO idler wave (2.28 μ m), θ = 62.22 $^{\circ}$	$0.5~\mathrm{kW},20~\mathrm{Hz},10~\mathrm{ns}$	[77And2]	
9–22 10–20	OPO (2–4 μ m) + radiation at $\lambda = 1.4$ –2.13 μ m OPO signal and idler waves, $\theta = 70$ °, eoo	10–100 W, 8 ns 50 % (quantum), 5–40 μJ, 10 ps	[86Bet] [95Dhi]	

(l) Crystal: Te

$\frac{\lambda}{[\mu \mathrm{m}]}$	Sources of interacting radiations, crystal parameters	Conversion efficiency, energy, power, $\tau_{\rm p}$	Ref.
10.9–11.1	CO ₂ laser (10.2 µm) + cw spin-flip laser (5.3 µm), $\theta_{\rm eeo} = 14$ $^{\circ}$	10 μW	[75Bri]

Table 4.1.29. Difference frequency generation in the far IR region.

Pump sources	Crystal	$\nu \ [\mathrm{cm}^{-1}]$	$\lambda \; [\mathrm{mm}]$	Power, energy	Ref.
Nd:glass (1.06 μm)	LiNbO ₃	100	0.1	_	[65Zer]
Ruby laser $(0.694 \mu m)$	$LiNbO_3$	29	0.33	_	[69Yaj]
Two ruby lasers (0.694 µm), 1 MW,	Quartz,	1.2 - 8.0	1.25 - 8.33	$20~\mathrm{mW}$	[69Far]
30 ns	$LiNbO_3$				
Nd:glass $(1.06 \mu m)$, $50 mJ$, $10 ps$	ZnTe,	8-30	0.33 – 1.25	20 mW/cm^{-1}	[71Yaj]
	$LiNbO_3$				
Nd:glass $(1.06 \mu m)$, 10 ps	$LiIO_3$	_	_	_	[72Tak]
Dye laser $(0.73-0.93 \mu m)$,	ZnTe, ZnSe,	5 - 30	0.33 – 2.00	1 W (ZnTe)	[73Mat]
11–15 ns, 4–13 MW	$LiNbO_3$				
Nd:glass $(1.064 \mu m)$, 10 ps	$LiNbO_3$	0.4 – 2.5	4-25	60 W	[76Ave]
Two ruby lasers $(0.694 \mu m)$, 20 ns	$LiNbO_3$	1-3.3	3-10	$0.5~\mathrm{W}$	[79Ave]
Ruby laser $(0.694 \mu m)$	$LiNbO_3$	1.67 - 3.3	3-6	_	[80Mak]
Two dye lasers:	$LiNbO_3$	20 – 200	0.05 – 0.5	$3 \mathrm{~nJ}$	[85Ber]
$\tau_1 = 1-2 \text{ ps}, \lambda_1 = 589 \text{ nm},$					
$E_1 = 0.2 \text{ mJ}; \tau_2 = 20 \text{ ns},$					
$\lambda_2 = 590-596 \text{ nm}, E_2 = 20 \text{ mJ}$					
Nd:YAG laser (45 ps) + OPO (35 ps)	$LiNbO_3$	10 - 200	0.05 - 1	10 kW	[95Qiu]
CO ₂ laser at two frequencies	GaAs	2-100	0.1 – 5.0	_	[85Rya]
Two CO_2 lasers	$ZnGeP_2$	70 - 110	0.09 – 0.14	$1.7~\mu\mathrm{W}$	[72Boy]
Two CO_2 lasers	${\rm ZnGeP_2}$	99-100	0.1 – 0.11	$3.6~\mu J$	[96Apo]
Nd:YAG (1.064 μ m), 30 ns	GaP	0.33-1	10-30	1 mW	[87Len]

(continued)

4.1.7 Optical parametric oscillation

Table 4.1.30. Continuous wave (cw) and nanosecond OPO in the UV, visible, and near IR regions.

Crystal	$\theta_{\rm pm}$, type of interaction	$\lambda_{ m pump} \ [\mu m m]$	$I_{ m thr}$ [MW cm ⁻²]	$\lambda_{ m OPO} \ [\mu { m m}]$	$ au_{ m p}$ [ns]	η [%]	Ref.	Notes
KDP	eoe	0.532	1000-2000	_	_	40-42 a	[86Bar]	TWOPO, $L_1 = 4 \text{ cm}, L_2 = 6 \text{ cm}, E = 2 \text{ J}$
	eoe	0.35	1000	0.45 – 0.6	0.5	41 ^a	[87Beg]	TWOPO, $L_1 = 2$ cm, $L_2 = 6$ cm, $E = 0.35$ J, $I_0 = 6-8$ GW cm ⁻²
ADP	_	0.527	1500	0.93 - 1.21	_	37 ^a	[84Akh]	TWOPO, $E = 2.3 \text{ J}$, $I_0 = 10 \text{ GW cm}^{-2}$
	ooe	0.266	_	0.42 – 0.73	2	25	[71Yar]	TWOPO, $T = 50-105$ °C
	ooe	0.266	250	_	14	30	[75Zhd]	$L = 6 \text{ cm}, I_0 = 1 \text{ GW cm}^{-2}$
$LiIO_3$	$\theta_{\rm ooe} = 24$ °	1.06	50	2.5 – 3.2	40	15	[84Ash]	SROPO, $L=6$ cm, $E=0.1$ J
	$\theta_{\rm ooe} = 23.1 22.4$ $^{\circ}$	0.694	5	1.15–1.9	20	50 ^a	[71Cam, 72Cam]	DROPO, $L=0.85$ cm, $P=10$ kW
	$\theta_{ m ooe} = 2530~^\circ$	0.53	10	0.68 – 2.4	15	8	[70Izr]	SROPO, $L = 1.6$ cm
	$\theta_{\rm ooe} = 2330~^\circ$	0.532	10	0.63 – 3.35	30	20	[77Dzh]	SROPO
${\rm LiNbO_3}$	$\theta_{\rm ooe} = 90$ °	1.06	_	2.13	100	8	[69Amm]	DROPO, $L = 3 \text{ mm}$
	$\theta_{\rm ooe} = 90$ °	1.06	_	1.4 – 4.45	20	15	[74 Her]	SROPO, $I_0 = 10 \text{ MW cm}^{-2}$
	43.3 °	0.93	$8 \mathrm{mJ}$	1.48-1.8;	16	9.7	[97Raf]	SROPO, $L = 50$ mm, broad spectral bandwidth
				1.95 – 2.55				$(\Delta \lambda = 320 \text{ nm})$
	$\theta_{\rm ooe} = 90$ °	0.473 - 0.659	_	0.55 – 3.65	130–700	$46 (67^{a})$	[70Wal]	SROPO, $T = 110-430$ °C, $P_{av} = 105 \text{ mW}$
LiNbO ₃ :MgO	$\theta_{\rm ooe} = 90$ °	1.06	$0.4~\mathrm{mW}$	1-1.14	cw	_	[93Sch]	Quadruply resonant OPO
	$\theta_{\rm ooe} = 90$ °	0.532	$35~\mathrm{mW}$	1.01-1.13	cw	$40 (60^{a})$	[89Koz]	DROPO, $T = 107-110 ^{\circ}\text{C}$
	$\theta_{\rm ooe} = 90$ $^{\circ}$	0.532	$12~\mathrm{mW}$	1.007 – 1.129	cw	$34 (78^{a})$	[89Nab]	DROPO, $T = 107-111 ^{\circ}\text{C}, P = 8.15 \text{mW}$
	$\theta_{\rm ooe} = 90$ °	0.532	$13~\mathrm{mW}$	0.966 – 1.185	cw	$38 (73^{a})$	[93Ger]	DROPO, $T = 113-126$ °C, $L = 15$ mm,
	0 00 °	0.530	00 W	10 110		01	[0FD]	P = 100 mW
	$\theta = 90^{\circ}$	0.532	28 mW	1.0-1.12	cw	81	[95Bre]	DROPO, $P = 105 \text{ mW}, L = 7.5 \text{ mm}$
	$\theta_{\rm ooe} = 90$ °	0.532	80 mW	0.788 - 1.640	cw	_	[98Tsu]	DROPO, $T = 80-180 ^{\circ}\text{C}$, $L = 15 \text{mm}$

Ref. p. 187]

Crystal	$\theta_{\rm pm}$, type of interaction	$\lambda_{ m pump} \ [\mu{ m m}]$	${I_{\rm thr} \atop [{\rm MW~cm^{-2}}]}$	$\lambda_{ m OPO} \ [\mu m]$	$ au_{ m p} \ [m ns]$	$\eta \ [\%]$	Ref.	Notes
BBO	$\theta_{\rm ooe} = 21.7 - 21.9$	0.532	278	0.94-1.22	12	10	[89Fan]	SROPO, $L = 9 \text{ mm}, E = 1 \text{ mJ}$
	ooe	0.355	130	0.45 - 1.68	8	9.4	[88Che]	SROPO, $L = 11.5 \text{ mm}, E = 15 \text{ mJ}$
	$\theta_{ m ooe} = 2433$ $^{\circ}$	0.355	20	0.412 – 2.55	2.5	24	[88Fan]	SROPO, $L = 12 \text{ mm}$, $P_{\text{av}} = 140 \text{ mW}$
	ooe	0.355	27	0.42 – 2.3	8	32	[89Bos]	SROPO, $L_1 = 11.5 \text{ mm}$, $L_2 = 9.5 \text{ mm}$, $\Delta \lambda = 0.03 \text{ nm}$
	$\theta_{\rm ooe} = 33.744.4$ °	0.355	38	0.48-0.63; 0.81-1.36	8	12	[90Bos]	SROPO, $L_1 = 17 \text{ mm}$, $L_2 = 10 \text{ mm}$, $\Delta \lambda = 0.05-0.3 \text{ nm}$
	$\theta_{ m ooe} = 2333~^\circ$	0.355	20-40	0.402 – 3.036	7	40-61	[91Fix, 93Fix]	SROPO, $L = 15$ mm, $E = 0.1-0.2$
	$\theta=28^\circ$	0.355	_	0.453 – 2.3	6	7–9	[95Joh]	SROPO, $\Delta \nu = 0.2 \text{ cm}^{-1}$, $E = 100 \text{ mJ}$, SHG i KDP and BBO (220–450 nm)
	$\theta=2333^\circ$	0.355	20	0.465 - 1.5	10	40	[94Glo]	,
	$\theta = 33^{\circ}$	0.355	$3.2~\mathrm{mJ}$	0.44 – 1.76	10	37	[97Oie]	SROPO, $L = 12$ mm, collinear and noncollinear geometries
	$\theta = 35.9^{\circ}$	0.355	-	0.5 – 0.7	10	_	[97Wan]	Broad spectral bandwidth OPO ($\Delta \lambda > 100$ nm with noncollinear geometry, $L = 18$ mm
	$\theta_{ m ooe} = 35.5 37~^\circ$	0.308	150	0.422 – 0.477	8	10	[88Kom]	
	ooe	0.308	18	0.354 - 2.37	17	64^{a}	[91Rob]	
	ooe	0.308	-	0.4 – 0.56	17	15	[93Rob]	SROPO, $L = 20 \text{ mm}$, $\Delta \nu = 0.07 \text{ cm}^{-1}$ (wit intracavity etalon)
	$\theta_{\mathrm{ooe}} = 3048$ $^{\circ}$	0.266	_	0.302 - 2.248	7	6.3	[91Fix]	SROPO
	$\theta_{ m ooe} = 38.3$ $^{\circ}$	0.266	58	0.3 – 2.34	4.5	15	[00Kon]	$L=14~\mathrm{mm}$
LBO	$\theta=90^{\circ},\varphi=0^{\circ}$	0.78 - 0.81	$360~\mathrm{mW}$	1.49 – 1.70	cw	40^{a}	[94Col1]	DROPO, $L = 2 \text{ cm}, T = 130185 ^{\circ}\text{C}, P = 30 \text{mV}$
	$\theta = 90^{\circ}, \varphi = 0^{\circ}$	0.5235	700	0.924 – 1.208	12	45	[93Hal2]	DROPO, $L = 12 \text{ mm}, T = 156-166 ^{\circ}\text{C}$
	$\theta = 90^{\circ}, \varphi = 0^{\circ}$	0.5145	50 mW	0.966-1.105	cw	10	[93Col1]	TROPO, $L = 20 \text{ mm}$, $T = (183 \pm 3) ^{\circ}\text{C}$, $P = 90 \text{ mW}$
	$\theta=0^{\circ}, \varphi=0^{\circ}$	0.5145	$1 \mathrm{~W}$	0.93 – 0.946	cw	15	[94Rob2]	SROPO, $P = 0.5 \text{ W}, L = 25 \text{ mm}$
	$\theta = 0^{\circ}, \ \varphi = 90^{\circ}$	0.364	115 mW	0.494-0.502; 1.32-1.38	cw	9.4	[93Col2, 94Col2]	SROPO and DROPO, $L=20$ mm, $T=18-86$ °C $P=103$ mW
	$\theta=90^{\circ},$ $\varphi=2442^{\circ}$	0.355	14	0.435 - 1.922	10	22		DROPO, $I_0 = 40 \text{ MW cm}^{-2}$, $E = 2.7 \text{ mJ}$

Table 4.1.30 continued.

Crystal	$\theta_{\rm pm}$, type of interaction	$\lambda_{ m pump} \ [\mu { m m}]$	$I_{ m thr} \ [{ m MW~cm^{-2}}]$	$\lambda_{ m OPO} \ [\mu { m m}]$	$ au_{ m p} \ [m ns]$	η [%]	Ref.	Notes
LBO	$\theta = 0^{\circ}, \varphi = 0^{\circ}$	0.355	15	0.48-0.457; 1.355-1.59	12	27	[92Cui]	SROPO, $T = 20-200$ °C
	$ heta=90^\circ, \ arphi=2742^\circ$	0.355	60	$0.455-0.655; \ 0.76-1.62$	10	35 ^a	[93Cui]	SROPO, $L = 16 \text{ mm}$
	$\theta=90^{\circ},$ $\varphi=20.1$ –42.1 $^{\circ}$	0.355	50	0.414 - 2.47	5	45	[94Sch]	SROPO, $L = 15 \text{ mm}$
	$\theta = 90^{\circ},$ $\varphi = 26-52^{\circ}$	0.308	26	$0.355-0.497; \ 0.809-2.34$	17	$28-40^{\text{ a}}$	[91Rob, 92Rob]	SROPO, $L = 15 \text{ mm}$
	type II in XZ and YZ planes, $\theta = 0-9$ °	0.308	30	0.381–0.387; 1.5–1.6	5	35	[91Ebr2]	$L = 16 \text{ mm}, I_0 = 0.1 \text{ GW cm}^{-2}$
	$\theta = 0^{\circ}, \varphi = 0^{\circ}$	0.266	10	0.314; 1.74	10	10	[92Tan]	SROPO, $L = 16 \text{ mm}, T = 20 ^{\circ}\text{C}$
	$\theta = 90^{\circ},$ $\varphi = 37-47^{\circ}$	0.266	-	0.307-0.325	4	_	[94Sch]	SROPO, $L = 15 \text{ mm}$
KTP	$\theta = 5058$ °, $\varphi = 0$ °	1.064	_	1.8 – 2.4	10	10	[90Lin1]	DROPO, $E = 0.1$ –0.5 mJ
	$\theta = 90^{\circ}, \varphi = 53^{\circ}$	1.064	80	3.2	10	5	[91Kat]	SROPO, $L = 15 \text{ mm}, P = 0.2 \text{ W}$
	$\theta = 90^{\circ}, \varphi = 0^{\circ}$	1.06	_	1.61	15	47 (66 a)) [93Mar1]	Diode-pumped Nd:YAG laser
	_	1.047	$0.5~\mathrm{mJ}$	$1.54;\ 3.28$	18	20	[94Ter]	
	$\theta = 63.4^{\circ}, \varphi = 0^{\circ}$		$0.6~\mathrm{mJ}$	1.58 – 1.84	10	40	[97Tan]	NC SROPO, $L = 25 \text{ mm}$
	$\theta = 90^{\circ}, \varphi = 0^{\circ}$	0.7 – 0.95	70	1.04-1.38; $2.15-3.09$	10	20	[92Kat]	SROPO, $L = 15 \text{ mm}$
	$\theta = 90^{\circ}$	0.7 – 0.9	_	1.03–1.28; 2.18–3.03	20	55	[94Zen]	$E=49~\mathrm{mJ},L=15~\mathrm{mm}$
	$\theta = 90^{\circ}, \varphi = 0^{\circ}$	0.769	$6~\mathrm{mW}$	1.1; 2.54	cw	_	[95Sch]	TROPO, $L = 12 \text{ mm}$
	$\theta = 54^{\circ}, \varphi = 0^{\circ}$		_	1.38 - 1.67	cw	0.001		$L = 10 \text{ mm}, P = 2 \mu\text{W}$
	$\theta = 90^{\circ}, \varphi = 0^{\circ}$		1.4 W, SROPO; 30 mW, DROPO	1.039; 1.09	cw	35		SROPO and DROPO, $L = 10$ mm, $P = 1.07$ W

Crystal	$\theta_{\rm pm}$, type of interaction	$\lambda_{ m pump} \ [\mu { m m}]$	${I_{\rm thr} \atop [{\rm MW~cm^{-2}}]}$	$\lambda_{ m OPO} \ [\mu { m m}]$	$ au_{ m p} \ [m ns]$	η [%]	Ref.	Notes
KTP	$\varphi = 0$ °	0.532	80	0.7-0.9; 1.3-2.2	3.5	12	[93Bos, 92Bos]	$L = 15 \text{ mm}, E = 3 \text{ mJ}, \Delta \nu = 0.02 \text{ cm}^{-1}$
	$\theta=69^{\circ},\varphi=0^{\circ}$	0.532	_	0.75 - 1.04	4-6	27	[95Sri]	OPO-OPA, $L_1 = L_2 = 10 \text{ mm}, E = 45 \text{ mJ}$
	$\theta = 60^{\circ}, \varphi = 0^{\circ}$	0.532	_	0.75-0.87; 1.83-1.37	4	_	[95Hui]	$\Delta \nu = 200$ MHz (with Fabry-Perot etalon), E = 0.6 mJ, $L = 16$ mm
	$\theta = 90^{\circ}, \varphi = 0^{\circ}$	0.532	$4.3~\mathrm{W}$	1.09; 1.039	cw	28 (64 a)) [94Yan]	SROPO, $P = 1.9 \text{ W}$, $L = 15 \text{ mm}$
	$\theta = 90^{\circ},$ $\varphi = 25.3^{\circ}$	0.531	$40~\mathrm{mW}$	1.0617	cw	30	[93Lee]	DROPO, $L = 8 \text{ mm}$
	$\theta=69^{\circ},\varphi=0^{\circ}$	0.532	7	0.76 – 1.04	6	30	[93Mar2]	$L=15$ mm, ICSHG in BBO with $\eta=40\%$ (380–520 nm)
KTA	$\theta = 53^{\circ}, \varphi = 0^{\circ}$	0.773 – 0.792	_	1.45; 1.7	300	0.3	[92Jan]	DROPO, $L = 7 \text{ mm}$
	type II	_	_	1.11 - 1.20;	cw	90	[98Edw]	Intracavity (Ti:Sa) SROPO, $P = 1.46$ W,
				2.44 - 2.86				L = 11.5 mm
RTA	$\theta = 90^{\circ}, \varphi = 0^{\circ}$	0.77-0.83	$70~\mathrm{mW}$	1.21-1.26; $2.1-2.4$	cw	_	[97Sch]	SROPO, $L=12$ mm, $P=84$ mW, $\Delta\nu<10$ MHz
Banana	$\theta_{\rm ooe} = 90$ $^{\circ}$	0.532	_	0.75 – 1.82	10	5	[80Bar3]	SROPO, $T = 80220 ^{\circ}\text{C}$
KNbO_3	along the b axis	0.532	3.5	0.88 – 1.35	10	32	[82Kat]	DROPO, $T=180200\ ^{\circ}\mathrm{C},P=12\ \mathrm{MW}$
Urea	$\theta_{\mathrm{oeo}} = 8190$ °	0.355	55 (45 mW)	0.5-0.51; $1.17-1.22$	7	20	[84Don]	SROPO, $L = 12.7 \text{ mm}, I_0 = 90 \text{ MW cm}^{-2}$
	$\theta_{ m oeo} = 50$ –90 $^{\circ}$	0.355	_	0.5 – 1.23	7	23	[85Ros2, 85Ros1]	SROPO, $L = 23 \text{ mm}$
	$\theta_{\rm oeo} =$ 64–90 $^{\circ}$	0.308	16-20	0.537 – 0.72	4-6	37		$L=15~\mathrm{mm}$
NPP	eeo	0.266	_	0.33 – 0.42	7	_	[85Ros1]	
	$\theta = 9.5 13^{\circ},$	0.5927	30	0.9 – 1.7	1	5	[92Jos,	L = 1.9 mm
	$\varphi = 0$ ° $\theta = 30$ °	0.583-0.59	0.5	1-1.5	7	_	93Dou] [95Kho]	

^a Pump depletion.

Table 4.1.31. Picosecond OPO in the UV, visible, and near IR regions.

Crystal	$\theta_{\rm pm}$, type of interaction	$\lambda_{ m pump} \ [\mu { m m}]$	$I_{ m thr}$ [MW cm ⁻²]	$\lambda_{ m OPO} \ [\mu { m m}]$	$ au_{ m p} \ [m ps]$	η [%]	Ref. Notes
KDP	eoe	0.532	_	0.8 – 1.67	40	25	[78Kry] TWOPO, $E = 1 \text{ mJ}$, $L_1 = L_2 = 4 \text{ cm}$
	eoe	0.532	-	0.9 – 1.3	30	51	[78Dan2, TWOPO, $\Delta \nu \Delta \tau = 0.7$, $L_1 = 4$ cm, $L_2 = 6$ cm 79Kab] $I_0 = 15-20$ GW cm ⁻²
	eoe	0.527	_	0.82 - 1.3	0.3 – 0.5	2	[83Bar, SP OPO, $E = 20 \mu J$ 82Dan]
	eoe	0.355	-	$0.45 - 0.64, \\ 0.79 - 1.69$	45	15	[78Dan3] TWOPO, $L_1 = L_2 = 4 \text{ cm}$
ADP	$\theta_{\rm ooe} = 90$ $^{\circ}$	0.266	_	0.44 – 0.68	10	10	[76Mas2] TWOPO, $T=50$ –110 °C, $L_1=L_2=5$ cm
CDA	$\theta_{\rm ooe} = 90$ °	0.532	_	0.854-1.41	10	30–60	[74Mas] $L = 3$ cm, $T = 50-70$ °C, $I_0 = 0.3$ GW cm ⁻²
	$\theta_{\rm ooe} = 90$ °	0.53	1000	0.8 – 1.3	10	12.5	[87Ion] SP OPO, $L = 4 \text{ cm}$, $I_0 = 3 \text{ GW cm}^{-2}$
$LiIO_3$	ooe	0.532	_	0.61 – 4.25	6	4	[77Dan, TWOPO, $L_1 = 1$ cm, $L_2 = 2.5$ cm, 78Dan1] $I_0 = 2$ GW cm ⁻²
	$\theta_{\rm ooe} = 2530~^\circ$	0.53	3000	0.68 – 2.4	_	5	[77Kry] TWOPO, $L_1 = L_2 = 4 \text{ cm}$, $I_0 = 6 \text{ GW cm}^{-2}$
${\rm LiNbO_3}$	ooe	1.06	_	1.4-4.0	3.5	10	[78Sei] TWOPO, $\Delta \nu = 6.5 \text{ cm}^{-1}$, $I_0 = 1 \text{ GW cm}^{-2}$
	4551 $^{\circ}$	1.064	_	1.37 - 4.83	40	17	[77Iva] TWOPO
	$47~^{\circ}$	1.054	100	1.35 – 2.11	0.5	15	[90Lae1] SP OPO, $L = 18 \text{ mm}$, $I_0 = 0.14 \text{ GW cm}^{-2}$
	84 °	0.53	_	0.66 – 2.7	40	17	[77Iva] TWOPO, $T = 46-360$ °C
	90 °	0.532	_	0.68 – 0.76	20	9	[79Liu] SP OPO
	90 °	0.532	8	0.85 – 1.4	15	17.5	[86Pis] SROPO, $P = 30 \text{ kW}, f = 10 \text{ kHz}$
	90 °	0.532	< 30	0.65 – 3.0	10	7.2	[87Ion] SP OPO, $L = 25 \text{ mm}$
LiNbO ₃ :MgO		0.75 – 0.84	4000	2.6 – 4.5	2-3	18	[96Lin] $L_1 = L_2 = 20 \text{ mm}$
	$\theta_{\rm ooe} = 60$ –84 $^{\circ}$	0.532	_	0.7 – 2.2	30	5.4	[91He] TWOPO, $\Delta \lambda = 0.3$ nm (0.7 μ m) and 1.4 nm (2 μ m)

Table 4.1.31 continued.

Crystal	$\theta_{\rm pm}$, type of interaction	$\lambda_{ m pump} \ [\mu { m m}]$	${I_{\rm thr} \atop [{\rm MW~cm}^{-2}]}$	$\lambda_{ m OPO} \ [\mu { m m}]$	$ au_{ m p} \ [m ps]$	η [%]	Ref.	Notes
BBO	$\theta_{\rm ooe}=20.722.8^\circ$	0.532	_	0.67 - 2.58	18	13	[92Zhu]	TWOPO, $L_1 = L_2 = 9 \text{ mm}$, $I_0 = 2.5 - 3.8 \text{ GW cm}^{-2}$, $E = 0.1 - 0.5 \text{ mJ}$
	ooe	0.53	_	0.63 – 3.2	1.3	25	[93Dan]	TWOPO-OPA, $L_1 = L_2 = 8 \text{ mm}$
	ooe	0.36	500	0.406 – 3.17	20	30	[90Bur]	SP OPO, $L = 12 \text{ mm}$, $I_0 = 2 \text{ GW cm}^{-2}$, $E = 3 \text{ mJ}$, $\Delta \lambda = 0.24 \text{ nm}$
	$\theta_{ m ooe} = 2633$ $^{\circ}$	0.355	_	0.4 – 2.0	15	30	[90Hua]	OPO-OPA, $L_1 = 12$ mm, $L_2 = 6$ mm, $L_3 = 15$ mm, $I_0 = 3$ GW cm ⁻² , $\Delta \lambda = 0.3$ nm
	ooe	0.355	_	0.4 – 2.86	24	6.5	[90Suk]	TWOPO, $L_1 = L_2 = L_3 = 8 \text{ mm},$ $I_0 = 5 \text{ GW cm}^{-2}, \Delta \nu = 10 \text{ cm}^{-1}$
	ooe	0.355	_	0.43 – 2.1	15	30	[93Zha1]	Injection seeding, $L = 15 \text{ mm}$
	$\theta_{ m ooe} = 33$ $^{\circ}$	0.3547	-	0.42 – 2.8	30	61	[94Hua]	OPO-OPA, $P = 51 \text{ MW}$
LBO	ooe, $\theta = 90^{\circ}$, $\varphi = 0^{\circ}$	0.8	700 (400 mW)	1.15 – 2.26	1 - 2.2	27 (48 ^a)	[95Ebr1]	SP OPO, $P=325$ mW, $L=30$ mm, $T=120-230~^{\circ}\mathrm{C}$
	ooe, $\theta = 90^{\circ}$, $\varphi = 0^{\circ}$	0.8	320	1.374–1.53; 1.676–1.828	0.52	7.5	[95Ebr3]	SP OPO, $P = 90$ mW, $L = 16$ mm
	type I	0.77 – 0.8	350	1.16 - 2.26	1	34	[96Fre]	SP SROPO, $P = 580 \text{ mW}$, $L = 16 \text{ mm}$
	$\theta=81^{\circ},\varphi=5^{\circ}$	0.57 – 0.63	_	1.2 – 1.5	0.58	10	[91Bay]	Injection seeding by 1.08 μm
	$\theta=85^{\circ},\varphi=9^{\circ}$	0.57 – 0.63	_	1.2 – 1.5	0.4	25	[92Akh]	Injection seeding by 1.08 μ m (40 ps), $L = 9$ mm $I_0 = 1$ TW cm ⁻²
	$\theta = 90^{\circ}, \varphi = 0^{\circ}$	0.532	_	0.75 - 1.8	35	20	[91Hua]	Injection seeding OPO, $T=106.5$ –148.5 °C
	$\theta = 90^{\circ}$, ooe	0.53	_	0.65 – 2.5	15	24	[91Lin]	OPA, angle ($\varphi = 8.715.9^{\circ}$) and temperature tuning ($T = 103210^{\circ}\text{C}$), $E = 0.45 \text{ mJ}$
	$\theta = 90^{\circ}, \varphi = 0^{\circ}$	0.532	1500	0.77 – 1.7	100	30	[93Zho]	SP SROPO, $L=15$ mm, $T=105$ –137 °C, $\Delta \lambda = 0.14$ nm
	ooe, $\theta = 90^{\circ}$, $\varphi = 11.5^{\circ}$	0.532	_	0.68 – 2.44	25	8	[95Liu]	$L_1 = L_2 = 15 \text{ mm}$
	$\theta = 0^{\circ}, \varphi = 0^{\circ}$	0.532	_	0.75 - 1.8	15	20	[95Wal]	P = 200 mW, L = 15 mm
	$\theta = 90^{\circ}, \varphi = 0^{\circ}$	0.5235	2500 (10 mW)	0.652 - 2.65	12	13	[92Ebr2, 93Ebr, 93Hal1]	SROPO, $L=12$ mm, $T=125$ –190 °C

(continued)

Table 4.1.31 continued.

Crystal	$\theta_{\rm pm}$, type of interaction	$\lambda_{ m pump} \ [\mu{ m m}]$	$I_{\rm thr}$ [MW cm ⁻²]	$\lambda_{ m OPO} \ [\mu { m m}]$	$ au_{ m p} \ [m ps]$	η [%]	Ref.	Notes
LBO	$\theta = 90^{\circ}, \varphi = 0^{\circ}$	0.5235	1100 (4.5 mW)	0.909-1.235	33	50	[93Hal1, 92Ebr1]	DROPO, $T = 167-180 ^{\circ}\text{C}$
	$\theta = 90^{\circ}, \varphi = 0^{\circ}$	0.5235	15 (30 mW)	0.65 – 2.65	1.7	50	[93Hal1]	DROPO, $L = 12$ mm, $P = 0.21$ W
	ooe, $\theta = 90^{\circ}$, $\varphi = 0^{\circ}$	0.5235	$47~\mathrm{mW}$	0.839 - 1.392	1.8	70 ^a	[94Rob1]	SP SROPO, $P = 88$ mW, $L = 3$ mm
	ooe, $\theta = 90^{\circ}$, $\varphi = 0^{\circ}$	0.5235	$170~\mathrm{mW}$	0.65 – 2.7	1.63	75 ^a	[95But]	SP OPO, $P=210$ mW, $L=15$ mm
	$\theta = 90^{\circ}, \varphi = 0^{\circ}$	0.523	100	0.72 – 1.91	1	34	[93McC1] SP SROPO, $L=13$ mm, $T=125$ –175 °C, $P_{\rm av}=89$ mW
	$\theta = 90^{\circ}, \varphi = 0^{\circ}$	0.523	80 (70 mW)	0.8 – 1.5	1.2 – 1.5	27 (75 a)) [93But]	SROPO, $L = 12 \text{ mm}, P_{\text{av}} = 78 \text{ mW}$
	$\theta = 90$ °, $\varphi = 27-43$ °	0.355	_ ` ,	0.46 – 1.6	15	30	[91Zha]	Injection seeding from OPO, $L = 16$ mm, $I_0 = 2.8$ GW cm ⁻² , $E = 0.3$ mJ
	$\overset{\cdot}{\theta} = 90^{\circ},$ $\varphi = 18-42^{\circ}$	0.355	_	0.403 – 2.58	12	28	[92Kra]	TWOPO, $L_1 = L_2 = 15 \text{ mm}$, $I_0 = 5 \text{ GW cm}^{-2}$, $E = 0.1-1 \text{ mJ}$
	$\overset{\cdot}{\theta} = 0^{\circ}, \varphi = 0^{\circ}$	0.355	2300	0.4159 - 0.4826	30	38	[92Hua, 93Hua]	TWOPO, $L = 10 \text{ mm}$, $T = 21-450 ^{\circ}\text{C}$, $I_0 = 18 \text{ GW cm}^{-2}$, $\Delta \lambda = 0.15 \text{ nm}$
	$\theta = 90^{\circ},$ $\varphi = 3042^{\circ}$	0.355	1000	0.452 - 1.65	9	26	[93Agn]	DROPO, $L = 10.5$ mm, $E = 0.15$ mJ
KTP	$\theta = 82 90 ^{\circ},$ $\varphi = 0 ^{\circ}$	1.064	0.8 W	1.57–1.59; 3.21–3.30	2-3	15	[93Chu]	SROPO, $L=10$ mm, $f=75$ MHz, $\Delta \lambda = 1.5$ nm
	_	1.064		2.128	100	25	[93Lot]	SP OPO with 6 KTP (total length 58 mm), $P = 14 \text{ W}$
	$\theta=81$ –90 °, $\varphi=0$ °	1.053	5.8 W	1.55–1.56; 3.22–3.28	12	21	[93Gra]	SP OPO, $L=6$ mm, $P=2$ W
	$\theta = 40.6 \text{-} 45.2^{\circ},$ $\varphi = 0^{\circ}, \text{ oeo}$	0.8	_	1.02-1.16; $2.6-3.7$	2.6	_	[95Gra]	OPA, $E = 0.04 \text{ mJ}$
	$\theta = 53^{\circ}, \varphi = 0^{\circ},$ oeo	0.72 – 0.85	0.8 W	1.44–1.64	1.1	20	[96Qia]	SP SROPO, $P=200$ mW, $L=7$ mm

Crystal	$\theta_{\rm pm}$, type of interaction	$\lambda_{ m pump} \ [\mu{ m m}]$	$I_{ m thr} \ [{ m MW~cm^{-2}}]$	$\lambda_{ m OPO} \ [\mu { m m}]$	$ au_{ m p} \ [m ps]$	η [%]	Ref.	Notes
KTP	$\theta = 90^{\circ}, \varphi = 0^{\circ}$	0.72 – 0.853	150	1.052–1.214; 2.286–2.871	1.2	42	[93Neb]	SP OPO, $L = 6$ mm, $P = 0.7$ W
	$\theta = 54^{\circ}, \varphi = 0^{\circ},$	0.532	250	0.614-4.16	0.39	12	[94Umb]	$L=14~\mathrm{mm}$
	eoe, $\theta = 90^{\circ}$, $\varphi = 35^{\circ}$	0.527	0.9 W	0.851-0.938; 1.2-1.381	2.4 – 3.2	13	[95Che]	SROPO, $P=80280$ mW, $L=5$ mm
	$\overset{\circ}{\theta} = 40 70 \overset{\circ}{\circ},$ $\varphi = 90 \overset{\circ}{\circ}$	0.526	_	0.6 – 2.0	30	10	[88Van]	$L=20~\mathrm{mm}$
	$\theta = 40-80^{\circ},$ $\varphi = 0^{\circ}$	0.526	_	0.6 – 4.3	30	10	[88Van]	$L=20~\mathrm{mm}$
	$\theta = 90^{\circ},$ $\varphi = 10-35^{\circ}$	0.523	57 (61 mW)	1.002 – 1.096	2.2	16 (79 a)) [92McC]	SP OPO, $L = 5$ mm, $P = 42$ mW
	_	0.5235	1000 (2 mW)	0.946–1.02; 1.075–1.172	8	10 (56 a) [91Ebr1, 93Hal1]	SP SROPO, $L=5$ mm, $P=2$ mW
	$\theta = 90^{\circ}$	0.523	, ,	0.938 - 1.184	1-2	16		SROPO, $L = 5$ mm, $f = 125$ MHz, $P = 40$ mW
	$\theta=90^{\circ},$ $\varphi=0$ –33 $^{\circ}$	0.526	0.5 W	1.01-1.1	14	44		SP OPO, $L = 6 \text{ mm}, P = 0.58 \text{ W}$
KTA	$\theta=90^\circ,\varphi=0^\circ$	1.064	_	1.54; 3.47	7	75	[98Ruf]	SP OPO, $L=15~\mathrm{mm}$
Banana	$\theta_{\rm ooe} = 90$ °	0.532	5	0.8 – 1.6	10	25	[83Oni]	SP OPO
	$\theta_{\rm ooe} = 90$ °	0.53	50	0.65 - 3	10	5.3	[87Ion]	SP OPO, $I_0 = 250 \text{ MW cm}^{-2}$
	$\theta_{ m ooe} = 90$ °	0.532	7–9	0.672 - 2.56	15–45	8.1	[89Pis, 90Pis]	SP SROPO, $L=10$ mm, $f=139$ MHz, $T=75350~^{\circ}\text{C}$
$lpha$ –HIO $_3$	eoe eoe	$0.532 \\ 0.527$	- 60	0.7 – 2.2	30–45 5–6	10–12 10	[77Dan] [80Bar2]	TWOPO, $L_1=L_2=2$ cm, $I_0=4$ –5 GW cm $^{-2}$ SP OPO, $\Delta \nu \Delta \tau=0.7$

^a Pump depletion.

Table 4.1.32. Femtosecond OPO in the UV, visible, and near IR regions.

Crystal	$\theta_{\rm pm}$, type of interaction	$\lambda_{ m pump} \ [\mu { m m}]$	$I_{ m thr}$ [MW cm ⁻²]	$\lambda_{ m OPO} \ [\mu { m m}]$	$ au_{ m p}$ [fs]	η [%]	Ref.	Notes
BBO	$\theta_{\rm eoe} = 28$ °	0.78	_	1.1-2.6	60	35	[94Nis]	TWOPO, $L_1 = L_2 = 4.8 \text{ mm}, E = 0.15 \text{ mJ}$
	$\theta_{ m ooe} = 20$ $^{\circ}$	0.8	_	1.2 – 1.3	70	5	[94Sei1]	TWOPO, $L = 4 \text{ mm}$
	ooe	0.62	_	0.45 – 2.8	200	15	[91Joo]	TWOPO, $L_1 = 5 \text{ mm}, L_2 = 7 \text{ mm}, E = 20 \mu\text{J}$
	ooe, eoe	0.6	_	0.75 – 3.1	180-250	23	[93Dan]	TWOPO-OPA, $L_1 = L_2 = 8 \text{ mm}$, $I_0 = 70 \text{ GW cm}^{-2}$
	$\theta_{ m ooe} =$ 19.5–21 $^{\circ}$	0.53	2200	0.68 – 2.4	75	30	[88Bro]	SP SROPO, $L = 7.2 \text{ mm}$, $I_0 = 2.2 \text{ GW cm}^{-2}$, $E = 2 \text{ mJ}$
	006	0.527	_	0.7–1.8	65–260	3	[90Lae2, 91Lae, 93Lae]	SP SROPO, $L = 5.8 \text{ mm}$
	ooe	0.527	_	1.04 – 1.07	70	_		OPA with gain ratio 2×10^4
	$\theta_{ m ooe} = 32$ $^{\circ}$	0.4	_	0.566 - 0.676	30	10	[94Dri]	SP OPO, $P = 100 \text{ mW}$
	ooe	0.4	_	0.59 – 0.666	13	50	[95Gal2]	SP OPO, $P = 130 \text{ mW}$
	ooe	0.395	_	0.55 – 0.69	14	_	[98Shi]	NC OPA, seeding with white light continuum
	$\theta_{\rm ooe} = 32$ °	0.39	=	0.5 – 0.7	11	_	[97Cer]	OPA, seeding with white light continuum, $L = 1 \text{ mm}$
LBO	_	0.8	_	1.1-2.4	40	38	[95Kaf]	SP OPO, $L = 6$ mm, $P = 550$ mW
	-	0.77 – 0.8	320	1.374–1.530; 1.676–1.828	720	7.5	[95Ebr2]	SP OPO, $P = 90 \text{ mW}$
	$\theta = 90^{\circ}, \varphi = 0^{\circ}$	0.605	_	0.85 - 0.97; 1.6 - 2.1	200	10–15		TWOPO, $L_1 = L_2 = L_3 = 15 \text{ mm}$, $T = 30-85 ^{\circ}\text{C}$, $I_0 = 25 \text{ GW cm}^{-2}$
KTP	$\theta = 43^{\circ}, \ \varphi = 0^{\circ},$	0.83	$325~\mathrm{mW}$	1.05–1.16; 2.9–4.0	175	15 ^a	[95McC]	SP OPO, $L=2 \text{ mm}$
	$\theta = 90^{\circ}, \ \varphi = 0^{\circ},$	0.816	_	2.5–2.9	160	55 ^a	[95Hol]	SP OPO-OPA, $L_1 = L_2 = 0.9 \text{ mm}, E = 0.55 \mu\text{J}$
	$\varphi=0$ °	0.765-0.815	_	$1.22-1.37; \\ 1.82-2.15$	57-135	55 ^a	[92Pel]	L=1.15 mm, $f=90$ MHz, $P=340$ mW (135 fs) and 115 mW (57 fs)

Table 4.1.32 continued.

Crystal	$\theta_{\rm pm}$, type of interaction	$\lambda_{ m pump} \ [\mu { m m}]$	$I_{\rm thr}$ [MW cm ⁻²]	$\lambda_{\mathrm{OPO}} \ [\mu\mathrm{m}]$	$ au_{ m p} \ [m fs]$	η [%]	Ref.	Notes
KTP	$\theta=67^{\circ},\varphi=0^{\circ}$	0.765	40000; (180 mW)	1.2–1.34; 1.78–2.1	62	_	[92Fu]	$\mathrm{SP}\ \mathrm{OPO}, L = 1.5\ \mathrm{mm}, f = 76\ \mathrm{MHz}, P = 175\ \mathrm{mW}$
	_	0.745	100 mW	0.53 – 0.585	200	29	[97Kar]	SP OPO with ICSHG (self-doubling OPO)
	$\theta=45^{\circ},\varphi=0^{\circ}$	0.68	_	1.16-2.2; $0.58-0.657$	57	60 ^a	[93Pow1]	$L = 1.5$ mm, $P = 0.68$ W, ICSHG in BBO ($L = 47 \mu \text{m}$)
	$\varphi = 0$ $^{\circ}$	0.645	$110~\mathrm{mW}$	1.2 – 1.34	220	13	[92Mak]	SP OPO, P = 30 mW
	$\theta = 53^{\circ}, \varphi = 0^{\circ}$	0.61	_	0.755-1.04; 1.5-3.2	105–120	_	[90Wac, 91Wac]	SP OPO in CPM dye laser cavity, $L=1.4~\mathrm{mm}$
	$\theta = 62^{\circ}, \varphi = 0^{\circ},$ eoo	0.524	2000	1.2 - 1.6	260	10	[95Rau]	SP OPO, $L = 3 \text{ mm}$
	$\theta=62^\circ,\varphi=0^\circ$	0.5235	_	1.2 – 1.7	300	_	[98Lae]	SP OPO, $L=6$ mm, $E=10$ nJ
KTA	$\varphi=0$ °, oeo	0.78	_	1.29–1.44; 1.83–1.91	85-150	10-15	[93Pow2]	$L=1.47~\mathrm{mm},P=75~\mathrm{mW}$
RTA	$\theta = 53^{\circ}, \varphi = 0^{\circ},$ eoo	0.76 – 0.82	_	1.03–1.3; 2.15–3.65	58	25	[94Pow]	SP SROPO, $L=1.8~\mathrm{mm},P=250~\mathrm{mW}$
	$\theta = 90^{\circ}, \varphi = 0^{\circ},$	0.78 – 0.86	$50~\mathrm{mW}$	1.33	70	32	[95Rei]	$L=2~\mathrm{mm},P=185~\mathrm{mW}$
	_	Ti:Sa	-	$1.25;\ 2.25$	78	33	[97Rei]	SP OPO, $f=344~\mathrm{MHz},P=0.6~\mathrm{W}$
KNbO_3	$\theta = 38^{\circ}, \varphi = 90^{\circ}$	0.78	_	2.3 – 5.2	60-90	23	[95Spe, 96Spe]	$L=1~\mathrm{mm},~P=170300~\mathrm{mW}$
NPP	_	0.62	_	0.8 – 1.6	150-290	_	[86Led, 87Led]	$L=1.5~\mathrm{mm}$

^a Pump depletion.

 ${\bf Table~4.1.33.~Optical~parametric~oscillation~in~the~mid~IR~region.}$

Crystal	$\lambda_{ m pump} \ [\mu { m m}]$	λ _{OPO} [μm]	$ au_{ m p}$	Conversion efficiency [%]	Ref.
$\overline{\mathrm{Ag_3AsS_3}}$	1.065	1.82-2.56	26 ns	1	[72Han]
	1.064	1.2-8	8 ps	0.01-1	[83Els]
$AgGaS_2$	1.064	1.2–10	8 ps	0.1 – 10	[84Els]
	1.06	1.4 – 4.0	18 ns	16	[84Fan]
	1.064	4.5 - 8.7	15-20 ps	5.4	[91Bak]
	1.064	1.16 – 12.9	19 ps	25	[93Kra]
	1.064	1.319; 5.505	45–80 ps	63 ^a	[94Che]
	1.047	2.6-7	0.5 - 2.6 ps	_	[98Lae]
	0.845	1.267; 2.535	cw	2	[98Dou]
	0.74 – 0.85	3.3–10	160 fs	20	[94Sei2]
$AgGaSe_2$	2.05	2.65 - 9.02	30 ns	> 18	[86Eck]
J	2.06	~ 4.1	$\sim 30~\mathrm{ns}$	23	[93Bud]
	1.57	6-14	6 ns	20	[97Cha]
	1.34	1.6 - 1.7; 6.7 - 6.9	30 ns	> 18	[86Eck]
${\rm ZnGeP_2}$	2.94	5.51–5.38; 6.29–6.46	80 ps	5.3	[85Vod]
	2.94	5-5.3; 5.9-6.3	150 ps	17	[87Vod]
	2.79	5.3; 5.9	$\sim 100 \mathrm{\ ps}$	10	[93Vod2]
	2.8; 2.94	4–10	$\sim 100 \text{ ps}$	1–18	[91Vod, 93Vod1, 95Vod2]
GaSe	2.8; 2.94	3.5–18	$\sim 100~\mathrm{ps}$	1	[91Vod, 93Vod1, 95Vod1]
CdSe	1.833	9.8–10.4; 2.26–2.23	300 ns	40	[72Her]
	2.36	7.9 - 13.7	40 ns	15	[72Dav, 73Dav]
	2.87	4.3 - 4.5; 8.1 - 8.3	140 ns	15	[74Wei]
	2.87	14.1 - 16.4	_	_	[76Wen]

^a Pump depletion.

4.1.8 Picosecond continuum generation

Table 4.1.34. Picosecond continuum generation in crystals.

Crystal	$\lambda_{ m pump} \ [\mu { m m}]$	I_{pump} [10 ⁹ W cm ⁻²]	$\lambda_{ m cont} \ [\mu { m m}]$	η [%]	Cut angle of crystals	Ref.
KDP	1.054	50	0.3–1.1	10	$\theta = 49^{\circ}$	[83Mur]
KDP	0.527	30-40	0.84 – 1.4	15	$\theta=42^\circ$	[82Bar]
$LiIO_3$	0.355	_	0.46 – 1.55	_	$\theta = 90^{\circ}$	[85Pok]
$LiIO_3$	0.532	0.3	0.67 - 2.58	-	$\theta = 90^{\circ}$	[85Pok]
$LiIO_3$	1.064	_	1.72 – 3.0	-	$\theta = 90^{\circ}$	[85Pok]
$LiNbO_3$	1.064	_	1.92 – 2.38	3	$ heta=44.7^\circ$	[75Cam]
GaAs	9.3	100	3-14	-	_	[85Cor]

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4.2 Frequency conversion in gases and liquids

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4.2.1 Fundamentals of nonlinear optics in gases and liquids

This chapter covers the properties of a nonlinear medium having spherical symmetry like gases and liquids. They therefore clearly differ from the properties of most solids (see Chap. 4.1).

Lasers have become so powerful these days that one can easily generate various kinds of optical

$$\omega_{o} = \sum_{i,q} n_{i} \cdot \omega_{i} \pm k_{q} \cdot \omega_{res,q} > 0 , \qquad (4.2.1)$$

where n_i and k_q are some integer (including $n_i = 0$ or $k_q = 0$), using a suitable nonlinear medium with eigenfrequencies $\omega_{{\rm res},q}$ and an incident laser frequency ω_i (conservation of energy).

In case of frequency conversion in gases one generally has $k_q = 0$ and deals with sum or difference frequency mixing

$$\omega_{\rm s} = \omega_i \pm \sum_j \omega_j > 0 \tag{4.2.2}$$

which may be enhanced by exploiting suitable resonances of the atomic or molecular gas.

In case of stimulated scattering one generally has $n_i = 1$. Then $\omega_{res,q}$ is a suitable manifold of atomic or molecular (rotational or vibrational) resonances of the gaseous or liquid scattering medium numbered by the index q. Like in classical spectroscopy the plus sign stands for Stokes processes, whereas the minus sign is responsible for Anti-Stokes processes.

4.2.1.1 Linear and nonlinear susceptibilities

Linear and nonlinear susceptibilities are discussed in [87Vid].

The complex linear susceptibility is given by

$$\chi^{(1)} = \bar{\chi}^{(1)} + i\,\tilde{\chi}^{(1)} = \frac{1}{\hbar} \sum_{a} \frac{|\mu_{ag}|^2}{(\Omega_{ag} - \omega)}$$
(4.2.3)

with the complex transition frequency

$$\Omega_{ag} = \omega_{ag} - i \Gamma_{ag} \tag{4.2.4}$$

and the dipole moment matrix elements μ_{ag} between the states $|a\rangle$ and $|g\rangle$. The nonlinear polarization is

$$\boldsymbol{P}^{\mathrm{NL}} = \sum_{n=2}^{\infty} \boldsymbol{P}^{(n)} . \tag{4.2.5}$$

The definition of the electric field amplitude is given by

$$\boldsymbol{E}(r,t) = \frac{1}{2} \sum_{j} \boldsymbol{e}_{j} \hat{E}(r,\omega_{j}) \exp(\mathrm{i} \, k_{j} r - \mathrm{i} \, \omega_{j} t) + c.c. , \qquad (4.2.6)$$

resulting in the definition of the total polarization

$$\mathbf{P}(r,t) = \frac{1}{2} \sum_{j} \mathbf{e}_{j} P(r,\omega_{j}) \exp(-\mathrm{i}\,\omega_{j}t) + c.c. , \qquad (4.2.7)$$

where the n th-order polarization is given by

$$\boldsymbol{P}_{\alpha_{s}}^{(n)}(r,\omega_{s}) = \frac{n!N}{2^{n-1}} \epsilon_{0} \sum_{\alpha_{1}...\alpha_{n}} \chi_{\alpha_{s}\alpha_{1}...\alpha_{n}}^{(n)}(-\omega_{s};\omega_{1}...\omega_{n}) \, \boldsymbol{E}_{\alpha_{1}}(r,\omega_{1}) ... \boldsymbol{E}_{\alpha_{n}}(r,\omega_{n}) .$$
(4.2.8)

The α_s are the unit vectors of the spatial coordinates, which may be cartesian, cylindrical, or spherical. The polarization can be expressed in terms of the density matrix [71Han]

$$\langle \mathbf{P}(t) \rangle = N \operatorname{Tr} \left[\boldsymbol{\rho}(t) \, \boldsymbol{\mu} \right] = N \sum_{mn} \rho_{mn}(t) \, \mu_{mn} ,$$
 (4.2.9)

whose elements are given by $i\hbar \dot{\rho}_{mn} = [H,\rho]_{mn}$, where the Hamiltonian $H = H^0 + H'$ contains $H' = -\mu E(t)$. From a perturbation approach one obtains

$$\chi_{\alpha_{1},\alpha_{2}...\alpha_{n}}^{(n)}(-\omega_{s};\omega_{1}...\omega_{n}) = \frac{1}{n! \,\hbar^{n}} \sum_{gb_{1}...b_{n}} \rho(g) \, \frac{\langle g|\boldsymbol{e}_{s}\boldsymbol{\mu}|b_{1}\rangle\langle b_{1}|\boldsymbol{e}_{1}\boldsymbol{\mu}|b_{2}\rangle...\langle b_{n}|\boldsymbol{e}_{n}\boldsymbol{\mu}|g\rangle}{(\Omega_{b_{1}g} - \omega_{1} - \cdots - \omega_{n})(\Omega_{b_{2}g} - \omega_{2} - \cdots - \omega_{n})....(\Omega_{b_{n}g} - \omega_{n})} \,.$$
(4.2.10)

4.2.1.2 Third-order nonlinear susceptibilities

These processes are responsible for the lowest-order frequency conversion in gases such as sum or difference frequency mixing, stimulated scattering processes and photorefraction. For the *degenerate* case the dominant terms in a system of spherical symmetry are [71Han]:

$$\chi^{(3)}(-3\,\omega;\omega,\,\omega,\,\omega) = \chi_{\rm T}^{(3)}(3\,\omega) = \hbar^{-3}\,\sum_{abc} \frac{\langle g|\boldsymbol{e}_{\rm s}\boldsymbol{\mu}|a\rangle\langle a|\boldsymbol{e}_{1}\boldsymbol{\mu}|b\rangle\langle b|\boldsymbol{e}_{2}\boldsymbol{\mu}|c\rangle\langle c|\boldsymbol{e}_{3}\boldsymbol{\mu}|g\rangle}{(\Omega_{ag}-\omega)(\Omega_{bg}-2\,\omega)(\Omega_{cg}-3\,\omega)}\,,\tag{4.2.11}$$

where the index T stands for the third harmonic generation.

For the *nondegenerate* case we have the general third-order nonlinear susceptibility [62Arm, 71Han]

$$\chi_{\alpha_{1},\alpha_{2},\alpha_{3},\alpha_{s}}^{(3)}(-\omega_{s};\omega_{1},\omega_{2},\omega_{3}) = \frac{1}{6\,\hbar^{3}} \sum_{gabc} \rho(g) \frac{\langle g|\boldsymbol{e}_{s}\boldsymbol{\mu}|a\rangle\langle a|\boldsymbol{e}_{1}\boldsymbol{\mu}|b\rangle\langle b|\boldsymbol{e}_{2}\boldsymbol{\mu}|c\rangle\langle c|\boldsymbol{e}_{3}\boldsymbol{\mu}|g\rangle}{(\Omega_{ag}-\omega_{1}-\omega_{2}-\omega_{3})(\Omega_{bg}-\omega_{2}-\omega_{3})(\Omega_{cg}-\omega_{3})}$$

$$(4.2.12)$$

obeying the conservation of energy

$$\omega_{\rm s} = \omega_1 + \omega_2 + \omega_3 \ . \tag{4.2.13}$$

4.2.1.3 Fundamental equations of nonlinear optics

Maxwell's equations in SI units [62Jac, 87Vid] are given by (1.1.4)–(1.1.7) and the material equations (1.1.8) and (1.1.9), see Chap. 1.1.

With the following three approximations

- 1. magnetization $\mathbf{M} = 0$: $\mu_0 \mathbf{H} = \mathbf{B} \rightarrow \mu = 1$,
- 2. source-free medium: $\rho = 0$,
- 3. currentless medium: j = 0

we get the simplified Maxwell equations

$$\nabla \times \mathbf{E} = -\mu_0 \frac{\partial \mathbf{H}}{\partial t} \tag{4.2.14}$$

$$\nabla \times \boldsymbol{H} = \epsilon_0 \frac{\partial \boldsymbol{E}}{\partial t} + \frac{\partial \boldsymbol{P}}{\partial t} \tag{4.2.15}$$

resulting in the wave equation

$$\Delta \mathbf{E} - \frac{1}{c^2} \frac{\partial^2 \mathbf{E}}{\partial t^2} = \frac{1}{\epsilon_0 c^2} \frac{\partial^2 \mathbf{P}}{\partial t^2}$$
(4.2.16)

with the polarization $P = P^{L} + P^{NL}$. This gives the driven wave equation

$$\Delta \mathbf{E} - \frac{n^2}{c^2} \frac{\partial^2 \mathbf{E}}{\partial t^2} - i \frac{\epsilon_0}{c^2} \frac{\partial^2 \mathbf{E}}{\partial t^2} = \frac{1}{\epsilon_0 c^2} \frac{\partial^2 \mathbf{P}^{NL}}{\partial t^2} . \tag{4.2.17}$$

With the plane-wave approximation $\hat{E}(r,\omega) = \hat{E}(z,\omega)$ and the slow-amplitude approximation

$$\frac{\partial \hat{E}_j}{\partial t} \ll \omega \, \hat{E}_j \,, \quad \frac{\partial \hat{E}_j}{\partial z} \ll k \, \hat{E}_j \,,$$
 (4.2.18)

we get the fundamental equations of nonlinear optics

$$\frac{\mathrm{d}\,\hat{E}_j}{\mathrm{d}\,z} = \mathrm{i}\,\frac{\omega_j}{2\,\epsilon_0\,c\,n_j}\,P_j^{\mathrm{NL}}\,\exp(-\mathrm{i}\,k_jz) - \frac{\kappa_j}{2}\hat{E}_j\;,\tag{4.2.19}$$

where κ is the absorption coefficient and where the total derivative is given by the partial derivatives

$$\frac{\mathrm{d}\,\hat{E}_j}{\mathrm{d}\,z} = \frac{\partial\hat{E}_j}{\partial z} + \frac{n_j}{c}\frac{\partial\hat{E}_j}{\partial t}\,\,,\tag{4.2.20}$$

and \hat{E}_i is a slowly-varying-envelope function in space and time.

4.2.1.4 Small-signal limit

In this case the only nonlinear polarization for a medium of density N is given by

$$P_{\rm s}^{(3)}(\omega_{\rm s}) = \frac{3}{2} \epsilon_0 N \chi_{\rm T}^{(3)}(-\omega_{\rm s}; \omega_1, \, \omega_2, \, \omega_3) E_1 E_2 E_3 \,. \tag{4.2.21}$$

Within the plane-wave approximation one obtains

$$\frac{\mathrm{d}\,\hat{E}_{\mathrm{s}}}{\mathrm{d}\,z} = \mathrm{i}\,\frac{3\,\pi\,\omega_{\mathrm{s}}}{c\,n_{\mathrm{s}}}\,N\,\chi_{\mathrm{T}}^{(3)}\,E_{10}E_{20}E_{30}\exp\left\{\left(-\frac{\kappa_{1}+\kappa_{2}+\kappa_{3}}{2}-\mathrm{i}\,\Delta\,k\right)z\right\}\,\,,\tag{4.2.22}$$

where the wave-vector mismatch is given by the conservation of momenta

$$\Delta k = k_{\rm s} - k_1 - k_2 - k_3 \ . \tag{4.2.23}$$

The wave vector k_j of the j th wave is given by the refractive index n_j

$$k_j = \frac{\omega_j \, n_j}{c} \,. \tag{4.2.24}$$

With the optical depth $\tau_j = \kappa_j L = \sigma_j^{(1)}(\omega_j) NL$ and the length L of the nonlinear medium we have

$$\hat{E}_{s}(L) = i \frac{3\pi\omega_{s}}{c n_{s}} NL \epsilon_{0} \chi_{T}^{(3)} E_{10} E_{20} E_{30} \frac{\exp\left(-\frac{\tau_{s}}{2}\right)}{\frac{\tau_{s} - \tau_{0}}{2} - i \Delta k L} \left\{ \exp\left[\frac{\tau_{s} - \tau_{0}}{2} - i \Delta k L\right] - 1 \right\},$$
(4.2.25)

where the total optical depth $\tau_0 = \tau_1 + \tau_2 + \tau_3$. With the intensity

$$\Phi_j = \frac{\epsilon_0 \, n_j \, c}{2} \, |E_j|^2 \tag{4.2.26}$$

the intensity conversion is given by

$$\frac{\Phi_{\rm s}}{n_{\rm s}} = \left[\frac{6\pi\omega_{\rm s}}{c^2 n_{\rm s}} N L \chi_{\rm T}^{(3)}(-\omega_{\rm s}; \omega_1, \omega_2, \omega_3) \right]^2 \frac{\Phi_{10}\Phi_{20}\Phi_{30}}{n_1 n_2 n_3} F(\Delta k L, \tau_0, \tau_{\rm s}) , \qquad (4.2.27)$$

containing the general phase-matching factor

$$F(\Delta k L, \tau_0, \tau_s) = \frac{\exp(-\tau_0) + \exp(-\tau_s) - 2\exp(-\frac{\tau_0 + \tau_s}{2})\cos(\Delta k L)}{\left(\frac{\tau_s - \tau_0}{2}\right)^2 + (\Delta k L)^2} < 1.$$
 (4.2.28)

4.2.1.5 Phase-matching condition

Maximum conversion efficiency is achieved for conservation of momenta k_j where $\Delta k = 0$

$$\omega_1 \, n_1 + \omega_2 \, n_2 + \omega_3 \, n_3 = \omega_s \, n_s \, . \tag{4.2.29}$$

In case of the third harmonic generation this gives $n_1 = n_{\rm s}$. Frequency mixing in a two-component system results in

$$\frac{N_a}{N_b} = \frac{\omega_s \,\bar{\chi}_b^{(1)}(\omega_s) - \sum_{j=1}^3 \omega_j \,\bar{\chi}_b^{(1)}(\omega_j)}{\sum_{j=1}^3 \omega_j \,\bar{\chi}_a^{(1)}(\omega_j) - \omega_s \,\bar{\chi}_a^{(1)}(\omega_s)} \,. \tag{4.2.30}$$

For the third harmonic generation in a two-component system we have:

$$\frac{N_a}{N_b} = \frac{\bar{\chi}_b^{(1)}(3\,\omega) - \bar{\chi}_b^{(1)}(\omega)}{\bar{\chi}_a^{(1)}(\omega) - \bar{\chi}_a^{(1)}(3\,\omega)} \,. \tag{4.2.31}$$

The frequency mixing in a one-component system is given by:

$$\omega_{\rm s}\,\bar{\chi}^{(1)}(\omega_{\rm s}) = \sum_{j=1}^{3} \omega_{j}\,\bar{\chi}^{(1)}(\omega_{j}) \,. \tag{4.2.32}$$

4.2.2 Frequency conversion in gases

The following conditions have to be met for large conversion efficiencies:

- 1. a large nonlinear susceptibility $\chi_{\rm T}^{(3)}$ which may be enhanced by a proper two-photon resonance,
- 2. large column densities with a proper phase matching,
- 3. small optical depths for the incident and generated waves to avoid reabsorption.

4.2.2.1 Metal-vapor inert gas mixtures

Metal-vapor inert gas mixtures are generally generated in concentric heat pipes because for efficient frequency mixing the phase matching can be accurately and independently adjusted through the partial pressures in the heat pipe [71Vid, 87Vid, 96Vid].

Tables of the multi-wave mixing experiments in different gaseous nonlinear media are arranged according to the elements, Table 4.2.1. For every element the wavelength is given together with the method of generation. The method of generation is indicated where $(\omega_1 + \omega_1)_{\rm Res} + \omega_2$, for example, indicates a two-photon resonance of ω_1 in the particular atomic or molecular medium and the additional wave ω_2 can make the resulting radiation tunable.

4.2.2.2 Mixtures of different metal vapors

The modified concentric heat pipe [71Vid, 87Vid, 96Vid] is used for phase matching with small partial pressures avoiding strong homogeneous broadening.

In Table 4.2.2 mixtures of different metal vapors are listed.

4.2.2.3 Mixtures of gaseous media

For $\lambda > 106$ nm one prefers gas cells with lithium fluoride windows [87Vid]. For $\lambda < 106$ nm one should use either pulsed nozzle beams [87Bet] without windows or gas cells with a fast shutter [85Bon].

In Table 4.2.3 mixtures of gaseous media are given.

 ${\bf Table\ 4.2.1.\ Metal-vapor\ inert\ gas\ mixtures.}$

Vapor	Wavelength [nm]	Method	Ref.
Na	354.7	$3 \cdot \omega_1$	[75Blo1, 75Blo2, 76Oha]
Na	330.5	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$	[74Blo]
Na	268 (cw)	$(\omega_1 + \omega_2)_{\mathrm{Res}} + \omega_3$	[84Bol]
Na	231	$(\omega_1 + \omega_2)_{\mathrm{Res}} + \omega_3$	[76Bjo]
Na	151.4	$7\cdot\omega_1$	[77Gro2, 79Mit]
Na	117.7	$9 \cdot \omega_1$	[77Gro1, 79Mit]
Rb	354.7	$3 \cdot \omega_1$	[71You, 75Blo1, 76Pue, 76Oha]
Cs	213.4	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_1$	[74Leu, 75War]
Be	121–123	$(\omega_1 + \omega_1)_{\mathrm{Res}}^{\mathrm{rtes}} + \omega_2$	[79Mah]
Mg	173.5	$2 \cdot \omega_1 + \omega_1 + \omega_1$	[85Hut]
Mg	140-160	$(\omega_1 + \omega_1)_{\text{Res}} + \omega_2$	[76Wal, 80Jun, 96Ste]
Mg	143.6 (cw)	$(\omega_1 + \omega_1)_{\text{Res}}^{\text{Res}} + \omega_1$	[83Tim]
Mg	115, 121.2, 127	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$	[81Car, 85Car]
Mg	121–129	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$	[78McK]
Mg^{+}	123.6	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$ $(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$	[85Leb]
Ca	200	$3 \cdot \omega_1$	[76Fer]
Ca ⁺	127.8	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$	[75Sor]
Zn	106–140	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$ $(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$	[82Jam]
Cd	118.2, 152, 177.3	$(\omega_1 + \omega_1)_{\text{Res}} + \omega_2; 3 \cdot \omega_1$	[72Kun]
Cd	128.7–135.3		[84Miy]
Cd	145.3–171.1	$(\omega_1 + \omega_1)_{\text{Res}} + \omega_2$	[86Miy]
Sr	155.3, 166.7, 169.7, 173.5, 183.5 (cw)	$(\omega_1 + \omega_1)_{\text{Res}} - \omega_2$	[90Nol1]
Sr		$(\omega_1 + \omega_2)_{\text{Res}} + \omega_3$	
Sr	184.6, 185.7, 192.0, 195.8, 208.8, 217.9 (cw)	$(\omega_1 + \omega_2)_{\text{Res}} + \omega_3$	[90Nol1]
51	171.2, 168.3, 169.7, 190 (cw)	$(\omega_1 + \omega_2)_{\text{Res}} + \omega_3$	[77Bjo, 78Fre]
Ca	152 0 150 5 161 2 162 2 167 0 (cm)	$(\omega_1 + \omega_1)_{\text{Res}} + \omega_2$	[90Nol2]
Ca	153.0, 159.5, 161.3, 163.3, 167.0 (cw)	$(\omega_1 + \omega_2)_{\text{Res}} + \omega_3$	[90Nol1]
Ca	169.7, 170.9, 172.3, 173.1, 176.9 (cw)	$(\omega_1 + \omega_2)_{\text{Res}} + \omega_3$	[90Nol1]
Zn	134.5–141.6 (cw)	$(\omega_1 + \omega_2)_{\text{Res}} + \omega_3$	[90Nol1]
Cd	138.1–140.3	$(\omega_1 + \omega_1)_{\text{Res}} + \omega_2$	[88Sch]
Sr	177.8–195.7	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$	[74Hod, 75Sor, 76Sor]
Sr	165–166	$(\omega_1 + \omega_2)_{\text{Res}} + \omega_3$	[78Eco]
Sr	192.3	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_1$	[80Pue, 81Egg]
Sr	171.2	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$	[80Eco]
Ba	190–200	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$	[80Hei]
Hg	109–196	$(\omega_1 + \omega_1)_{\mathrm{Res}} \pm \omega_2$	[83Hil2]
Hg	184.9, 143.5, 140.1, 130.7, 125.9, 125.0	$(\omega_1 + \omega_1)_{\mathrm{Res}} \pm \omega_2$	[81Bok]
Hg	125.1, 183.3, 208.5	$(\omega_1 + \omega_1)_{\mathrm{Res}} \pm \omega_2$	[81Tom]
Hg	124.7 - 125.5, 122.8 - 123.5, 117.4 - 122	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$	[82Mah, 82Tom]
$_{\mathrm{Hg}}$	120.3	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$	[76Hsu]
Hg	87.5–105, 99.1–126.8	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$	[85Her, 83Hil2]
Hg	89.6	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_1$	[78Sla]
Hg	132–185	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$	[86Hil2]
Tl	195.1	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_1$	[75Wan]
Eu, Yb	185.5, 194	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$	[75Sor, 76Sor]

 ${\bf Table~4.2.2.~Mixtures~of~different~metal~vapors.}$

Mixture	Wavelength [nm]	Method	Ref.
$\overline{Na + K}$ $Na + Mg$	2–25 μm 354.7	$\left(\omega_1 - \omega_2\right)_{\mathrm{Res}} - \omega_3 \ 3 \cdot \omega_1$	[74Wyn] [75Blo2]

 ${\bf Table~4.2.3.~Mixtures~of~gaseous~media.}$

Gas	Wavelength [nm]	Method	Ref.
He, Ne, Ar, Kr, Xe	231.4	$3 \cdot \omega_1$	[67New, 69War]
Не	53.2	$5\cdot\omega_1$	[76Rei, 77Rei, 77She,
			78Rei2, 78Rei1]
Не	38	$7\cdot\omega_1$	[77Rei, 78Rei2]
Не	82.8, 50, 35.5	$3 \cdot \omega_1; 5 \cdot \omega_1; 7 \cdot \omega_1$	[83Bok]
He, Ne	106.4	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$	[78Rei2]
He, Ne	88.7	$3\cdot\omega_1$	[78Rei2, 78Rei1]
He, Ne	76, 70.9, 62.6, 59.1	$4 \cdot \omega_1 \pm \omega_2$	[77She, 78Rei2]
He, Xe	$49.7,\ 35.5$	$5 \cdot \omega_1; 7 \cdot \omega_1$	[83Bok]
Ne	53.2, 118.2	$5 \cdot \omega_1; (\omega_1 + \omega_1)_{\text{Res}} + \omega_2$	[76Rei, 77She, 78Rei2]
Ne	72.05 - 73.58, 74.3 - 74.36	$3\cdot\omega_1$	[84Hil]
Ar	120.4	$(\omega_1 + \omega_1)_{\text{Res}} + 3 \cdot \omega_1$	[80Din]
Ar	85.7 - 87.0, 97.4 - 104.8	$3\cdot\omega_1$	[83Hil3, 83Mar]
Ar	106.7	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_1$	[82Mil]
Ar	102.6 – 102.8	$3\cdot\omega_1$	[79Rei, 80Rei]
Ar, Kr	61.6, 53.2	$5 \cdot \omega_1$	[78Rei2, 81Rei]
Ar	57	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_1$	[76Hut]
Kr	131.2, 92.3, 92.8, 94.2	$(\omega_1 + \omega_1)_{\mathrm{Res}} \pm \omega_2$	[85Bon]
Kr	121.6	$3\cdot\omega_1$	[81Hil, 80Lan, 81Bat]
Kr	123.6	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_1$	[82Mil]
Kr	112.4, 120.3–123.6	$(\omega_1 + \omega_1)_{\mathrm{Res}}^{\mathrm{Res}} + \omega_2$	[79Cot1, 79Cot2]
Kr	131.2	$(\omega_1 + \omega_1)_{\mathrm{Res}} - \omega_2$	[85Bon]
Kr	72.5-83.5; 127-180	$(\omega_1 + \omega_1)_{\mathrm{Res}} \pm \omega_2$	[87Hil]
Kr	121-200	$(\omega_1 + \omega_1)_{\mathrm{Res}}^{\mathrm{Res}} - \omega_2$	[90Mar]
Kr, Xe	71 - 92	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$	[89Miy]
Kr, Xe	110-210	$(\omega_1 + \omega_1)_{\mathrm{Res}}^{\mathrm{rtes}} \pm \omega_2$	[82Hil]
Xe	84.6–109.5, 155–220	$(\omega_1 + \omega_1)_{\mathrm{Res}}^{\mathrm{rtes}} \pm \omega_2$	[82Hil, 82Hag, 83Hil1]
Xe	155	$(\omega_1 + \omega_1)_{\mathrm{Res}}^{\mathrm{Res}} - \omega_2$	[83Hut]
Xe	147	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_1$	[82Mil]
Xe	140.3-146.9	$3 \cdot \omega_1; (\omega_1 + \omega_1)_{\text{Res}} + \omega_2$	[81Hil, 83Val]
Xe	74.8, 75, 75.2	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_1$	[82Mui]
Xe	125.4, 125.9, 126.1	$(\omega_1 + \omega_1)_{\mathrm{Res}}^{\mathrm{Res}} - \omega_2$	[82Mui]
Xe	101.5, 101.8, 13.0	$(\omega_1 + \omega_1)_{\mathrm{Res}} + \omega_2$	[82Mui]
Xe	163.1 – 194.6	$(\omega_1 + \omega_1)_{\mathrm{Res}}^{\mathrm{Res}} \pm \omega_2$	[74Kun]
Xe	118.2	$3\cdot\omega_1$	[83Kun, 76Kun, 82Gan,
			83Bok]
Ar, Kr, Xe, CO, N_2	72; 90.4–102.5	$3\cdot\omega_1$	[87Pag]
Ne, Ar, Kr, Xe, Hg	60-200	$(\omega_1 + \omega_1)_{\mathrm{Res}} \pm \omega_2$	[86Hil1]
Ar, Xe, CO	74.2, 80.4, 95.1, 98.2, 100.1,	$\omega_1 + \omega_1 + \omega_2; \ 3 \cdot \omega_1$	[89Cro]
	116.5, 117.8, 123.6		-

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4.3 Stimulated scattering

A. Laubereau

4.3.1 Introduction

The first example of stimulated scattering was incidentally discovered in 1962 as a "new laser line in the emission of a ruby laser" [62Woo]. The phenomenon occurred when the laser was equipped with a nitrobenzene cell for Q-switching operation. The emitted frequency component was identified as an amazingly intense Raman line [62Eck] due to stimulated Raman scattering predicted theoretically in 1931 [31Goe]. Hundreds of papers appeared since then on the novel phenomenon. Compared to the wealth of experimental evidence full quantitative information about the individual scattering processes however is rather scarce since many publications confine themselves to reported frequency shifts. A quantitative analysis is also often impeded by competing nonlinear effects and by the not too well known properties of the applied laser pulses. Three cases were investigated in detail: Stimulated Raman Scattering (SRS), Stimulated Brillouin Scattering (SBS), and stimulated Rayleigh scattering.

This chapter, Chap. 4.3, follows the discussions given by Maier and Kaiser [72Mai], by Maier [76Mai], and by Penzkofer et al. [79Pen]. Circular frequencies are denoted in the following by ω_i while the corresponding frequency values are represented by $\nu_i = \omega_i/2\pi$. The term "circular" is often omitted in context with the ω_i 's.

4.3.1.1 Spontaneous scattering processes

Fluctuations of the molecular polarizability and of the number density of atoms or molecules give rise to various scattering processes when light passes a transparent medium. The scattering is characterized by the frequency $\nu_{\rm sc}$ of the scattered light relative to the incident laser frequency $\nu_{\rm L}$, the linewidth $\delta\nu$, its polarization properties, and the scattering intensity. Here we introduce the scattering cross section ${\rm d}\sigma/{\rm d}\Omega$ relating the power $P_{\rm sc}$ of the light scattered into a solid angle $\Delta\Omega$ to the incident laser power $P_{\rm L}$:

$$P_{\rm sc} = N \frac{\mathrm{d}\sigma}{\mathrm{d}\Omega} \ell P_{\rm L} \Delta\Omega \tag{4.3.1}$$

with number density N of the (quasi-)particles generating the scattering. The interaction length is denoted by ℓ . $\mathrm{d}\sigma/\mathrm{d}\Omega$ is the differential cross section with respect to solid angle but integrated over the spectral lineshape. The spectrum of four scattering processes is depicted schematically in Fig. 4.3.1a. Two unshifted components are indicated: the narrow Rayleigh line scattered from non-propagating entropy (temperature) fluctuations and the broader Rayleigh-wing line due to orientation fluctuations of anisotropic molecules. The lines are accompanied by the Brillouin doublet representing scattering from propagating isentropic density fluctuations. In the quantum-mechanical approach the Brillouin lines are related to the annihilation (frequency up-shifted anti-Stokes component) and creation (down-shifted Stokes line) of acoustic phonons with conservation of quantum energy and (pseudo-)momentum:

218 4.3.1 Introduction [Ref. p. 232

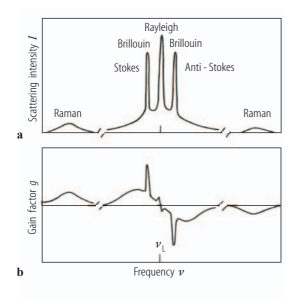


Fig. 4.3.1. (a) Schematic of the spectral intensity distribution of spontaneous light scattering in condensed matter with unshifted Rayleigh and Rayleigh-wing lines (quasi-elastic scattering) as well as Stokes- and anti-Stokes-shifted Brillouin and Raman lines (inelastic light scattering). (b) Frequency dependence of the corresponding gain factors of stimulated scattering (see text).

$$h \nu_{\rm L} = h \nu_{\rm sc} \pm h \nu_{\rm o} , \qquad (4.3.2)$$

$$\hbar \, \mathbf{k}_{\rm L} = \hbar \, \mathbf{k}_{\rm sc} \pm \hbar \, \mathbf{k}_{\rm o} \tag{4.3.3}$$

with Planck's constant h and wavevector k, $\hbar = h/2\pi$. Subscript "o" refers to the material excitation, i.e. acoustic phonons. The positive sign in (4.3.2) and (4.3.3) corresponds to the Stokes process (sc = S), while the negative sign applies for anti-Stokes scattering (sc = A). Due to the dispersion relation of acoustic phonons (phase velocity v of sound waves) the frequency shift is given by

$$\nu_{\rm o} = \frac{vk_{\rm o}}{2\pi} = 2v \frac{\nu_{\rm L}n}{c} \sin\left(\frac{\theta}{2}\right). \tag{4.3.4}$$

Here c/n denotes the speed of light in the medium; θ is the scattering angle between wave vectors $\mathbf{k}_{\rm L}$ and $\mathbf{k}_{\rm sc}$ ($k_{\rm sc} \cong k_{\rm L}$, since $\nu_{\rm o} \ll \nu_{\rm L}$). Equation (4.3.4) refers to isotropic media, e.g. gases and liquids. For anisotropic solids three Brillouin doublets occur in the general case and (4.3.4) has to be modified according to the considered transverse or longitudinal acoustic phonon branch and the respective orientation-dependent sound velocity in the crystal. As a consequence of (4.3.3), $k_{\rm o} \leq 2 \, k_{\rm L}$, so that only acoustic phonons close to the center of the first Brillouin zone are involved (note $k_{\rm L} \sim 10^5~{\rm cm}^{-1}$).

Figure 4.3.1a also schematically shows the Stokes and anti-Stokes line of Raman scattering off a molecular vibration or off an optical phonon branch, displaying a larger frequency shift. As before, only phonons of relatively small k_0 are involved. Polyatomic molecules display a variety of such vibrational Raman lines. In gases many vibration-rotation Raman lines occur in addition and also rotational lines with small frequency shifts. In ionic crystals the relevant material excitation is of mixed phonon-photon character and termed polariton. Since the excited states of molecular vibrations and optical phonons are weakly populated, the anti-Stokes line intensity is also small compared to the corresponding Stokes line.

A further unshifted scattering component in liquids, the Mountain line [76Ber], is only mentioned here since it was not yet observed in stimulated scattering because of its weakness and broad width. Typical values for the frequency shift $\nu_{\rm o}/c$ and the linewidth $\delta\nu/c$ (FWHM) in wavenumber units of the various processes are given in Tables 4.3.1–4.3.5. Some scattering cross sections for the Raman interaction are listed in Table 4.3.2. The scattered light intensity is small. Even for the large

number density of condensed matter of $\sim 10^{22}~\rm cm^{-3}$ a small fraction $< 10^{-5}$ of the incident light is distributed into the whole solid angle 4π per cm interaction length by spontaneous scattering.

4.3.1.2 Relationship between stimulated Stokes scattering and spontaneous scattering

The elementary interaction for Stokes scattering is illustrated in Fig. 4.3.2a (solid arrows). The process involves a transition from an initial to a final energy level of the medium (horizontal lines). The relationship between the stimulated and the spontaneous process is close and originates from the Boson character of photons, i.e. the analogy of the eigenmodes of the electromagnetic field with the harmonic oscillator, the transition probability of which increases with occupation number. As a result the rate of photons scattered into an eigenmode of the Stokes field (subscript "S") depends on the occupation number $n_{\rm S}$ of this mode. Under steady-state conditions we have:

$$\frac{\mathrm{d}\,n_{\mathrm{S}}}{\mathrm{d}\,t} = \mathrm{const.}\,n_{\mathrm{L}}\left(1 + n_{\mathrm{S}}\right). \tag{4.3.5}$$

The first term in the bracket on the right-hand side of (4.3.5) represents spontaneous scattering depending linearly on incident photon number $n_{\rm L}$ or laser power, compare (4.3.1), as long as $n_{\rm S} \ll 1$, i.e. a negligible number of scattered photons per mode of the radiation field is present. The second term on the right-hand side of (4.3.5) describes stimulated scattering that dominates for $n_{\rm S} > 1$ and requires sufficiently high laser intensities. In this regime an avalanche build-up of scattered photons can occur.

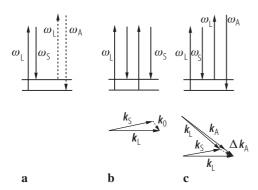


Fig. 4.3.2. (a) Schematic of the elementary scattering process of spontaneous scattering involving two energy levels (horizontal bars) of the medium with transition frequency ω_0 ; the Stokes (full arrows) and anti-Stokes (dashed arrows) processes are indicated. Corresponding diagrams for (b) stimulated Stokes scattering and (c) stimulated Stokes—anti-Stokes coupling in the stimulated scattering. Vertical arrows represent photons that are annihilated (upwards) or generated (downwards) in the interaction. The k-vector geometries of the stimulated processes are depicted in the lower part of the figure (see text).

4.3.2 General properties of stimulated scattering

4.3.2.1 Exponential gain by stimulated Stokes scattering

Integration of (4.3.5) yields exponential growth of Stokes-scattered photons, $n_S \propto \exp(\text{const. } n_L t)$, or equivalently for forward scattering in the z-direction:

$$I_{\rm S}(z) = I_{\rm S}(0) \exp(g I_{\rm L} z)$$
 (4.3.6)

Here we have replaced in the argument of the exponential the product "const. $n_{\rm L}$ t" by a more familiar term with laser intensity $I_{\rm L}$, the gain factor g for stimulated Stokes scattering, and the interaction length z. Equation (4.3.6) indicates exponential amplification of an initial signal $I_{\rm S}(0)$ that may be supplied by spontaneous scattering or by an additional input beam. The exponential growth of the scattered light is only limited by the energy conservation of (4.3.2), since for every scattered photon one incident laser photon has to be annihilated. The corresponding laser depletion leads to gain saturation not included in (4.3.6). Conversion efficiencies above 50% have been observed for stimulated scattering in a number of cases. Equation (4.3.6) refers to steady state.

The gain factor g is an important material parameter for stimulated scattering. The dependence of g on the frequency shift of the scattering is indicated in Fig. 4.3.1b. Maximum gain occurs in the center of the down-shifted Brillouin and Raman lines (Stokes process). For stimulated Rayleigh scattering the peak gain occurs for a Stokes shift equal to half of the full width, $\delta\nu/2$, of the respective line. The negative gain values in Fig. 4.3.1b indicate loss via stimulated scattering on the anti-Stokes side.

Typical values of the peak gain factors are listed in Tables 4.3.2–4.3.5. Under steady-state conditions stimulated Brillouin scattering often represents the dominant interaction. In absorbing media additional mechanisms occur. The corresponding processes, stimulated thermal Brillouin and stimulated thermal Rayleigh scattering, are discussed below.

4.3.2.2 Experimental observation

Stimulated scattering was studied using the following three different experimental approaches:

- 1. generator setup,
- 2. oscillator setup,
- 3. stimulated amplification setup.

4.3.2.2.1 Generator setup

Here only an intense laser beam is directed into the sample. The kind of stimulated scattering is selected by the material and laser beam properties. As a general rule, a large gain of $g I_L z \cong 30$ is required under steady-state conditions for the traveling-wave situation with a single pass through the medium (length z), in order to observe the respective stimulated process. The scattering occurs in forward and/or backward direction because of a simple geometrical argument (maximum interaction length in these directions). The process builds up from an equivalent noise input $I_S(0)$, see (4.3.6), that can be estimated from zero point fluctuations of the electromagnetic field [79Pen]. The growth of the Stokes component is finally limited by the simultaneous decrease of incident laser radiation. The observations are difficult to analyze because of the competition of nonlinear interactions including optical self-focusing. The latter is often involved in liquid media. The observed frequency shift of the stimulated process may slightly deviate from the value known from spontaneous scattering (up to a few cm⁻¹ in SRS) because of simultaneous self-phase modulation in the medium.

4.3.2.2.2 Oscillator setup

An optical resonator made up by mirrors or reflecting surfaces can provide feedback of the stimulated Stokes radiation so that the effective interaction length is increased by multiple passes through the medium. As a result the laser intensity requirements are lowered. The scattering angle is controlled by the cavity axis, so that off-axis emission is possible relative to the laser beam. The frequency-dependent feedback and the lower intensity level of the setup can be sufficient to select a specific stimulated scattering process. Among different Raman transitions only the one with largest gain factor q shows up in SRS in general.

4.3.2.2.3 Stimulated amplification setup

Two well defined beams representing the laser component and the incident Stokes radiation are directed into the scattering medium. Scattering angle and mechanism are determined by the direction and frequency shift of the incident Stokes beam. A second tunable laser is used for the latter in general. The pump intensity $I_{\rm L}$ is smaller by one or more orders of magnitude compared to the generator case, so that self-focusing and other competing effects including secondary scattering processes can be avoided. Quantitative information on the amplitude and/or frequency dependence of the gain factor $g(\nu_{\rm S})$ may be deduced from careful measurements of the amplification factor.

An example for the technique is Raman gain spectroscopy that is often applied in the low-intensity limit $g I_{\rm L} z \ll 1$. An alternative is Raman loss spectroscopy of the transmitted laser component, since the production of Stokes photons corresponds to the annihilation of the same number of laser photons.

4.3.2.3 Four-wave interactions

4.3.2.3.1 Third-order nonlinear susceptibility

Stimulated Stokes scattering can be treated as a four-photon (or four-wave) interaction involving the third-order nonlinear susceptibility $\chi^{(3)}(-\omega_S;\omega_L,-\omega_L,\omega_S)$. The interaction is illustrated by the energy level scheme of Fig. 4.3.2b. The two waves are resonantly coupled via a difference frequency resonance, $\omega_L - \omega_S = \omega_o$, to the relevant material excitation. The latter is enhanced by the scattering thus increasing the coupling strength. The photons at frequencies ω_L and ω_S enter the process twice (see Fig. 4.3.2a). Stimulated amplification is provided in the resonant case by the imaginary part χ_3'' of $\chi^{(3)}$, while the real part leads to frequency modulation. The gain factor is related to the imaginary part by:

$$g \propto |\chi_3''|^2 \,. \tag{4.3.7}$$

Outside difference frequency resonances the real part of $\chi^{(3)}$ is also important for stimulated amplification. The general case of stimulated 4-photon amplification is treated in [79Pen]. The (fourth-rank) tensor character of $\chi^{(3)}$ is omitted here for brevity considering only parallel polarization of the light field components.

The corresponding wave-vector diagram is shown in the lower part of Fig. 4.3.2b. The general case with off-axis geometry is considered. The scattering couples to a material excitation with wave vector $\mathbf{k}_{\rm o}$. The effective scattering angle is strongly influenced by interaction-length arguments. Because of the maximum interaction length, geometries with approximate forward and backward scattering are most important. In cases where the corresponding frequency shift $\omega_{\rm o}$ vanishes, e.g. SBS, stimulated scattering exactly in forward direction is not possible. For backward scattering of short pulses, e.g. SRS of a picosecond laser, the interaction length ℓ may be governed by the duration $t_{\rm p}$ (FWHM of intensity envelope) of the incident laser pulse setting an upper limit of $\ell = t_{\rm p}/2\,v_{\rm g}$ ($v_{\rm g}$: group velocity). In forward direction a less stringent limitation is set by group velocity dispersion between laser and Stokes pulses, $\ell = t_{\rm p}\,\Delta(1/v_{\rm g})$. As a result SRS of picosecond pulses preferentially occurs in forward direction.

4.3.2.3.2 Stokes-anti-Stokes coupling

The stimulated Stokes scattering can be impeded by simultaneous anti-Stokes scattering, $\omega_{\rm A}=\omega_{\rm L}+\omega_{\rm o}$. The anti-Stokes process is depicted in Fig. 4.3.2a (dashed arrows) and "consumes" material excitation, so that (4.3.6) is not applicable. The corresponding four-wave interaction via $\chi^{(3)}(-\omega_{\rm A};\omega_{\rm L},\omega_{\rm L},-\omega_{\rm S})$ is termed Stokes-anti-Stokes coupling and depicted in Fig. 4.3.2c. The significance of the process is determined by its wave vector mismatch $\Delta k_{\rm A}$, depicted in the lower part of Fig. 4.3.2c, and the initial intensity ratio $I_{\rm A}(0)/I_{\rm S}(0)$ ($I_{\rm A}$: anti-Stokes intensity). $\Delta k_{\rm A}$ is governed by the scattering angle and the color dispersion of the refractive index $n(\omega)$ of the medium since

$$k_i = n(\omega_i) \frac{\omega_i}{c}$$
; $(i = A, L, S)$. (4.3.8)

For a collinear geometry we simply have $\Delta k_{\rm A}=k_{\rm A}+k_{\rm S}-2k_{\rm L}$. For $\Delta k_{\rm A}=0$ and $I_{\rm A}/I_{\rm S}=1$, the inverse process of anti-Stokes scattering fully inhibits stimulated Stokes scattering. An example in this context is exact forward scattering in gases, where $\Delta k_{\rm A}$ is small, so that the observed weakness of SRS in exact forward direction is explained in this way. For a large mismatch, $|\Delta k_{\rm A}|>3\,g\,I_{\rm L}$, on the other hand, the Stokes-anti-Stokes coupling is negligible. This condition is always fulfilled for backward scattering so that simultaneous anti-Stokes scattering cannot perturb the stimulated Stokes process notably. For $I_{\rm A}\ll I_{\rm S}$, the perturbation of Stokes scattering by anti-Stokes production is negligible, too. In this case the process of Fig. 4.3.2c is also called Coherent Anti-Stokes Raman Scattering, CARS, an important nonlinear spectroscopy (preferentially applied for phase-matching geometries, $\Delta k_{\rm A}\cong 0$).

Outside Raman resonances the properties of Stokes—anti-Stokes coupling differ notably from the near-resonant case considered here.

4.3.2.3.3 Higher-order Stokes and anti-Stokes emission

For high conversion efficiency of the stimulated scattering the Stokes intensity $I_{\rm S}$ becomes comparable to the incident radiation $I_{\rm L}$, and the material excitation is significant. As a consequence secondary processes show up, generating a cascade of higher-order Stokes and anti-Stokes lines with relative frequency shift $\omega_{\rm o}$ and decreasing intensity levels. Two mechanisms are relevant here:

- 1. stimulated Stokes scattering where the intense first-order Stokes component serves as the pump radiation for generating the second-order line and so forth;
- coherent Stokes or anti-Stokes scattering off the material excitation generated by the primary Stokes scattering producing new frequency-shifted lines. The mechanism is effected by wavevector mismatches of the individual processes.

The Stokes—anti-Stokes coupling discussed above is responsible for the generation of the first-order anti-Stokes component. Higher-order Stokes scattering limits the energy conversion efficiency of first-order Stokes production. The higher-order stimulated scattering should be distinguished from higher-order spontaneous scattering since only a fundamental material transition is involved in the former case.

4.3.2.4 Transient stimulated scattering

The build-up of a material excitation in stimulated scattering involves the response time T_2 (dephasing time) of the medium. When the pulse duration t_p of the incident laser is comparable to or smaller than T_2 , the interaction becomes less efficient and the actual gain of the stimulated Stokes

process is smaller than in the steady state. Equation (4.3.6) for the stationary case is not valid for $t_{\rm p}/T_2 < 10$. The smaller transient gain for a given input situation may be overcome experimentally by increased pump intensities. For details the reader is referred to the literature [78Lau]. Here only three remarks are given:

1. For homogeneous broadening of the material transition ω_0 involved in the stimulated scattering the relaxation time can be simply derived from the linewidth $\delta\nu$ (FWHM)

$$T_2 = (\pi \,\delta \nu)^{-1} = \frac{1}{\Gamma} \,.$$
 (4.3.9)

For inhomogeneous broadening (4.3.9) may be also used to estimate an effective T_2^* from the line broadening that may be sufficient for a semi-quantitative discussion of the transient scattering. For the competition among different Raman transitions in transient SRS both gain factor g and dephasing time T_2 are relevant.

2. For frequency-modulated laser pulses the temporal behavior is not fully described by the duration $t_{\rm p}$ of the pulse envelope. Because of intensity fluctuations the effective duration of the pulse can be estimated to be $t_{\rm p}^* \cong (2 \ \delta \nu_{\rm L})^{-1} < t_{\rm p} \ (\delta \nu_{\rm L}$: frequency width (FWHM) of the laser pulse). To ascertain steady-state conditions the condition

$$\frac{t_{\rm p}^*}{T_2} > 10 \tag{4.3.10}$$

should be fulfilled.

3. Choice of a short t_p may allow to suppress stimulated scattering of transitions with longer T_2 that would have to occur in a less favorable transient situation. An example is SRS in liquids in forward direction with picosecond pulses that is observed in spite of the larger stationary gain factor of SBS. Here the different interaction lengths of forward (SRS) and backward scattering (SBS) also play a role.

4.3.3 Individual scattering processes

4.3.3.1 Stimulated Raman scattering (SRS)

The gain constant for stimulated amplification of the first Stokes component (4.3.6) at resonance, $\omega_S = \omega_L - \omega_o$ is given by

$$g_{\rm S} = \frac{4\pi^2 N \left(\partial \alpha / \partial q\right)^2 \omega_{\rm S}}{n_{\rm L} n_{\rm S} c^2 m \omega_{\rm o} \Gamma} \,. \tag{4.3.11}$$

Here N denotes the molecular number density. A highly polarized vibrational Raman line with halfwidth Γ (HWHM, isotropic scattering component) is considered. $(\partial \alpha/\partial q)$ is the isotropic part of the Raman polarizability (derivative of the molecular polarizability with respect to the vibrational coordinate q of transition ω_o). m represents the reduced mass of the molecular vibration. n_i (i=L,S) is the refractive index at frequency ω_i . $(\partial \alpha/\partial q)$ is connected to the Raman scattering cross section by the relation:

$$\frac{\mathrm{d}\sigma}{\mathrm{d}\Omega} = \frac{(\partial \alpha/\partial q)^2 \,\omega_{\mathrm{S}}^4 \, h \, n_{\mathrm{S}}}{4 \,\pi \, c^4 \, m \, \omega_{\mathrm{O}} \, n_{\mathrm{L}}} \,. \tag{4.3.12}$$

The frequency dependence of the gain factor is given by:

$$g(\omega_{\rm S}) = \frac{g_{\rm S} \Gamma^2}{(\omega_{\rm S} - \omega_{\rm L} + \omega_{\rm o})^2 + \Gamma^2} . \tag{4.3.13}$$

A Lorentzian lineshape is assumed in (4.3.13) that holds well in gases at sufficiently high pressure, weakly associated liquids and solids. SRS of notably depolarized Raman lines is discussed in [78Lau]. Frequency shifts observed for SRS in the generator setup are compiled in Table 4.3.1. A list of gain factors $g_{\rm S}$ and other parameters is presented in Table 4.3.2. The relaxation time T_2 in condensed matter is in the range 10^{-12} to 10^{-10} s.

Table 4.3.1. Frequency shifts (in wavenumber units) observed in stimulated Raman scattering of various materials.

(a) Liquids

Medium	Stokes shift $\nu_0/c \ [\mathrm{cm}^{-1}]$	Excitation wavelength [nm]	Reference
Acetic acid	2944		[84Kru]
Acetone	2925	527	[68Bre, 69Col]
Aniline	997	694	[66Eck]
Benzaldehyde	1001	694	[66Bar]
Benzene	992	527	[67Sha, 68Bre, 69Col, 70Alf]
Benzene	3064	694	[66Eck]
Benzene-d ₆	944	694	[67Blo]
Benzonitrile	2229	694	[66Eck]
Bromobenzene	998, 1000	527, 694	[66Eck, 67Sha]
Bromopropane	2962	694	[66Bar]
2-Bromopropane	2920	694	[66Bar]
1-Bromopropane	2935	694	[66Bar]
Butyl-benzene (tert.)	1000	694	[66Bar]
Carbondisulfide	656	527	[67Sha, 68Bre, 69Col, 70Alf]
Carbontetrachloride	460	694	[66Eck]
Chlorobenzene	1002	527	[67Sha, 69Col]
Chloromethylbutane	2927	694	[66Bar]
Chloroform	663	694	[66Eck]
Cyclohexane	2825		[84Kru]
Cyclohexanone	2683	694	[66Eck]
1,3-Dibromobenzene	992	694	[66Bar]
1,2-Dichloroethane	2958		[84Kru]
Dichloromethane	2989		[75Lau]
2,2-Dichlorodiethylether	2938	527	[83Tel]
1,2-Diethylbenzene	2934	694	[66Bar]
1,2-Dimethylcyclohexane	2853, 2921	694	[66Bar]
1,4-Dimethylcyclohexane	2876	694	[66Bar]
Dimethylhexadiene	2910	694	[66Bar]
1,4-Dioxane	2967		[84Kru]
DMSO, dimethylsulfoxide	2911		[95Go]
Ethanol	2928	527	[69Col, 71Lin]
Ethyl-Benzene	1002	694	[66Eck]
1-Fluoro-2-chlorobenzene	1034	694	[66Bar]
Fluorobenzene	1009	694	[66Eck]
Fluoromethane	2970	694	[78Map]
Isopropanol	2882	527	[69Col]
Methanol	2835	527	[69Col, 70Alf]
$Methanol-d_4$	2200	527	[73Lau]
3-Methylbutadiene	1638	694	[66Eck]

 $({\rm continued})$

Table 4.3.1a continued.

Medium	Stokes shift $\nu_0/c \text{ [cm}^{-1}\text{]}$	Excitation wavelength [nm]	Reference
Nitrobenzene	1344	527, 694	[67Blo, 67Sha, 69Col]
2-Nitropropane	2945	694	[66Bar]
Nitrogen $(T = 77 \text{ K})$	2326	527	[74Lau]
1,3-Pentadiene	1655	694	[66Eck]
Piperidine	2933	694	[66Eck]
Pyridine	992	694	[66Eck]
Siliciumtetrachloride	425	527	[71Lau]
Styrene	1315, 1631, 3056	694	[66Eck]
Tintetrabromide	221	527	[78Lau]
Tintetrachloride	368	527	[78Lau]
Tetrachloroethane	2984	694	[66Eck]
Tetrachloroethylene	448, 2939	527	[69Col, 72Lau, 66Bar]
Tetrahydrofuran	2849	694	[66Eck]
Toluene	1004	527, 694	[67Blo, 67Sha]
Water	3450	527	[68Bre, 69Col, 69Rah]
m-Xylene	2933	694	[66Eck]
o-Xylene	2913	694	[66Eck]
p-Xylene	2998	694	[66Eck]

(b) Solids

Medium	Stokes shift $\nu_0/c \; [\mathrm{cm}^{-1}]$	Excitation wavelength [nm]	Reference
$\overline{\mathrm{Al_2O_3}}$	416	532	[97Kam2]
1-Bromonaphthalene	1363	694	[66Eck]
Calcite	1086	527	[69Col]
1-Chloronaphthalene	1368	694	[66Eck]
Diamond	1332	527	[71Lau]
2-Ethylnaphthalene	1382	694	[66Bar]
$Gd_2(MoO_4)_3$	960	532	[97Kam1]
$\text{LiHCOO} \cdot \text{H}_20$	104, 1372	694	[90Lai]
$NaClO_3$	936	532	[97Kam3]
Naphthalene	1380	694	[66Eck]
Polydiacetylene	1200		[94Yos]
Sulfur	216, 470	694	[66Eck]

(c) Gases

Medium	Stokes shift $\nu_0/c \; [\mathrm{cm}^{-1}]$	Excitation wavelength [nm]	Reference
Ammonia	3339		[72Car]
Barium vapor	11395	552	[83Sap, 87Glo]
Ethylene (55 atm)	1344	694	[70Mac]
Butane (90 atm)	2920	694	[70Mac]
Carbondioxide (20–50 atm)	1385	694	[70Mac, 78Map]
Carbonmonoxide	2145	694	[72Car]

(continued)

Table 4.3.1c continued.

Medium	Stokes shift $\nu_0/c \; [\mathrm{cm}^{-1}]$	Excitation wavelength [nm]	Reference
Cesium vapor	14597		[84Har]
Chlorine	556	694	[72Car, 78Map]
Deuterium	2991	694	[67Blo]
Hydrogen	4160	694	[67Blo, 75Cha]
Hydrogenbromide (20 atm)	2558	694	[70Mac, 78Map]
Hydrogenchloride (35 atm)	2883	694	[70Mac, 78Map]
Methane (10 atm)	2917		[70Mac]
Nitrogen (55–100 atm)	2330	694	[70Mac, 75Cha]
N_2O (50 atm)	774	694	[70Mac, 78Map]
NO	1877	694	[72Car]
Oxygen (50–100 atm)	1550	694	[70Mac]
SF ₆ (15–20 atm)	1551	694	[70Mac]
SF ₆ (18 atm)	775	694	[72Car]

Table 4.3.2. Gain factor and other parameters of stimulated Raman scattering.

(a) Liquids

Medium	Stokes shift ν_0/c [cm ⁻¹]	Scattering coefficient $N \times d\sigma/d\Omega$ [10 ⁷ m ⁻¹ sr ⁻¹]	Linewidth $\delta \nu/c$ [cm ⁻¹]	Gain factor $g_{\rm S}$ [10 ¹² m/W]	Excitation wavelength [nm]	Ref.
Acetone	2925		17.4	12	530	[69Col]
Benzene	992		2.2	28	694	[72Mai]
Bromobenzene	1000	15	1.9	15	694	[72Mai]
Carbondisulfide	655	75	0.50	240	694	[72Mai]
Chlorobenzene	1002	15	1.6	19	694	[72Mai]
Ethanol	2928		17.4	51	530	[69Col]
Isopropanol	2882		26.7	9.2	530	[69Col]
Methanol	2834		18.7	23	530	[69Col]
Methanol	2944		26.5	18	530	[69Col]
Nitrogen	2326	2.9	0.067	170	694	[72Mai]
Nitrobenzene	1345	64	6.6	21	694	[72Mai]
Oxygen	1552	4.8	0.117	140	694	[72Mai]
Tetrachloroethylene	447			17	598	[76Mai]
Toluene	1003	11	1.9	12	694	[72Mai]
1,1,1-Trichloroethane	2939		5.2	51	530	[69Col]
Water	3450	430		1.4	530	[69Col]

/1 \	O 1: 1
(\mathbf{b})	Solids

Medium	Stokes shift ν_0/c [cm ⁻¹]	Linewidth $\delta \nu/c \ [{ m cm}^{-1}]$	Gain factor g_s [10 ¹² m/W]	Excitation wavelength [nm]	Ref.
Ba ₂ NaNb ₅ O ₁₅	650		67	694	[72Mai]
Calcite	1086	1.1	1.4	530	[69Col]
$CuAlS_2$	314		21,000	514	[97Bai]
GaP	403		19,000	632	[97Bai]
$^6\mathrm{LiNbO_3}$	256		180	694	[72Mai]
$^{7}\mathrm{LiNbO_{3}}$	256		89	694	[72Mai]
$^6\mathrm{LiTaO_3}$	600		43	694	[72Mai]
Quartz	467		0.15	527	[67Wig]

(c) Gases

Medium	Stokes shift ν_0/c [cm ⁻¹]	Differential scattering cross section $d\sigma/d\Omega$ [10 ³⁶ m ² sr ⁻¹]	Dephasing time T_2 [ps]	Gain factor $g_{\rm S}$ [10 ¹² m/W]	Excitation wavelength [nm]	Ref.
$\overline{\mathrm{H}_2,\mathrm{Q}(1)}$	4155	1.2	208	9.7	1064	[86Han]
$H_2, Q(1)$	4155	79	208	27.6	532	[86Han]
$D_2, Q(2)$	2987	2.0	150	3.7	1064	[86Han]
$D_2, Q(2)$	2987	8.0	150	10	532	[86Han]
Methane, Q	2917	7.0	16	3.3	1064	[86Han]
Methane, Q	2917	270	16	8.6	532	[86Han]

4.3.3.2 Stimulated Brillouin scattering (SBS) and stimulated thermal Brillouin scattering (STBS)

Stimulated Brillouin scattering was extensively studied in liquids, solids, and gases. In many substances it is the dominant process under stationary conditions and occurs generally in backward direction. The scattering originates from two coupling mechanisms between the electromagnetic field and the medium: electrostriction and absorption. In transparent media only electrostriction is relevant. In absorbing media the second contribution called Stimulated Thermal Brillouin Scattering (STBS) is caused by absorption-induced local temperature changes leading to propagating density waves. The frequency dependencies of the gain factors for the two mechanisms are different. The peak values of stimulated gain are given by:

$$g_{\rm B}^{\rm e} = \frac{(\partial \varepsilon / \partial \rho)_T^2 \,\omega_{\rm S}^2 \,\rho_{\rm o}}{2 \,c^3 \,n_{\rm S} \,v \,\Gamma_{\rm B}} \tag{4.3.14}$$

for the electrostrictive contribution (superscript "e"), and by

$$g_{\rm B}^{\rm a} = \frac{\alpha \left(\partial \varepsilon / \partial \rho\right)_T \,\omega_{\rm S} \,\beta_T}{4 \,c \,n_{\rm S} \,C_p \,\Gamma_{\rm B}} \tag{4.3.15}$$

for STBS. Here $(\partial \varepsilon/\partial \rho)_T$ is the change of the relative dielectric constant with mass density ρ at constant temperature T. ρ_0 is the equilibrium density value. v denotes the sound velocity at

frequency $\omega_{\rm o}=\omega_{\rm L}-\omega_{\rm S}$ (see (4.3.4) for backward scattering, $\theta=180^{\circ}$). The parameters in (4.3.15) are the absorption coefficient of the laser intensity α and the relative volume expansion coefficient β_T . The half-width $\Gamma_{\rm B}=\pi~\delta\nu$ of the corresponding spontaneous Brillouin line that displays an approximately quadratic frequency dependence also enters the expressions above. For liquids one can write:

$$\frac{\Gamma_{\rm B}}{\omega_{\rm o}^2} = \frac{\frac{4}{3}\eta_{\rm S} + \Lambda \left(\frac{1}{C_{\rm V}} - \frac{1}{C_{\rm p}}\right) + \eta_{\rm V}}{2\rho_{\rm o} v^2} \,,\tag{4.3.16}$$

where $\eta_{\rm S}$ and $\eta_{\rm V}$, respectively, denote the shear and volume viscosity; the latter is to some extent frequency-dependent via relaxation phenomena. Λ is the thermal conductivity. $C_{\rm V}$ and C_p are the specific heat per unit mass at constant volume and pressure, respectively. The phonon lifetime τ of the involved acoustic phonons with circular frequency $\omega_{\rm o}$ is related to the linewidth by $\tau = T_2/2 = 1/(2\,\Gamma_{\rm B})$. The peak gain value $g_{\rm B}^{\rm a}$ increases proportional to α and is of same order of magnitude as $g_{\rm B}^{\rm e}$ for $\alpha\approx 1~{\rm cm}^{-1}$.

The total frequency-dependent gain factor for the (first-order) Stokes component of SBS including STBS is given by

$$g(\omega_{\rm S}) = \frac{g_{\rm B}^{\rm e} \, \Gamma_{\rm B}^2}{(\omega_{\rm S} - \omega_{\rm L} + \omega_{\rm o})^2 + \Gamma_{\rm B}^2} - \frac{g_{\rm B}^{\rm a} \, 2 \, \Gamma_{\rm B} \, (\omega_{\rm S} - \omega_{\rm L} + \omega_{\rm o})}{(\omega_{\rm S} - \omega_{\rm L} + \omega_{\rm o})^2 + \Gamma_{\rm B}^2} \,. \tag{4.3.17}$$

The maximum contribution of STBS is red-shifted relative to the Brillouin line and occurs at $\omega_{\rm S} = \omega_{\rm L} - \omega_{\rm o} - \Gamma_{\rm B}$. In the blue wing of the Brillouin Stokes line the mechanism produces stimulated loss. Equation (4.3.17) states that the Stokes shift observed in the stimulated Brillouin scattering of absorbing media in the generator or oscillator setup – occurring at the peak value of $g(\omega_{\rm S})$ – is modified compared to the spontaneous Brillouin line.

A list of frequency shifts observed in SBS of transparent media is presented in Table 4.3.3 where values for the Brillouin linewidth $\delta\nu$ and the gain parameters $g_{\rm B}^{\rm a}/\alpha$ and $g_{\rm B}^{\rm e}$ are also compiled. The relaxation time T_2 (= $1/\pi$ $\delta\nu$) in condensed matter is in the order of 10^{-9} s so that SBS is close to steady state for giant laser pulses with $t_{\rm p}\approx 10^{-8}$ s (if self-focusing is avoided), but is of transient character in the subnanosecond time domain.

4.3.3.3 Stimulated Rayleigh scattering processes, SRLS, STRS, and SRWS

Three mechanisms can be distinguished:

- 1. Stimulated Rayleigh Line Scattering in transparent substances, SRLS, by electrostrictive coupling to non-propagating density changes,
- 2. Stimulated Thermal Rayleigh Scattering, STRS, by absorptive coupling similar to the STBS case, and
- 3. Stimulated Rayleigh Wing Scattering, SRWS, in liquids by orientational changes of anisotropic molecules.

The frequency shifts of the Stokes component of the first two cases are considerably smaller than for SBS. SRLS is difficult to observe because of the small gain factor and the relatively long relaxation time $T_2 \approx 10^{-8}$ s for backward scattering leading to transient scattering for nanosecond pulses.

Table 4.3.3. Stimulated Brillouin scattering in backward direction: frequency shift, linewidth, and gain factor.

Medium	Stokes shift ν_0/c [cm ⁻¹]	Linewidth $\delta \nu$ [MHz]	$egin{aligned} & { m Gain} \\ & { m coefficient} \\ & g_{ m B}^{ m a}/\alpha \\ & { m (calculated)} \\ & { m [cm^2~/~MW]} \end{aligned}$	$g_{ m B}^{ m e}$ (calculated) [cm / MW]	Gain factor $g_{\rm B}^{\rm e}$ (measured) [cm / MW]	Ref.
Acetic acid	0.152					[67Wig]
Acetone	0.154	180	0.022	0.017	0.020	[70Poh]
Aniline	0.259					[67Wig]
Benzaldehyde	0.224					[67Wig]
Benzene	0.211		0.024	0.024	0.018	[68Den]
Bromobenzene	0.188					[67Wig]
Carbondisulfide	0.194	75	0.213	0.197	0.068	[97Jo]
Carbontetrachloride	0.146	650	0.0134	0.0084	0.006	[68Den, 70Poh]
Chloroform	0.148					[67Wig]
Cyclohexane	0.180			0.007	0.0068	[68Den]
p-Dichlorobenzene	0.184					[67Wig]
Ethanol	0.152		0.010	0.012		[72Mai]
Fluorinert FC 72					0.006	[97Yos]
Fluorinert FC 75					0.005	[97Yos]
Glass BSC-2	0.866					[67Wig]
Glass DF-3	0.638					[67Wig]
Glycerol	0.386					[67Wig]
n-Hexane		220		0.027	0.026	[68Den, 70Poh]
InSb						[95Lim]
Methanol	0.142		0.013	0.013	0.013	[68Den, 70Poh]
Methylodide	0.166					[67Wig]
Nitrobenzene	0.228					[67Wig]
m-Nitrotoluene	0.229					[67Wig]
n-Nitrotoluene	0.217					[67Wig]
Octanol	0.194					[67Wig]
Pyridine	0.226					[67Wig]
Quartz	1.16				0.005	[89Agr]
Sulfurhexafluoride					0.0015	[93Fil, 97Jo]
(20 atm)						
Tetrabromomethane	0.173					[67Wig]
Toluene	0.193	480		0.013	0.013	[68Den, 70Poh]
Water	0.197		0.0008	0.0066	0.0048	[68Den, 77Rys, 94Yos]
p-Xylene	0.199					[67Wig]

1. and 2. The peak value of the stimulated gain for SRLS is given by:

$$g_{\rm RL}^{\rm e} = \frac{(\partial \varepsilon / \partial \rho)_T^2 \,\omega_{\rm S} \,\rho_{\rm o} \,(\gamma - 1)}{4 \,c^2 \,n_{\rm S}^2 \,v^2} \,, \tag{4.3.18}$$

where $\gamma=C_p/C_V$. It is interesting to notice that $g_{\rm RL}^{\rm e}$ does not depend on scattering angle $(\omega_{\rm S}\cong\omega_{\rm L})$. A finite optical absorption coefficient α of the medium gives rise to a second contribution with peak value:

$$g_{\rm RL}^{\rm a} = \frac{\alpha \left(\partial \varepsilon / \partial \rho\right)_T \,\omega_{\rm S} \,\beta_T}{2 \,c \,n_{\rm S} \,C_{\rm p} \,\Gamma_{\rm RL}} \,. \tag{4.3.19}$$

Medium	Gain factor	Gain coefficient	Linewidth	Reference
	$g_{ m RL}^{ m e}$	$g_{\mathrm{RL}}^{\mathrm{a}}(\mathrm{max.})/\alpha$	δu	
	$[\mathrm{cm}/\mathrm{MW}]$	$[{ m cm}^2/{ m MW}]$	[MHz]	
Acetone	2	0.47	21	[70Rot]
Benzene	2.2	0.57	24	[70Rot]
Carbondisulfide	6	0.62	36	[70Rot]
Ethanol		0.38	18	[70Rot]
Methanol	8.4	0.32	20	[70Rot]
Tetrachloromethane	2.6×10^{-4}	0.82	17	[70Rot]
Water	0.02	0.019	27.5	[70Rot]

Table 4.3.4. Stimulated Rayleigh scattering: gain factors and linewidth values.

Here $\Gamma_{\rm RL} = \pi \, \delta \nu$ is the halfwidth (HWHM) of the spontaneous Rayleigh line in circular frequency units that originates from the damping of entropy changes via thermal conductivity Λ :

$$\Gamma_{\rm RL} = \frac{4 k_{\rm L}^2 \Lambda \sin^2 (\theta/2)}{\rho_{\rm o} C_p} \,.$$
 (4.3.20)

Via $\Gamma_{\rm RL}$ the gain constant $g_{\rm RL}^{\rm a}$ strongly depends on scattering angle. The frequency dependencies of SRLS and STRS have opposite sign, leading to the total gain factor:

$$g(\omega_{\rm S}) = \frac{(g_{\rm B}^{\rm e} - g_{\rm B}^{\rm a}) \, 2 \, \Gamma_{\rm RL}(\omega_{\rm L} - \omega_{\rm S})}{(\omega_{\rm L} - \omega_{\rm S})^2 + \Gamma_{\rm RL}^2} \,. \tag{4.3.21}$$

For dominant coupling via electrostriction positive gain occurs on the Stokes side, $\omega_{\rm S} < \omega_{\rm L}$. Small absorption values, $\alpha > 10^{-3}~{\rm cm}^{-1}$, can be sufficient for dominant STRS that produces gain on the anti-Stokes side. For zero frequency shift, $\omega_{\rm S} = \omega_{\rm L}$, the gain factor vanishes in the steady state, (4.3.21), but not in the transient case. Stimulated scattering in forward scattering is particularly delicate since $\Gamma_{\rm RL} \to 0$, (4.3.20), and Stokes–anti-Stokes coupling has to be included. Values for g and $\delta \nu$ are listed in Table 4.3.4.

3. Stimulated Rayleigh wing scattering is connected with the overdamped rotational motion of liquid molecules in combination with an anisotropic polarizability tensor. The latter is also involved in the optical Kerr effect enhancing the nonlinear refractive index of the medium (optical self-focusing, self-phase modulation). The maximum steady-state gain factor for SRWS is given by:

$$g_{\rm RW} = \frac{16\,\pi^2\,N\,\omega_{\rm S}\,(\alpha_{\parallel} - \alpha_{\perp})^2}{45\,k_{\rm B}\,T_{\rm o}\,c^2\,n_{\rm S}^2} \ . \tag{4.3.22}$$

The difference of the molecular polarizability parallel and perpendicular to the (assumed) molecular axis of rotational symmetry is denoted by $\alpha_{\parallel} - \alpha_{\perp}$. $k_{\rm B}$ is the Boltzmann constant, $T_{\rm o}$ the sample temperature. The frequency dependence of the gain factor is analogous to the previous cases:

$$g(\omega_{\rm S}) = \frac{g_{\rm RW} 2 \Gamma_{\rm RW} (\omega_{\rm L} - \omega_{\rm S})}{(\omega_{\rm L} - \omega_{\rm S})^2 + \Gamma_{\rm RW}^2} . \tag{4.3.23}$$

Maximum gain of SRWS occurs for $\omega_{\rm S}=\omega_{\rm L}-\varGamma_{\rm RW}$. The halfwidth $\varGamma_{\rm RW}$ of the Rayleigh wing line may be taken from spontaneous scattering observations or from the reorientational time $\tau_{\rm or}$. The latter can be derived from spontaneous Raman spectroscopy, NMR, or time-resolved spectroscopy, e.g. transient optical Kerr effect observations: $\tau_{\rm or}=T_{2,{\rm RW}}=1/\varGamma_{\rm RW}$. An estimate of the halfwidth may be computed from shear viscosity and the size of the molecules using the Debye theory:

$$\Gamma_{\rm RW} = \frac{3 k_{\rm B} T_{\rm o}}{8 \pi R \eta_{\rm S}} \,.$$
(4.3.24)

Here R denotes an effective mean radius of the molecule. The proportionality $\Gamma_{\rm RW} \propto \eta_{\rm S}$ was demonstrated experimentally for numerous examples.

Equations (4.3.6), (4.3.22), and (4.3.23) hold for large scattering angles where Stokes–anti-Stokes coupling can be neglected. Close to forward direction simultaneous anti-Stokes scattering enhances the stimulated Stokes scattering, in contrast to SRS. Including the Stokes–anti-Stokes coupling maximum gain is predicted for an optimum scattering angle

$$\theta_{\text{opt}} = \left(\frac{2 g_{\text{RW}} I_{\text{L}} c}{n_{\text{L}} \omega_{\text{L}}}\right)^{\frac{1}{2}} . \tag{4.3.25}$$

The corresponding gain factor for stimulated amplification in the scattering direction $\theta_{\rm opt}$ without frequency shift, $\omega_{\rm S} = \omega_{\rm A} = \omega_{\rm L}$, amounts to:

$$g_{\text{opt}} = 2 g_{\text{RW}} ,$$
 (4.3.26)

where g_{RW} is given by (4.3.22). For more general cases the reader is referred to the literature, e.g. [72Mai]. Frequency shift and gain factor numbers are compiled in Table 4.3.5.

Table 4.3.5. Frequency shift and gain factor of stimulated Rayleigh wing scattering.

Medium	Frequency shift $\times c^{-1}$ [cm ⁻¹]	Gain factor $G_{\rm RW}$ [10 ¹² m/W]	Reference
Azoxybenzene	0.036		[68Fol]
Benzene		6	[72Mai]
Benzonitrol	0.198		[68Fol]
Benzoylchloride	0.184		[68Fol]
Benzylidenaniline	0.065		[68Fol]
Bromobenzene		14	[72Mai]
1-Bromonaphthalene	0.076		[68Fol]
Carbondisulfide		30	[72Mai]
Chlorobenzene		10	[72Mai]
Chloronaphthalene	0.100		[68Fol]
1,4-Dimethylnitrobenzene	0.090		[68Fol]
m-Dinitrobenzene	0.116		[68Fol]
2,4-Dinitrotoluene	0.098		[68Fol]
Naphthalene	0.5		[68Fol]
Nitroacetophenone	0.105		[68Fol]
o-Nitroaniline	0.107		[68Fol]
p-Nitroanisol	0.075		[68Fol]
Nitrobenzaldehyde	0.101		[68Fol]
Nitrobenzene	0.111	76	[72Mai]
o-Nitrophenol	0.078		[68Fol]
m-Nitrotoluene	0.097		[68Fol]
o-Nitrotoluene	0.133		[68Fol]
p-Nitrotoluene	0.145		[68Fol]
Styrene	0.4		[68Fol]
Toluene		20	[72Mai]

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4.4 Phase conjugation

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4.4.1 Introduction

Phase conjugation is a nonlinear optical process which generates a light beam having the same wavefronts as an incoming light beam but opposite propagation direction, see Fig. 4.4.1. Therefore phase conjugation is also called wavefront reversal. A nonlinear optical device generating a phase-conjugated wave is called a *phase conjugator* or *Phase-Conjugate Mirror* (*PCM*).

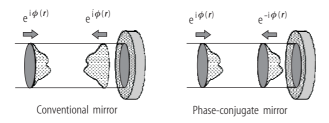


Fig. 4.4.1. Wavefront reflection at a conventional mirror and at a phase-conjugate mirror (PCM).

In Fig. 4.4.2 we consider the conjugation property of a PCM on a probe wave emanating from a point source. A diverging beam, after "reflection" from an ideal PCM, gives rise to a converging conjugate wave that precisely retraces the path of the incident probe wave, and therefore propagates in a time-reversed sense back to the same initial point source.

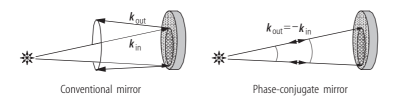


Fig. 4.4.2. Beam propagation after reflection at a conventional mirror and a PCM, both illuminated by a point source.

A phase conjugator reflects light, mostly laser beams, only if the incident power is high enough (self-pumped phase conjugator) or if the nonlinear material in the phase conjugator is pumped by additional laser beams, e.g. two additional beams in a degenerate four-wave mixing arrangement. In principle phase conjugation could be achieved also by a deformable mirror which is controlled by a wavefront sensor adapting the local mirror curvature to the incoming wavefront. Instead of a deformable mirror also a 2-dimensional phase modulator could be used. However, deformable mirrors and other phase modulators up to now are more complicated set-ups with longer reaction periods than nonlinear optical phase conjugators to solve practical problems requiring phase conjugation.

4.4.2 Basic mathematical description

The incoming wave $E_{\rm in}$ is given by (4.4.1) with frequency f, where the amplitude E_0 and phase Φ are combined to the complex amplitude A. The complex conjugate is denoted by c.c.

$$E_{\rm in}(x, y, z, t) = \frac{1}{2} E_0(x, y, z) e^{2\pi i (ft + \Phi(x, y, z))} + \text{c.c.} = A(x, y, z) e^{i\omega t} + \text{c.c.},$$
(4.4.1)

$$A(x,y,z) = \frac{1}{2} E_0(x,y,z) e^{2\pi i \Phi(x,y,z)}.$$
(4.4.2)

The phase-conjugated wave exhibits the same wavefronts, however the sign of the phase Φ is inverted due to the inverted propagation direction. Thus, the phase-conjugated wave $E_{\rm pc}$ can be written as (4.4.3):

$$E_{\rm pc}(x, y, z, t) = \frac{1}{2} E_0(x, y, z) e^{2\pi i (ft - \Phi(x, y, z))} + \text{c.c.} = A_{\rm pc}(x, y, z) e^{i\omega t} + \text{c.c.},$$
(4.4.3)

$$A_{\rm pc}(x,y,z) = \frac{1}{2} E_0(x,y,z) e^{-2\pi i \Phi(x,y,z)} = A^*(x,y,z) . \tag{4.4.4}$$

As can be seen A_{pc} equals the complex-conjugated A^* , which explains the term *phase conjugation*. From (4.4.1) and (4.4.3) we derive that the incident and phase-conjugated wave are also related to each other by

$$E_{\rm in}(x, y, z, -t) = E_{\rm pc}(x, y, z, t) . \tag{4.4.5}$$

Thus, the phase-conjugate wave $E_{\rm pc}$ propagates as if one would reverse the temporal evolution of the incident wave $E_{\rm in}$. Therefore the term "time-reversed replica" is sometimes used to describe the phase-conjugate wave.

An ideal PCM also maintains the polarization state of an incident wave after phase conjugation. As an example, a probe wave that is Right-Handed Circularly Polarized (RHCP) will result in a RHCP-reflected wave after conjugation. This is in contrast to a conventional mirror, which reflects an incident RHCP field to yield a Left-Handed Circularly Polarized (LHCP) wave [82Pep].

One should realize that an ideal phase-conjugated wave exhibits the same frequency f as the incident wave and reveals the same polarization state. Often, real phase conjugators do not have these properties. However, if an PCM maintains the polarization state it is called a "vector phase conjugator".

The nonlinear optical process which comes closest to yielding an ideal phase-conjugate wave is the backward-going, degenerate four-wave mixing interaction. Other classes of interaction (e.g. stimulated effects) result in nonideal conjugate waves due to frequency shifts, nonconjugated field polarization, etc. Although the application of stimulated effects, especially Stimulated Brillouin Scattering (SBS), yields to nonideal phase-conjugate mirrors they are used the most to solve practical problems requiring phase conjugation (e.g. compensation of phase distortions in high average power laser systems [99Eic]).

4.4.3 Phase conjugation by degenerate four-wave mixing

Four-wave mixing can be understood as a real-time holographic process, which facilitates phase conjugation. If the frequencies of the incoming wave, the two additionally required pump waves, and the phase-conjugated or reflected wave are equal the process is called $Degenerate\ Four-Wave\ Mixing\ (DFWM)$.

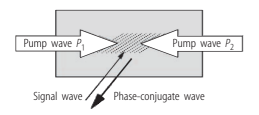


Fig. 4.4.3. Setup for phase conjugation by four-wave mixing.

In Fig. 4.4.3 the setup for phase conjugation by four-wave mixing is shown.

Interference of the incoming wave $E_{\rm in}(x,y,z,t)$ with the pump wave $P_1(x,y,z,t)$ results in a spatially periodic intensity pattern which modulates the absorption coefficient or refractive index of the optical material resulting in a dynamic or transient amplitude or phase grating. The other pump $P_2(x,y,z,t)$ is diffracted at this grating producing the phase-conjugated wave. This corresponds to the conventional holographic process where the read-out wave is replaced by the second pump wave counterpropagating to the first pump or reference wave.

Recording of a hologram is the first step in phase conjugation and leads to a transmission function t in the hologram plane (variables will not be noted furthermore to simplify the readability):

$$t \propto |P_1 + E_{\rm in}|^2 = \dots = |P_1|^2 + P_1 E_{\rm in}^* + P_1^* E_{\rm in} + |E_{\rm in}|^2$$
 (4.4.6)

During the read-out the phase-conjugate wave can be generated. Therefore, the hologram is illuminated with a second pump wave P_2 , propagating in the opposite direction to P_1 . This is in contrast to standard holography. Since P_2 precisely retraces the path of P_1 in the opposite propagation direction, P_2 equals P_1^* . This means, that the two pump beams should be phase-conjugated to each other, so that their spatial phases cancel and do not influence the phases of the reflected beam.

In the hologram plane we obtain a field strength distribution as follows:

$$P_2 t = P_1^* t \propto P_1^* |P_1|^2 + |P_1|^2 E_{\text{in}}^* + (P_1^*)^2 E_{\text{in}} + P_1^* |E_{\text{in}}|^2 .$$
(4.4.7)

The second term $|P_1|^2 E_{\text{in}}^*$ corresponds to the phase-conjugate wave of E_{in} . The other expressions lead to three additional waves which are not of interest here. They can be suppressed in thick nonlinear media in case of Bragg diffraction.

Common dynamic grating materials for phase conjugation are:

- photorefractive crystals (LiNbO₃, BaTiO₃, ...),
- liquid crystals (molecular reorientation effects),
- laser crystals (spatial hole-burning, excited-state absorption),
- saturable absorbers,
- absorbing gases and liquids (thermal gratings),
- semiconductors (Si, GaAs, ...).

The disadvantage of phase conjugation by four-wave mixing is the requirement of two additional pump waves for the nonlinear medium. However, this facilitates amplification of the phase-conjugate wave in the nonlinear medium at the same time. Vector phase conjugation is not achieved by this simple DFWM scheme, but requires polarization-dependant interactions.

4.4.4 Self-pumped phase conjugation

Self-pumped phase conjugation of continuous-wave laser beams in the lower power range (mW ... W) can be realized in Four-Wave Mixing (FWM) loop arrangements using photore-fractive media, see Sect. 4.4.6 for detailed discussion.

SBS medium	Brillouin gain coefficient g [cm/GW]	Phonon lifetime τ [ns]
$\overline{\mathrm{SF}_6 \ (20 \ \mathrm{bar})}$	25	15
Xe (50 bar)	90	33
C_2F_6 (30 bar)	60	10
CS_2	130	5.2
CCl_4	6	0.6
Acetone	20	2.1
Quartz	2.4	5

Table 4.4.1. Brillouin gain coefficient g and phonon lifetime τ for different SBS media.

For pulsed lasers, self-pumped phase conjugation is achieved by stimulated scattering. For practical application, stimulated Brillouin scattering [72Kai] in

- gases (SF₆, Xe, C_2F_6 , CH_4 , N_2 , ...) under high pressure,
- liquids (CS₂, CCl₄, acetone, freon, GeCl₄, methanol, ...), and
- solids (bulk quartz glass, glass fibers)

is used.

Table 4.4.1 shows the Brillouin gain coefficient g and the phonon lifetime τ for different gaseous, liquid, and solid-state SBS media.

A phase-conjugate mirror consists simply of a gas or liquid cell or a fiber piece. The incoming wave is focused into the material where an oppositely traveling wave is generated initially by spontaneous scattering. This wave interferes with the incoming wave and induces a sound wave or another type of phase grating reflecting the incoming beam similarly as a dielectric multilayer mirror. The induced density variations have the frequency of the initial sound wave, which is amplified therefore and reinforces the backscattering. A detailed discussion of stimulated Brillouin scattering is given in Sect. 4.3.3.2.

The amplification depends strongly on the extension of the interference area. Therefore the phase-conjugated backscattered part dominates, leading to an exponential rise of the reflected phase-conjugated signal. The wavefronts of the sound-wave grating match the wavefronts of the incoming beam. Any disturbance of the incident wavefront will result in a self-adapted mirror curvature with response times in the ns range.

For applications the "threshold", reflectivity, and conjugation fidelity are the most important parameters that characterize the performance of a Brillouin-scattering phase-conjugate mirror. A sharply defined threshold does not exist for the nonlinear SBS process. However, after exceeding a certain input energy a steep increase of reflectivity can be observed. Often this is called the energy threshold of the phase-conjugate medium. For long pulses as compared to the phonon lifetime (typically several ns) the SBS is expected to become stationary. In this case the energy threshold can be substituted by a power threshold. Well above this threshold, the reflectivity is not stationary but exhibits statistical fluctuations because SBS starts from noise.

It is important to emphasize that the power and not the intensity determines the "threshold" in case of strongly monochromatic input waves. Slight focusing leads to lower intensity, but also to a longer Rayleigh length and a larger interaction area. Stronger focusing reduces the interaction length, but results in stronger refractive-index modulation. Both effects compensate each other if the interaction length is not limited by the coherence length.

Practically, for most laser sources the coherence length is rather short. Here the interaction length should be short compared to the coherence length. This requires adequate focusing of the beam into the SBS medium. Focal length and scattering material have to be chosen suitable to achieve a high SBS reflectivity and a good reproduction of the wavefront. Side effects in the material like absorption, optical breakdown, or other scattering processes have to be avoided. Figure 4.4.4

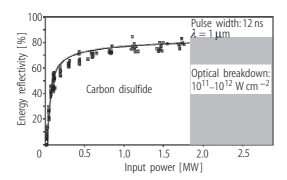


Fig. 4.4.4. Commonly used carbon disulfide (CS₂) shows an SBS threshold of about 18 kW (pulse peak power) under stationary conditions.

Table 4.4.2. SBS threshold, max. reflectivity, far-field fidelity, M^2 -limit, and power limit for different fiber phase conjugators, coherence length 1.5 m. The reflectivity is corrected with respect to Fresnel and coupling losses.

$\begin{array}{c} \hline \\ \text{Core diameter} \\ [\mu m] \end{array}$	SBS threshold [kW]	Maximum reflectivity [%]	Far-field fidelity [%]	M^2 -limit	Power limit [kW]
200	17	80	93	63	160
100	6.4	80	91	31	40
50	2.0	88	70	16	10
25	0.3	86	_	8	2.5

shows the energy reflectivity of carbon disulfide as a function of the input power at 1 μ m wavelength. Carbon disulphide shows one of the smallest power thresholds for liquids of about 18 kW. Applying gases as SBS media, the power thresholds are about one order of magnitude higher. A saturation of energy reflectivity close to 80 % is a typical value for liquid SBS media, although reflectivities up to 96 % had been demonstrated [91Cro].

For high-power input pulses bulk solid-state media like quartz are investigated as SBS media, too [97Yos]. To reduce the power threshold of SBS a waveguide geometry can be applied [95Jac]. The beam intensity inside the waveguide is high within a long interaction length resulting in low power thresholds. To avoid toxic liquids and gases under high pressure multimode quartz fibers can be used [97Eic]. The lower Brillouin gain of quartz glass compared to suitable SBS gases and liquids can be overcome using fibers with lengths of several meters resulting in SBS thresholds down to 200 W peak power [98Eic].

The power threshold P_{th} can be estimated from (4.4.8), where A_{eff} is the effective mode field area inside the fiber core, L_{eff} the effective interaction length, which depends on the coherence length, and g is the Brillouin gain coefficient; for quartz g is about 2.4 cm/GW [89Agr].

$$P_{\rm th} = \frac{21 \ A_{\rm eff}}{L_{\rm eff} \ g} \ .$$
 (4.4.8)

Table 4.4.2 shows the power threshold, the maximum energy reflectivity, the far-field fidelity, the M^2 -limit (see below), and an approximated power limit of fiber phase conjugators with different core diameters. The used quartz-quartz fibers had a step-index geometry and a numerical aperture of 0.22. They were investigated with an Nd:YAG oscillator amplifier system generating pulses of 30 ns (FWHM) at 1.06 μ m wavelength. Regarding applications it is important to couple also spatially aberrated beams into the fiber. The upper limit for the beam parameter product is due to the finite numerical aperture and the core diameter of the fiber. This can by expressed by a "times diffraction limit value" M^2 , see Chap. 2.2 for further information about beam characterization. The upper power limit is approximated assuming a damage threshold above 500 MW/cm² for ns pulses.

An important feature of a fiber phase conjugator is the threshold behavior for different M^2 -values of the incoming beam. In case of a fiber the SBS threshold is nearly independent of the incoming beam quality. This is caused by mode conversion inside the fiber resulting in homogeneous illumination and therefore in constant SBS reflectivity. In case of a Brillouin cell the reflectivity depends on the far-field distribution of the incoming beam. Here phase distortions result in amplitude fluctuations in the focal region. A comparison between a diffraction-limited beam $(M^2 = 1.0)$ and a highly distorted beam $(M^2 = 10)$ showed an increase of the SBS threshold of 300% in case of the Brillouin cell. For the fiber phase conjugator no remarkable changes of the power threshold were observed [97Eic].

Practically, the reproduction of the initial wavefront is not perfect after phase conjugation. To characterize the deviation with respect to the reference wave the term fidelity F is introduced [77Zel]:

$$F = \frac{\left| \int E_{\rm in} E_{\rm p}^* \, \mathrm{d}^2 \, r \right|^2}{\int \left| E_{\rm in} \right|^2 \, \mathrm{d}^2 \, r \cdot \int \left| E_{\rm p} \right|^2 \, \mathrm{d}^2 \, r} \,. \tag{4.4.9}$$

The fidelity equals unity in case of perfect wavefront reproduction and is smaller than unity for practical cases. To calculate the fidelity, the electric field distribution of the incident signal $E_{\rm in}$ and the not perfectly phase-conjugated wave $E_{\rm p}$ – the perfectly phase-conjugated wave is denoted $E_{\rm pc}$ in Sect. 4.4.2 – has to be known. The determination requires sophisticated measurement equipment. In contrary, the *far-field fidelity* can be measured with less effort and is therefore often used. The transmission through an aperture of the phase-conjugate signal is compared with the transmission of the input signal. The ratio is called far-field fidelity, because the aperture is placed in the focal plane of a focusing lens.

4.4.5 Applications of SBS phase conjugation

Phase conjugation generates a wave which retraces the incoming wave in a time-reversed way. Thereby it is possible to eliminate phase distortions in optical systems. For example, in a solid-state laser amplifier, the incoming beam is not only amplified but suffers also from phase distortions due to thermal refractive-index changes in the laser crystal. After passing this amplifier crystal, the beam is reflected by a phase conjugator and passes the crystal a second time. As the wavefronts are inverted with respect to the propagation direction, the refractive-index changes reduce the phase distortions and after the second passage, these distortions disappear so that the beam quality of the incoming wave is reproduced. In Fig. 4.4.5 a double-pass scheme with phase-conjugate mirror to compensate for phase distortions is shown.

Typically, phase conjugators are applied in $Master\ Oscillator\ Power\ Amplifier\ (MOPA)$ setups, where a nearly diffraction-limited master oscillator beam is increased in power within an amplifier

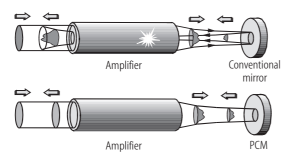


Fig. 4.4.5. Double-pass scheme with phase-conjugate mirror to compensate for phase distortions.

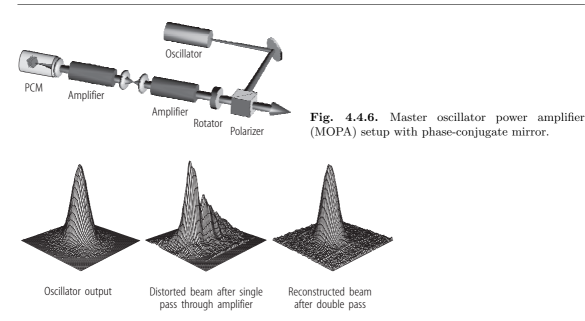


Fig. 4.4.7. Far-field intensity distributions of the oscillator beam, the distorted beam after single-pass amplification, and the highly amplified beam after double-pass amplification with phase conjugation.

arrangement, see Fig. 4.4.6. After the first amplification pass the beam quality is reduced due to thermally induced phase distortions. The spatial-distorted beam enters the SBS mirror and becomes phase-conjugated. The initial beam quality of the master oscillator can be roughly reproduced after the second amplification pass. The amplified beam is extracted with an optical isolation, which consists in this case of a Faraday rotator and a polarizer.

Figure 4.4.6 shows a MOPA system producing up to 210 W average output power at 2 kHz average repetition rate (1.08 µm wavelength). The system is part of an advanced setup yielding up to 520 W average output power [99Eic]. The oscillator beam has a nearly diffraction-limited beam quality ($M^2 < 1.2$) which is already reduced in front of the first amplifier ($M^2 \cong 1.5$). This results from optical components between oscillator and amplifier which introduce phase distortions. After single-pass amplification the beam quality decreases to $M^2 \cong 5$ due to phase distortions introduced by both pumped amplifier rods at 6.5 kW pumping power for each amplifier. After phase conjugation and double-pass amplification the initial beam quality can be nearly reproduced ($M^2 < 1.9$). Differences between the initial and final beam quality are caused by a fidelity smaller than unity and diffraction at several apertures in the amplifier chain.

The performance of the phase-conjugate mirror can be illustrated by far-field intensity profiles recorded at different positions in the setup. In Fig. 4.4.7 the oscillator output beam exhibits a smooth Gaussian profile corresponding to the nearly diffraction-limited beam quality. After single-pass amplification the reduction of beam quality is confirmed by a strongly aberrated far-field profile. After phase conjugation and double-pass amplification the initial intensity distribution can be nearly reproduced. In this example the average power of the master oscillator beam (approx. 1 W) was increased to 130 W after double-pass amplification.

Presently, phase distortion elimination in double or multipass laser amplifiers is the most often application of phase conjugation. In addition phase conjugators are useful as mirrors in laser oscillators replacing one of the conventional mirrors. Again, the phase conjugator eliminates phase distortions in the laser medium induced by optical or discharge pumping. For recent advances and applications of SBS-phase-conjugation see [02Eic, 03Rie, 04Rie].

4.4.6 Photorefraction

The photorefractive effect belongs to the nonlinear optical effects with the highest sensitivity for operation at low optical intensity levels. Photorefractive phase conjugators are able to operate using intensities of only mW/cm^2 . The price paid of the low intensity performance is diminished speed. The response times of recent photorefractive phase conjugators span in the range of milliseconds to several minutes.

The photorefractive effect describes light-induced refractive-index changes in the material when the incident light is spatially nonuniform [88Gue, 93Yeh, 95Nol, 96Sol]. The spatial nonuniformity distinguishes the photorefractive effect from other common nonlinear optical effects that occur under spatial uniform intensity. The maximum refractive-index change induced in a photorefractive material does not occur necessarily locally where the light intensity is a maximum. The nonlocal response occurs because electric charges move and are stored inside the material. In case of classical nonorganic bulk photorefractive materials, such as ferroelectric oxides (BaTiO₃, LiNbO₃, KNbO₃), sillenites (Bi₁₂SiO₂₀, Bi₁₂TiO₂₀, Bi₁₂GeO₂₀) or semi-insulating semiconductors (GaAs, InP, CdTe), electrons (or holes) are photoexcited from localized impurity centers or defect sites, which are energetically located deep in the band gap of the material, into the conduction (or valence) band.

The energy of the exciting photons is smaller than the band-gap energy. Free carriers excited in bright crystal regions move due to diffusion and drift into dark crystal regions where they are trapped by empty defect sites, see Fig. 4.4.8. As a consequence of separated and trapped electric charges the formation of space-charge electric fields occurs. These electric fields change the refractive index of the material by electrooptics effects, usually the Pockels effect.

Nonuniform illumination occurs when two coherent laser beams interfere in the crystal. The intersecting beams create a periodical interference pattern. The formation of a photorefractive index grating due to a sinusoidal intensity pattern is shown in Fig. 4.4.9. When diffusion is the main effect for the transport of the excited charge carriers (there is no external electric field applied on the crystal) the electric-field maxima are shifted by a quarter fringe spacing relative to the intensity maxima. This $\pi/2$ phase shift of the induced index grating plays a fundamental role in photorefractive non-linear optical wave mixing. It allows for an energy transfer between the two beams writing the grating in a process called two-wave mixing. One of the beams (called signal beam) is amplified at the expense of the other beam (called pump beam).

A phase-conjugate beam can be created by four-wave mixing processes. In this case the two-wave mixing arrangement is extended with a second pump beam which counterpropagates with respect to the first pump beam, see Fig. 4.4.3. In case of external pump beams, the phase-conjugation

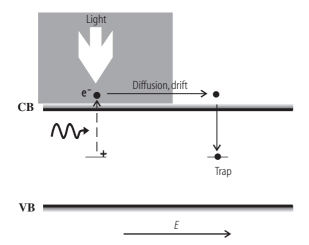


Fig. 4.4.8. Band transport model of photorefraction.

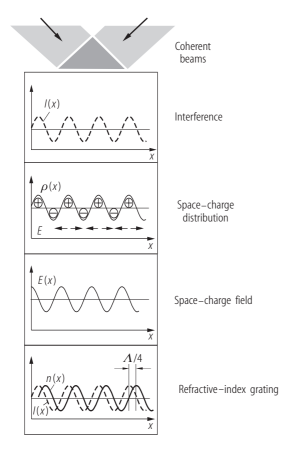


Fig. 4.4.9. Formation of a photorefractive index grating due to a sinusoidal intensity pattern.

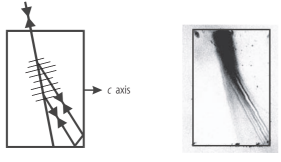


Fig. 4.4.10. Scheme of photorefractive total-internal-reflection phase conjugator (cat conjugator), left. The light propagation in the crystal can be seen due to scattering, right.

process may be highly efficient leading to large reflectivities well above $100\,\%$ in relation to the incoming power.

Self-pumped phase conjugators require only a single incident beam and because of their simplicity they are more advantageous for practical applications. The operation of photorefractive self-pumped phase conjugators is based on a non-linear optical process called beam fanning. When a single beam is incident on a photorefractive crystal, some light is scattered inside the crystal. This scattered light forms a set of gratings with the incident light and is amplified by two-wave mixing. This process was named fanning because a broad fan of scattered amplified light is generated emerging from the crystal.

Perhaps the most commonly used photorefractive self-pumped phase conjugator type is the so-called cat conjugator [82Fei]. In this case the first pump beam is generated from the incident beam by fanning, the second pump beam by backreflection on the crystal corner. Figure 4.4.10 shows a rhodium-doped barium titanate crystal which acts as a cat conjugator for an incident beam of $5~\mathrm{mW}$

244 4.4.6 Photorefraction [Ref. p. 245]

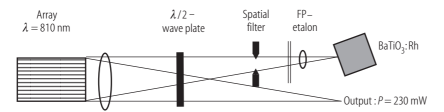


Fig. 4.4.11. Coherent diode laser array coupled by a phase-conjugating BaTiO₃:Rh crystal [98Lob].

optical power at 808 nm wavelength. The formed internal phase-conjugation loops can be observed in the lower right-hand corner of the crystal. Self-pumped phase-conjugate reflectivities as high as $60-80\,\%$ have been reported for visible and near-infrared wavelengths by numerous investigators using photorefractive crystals in various arrangements [85Gue, 86Pep, 95Mu, 94Wec, 97Huo].

The efficient operation of photorefractive phase conjugators at low and moderate power levels makes this type of device attractive especially for diode-laser applications. Free-running high-power diode laser arrays emit laser beams of poor spatial and spectral quality. Optical phase-conjugate feedback can increase both the spatial and the temporal coherence of the radiation. Figure 4.4.11 shows an external-cavity diode laser system comprising a photorefractive BaTiO₃ crystal as phase conjugator, a Fabry-Perot etalon, and a spatial filter forcing the laser diode array to operate in a single spatial and a single longitudinal mode [98Lob]. The coherence length of the phase-conjugate laser system has been increased by a factor of 70 and the output has become almost diffraction-limited. The output power is reduced from 440 mW to 230 mW.

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Index

Landolt-Börnstein New Series VIII/1A1

α -HIO ₃ 143, 183	Acetic acid 224, 229
α -InO ₃ 145, 165 α -Iodic Acid 143	Acetone 224, 226, 229, 230, 238
β -BaB ₂ O ₄ 142	Achromatic correction 97
μ -Dab ₂ O ₄ 142 π -pulse 36	Acoustic phonons 217
⁶ LiNbO ₃ 227	ACOUSTIC PHOHOIS 217 ADA 143, 165
6LiTaO ₃ 227	Adiabatic
-	
LiNbO ₃ 227	equations 20
1,1,1-Trichloroethane 226	pulse amplification 25
1,2-Dichloroethane 224	ADP 143, 160, 161, 164, 166, 168, 176, 180
1,2-Diethylbenzene 224	Ag_3AsS_3 142, 166, 171, 173, 186
1,2-Dimethylcyclohexane 224	Ag_3SbS_3 142
1,3-Dibromobenzene 224	$AgGaS_2$ 142, 166, 171, 173, 186
1,3-Pentadiene 225	$AgGaSe_2$ 142, 166, 174, 186
1,4-Dimethylcyclohexane 224	Air
1,4-Dimethylnitrobenzene 231	thick lens in 113
1,4-Dioxane 224	Airy's disc 92
1-Bromonaphthalene 225, 231	Al_2O_3 225
1-Bromopropane 224	Ammonia 225
1-Chloronaphthalene 225	Ammonium Dihydrogen Arsenate 143
1-Fluoro-2-chlorobenzene 224	Ammonium Dihydrogen Phosphate 143
2π -pulse 35	Amplification
2,2-Dichlorodiethylether 224	stimulated 221
2,4-Dinitrotoluene 231	Amplifier
2-Bromopropane 224	feed-back 3
2-Ethylnaphthalene 225	Angle
2-Nitropropane 225	birefringence 145
3-Methyl-4-Nitro-Pyridine-1-Oxide 142	Brewster's 101
3-Methylbutadiene 224	divergence 82
4-Hydroxy-3-Methoxy-Benzaldehyde (Vanillin)	walk-off 145
142	Angular
	moment 56
Abbe's number 97	-spectrum representation 87
ABCD	Aniline 224, 229
matrix 111	Annular aperture 92
transformation 120	Anti-Stokes
Aberration 127	emission, higher-order 222
lens 117	line 218
spherical 117	Raman scattering, coherent 222
third-order 119	scattering 222
Absorber	9
	Aperture
saturable 237	annular 92
Absorbing	circular 90
gas 237	length 151
liquid 237	rectangular 89
Absorption	Apodization
excited-state 237	Gaussian 114

Approach	Gauss
perturbation 206	-Hermite 81
Approximation	-Laguerre 83
dipole 12	-Schell model 53
far-field, Fraunhofer 87	Gaussian 53, 81
Fraunhofer 88	elliptical 57
far-field 87	general astigmatic 64, 121, 123
Fresnel's 85, 86	Hermite–Gaussian 53
Fresnel-Kirchhoff 86	higher-order 123
plane-wave 152	Laguerre–Gaussian 53
Rayleigh–Sommerfeld 86	ordinary 145
rotating-wave 14	partially coherent 54
Ar 211	positional stability 67
Argon laser 166	principal axis of 57
Astigmatic	propagation 120
beam	Gaussian 111
general $121, 123$	ratio, effective 60
simple $120, 121$	pseudo-symmetric 64
general 64	simple astigmatic 63, 120, 121
simple 63	stigmatic 62, 120, 121
system	waist 82
general 116	width 57
Astigmatism	long-term 69
inner 64	Benzaldehyde 224, 229
intrinsic 60,65	Benzene 224, 226, 229–231
Axis	Benzene-d ₆ 224
principal 58	Benzonitrile 224
of the beam 57	Benzonitrol 231
Azoxybenzene 231	Benzoylchloride 231
Do 910	Benzylidenaniline 231
Ba 210 Pa NaNh O 142 227	Bessel beam
$Ba_2NaNb_5O_{15}$ 142, 227	diffraction-free 79
Banana (Barium Sodium Niobate) 142, 158, 179, 183	real 80
	vectorial 80
Barium Sodium Niobate (Banana) 142, 158, 179, 183	Beta-Barium Borate 142
Barium vapor 225	$Bi_{12}GeO_{20}$ 242
Basov 3	$Bi_{12}SiO_{20}$ 242
BaTiO ₃ 237, 242, 244	$Bi_{12}TiO_{20}$ 242
BBO 142, 157–161, 163–166, 168, 172, 177, 181,	Biaxial crystal 105, 145
184	Birefringence 106
Be 210	angle 145
Beam	Bisectrix 147
Bessel	Black-body radiator 47
diffraction-free 79	Bloch vector 33
real 80	Bolometer 48
vectorial 80	Brewster angle 101
characterization 53	-tilted plate 76
classification 61	Brillouin
conversion 65	doublet 218
diameter	gain coefficient 238
generalized 57	lines 217
diffraction-free 79	scattering
elliptical Gaussian 57	stimulated $217, 227, 229, 236$
extraordinary 145	stimulated, thermal 220, 227
fanning 243	Broadening
fluctuation matrix 67	collision 28

Doppler 28	Chloroform 224, 229
homogeneous 29, 223	Chloromethylbutane 224
inhomogeneous 30, 223	Chloronaphthalene 231
line 26, 28	Circular aperture 90
pressure 28	Classification 104
saturation 29	beam 61
types of 29	
Bromobenzene 224, 226, 229, 231	CLBO 142, 157, 160, 161
, , ,	CO 211
Bromopropane 224 Butane 225	laser 166
	CO ₂ laser radiation
Butyl-benzene (tert.) 224	harmonic generation of 166
C II N O 140	up-conversion of 171
$C_{10}H_{11}N_3O_6$ 142	Coefficient
$C_{10}H_{13}N_3O_3$ 142	gain $18,229$
$C_{11}H_{14}N_2O_3$ 142	Miller 147
$C_2F_6 = 238$	nonlinear
$C_6H_6N_2O_3$ 142	effective 150
$C_8H_8O_3$ 142	scattering 226
Ca 210	Seebeck 48
Ca^{+} 210	Coherence length 238
Cadmium Germanium Arsenide 142	Coherent
Cadmium Selenide 142	anti-Stokes Raman scattering (CARS) 222
Calcite 225, 227	interaction 31
Calculus	partially 54
Jones 75	pulses, resonant 34
Mueller 77	÷ '
Calorimeter 50	Collins integral 126
Calorimetry	Collision broadening 28
photoacoustic 50	Coma 119
Carbondioxide 225	Complex
Carbondisulfide 224, 226, 229–231	notation 6
Carbonmonoxide 225	q-parameter 120
Carbontetrachloride 224, 229	refractive index 8, 103
Cardinal	susceptibility 10
plane 110	Concave
point 110	grating
CARS (coherent anti-Stokes Raman scattering)	Rowland 114
222	mirror
Cat conjugator 243	spherical 114
CBO 142, 159, 160	Condition
CCl ₄ 238	phase-matching 208
Cd 210	resonance 5
CDA 142, 156, 180	steady-state 5
$CdGeAs_2$ 142, 166, 167, 174	symmetry
CdSe 142, 171, 174, 186	Kleinman 150
CdTe 242	threshold 4
	Conductivity
Centered moment 55	electric 6
Cesium Borate 142	Conjugation fidelity 238
Cesium Dideuterium Arsenate 142	v o
Cesium Dihydrogen Arsenate 142	Conjugator cat 243
Cesium Lithium Borate 142	Conservation
Cesium vapor 226	
CH ₄ 238	of momenta 208
Characterization	Constant
beam 53	time 31
Chlorine 226	Construction
Chlorobenzene 224, 226, 231	Listing's 110

Landolt-Börnstein New Series VIII/1A1

	G 11
Continuous wave	Cyclohexane 224, 229
optical parametric oscillation 176	Cyclohexanone 224
Continuum generation	Cylindrical
picosecond 186	vector wave 78
in crystals 186	wave 79
Contribution	
electrostrictive 227	$D_2, Q(2) = 227$
Conversion	DAN 142, 157
beam 65	Data
efficiency	of crystals 108
quantum 153	of gases 108
factor for SI and CGS-esu systems 150	of infrared materials 108
frequency	of liquids 108
efficiency 151	of metals 108
	of negative-refractive-index materials 108
in gases 209 Correction	of optical glass 108
	of photonic crystals 108
achromatic 97	of polymeric materials 108
Coupling	of semiconductors 108
laser fiber 129	of solid state laser materials 108
Stokes-anti-Stokes 222	DCDA 142, 156, 161
waveguide 127, 128	Decay
Cross section 18	time T_1 15
differential 217	time T_2 16
Raman scattering 223	Degenerate four-wave mixing 235, 236
scattering 217	Degree
differential 227	of polarization 77
Raman 223	Delta formulation, Miller 147
Cross-spectral density 53	Density
Cryogenic radiometer 47, 50	cross-spectral 53
Crystal	current 5
biaxial 105, 145	distribution
data of 108	power 55
isotropic 105	power, far-field 55
laser 237	inversion 12, 13, 17
liquid 237	matrix 206
negative uniaxial 105	of electric charges 5
optical 105	Dephasing 227
optics 104	Depth Depth
photonic, data of 108	penetration 102
photorefractive 237	Detector
picosecond continuum generation in 186	pyroelectric 49
positive uniaxial 105	quantum 49
uniaxial 145	thermal 48
negative 146	Determination
positive 146	of the ten second-order moments 66
Cs 210	Deuterium 226
CS ₂ 238	DFWM (Degenerate Four-Wave Mixing) 236
CsB_3O_5 142	Diameter Diameter Diameter
CsD_2AsO_4 142	beam, generalized 57
CsH_2AsO_4 142	Diamond 225
$CsLiB_6O_{10}$ 142	Dichloromethane 224
$CuAlS_2$ 227	Dielectric medium 96
Cubic 105	Dielectrics Dielectrics
Current density 5	homogeneous 6
Curvature	isotropic 6
phase, generalized 59	linear 6
phase, generanzed 99	micai U

Difference frequency generation	144, 153, 172	Effective
generation of IR radiation by	172	beam propagation ratio 60
in the far IR region 175		length 151
Differential		nonlinear coefficient 150
cross section 217		nonlinearity 147
scattering		Efficiency .
cross section 227		conversion
Diffraction 84		quantum 153
efficiency 92		diffraction 92
figure		frequency-conversion 151
Fresnel's 93		Einstein coefficient
-free		of induced emission 21
beam 79		of spontaneous emission 21
Bessel beam 79		Electric
length 151		charges, density of 5
pattern		conductivity 6
Fraunhofer 89		displacement 5
scalar theory of 85		field 5
theory		polarization 6
Rayleigh–Sommerfeld–Debye	89	susceptibility 6
time-dependent 89	00	Electromagnetic field 5
vector theory of 85		Electrostrictive contribution 227
Diffractive optics 94		Ellipse
Diffuse emitter 47		variance 57
Dimethylhexadiene 224	1	Ellipsoid
Dimethylsulfoxide (DMSO) 224	Ł	index 105
Dipole		Ellipsometry 104
approximation 12		Elliptical Gaussian beam 57
Hertz's 8		Emission
moment 11, 15		anti-Stokes, higher-order 222
oscillating 8		induced
Disc		Einstein coefficient 21
Airy's 92		spontaneous $4, 15, 28$
Dispersion		Einstein coefficient 21
formula 97		Stokes, higher-order 222
-spreading length 151		Emitter
Displacement		diffuse 47
electric 5		Energy
Distance		flux 7
Rayleigh 82		radiant $45,46$
Distribution		relaxation 15
power density 55		Entropy relaxation 16
far-field 55		Equation
Wigner 53		adiabatic 20
Divergence angle 82		fundamental
DKB5 143, 165		of nonlinear optics 207
		Helmholtz 74, 79
· · · · · · · · · · · · · · · · · · ·		•
DKDP 143, 156, 159, 160, 162		Maxwell's 5
DMSO (dimethylsulfoxide) 224		Maxwell–Bloch 15, 16
Doped media		rate 20
propagation in 10		slowly varying envelope (SVE) 74,80
Doppler broadening 28		SVE (slowly varying envelope) 74,80
Doublet		wave 73,78
Brillouin 218		Ethanol 224, 226, 229, 230
Dye laser radiation		Ethyl-Benzene 224
second harmonic generation of	164	Ethylene 225

Fourth harmonic generation 153, 156 Eu 210 Excitance of Nd:YAG laser radiation 160 of Ti:sapphire laser radiation 164 radiant 46 Excited-state absorption Fraunhofer External reflection 101 approximation Extraordinary beam 145 far-field diffraction pattern 89 Factor far-field approximation 87 conversion Free-space propagation 61 for SI and CGS-esu systems Freon 238 gain 220, 226, 227 Frequency slit 92 conversion Far field 58 efficiency 151 approximation gases 205 Fraunhofer 87 in gases 209 fidelity 239 liquids 205 Fraunhofer approximation 87 mixing 208 power density distribution Rabi 33 Far IR region spatial 88 difference frequency generation in 175 Fresnel Faraday rotator 76 approximation 85,86 Feed-back amplifier 3 diffraction figure 93 Femtosecond optical parametric oscillation 184 formulae 98,99 Ferroelectric oxide 242 -Kirchhoff approximation 86 Feynman representation 32 number 88 Fiber Fundamental equations laser coupling 129 of nonlinear optics 207 phase conjugator 240 Fidelity 240 GaAs 175, 186, 242 conjugation 238 Gain far-field 239 coefficient 18, 229 Field Brillouin 238 electric 5 factor 220, 226, 227 electromagnetic 5 small-signal 11 far 58 Gallium Selenide 142 magnetic 5 GaP 175, 227 near 56 Gas 205 Fifth harmonic generation 153, 156 absorbing 237 of Nd:YAG laser radiation 161 data of 108 Fluctuation matrix -eous media, mixture of 209,211beam 67 frequency conversion in 205, 209Fluorinert laser radiation FC 72 229 second harmonic generation of 166 FC 75 229 mixture, metal-vapor inert 209, 210 Fluorobenzene Raman Fluoromethane 224parameters 227 Formula scattering 225 dispersion 97 scattering Fresnel's 98, 99 Raman 225 Sellmeier's 97 GaSe 142, 167, 174, 186Four-level system 24 Gauss Four-wave -Hermite beam 81 interaction 221 -Laguerre beam 83 mixing, degenerate 235, 236-Schell model 53 Fourier beam 53 optics 94 Gaussian transform 88 apodization 114

beam 53,81	DF-3 229
elliptical 57	optical 97
propagation 111	data of 108
imaging 108	Glycerol 229
line shape 27	Goos–Hänchen shift 103
matrix 111	Gradient-index lens 113
$Gd_2(MoO_4)_3$ 225	Grating 92
$GeCl_4$ 238	concave
General	Rowland 114
astigmatic	thermal 237
beam 64, 121, 123	Group
system 116	point 150
·	pomo 100
parabolic	$H_2, Q(1) = 227$
system 112	Half-wave plate 76
ray-transfer matrix 117	Harmonic generation 156
Generalized	of CO ₂ laser radiation 166
beam diameter 57	of high-power Nd:glass laser radiation 162
phase curvature 59	of iodine laser radiation 162
Generation	of Nd:YAG laser radiation 161
continuum	
picosecond 186	Harmonics
picosecond, in crystals 186	of high-power Nd:glass laser radiation 162
difference-frequency 144, 153, 172	of iodine laser radiation 162
generation of IR radiation by 172	of Nd:YAG laser radiation 161
in the far IR region 175	He 211
fifth harmonic 153, 156	He-Ne laser 166
of Nd:YAG laser radiation 161	Heat pipe 209
fourth harmonic 153, 156	Helmholtz equation 74,79
of Nd:YAG laser radiation 160	Hermite–Gaussian beam 53
of Ti:sapphire laser radiation 164	higher-order 123
harmonic 156	Hertz's dipole 8
of CO ₂ laser radiation 166	Hexagonal 105
of harmonics	Hg 210, 211
	$HgGa_2S_4 $ 143, 171
of high-power Nd:glass laser radiation 162	Higher order
of iodine laser radiation 162	anti-Stokes emission 222
of Nd:YAG laser radiation 161	Hermite–Gaussian beam 123
second harmonic 144, 152, 156	Stokes emission 222
in "nonlinear regime" 154	Hole-burning
of dye laser radiation 164	spatial 237
of gas laser radiation 166	Holography 237
of Nd:YAG laser radiation 156, 157, 159	Homogeneous 6
of Nd:YAG laser radiation, intracavity 158	broadening 29, 223
of ruby laser radiation 163	dielectrics 6
of semiconductor laser radiation 164	system 16
of Ti:sapphire laser radiation 163	Huygens' principle 8,85
sixth harmonic 156	Hydrogen 226
of Nd:YAG laser radiation 161	Hydrogenbromide 226
sum-frequency 144, 153, 167	Hydrogenchloride 226
of UV radiation 167–169	Hyperbolic 220
third harmonic 152, 156, 208	propagation law 60
of Nd:YAG laser radiation 159	propagation iaw 00
of Ti:sapphire laser radiation 163	Imaging
Geometric optical radiance 54	Imaging Gaussian 108
	Gaussian 108 Index
Geometrical optics 108	
Glass	ellipsoid 105
BSC-2 229	surface 105, 146

T 1:	
Indicatrix	generation by difference frequency generation
optical 145	172
Induced emission	Irradiance 46
Einstein coefficient 21	Isopropanol 224, 226 Isotropic 7
Induction	crystal 105
magnetic 5	dielectrics 6
Infrared material, data of 108	
Inhomogeneous broadening 30, 223	Jones
broadening 30, 223 system 16	calculus 75
Inner astigmatism 64	matrix 75
InP 242	vector 75
Input	
noise 220	K 210
InSb 229	$KB_5O_8 \ 4D_2O \ 143$
Integral	$KB_5O_8 \ 4H_2O \ 143$
Collins 126	KB5 143, 161, 162, 165, 166, 169
Intensity 8	KD_2AsO_4 143
radiant 46	KD ₂ PO ₄ 143
saturation 18	KDP 143, 156, 159–164, 166, 167, 176, 180, 186
Interaction 14	KH ₂ PO ₄ 143 Kleinman symmetry conditions 150
coherent 31	KNbO ₃ 143, 157, 158, 163–165, 179, 185, 242
four-wave 221	Kr 211
Hamiltonian 14	Kramers–Kronig relation 97
length	KTA 143, 173, 179, 183, 185
nonlinear 152	KTiOAsO ₄ 143
quasistatic 151	$KTiOPO_4$ 143
nonlinear	$\mathrm{KTP} 143, 156 – 159, 170, 173, 178, 179, 182 – 185$
length 152	
quasistatic	Laguerre
length 151	-Gaussian beam 53
three-wave 144	polynomial 83
Internal reflection 101	Lamb dip 26
Intracavity second harmonic generation of Nd:YAG	Lambert's cosine law 47
laser radiation 158	Lambertian radiator 47
Intrinsic astigmatism 60,65	Laser
Inversion density 12, 13, 17	argon 166 CO 166
Iodine laser radiation	crystal 237
harmonics of 162	energy meter 50
IR	fiber coupling 129
far	He-Ne 166
region, difference frequency generation in 175	NH ₃ 166
material, data of 108	oscillator 3
mid	theory, semiclassical 4
region, optical parametric oscillation in 186	Law
near	Planck's 21
region, cw optical parametric oscillation in	propagation, hyperbolic 60
176	LBO 143, 157–159, 161, 163, 169, 177, 178, 181,
region, femtosecond optical parametric	182, 184
oscillation in 184	Length
region, nanosecond optical parametric	aperture 151
oscillation in 176	coherence 238
region, picosecond optical parametric oscillation in 180	diffraction 151
radiation	dispersion-spreading 151 effective 151
rauration	enective 191

interaction	Raman
nonlinear 152	parameters 226
quasistatic 151	scattering 224
nonlinear interaction 152	scattering
quasistatic interaction 151	Raman 224
Rayleigh 61	Listing's construction 110
Lens	Lithium Fomate 143
aberration 117	Lithium Iodate 143
gradient-index 113	Lithium Niobate 143
shape factor of 117	Lithium Triborate 143
thermal 113	Long-term beam width 69
thick 110	Lorentzian line shape 27
in air 113	m-Dinitrobenzene 231
thin 113	
LFM 143, 156, 165	m-Nitrotoluene 229, 231 m-Xylene 225
$Li_2B_4O_7$ 160, 161	Magnetic 223
LiB ₃ O ₅ 143	field 5
LiCOOH H ₂ O 143	induction 5
Lifetime	polarization 6
upper-level, T_1 18	susceptibility 6
Light	Maiman 3
-induced refractive-index change 242	MAP 142, 157
partially polarized 77	Master oscillator power amplifier 240
pressure 50	Matching
LiHCOO 225	mode 128
LiIO ₃ 143, 156, 158, 159, 161, 163, 165, 166, 170,	phase 144
172, 175, 176, 180, 186	condition 208
LiNbO ₃ 143, 156, 158, 159, 161, 166, 170, 172, 175,	noncollinear 146
176, 180, 186, 237, 242 Linbornary 142, 150, 176, 180	Material
LiNbO ₃ :MgO 143, 159, 176, 180 Line	infrared, data of 108
anti-Stokes 218	negative-refractive-index, data of 108
broadening 26, 28	optical 95
Raman	photorefractive 242
vibration-rotation 218	polymeric, data of 108
vibrational 218, 223	solid-state-laser, data of 108
scattering	Matrix
Rayleigh, stimulated 228	ABCD 111
shape 26	density 206
Gaussian 27	fluctuation
Lorentzian 27	beam 67 Gaussian 111
normalization of 27	Jones 75
normalized 27	ray-transfer 111
spectral 17	general 117
Stokes 218	system 55
width 26	variance 56
Linear 7	Maxwell
dielectrics 6	-Bloch equations 15, 16
optics 73	equations 5
susceptibility 205	Medium
Linewidth 226, 227	dielectric 96
Liquid 205	Meissner 3
absorbing 237	Mercury Thiogallate 143
crystal 237	meta-Nitroaniline 143
data of 108	Metal
frequency conversion in 205	data of 108

optics of 98	Multimode oscillation 26
vapor	Mutual power spectrum 53
inert gas mixture 209, 210	
mixture of different 209, 210	$n\pi$ -pulse 35
Methane 226	N_2 211, 238
Methane, Q 227	N_2O 226
Methanol 224, 226, 229, 230, 238	N-(4-Nitrophenyl)-(L)-Propinol 142
Methanol-d ₄ 224	N-[2-(Dimethylamino)-5-Nitrophenyl]-Acetamide 142
Methyl N-(2,4-Dinitrophenyl)-L-Alaninate 142	n-Hexane 229
Methylodide 229	n-Nitrotoluene 229
Mg 210	Na 210
Mg ⁺ 210	NaClO ₃ 225
Mg:O-doped Lithium Niobate 143	Nanosecond optical parametric oscillation 176
MHBA 142, 157	Naphthalene 225, 231
Mid IR region	Nd:glass high-power laser radiation
optical parametric oscillation in 186 Miller	generation of harmonics 162
coefficient 147	harmonics 162
delta formulation 147	Nd:YAG laser radiation
Mirror	fifth harmonic generation of 161
concave	fourth harmonic generation of 160
spherical 114	harmonics of 161
phase-conjugate 235	intracavity second harmonic generation of 158
self-adapted 238	second harmonic generation of 156, 157, 159
spherical 113	second harmonic generation, intracavity, of 158
concave 114	sixth harmonic generation of 161
Misalignment 116	third harmonic generation of 159
Mixed moment 56	Ne 211 Near field 56
Mixing	Near field 56 Near IR
frequency 208	radiation
two-wave 242	up-conversion of 170
Mixture	region
metal-vapor inert gas 209, 210	cw optical parametric oscillation in 176
of different metal vapors 209, 210	femtosecond optical parametric oscillation in
of gaseous media 209, 211	184
mNA 143, 157	nanosecond optical parametric oscillation in
Mode	176
competition 26	picosecond optical parametric oscillation in
hopping 26	180
matching 128	Negative
Model Gauss-Schell 53	refractive index material, data of 108
Moment 55	uniaxial crystal 105, 146
angular 56	$(NH_2)_2CO$ 143
centered 55	NH ₃ laser 166
dipole 11, 15	$NH_4H_2AsO_4$ 143 $NH_4H_2PO_4$ 143
mixed 56	Nitroacetophenone 231
second-order 55	Nitrobenzaldehyde 231
determination of 66	Nitrobenzene 225, 226, 229, 231
radiation field, propagation of 111	Nitrogen 225, 226
spatial 56	NO 226
Momentum	$NO_2C_6H_4NH_2$ 143
conservation of 208	Nodal point 110
Monoclinic 105	Noise input 220
MOPA (Master Oscillator Power Amplifier) 240	Non-symmetrical optical system 112
Mueller calculus 77	Noncollinear phase matching 146

Nonlinear	Oscillating dipole 8
coefficient	Oscillation
effective 150	multimode 26
interaction length 152	optical parametric 144, 153, 176
optics	continuous wave 176
fundamental equations of 207	femtosecond 184
regime	in the mid IR region 186
second harmonic generation in 154	nanosecond 176
susceptibility 205	picosecond 180
third-order 206, 221	Oscillator
Nonlinearity	self-sustained 3
effective 147	Oxide
Normalization	ferroelectric 242
of line shapes 27	Oxygen 226
Normalized	
line shape 27	p-Dichlorobenzene 229
shape function 27	p-Nitroanisol 231
Notation	p-Nitrotoluene 231
complex 6	p-Xylene 225, 229
NPP 142, 179, 185	Parabolic system, general 112
Number	Paraboloid
Abbe's 97	phase 59
Fresnel 88	Paraxial range 108
	Partially
o-Nitroaniline 231	coherent beam 54
o-Nitrophenol 231	polarized light 77
o-Nitrotoluene 231	PCM (Phase-Conjugate Mirror) 235
o-Xylene 225	Penetration depth 102
Octanol 229	Permeability 6
Optical	Permittivity 6
crystal 105	Perturbation approach 206
glass 97	Phase
data of 108	-conjugate mirror 235
indicatrix 145	conjugation 235
material 95	self-pumped 237
parametric oscillation 144, 153, 176	conjugator 235
continuous wave 176	fiber 240
femtosecond 184	photorefractive 242
in the mid IR region 186	self-pumped $235, 243$
nanosecond 176	vector 236
picosecond 180	curvature, generalized 59
radiometry 45	matching 144
self-focusing 220	condition 208
system	noncollinear 146
non-symmetrical 112	paraboloid 59
symplectic 116	Phonons
Optics	acoustic 217
crystal 104	Photoacoustic calorimetry 50
diffractive 94	Photoconductor 49
Fourier 94	Photodiode 49
geometrical 108	Photometric quantities 45
linear 73	Photonic crystals, data of 108
nonlinear, fundamental equations of 207	Photorefraction 242
of metals 98	Photorefractive
of semiconductors 98	crystal 237
Ordinary beam 145	material 242
Orthorhombic 105	phase conjugator 242

Landolt-Börnstein New Series VIII/1A1

D: 1	D
Picosecond	Pressure
continuum generation 186	broadening 28
in crystals 186	light 50
optical parametric oscillation 180	Primary standards 47
Pipe	Principal
heat 209	axis 58 of the beam 57
Piperidine 225	value 145
Planar plate 113	Principle
Planck's law 21	-
Planckian radiation 47	Huygens' 8, 85 Process
Plane 113	
cardinal 110	pumping 16 Prokhorov 3
wave 7,79	Propagation 60
approximation 152	beam 120
representation 87	Gaussian 111
Plate	free-space 61
Brewster-angle-tilted 76	in doped media 10
half-wave 76	law
planar 113	hyperbolic 60
quarter-wave 76	of the second-order moments of the radiation
Point	field 111
cardinal 110	short-pulse 97
group 150	three-dimensional 127
nodal 110	two-dimensional 126
Polarization 12, 17, 75	Proustite 142
degree of 77	Pseudo-symmetric beam 64
electric 6	Pulse
magnetic 6	2π - 35
Polarized	amplification, adiabatic 25
partially	$n\pi$ - 35
light 77	π- 36
Polydiacetylene 225	resonant coherent 34
Polymeric materials, data of 108	short, propagation of 97
Polynomial	Pumping
Laguerre 83	process 16
POM 142, 157	schemes 22
Position	Pyrargyrite 142
waist 60	Pyridine 225, 229
Positional stability	Pyroelectric detector 49
beam 67	V
Positive uniaxial crystal 105, 146	q-parameter, complex 120
Potassium Dideuterium Arsenate 143	Quantities
Potassium Dideuterium Phosphate 143	photometric 45
Potassium Dihydrogen Phosphate 143	radiometric 45
Potassium Niobate 143	Quantum
Potassium Pentaborate Tetradeuterate 143	conversion efficiency 153
Potassium Pentaborate Tetrahydrate 143	detector 49
Potassium Titanyl Arsenate 143	Quarter-wave plate 76
Potassium Titanyl Phosphate 143	Quartz 175, 227, 229, 238
Power	Quasistatic interaction length 151
density distribution 55	-
radiant 46	Rabi frequency 33
spectrum 53	Radiance 46
mutual 53	geometric-optical 54
Poynting vector 7	spectral 45

Radiant	-Sommerfeld
energy $45,46$	approximation 86
excitance 46	–Debye diffraction theory 89
intensity 46	wing scattering 217
power 46	wing scattering, stimulated 228, 230, 231
Radiation	Rb 210
field	RbH_2AsO_4 143
propagation of the second-order moments 111	RbH_2PO_4 143
Planckian 47	RbTiOAsO ₄ 143
synchrotron 47	RDA 143, 156, 159, 163
Radiator	RDP 143, 156, 159, 163, 164
black-body 47	Real Bessel beam 80 Rectangular aperture 89
lambertian 47	Rectangular aperture 89 Reflection
Radiometer	external 101
cryogenic 47, 50	internal 101
Radiometric	total 101
quantities 45	Refraction 101
standards 47	at a sphere 114
Radiometry 45	Refractive index
optical 45	change, light-induced 242
Raman	complex 8, 103
	Region
vibration-rotation 218	X-ray 47
vibrational 218, 223	Relation
parameters gases 227	Kramers-Kronig 97
liquids 226	Relaxation
solids 227	energy 15
scattering	entropy 16
cross section 223	Replica
gases 225	time-reversed 236
liquids 224	Representation
solids 225	angular-spectrum 87
stimulated 217, 223, 226	plane-wave 87
Range	Resonance condition 5
paraxial 108	Resonant coherent pulses 34
Seidel's 109	Responsivity 48
Rate equations 20	Rotating wave approximation 14 Rotator
Ray	Faraday 76
surface 105	Rowland concave grating 114
tracing 55	RTA 143, 179, 185
transfer matrix 111	Rubidium Dihydrogen Arsenate 143
general 117	Rubidium Dihydrogen Phosphate 143
Rayleigh	Rubidium Titanyl Arsenate 143
distance 82	Ruby laser radiation
length 61	second harmonic generation of 163
line scattering 217	G
stimulated 228	Saturable absorber 237
scattering	Saturation
line 217	broadening 29
line, stimulated 228	intensity 18
stimulated 217, 228, 230	of the two-level transition 17
stimulated, thermal 220, 228	SBS (stimulated Brillouin scattering) 227, 236
wing 217	threshold 238, 239
wing, stimulated 228, 230, 231	Scalar diffraction theory 85

Scattering	Secondary standards 48
anti-Stokes 222	Seebeck coefficient 48
anti-Stokes Raman, coherent 222	Seidel's range 109
Brillouin	Self-adapted mirror 238
stimulated 217, 227, 229, 236	Self-focusing
stimulated, thermal 220, 227	optical 220
coefficient 226	Self-pumped
coherent	phase conjugation 237
anti-Stokes Raman 222	phase conjugator 235, 243
cross section 217	Self-sustained oscillator 3
differential 227	Sellmeier's formula 97
Raman 223	Semiclassical laser theory 4
Raman	Semiconductor
cross section 223	data of 108
gases 225	laser radiation
liquids 224	second harmonic generation of 164
solids 225	optics of 98
stimulated 217, 223, 226	$SF_6 = 226, 238$
Rayleigh	Shape
line 217	factor of a lens 117
line, stimulated 228	function, normalized 27
stimulated 217, 228, 230	line 26
stimulated, thermal 220, 228	Shift
wing 217	Goos-Hänchen 103
wing, stimulated 228, 230, 231	Stokes 224–227
spontaneous	Short-pulse propagation 97
Stokes 219	Siliciumtetrachloride 225
stimulated 205, 217	Sillenites 242
Brillouin 217, 227, 229	Silver Gallium Selenide 142
Brillouin, thermal 220, 227	Silver Thiogallate 142
Raman 217, 223, 226	Simple astigmatic beam 63, 120, 121
Rayleigh 217, 228, 230	Sixth harmonic generation 156 of Nd:YAG laser radiation 161
Rayleigh, line 228	Slit factor 92
Rayleigh, thermal 220, 228	
Rayleigh, wing 228, 230, 231 Stokes 219	Slowly varying envelope (SVE) 9
transient 222	approximation 9 for diffraction 9
Stokes	
	equation 74,80 Small signal
spontaneous 219 stimulated 219	gain factor 11
	_
Schawlow 3 Schrödinger equation 13	solutions 19 Solid
of optics 9	Raman
Second harmonic generation 144, 152, 156	parameters 227
in "nonlinear regime" 154	scattering 225
of dye laser radiation 164	scattering 223
of gas laser radiation 166	Raman 225
of Nd:YAG laser radiation 156, 157, 159	-state laser material, data of 108
of Nd:YAG laser radiation, intracavity 158	Solutions
of ruby laser radiation 163	small-signal 19
of semiconductor laser radiation 164	steady-state 17
of Ti:sapphire laser radiation 163	strong-signal 19
Second-order moment 55	Spatial 19
determination of 66	frequency 88
radiation field	hole-burning 237
propagation of 111	moment 56
rr08	

Spectral	Rayleigh, wing 228, 230, 231
line shape 17	Stokes 219
radiance 45	transient 222
Spectrum	Stokes scattering 219
angular, representation 87	thermal
Sphere	Brillouin scattering 220, 227
refraction at 114	Rayleigh scattering 220, 228
Spherical	Stokes
aberration 117	-anti-Stokes coupling 222
third-order 119	emission
concave mirror 114	higher-order 222
mirror 113	line 218
surface 113	scattering
vector wave 78	spontaneous 219
wave 8,79	stimulated 219
Spontaneous	shift 224–227
emission $4, 15, 28$	spontaneous scattering 219
Einstein coefficient 21	stimulated scattering 219
scattering	vector 77
Stokes 219	Stop 127
Stokes scattering 219	Strong-signal solutions 19
Sr 210	STRS (stimulated thermal Rayleigh scattering)
SRLS (stimulated Rayleigh line scattering) 228	228
SRS (stimulated Raman scattering) 223	Styrene 225, 231
SRWS (stimulated Rayleigh wing scattering) 228,	Sulfur 225
230, 231	Sulfurhexafluoride 229
Stability	Sum frequency generation 144, 153, 167
positional	of UV radiation 167–169
beam 67	Superradiance 37
Standards	Surface
primary 47	index $105, 146$
radiometric 47	ray 105
secondary 48	spherical 113
STBS (stimulated thermal Brillouin scattering)	Susceptibility 19
227	complex 10
Steady state	electric 6
condition 5	linear 205
solutions 17	magnetic 6
Steradian 47	nonlinear 205
Stigmatic beam 62, 120, 121	third-order $206, 221$
Stimulated	third-order
amplification 221	nonlinear 221
Brillouin scattering 217, 227, 229, 236	SVE (Slowly varying envelope) 9
thermal 220, 227	approximation 9
Raman scattering 217, 223, 226	for diffraction 9
Rayleigh scattering 217, 228, 230	SVE (slowly varying envelope)
line 228	equation 74,80
thermal 220, 228	Symmetry conditions
wing 228, 230, 231	Kleinman 150
scattering 205, 217	Symplectic optical system 116
Brillouin 217, 227, 229	Synchrotron radiation 47
Brillouin, thermal 220, 227	System
Raman 217, 223, 226	four-level 24
Rayleigh 217, 228, 230	general
Rayleigh, line 228	astigmatic 116
Rayleigh, thermal 220, 228	parabolic 112

homogeneous 16	Tintetrabromide 225
inhomogeneous 16	Tintetrachloride 225
matrix 55	Tl 210
optical	Tl_3AsSe_3 143, 167
non-symmetrical 112	Toluene 225, 226, 229, 231
symplectic optical 116	Total reflection 101
three-level 23	Townes 3
two-level 11	Tracing
	ray 55
Te 143, 166	Transform
Tellurium 143	Fourier 88
Tetrabromomethane 229	Transformation
Tetrachloroethane 225	ABCD 120
Tetrachloroethylene 225, 226	waist 124
Tetrachloromethane 230	Transient stimulated scattering 222
Tetragonal 105	Triclinic 105
Tetrahydrofuran 225	Trigonal 105
Thallium Arsenic Selenide 143	Twist 59
Theory	Two-dimensional propagation 126
scalar	Two-level
of diffraction 85	system 11
vector	transition
of diffraction 85	saturation intensity 17
Thermal	Two-wave mixing 242
detector 48	O
grating 237	Uniaxial
lens 113	crystal 145
scattering	negative 146
Brillouin, stimulated 220, 227	positive 146
Rayleigh, stimulated 220, 228	negative
Thermodynamic considerations 21	crystal 105
Thermopile 48	positive
Thick lens 110	crystal 105
in air 113	Up-conversion
Thin lens 113	of CO ₂ laser radiation 171
Third harmonic generation 152, 156, 208	of near IR radiation 170
of Nd:YAG laser radiation 159	Upper-level lifetime T_1 18
of Ti:sapphire laser radiation 163	Urea 143, 161, 165, 179
Third-order	UV
nonlinear susceptibility 206, 221	radiation
spherical aberration 119	sum frequency generation of 167–169
Three-dimensional propagation 127	region
Three-level system 23	cw optical parametric oscillation in 176
Three-wave interaction 144	femtosecond optical parametric oscillation in
Threshold	184
condition 4	nanosecond optical parametric oscillation in
SBS- 238, 239	176
Ti:sapphire laser radiation	picosecond optical parametric oscillation in
fourth harmonic generation of 164	180
second harmonic generation of 163	
third harmonic generation of 163	Value
Time	principal 145
constants 31	Vanillin (4-Hydroxy-3-Methoxy-Benzaldehyde)
decay, T_1 15	142
decay, T_2 16	Vapor
-dependent diffraction theory 89	barium 225
-reversed replica 236	cesium 226

Variance	Water 225, 226, 229, 230
ellipse 57	Wave
matrix 56	cylindrical 79
Vector	equation 73,78
Bloch 33	plane $7,79$
-ial Bessel beam 80	approximation 152
Jones 75	representation 87
phase conjugator 236	spherical 8,79
Stokes 77	vector
theory of diffraction 85	cylindrical 78
wave	spherical 78
cylindrical 78	Waveguide coupling 127, 128
spherical 78	Width
Vibration	beam 57
-al Raman line 218, 223	long-term 69
-rotation Raman line 218	Wigner distribution 53
Visible region	Wing scattering
cw optical parametric oscillation in 176	Rayleigh, stimulated 228, 230, 231
femtosecond optical parametric oscillation in	
184	X-ray region 47
nanosecond optical parametric oscillation in 176	Xe 211, 238
picosecond optical parametric oscillation in 180	Yb 210
Waist	Zinc Germanium Phosphide 143
beam 82	Zn 210
position 60	$ZnGeP_2$ 143, 166, 167, 171, 175, 186
transformation 124	ZnSe 175
Walk-off angle 145	ZnTe 175