AEROSOL EXTINCTION TRANSMISSOMETRY AT VISIBLE AND INFRARED WAVELENGTHS APPLIED TO SPECIALIZED PROBLEMS IN CB DEFENSE

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Aerosol Extinction Transmissometry at Visible and Infrared Wavelengths Applied to Specialized Problems in CB Defense

Complex calculations of optical scattering using high-speed electronic computers have become so commonplace that the technology has evolved into several specialty fields, e.g., extinction spectroscopy, angular scattering measurements and nephelometry. As a consequence, the overall technology and our ability to remotely characterize and quantitatively analyze atmospheric aerosols might be further advanced than is realized by workers in the individual disciplines who do not have the time to keep up with advances in other fields.
18. Subject Terms (continued)

Nephelometry  Polarization
Atmosphere    Spheres
Optical scattering Spectroscopy
Visible        Geometric regime
Lasers         Rayleigh regime
He:Ne          
Turbidity      
Infrared       

19. Abstract (continued)

This report examines techniques of transmission (extinction) and turbidity spectroscopy, polarization and angular scattering measurements, and related developments and concludes that computer programs can be written to optimize desired parameters in interdisciplinary investigations. An appendix discusses sampling, which is a complicated problem in aerosol physics. A technique described to monitor continuously the mass concentration of monodisperse test aerosols using a He:Ne laser. Good approximations are possible from Mie theory. The method is fast, accurate, and eliminates the need for precalibration of a standard sampler to monitor aerosol mass concentration over increments of time. It can be used for sampler efficiency measurements and whenever it is desired to monitor test aerosols of spherical droplets continuously, e.g., in studies of the human respiratory system for droplets larger than a few micrometers. A second appendix discusses the estimation of mass extinction coefficients for nonabsorbing spherical aerosol particles in the geometric scattering regime. The techniques presented should find wide application in military electro-optical technology.
PREFACE

The work described in this report was authorized under Project No. 1L161101A91A, In-House Laboratory Independent Research (ILIR). This work was started in January 1978 and completed in December 1982. It was updated by reference to the recent open literature and prepared for publication in December 1988.

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This report has been approved for release to the public.

Acknowledgments

Significant contributions to the work described in the Appendixes of this report were made by David V. Kimball and Robert J. Wright, who carried out the laboratory investigations described, and by Robert Frickel and Merrill E. Milham, who performed many Mie calculations in support of this research. Bernard V. Gerber suggested the polarization and angular scattering studies that are described.
## CONTENTS

1. INTRODUCTION AND BACKGROUND ............................................... 7  
2. AEROSOL EXTINCTION MEASUREMENTS ....................................... 8  
3. POLARIZATION AND ANGULAR SCATTERING MEASUREMENTS .................. 20  
4. SUMMARY AND CONCLUSIONS .................................................. 22  

LITERATURE CITED ................................................................. 23  

APPENDIX A: LASER MONITORING OF MASS CONCENTRATIONS OF  
MONODISPERSE TEST AEROSOLS .................................................. 27  

APPENDIX B: MASS EXTINCTION COEFFICIENTS ESTIMATED FOR  
NONABSORBING SPHERICAL AEROSOL PARTICLES IN THE  
GEOMETRIC SCATTERING REGIME .............................................. 37  

## LIST OF FIGURES

1. Values of $a_\lambda . D_H$ vs $D_H$ Calculated From the Mie Program for  
Water; the "Constant" Function Actually is a  
Damped Oscillation ...................................................................... 10  

2. Comparison of Computed Curves Relating Visible to Infrared  
Optical Transmittance of Water Fog to Experimental Data  
Points for Cooling, Steam-Generated Water Fog Clouds ............... 11  

3. See Figure 2; Curves for Three Infrared Wavelengths  
Comprising the Broadband Composite of Wavelengths  
Shown as the Abscissa of Figure 2 ............................................. 12  

4. Calculated Curves for Water Droplets Showing $a_\lambda$ vs $D_H$; Note  
Constancy of $a_{12.5}$ at All Wavelengths Up to About $D_H = 15 \mu m$ .... 14  

5. Computed Values of $a_\lambda$ for Liquid Droplet Aerosols Comprising  
Several Concentrations of Orthophosphoric Acid ($H_3PO_4$) in Water. 15  

6. Plot of Equation (3) for $n_\lambda$ values of 0.1-4.0 Shown on the  
Curves and $k_\lambda$ Values Shown on the Abscissa, for Rayleigh-  
Scattering Aerosols "Only" ......................................................... 17  

7. Fractional Transmittance for a $C \times L$ Product = 5.0 (Equation (1))  
of Water Fog vs Droplet Diameter for Wavelengths, Reading from  
Top to Bottom of the Left-Hand Ends of the Curves, of $\lambda = 0.63$,  
8.5, 10.5 and 4$\mu m$, Which Are the Wavelengths Also  
Considered in Figures 1-3 ......................................................... 18
8. \( \alpha_\lambda \) vs \( D_\mu \) Calculated From the Mie Program at \( \lambda = 10\mu m \) for Water Droplets; the Subscripts T, S and A Refer to Equation (4)........ 19

9. Polarization Ratio \( P(\theta) \) vs Scattering Angle \( \theta \) for DOP.............. 21

A1. Values of \( \alpha_{0.63} \) vs \( D_\mu \) Calculated From the Mie Theory for Liquids Listed in Table A1.......................................................... 30

A2. Values of \( \alpha_\lambda \) vs \( D_\mu \) Calculated From the Mie Theory for Dimethyl Phthalate (DMP), Using the Sodium D Line Refractive Index Measured at Its Wavelength of 0.589 \( \mu m \) (Dashed Curve) and Assuming the Same Index at the He:Ne Laser Wavelength \( \lambda = 0.63 \mu m \) (Solid Curve).... 31

A3. Values of \( \alpha_{0.63} \) vs \( D_\mu \) Calculated From Mie Theory for Dioctyl Phthalate (DOP)........................................ 32

A4. Schematic Representation of the Experimental Setup; the Chamber Was Stirred to Ensure Uniform Aerosol Mixing in the Laser Beam.. 33

A5. Typical Sampling Efficiencies for DOP Test Aerosols Are Shown by the Curves; the Methodology of Measurements Also Is Illustrated................................. 35

B1. Mie-Calculated Extinction Coefficient vs Spherical Droplet Diameter for the He:Ne Wavelength (\( \lambda = 0.63 \mu m \)); the Dashed Curve Is for Water Droplets, While the Solid Curves Are for Droplets of Phthalates (Dibutyl, Diethyl, Dimethyl and Dioctyl). 38

**TABLE**

Liquids Used to Generate Test Aerosols......................... 28
AEROSOL EXTINCTION TRANSMISSOMETRY AT VISIBLE AND INFRARED WAVELENGTHS
APPLIED TO SPECIALIZED PROBLEMS IN CB DEFENSE

1. INTRODUCTION AND BACKGROUND

With the increasing availability of high-speed electronic computers over the past two decades, complex calculations of optical extinction and scattering of electromagnetic radiation by atmospheric aerosols using the Mie theory have become commonplace.

The technology has advanced so swiftly that specialists measuring, e.g., cloud extinction of aerosols often are unaware of advances being made simultaneously by other specialists measuring, e.g., angular scattering patterns of aerosols with nephelometers. Such measurements and calculations can be done in numerous wavelength regions, but the technology which concentrated first upon the visible wavelengths now is expanding rapidly especially into the regions of relatively high atmospheric transparency called "windows" in the infrared.

This technological expansion can be measured, for example, by a sampling of papers appearing one journal in one year alone, discussing aerosol measurements or calculations of extinction, angular light scattering or nephelometry, lidar, and related aerosol technology including thermal emissivity and calibration of optical particle counters.

The author has noted several trends in all of this work. First, the utility and information content of simple extinction measurements has tended to be overlooked in the rush to exploit scattering techniques. Second, the polarization techniques do not seem to include applicable techniques, some new, that can be applied both to extinction and to polarization measurements. Third, that if taken in combination at this point in time, the overall technology might be further advanced than is realized by workers in the individual disciplines.

For example, remote characterization or quantitative analysis of tropospheric aerosols seems a distinct possibility, both by conventional transmission (extinction) spectroscopy and by angular scattering or backscattering techniques. This report examines these possibilities by first reviewing the technology of aerosol extinction spectroscopy.

The Beer-Lambert equation for atmospheric aerosols can be written:

\[- \ln T_\lambda = \alpha_\lambda CL, \quad (1)\]
where for a given wavelength (μm), $T_\lambda$ is the fractional transmittance, $\alpha_\lambda$ is the optical mass extinction coefficient of the aerosol (m$^2$g$^{-1}$), $C$ is the aerosol mass concentration (g m$^{-3}$), and $L$ is the optical path length (m). Some form of Equation (1) has been used by spectroscopists virtually since the birth of spectroscopy and the equation has become so well known that the power of simple extinction measurements of aerosols often is not fully appreciated.

For example, using suitable multi-wavelength techniques, extinction measurements can contain sufficient information to allow direct determination of the mean size of a spherical droplet aerosol distribution, or even remotely to characterize chemical reactions in cloud droplets at isosbestic wavelengths without the use of polarized radiation or other more sophisticated techniques.

It is not surprising, however, that polarization techniques have received heavy emphasis since their exploitation in the early 1940s to characterize military fog-oil smoke mean droplet diameters based, in part, in earlier work at the U.S. Bureau of Standards by Stratton and Houghton using the Mie theory that was first put forth in 1908 but had to await the availability of high-speed computers for exploitation.

Stratton and Houghton measured the polarization ratio only for transparent spheres for which, if the complex index of refraction $n_\lambda$ is given by $(n - ik)_\lambda$ where $n_\lambda$ is the real index and $k_\lambda$ is the imaginary index (closely related to the absorption coefficient), then $k_\lambda = 0$.

Polarization measurements can be used to determine particle size distribution in addition to mean size of aerosol droplet distributions. This was recognized and the technique was developed in the 1960s. By 1969, Kerker and his co-workers had extended the techniques of particle size distribution measurements to the extent discussed in his classical book on light scattering that gives an excellent review of the technology as it existed then.

In recent years, there is an increased awareness that aerosol particle absorption, e.g., in the infrared where $k_\lambda$ is not zero, can be interpreted in new ways using not only extinction measurements or polarization measurements, but the two together. The author concurs that this is a promising field for new research.

2. AEROSOL EXTINCTION MEASUREMENTS

It is known that the variation of $k_\lambda$ from values near zero in the visible wavelengths to significant ones in the infrared wavelengths provides a method for the determination of aerosol mean droplet size $\mu$ mass concentration from simple extinction measurements if two or more wavelengths are selected properly for observation.

For example, at the He-Ne laser wavelength $\lambda = 0.63$ μm, $k_\lambda$ is near zero and typical water fog droplets have diameters $D_\mu$ such that $D_\mu >> \lambda$ or, to use Kerker's criterion, the size parameter $\pi D_\mu/\lambda \geq 2.0$ so that the conditions for geometric scattering exist.
This leads to a constancy of the product $a_{\lambda}D_u \rightarrow 3.0-3.2$, where $a_{\lambda}$ is defined in Equation (1). At the same time, the product $a_{\lambda}D_u$ decidedly is not constant with $D_u$ at wavelengths where $k_{\lambda}$ is not equal to zero in the infrared. This is illustrated in Figure 1, calculated from the Mie theory.

In Figure 1, curves are shown for the infrared wavelengths of 8.5, 10.5 and 12.57 (or 4$\pi$) $\mu$m, as well as for a composite of wavelengths in the 8.5-12.57 $\mu$m region where extinction can be measured, e.g., by a simple broadband transmissometer operating at 8-13 $\mu$m in the atmospheric window region there.

While operation at a specific wavelength such as the 10.6 $\mu$m CO$_2$ laser line for comparison to a visible wavelength such as the He:Ne laser line at $\lambda = 0.63$ $\mu$m has advantages, a broadband infrared reference band also can be used. In fact, related findings using a filtered light source in the visible at $\lambda = 0.515$ $\mu$m and a 9-12 $\mu$m broadband transmissometer to study the formation and dissipation of water fogs were reported at least as early as 1970 by Carlon.$^3$4

When the techniques of Carlon, et al.$^{25,27}$ are extended to use the Mie theory to calculate functions of the transmittances $T_{\lambda}$ at $\lambda = 0.63$ $\mu$m and the broadband or composite wavelength band 8.5-12.57 $\mu$m vs. water droplet diameter $D_u$, curves like those shown in Figure 2 are obtained. It is seen that the curves are widely separated.

Thus, a sensitive method is provided for the determination of mean droplet diameter. Experimental points for a steam-generated water fog are shown in Figure 2; these give excellent agreement with calculated curves for example at $D_u = 8 \mu$m. This result was cited by Weinman, et al.$^{35}$ as approximating Deirmendjian's 1969 Model C-1 (or vice-versa),$^{36}$ where $D_u$ is taken as the mass median diameter.

What is uncertain is the kind of distribution to which this diameter is most applicable. Kerker$^{33}$ has given an informative discussion of particle size distributions, and of transmission (extinction) and turbidity measurements. In this report, turbidity can be taken as the product $a_{\lambda}C$ in Equation (1), having the units $m^{-1}$.

Justification for the use of a composite of wavelengths (Figure 1 and abscissa, Figure 2) can be found by examining calculated curves and steam-generated fog data points for the individual wavelengths $\lambda = 8.5, 10.5$ and 12.57 (or 4$\pi$) $\mu$m and plotting curves like Figure 2 as is done in Figure 3.

Viewing the plots from top to bottom in Figure 3, it can be seen that the fog data gave the best agreement with calculation for a mass median diameter (MMD) of 6 $\mu$m at $\lambda = 8.5$ $\mu$m, MMD = 10 $\mu$m at $\lambda = 10.5$ $\mu$m, and MMD = 6 $\mu$m at $\lambda = 4\pi$ $\mu$m.

In his discussions of particle size distribution functions, Kerker$^{33}$ cautions that mean particle diameters obtained by using methods like those discussed in the present report may be adequate for most purposes if the distribution is sufficiently narrow, but strictly are valid only for monodisperse systems.
Figure 1. Values of $a_\lambda D_\mu$ vs $D_\mu$ Calculated From the Mie Program for Water; the "Constant" Function Actually is a Damped Oscillation.
Figure 2. Comparison of Computed Curves Relating Visible to Infrared Optical Transmittance of Water Fog to Experimental Data Points for Cooling, Steam-Generated Water Fog Clouds.
Figure 3. See Figure 2; Curves for Three Infrared Wavelengths Comprising the Broadband Composite of Wavelengths Shown as the Abscissa of Figure 2.
By using a composite of wavelengths to obtain a mean diameter, one is averaging both over wavelength intervals and over particle size distributions. But the results indicate that the technique works very well, especially under dynamic droplet growth or evaporation conditions where what is wanted most is a real-time indication of approximate droplet size, e.g., in studies of developing or dissipating water fogs.34,37

The dependence of $\alpha_\lambda$ upon $\lambda$ is, in fact, highly variable due to variations in $(n - ik)_\lambda$ in addition to which $\alpha_\lambda$ varies with $D_m$ as was discussed previously. This is illustrated in Mie calculations for the extinction coefficient of water droplets ($\alpha_\lambda$, ordinate) in Figure 4 vs $D_m$ for several wavelengths. At certain wavelengths, e.g., $\lambda = 12.5$ μm, the combined contributions of refractive ($n_\lambda$) and absorptive ($k_\lambda$) components of water droplet extinction lead to values of $\alpha_\lambda$ that virtually are independent of droplet size.

Experiments27 seem to confirm for example that $\alpha_{12.5}$ is nearly constant with $D_m$, suggesting that the liquid water content C in Equation (1) of a tropospheric optical path could be monitored by a simple 12.5 μm transmissometer with good precision for droplet sizes up to about 15 μm.38 In this application, however, care must be taken to account for absorption due to hydrogen-bonded molecular clusters (water clusters) in the vapor leading to absorption easily confused with droplet absorption.5

At wavelengths called "isosbestic points"28 the complex indices $(n - ik)_\lambda$ of some liquids are such that $\alpha_\lambda$ remains constant even though the liquid solutions comprising the droplet aerosol vary widely in their chemical composition, e.g., for droplets of orthophosphoric and related acids that might be formed by burning phosphorus in air of varying relative humidity (RH) such that solute (acid) concentrations vary from near zero to 85% by weight.

In Figure 5, $\lambda = 11.4$ μm is an isosbestic point calculated from refractive index measurements of pure orthophosphoric acid ($H_3PO_4$) in water.39,40 Real atmospheric smokes obtained by burning phosphorus may contain mixtures of acids depending on rates of combustion in the presence of varying humidities and so may not give experimental spectra like those for pure $H_3PO_4$ in Figure 5 where $\alpha_{11.4}$ remains essentially constant at 0.13-0.15 m²g⁻¹, while acid concentration varies from near zero to 85% by weight. This will be investigated.

Because of the constancy of $\alpha_{11.4}$ compared to the wide ranges of, say, $\alpha_{9.7}$ (see Figure 5) with solute concentration, remote characterization of the aerosol becomes a possibility if something is known about the droplet constituents. It can be shown that:

$$\alpha_\lambda \sim 4\pi k_\lambda f(m_\lambda)/\lambda \rho,$$

where $\rho$ (g cm⁻³) is the droplet acid (solution) density, i.e., mass density, and:
Figure 4. Calculated Curves for Water Droplets Showing $\alpha_\lambda$ vs $D_\mu$; Note Constancy of $\alpha_{12.5}$ at All Wavelengths Up to About $D_\mu = 15 \mu m$. 

\[
\lambda = 0.63 \quad 1.06 \quad 12.5 \\
10.0 \quad 8.6 \quad 4.0 \\
0.01 \quad 0.02 \quad 0.03 \quad 0.04 \quad 0.05 \quad 0.06 \quad 0.07 \quad 0.1 \quad 0.15 \quad 1.0 \quad 2.0 \quad 3.0 \quad 5.0 \quad 70.0 \quad 100.0
\]
Figure 5. Computed Values of $\alpha_\lambda$ for Liquid Droplet Aerosols Comprising Several Concentrations of Orthophosphoric Acid ($H_3PO_4$) in Water.
\[ f(m_\lambda) = \frac{9 \eta_\lambda}{(n_\lambda^2 + k_\lambda^2)^2 + 4(n_\lambda^2 - k_\lambda^2) + 4} \]  

where \( m_\lambda = (n - ik)_\lambda \) and the function of Equation 3 is plotted in Figure 6 for values of \( n_\lambda \) marked on the curves and ranging from 0.1 to 4.0, and for values of \( k_\lambda \) shown on the abscissa. It is emphasized that the above discussion applies to Rayleigh-scattering smokes where \( D_\mu \ll \lambda \) as clearly is the case for orthophosphoric or other acid aerosols (smokes) produced by burning phosphorus in moist air.

Often it is useful to plot extinction coefficients or fractional transmittances vs \( D_\mu \) rather than vs the size parameter \( \pi D_\mu / \lambda \), for specific wavelengths. Two such calculated plots\(^ {41} \) are shown for water droplets in Figures 7 and 8. In Figure 7, the curves are calculated for a product \( \Gamma x L = 5.0 \) (Equation (1)) and for wavelengths, reading from top to bottom of the left-hand ends of the curves, of \( \lambda = 0.63, 8.5, 10.5 \) and \( 4 \mu \text{m} \), i.e., the same wavelengths considered in Figures 1-3. In Figure 8, the curves are calculated for \( \lambda = 10 \mu \text{m} \) to show the functions vs \( D_\mu \) of total extinction, and of the scattered and absorbed components that are summed to give total extinction, i.e.,

\[ \alpha_T = \alpha_S + \alpha_A. \]  

where the subscripts indicate total (T), scattering (S), and absorptive (A) extinction.

Sassen\(^ {26} \) recognized that aerosol cloud extinction measured in conjunction with angular scattering measurements could lead to remote sensing of cloud composition by using two wavelengths like the laser wavelengths \( \lambda = 0.63 \) and \( 10.6 \mu \text{m} \). But the present discussion indicates that such remote characterization could result from cloud extinction measurements alone at two closely-spaced infrared wavelengths, one of them an isosbestic point.

Additional data might be obtained by combining extinction and angular scattering measurements at an isosbestic wavelength. Optical parameters can carry only so much information, but the idea is provocative enough to warrant further investigation.

Other investigations of remote characterization of tropospheric aerosol clouds are possible, such as those based on effects peculiar to Christiansen wavelengths.\(^ {42} \) A complete discussion is given by Carlon,\(^ {4,5,42} \) where Reference 5 discusses aerosols ranging in size from molecular clusters to water cloud droplets.
Figure 6. Plot of Equation (3) for \( n_{\lambda} \) values of 0.1-4.0 Shown on the Curves and \( \kappa_{\lambda} \) Values Shown on the Abscissa, for Rayleigh-Scattering Aerosols “Only”.
Figure 7. Fractional Transmittance for a C x L Product = 5 0 (Equation (1)) of Water Fog vs Droplet Diameter for Wavelengths, Reading from Top to Bottom of the Left-Hand Ends of the Curves, of $\lambda = 0.63$, 6.5, 10.5 and 4$\pi$ $\mu$m, Which Are the Wavelengths Also Considered in Figures 1-3.
WATER, $\lambda = 10\mu m$

Figure 8. $\alpha_\lambda$ vs $D_\mu$ Calculated From the Mie Program at $\lambda = 10\mu m$ for Water Droplets; the Subscripts T, S and A Refer to Equation (4).
3. POLARIZATION AND ANGULAR SCATTERING MEASUREMENTS

Kerker has given an excellent review of techniques used to determine particle size distributions using measurements like these, and the present author will not elaborate except to raise a few pertinent points.

Sassen verified that angular scattering patterns calculated from the Mie theory can be observed rather closely by experiment not only for spherical particles, but for irregularly-shaped ones as well. Thus polarization or angular scattering techniques combined with some of the recent advances in extinction measurement technology discussed here should provide new combinations for the remote characterization of tropospheric aerosols.

The polarization method for evaluating the distribution of sphere sizes, utilizes polarization of scattered radiation obtained from a monochromatic light source at various angles of observation. The intensity of the components of the scattered light whose electric vector vibrates perpendicular and parallel to the plane of observation, \( I_1(\theta) \) and \( I_2(\theta) \), is measured at a number of angles, where \( \theta \) is the scattering angle measured from the direction of the incident light beam. The polarization ratio, \( P(\theta) \), is defined such that:

\[
P(\theta) = \frac{I_2(\theta)}{I_1(\theta)}.
\]

For example, \( P(\theta) \) can be calculated as a function of \( \theta \) for any assumed particle distribution such as the zeroth order logarithmic distribution (ZOLD) and for any standard deviation, \( \sigma_0 \), ranging from \( \sigma_0 = 0 \) (i.e., a monodisperse aerosol), to values of 0.3 or more where the curves flatten and the method approaches its limit of usefulness.

Figure 9 shows the family of curves calculated for the He: Cd laser wavelength \( \lambda = 0.4416 \) \( \mu \)m, \( D_0 = 0.3 \) \( \mu \)m, \( n_\lambda = 1.484 \) and \( k_\lambda = 0 \) so that \( m_\lambda = (1.484 - i.0)_\lambda \) and the density is \( \rho = 0.98 \) g cm\(^{-3}\). The curves are for dioctyl phthalate (DOP), a liquid that is commonly used for test aerosols. Agreement between calculation and measurement using the polarization method is good to excellent, often within a few percent.

The effect of the wavelength of the monochromatic light upon the functions shown in Figure 9 is of special interest. This has been investigated for visible and near-infrared wavelengths, including cases such as vanadium pentoxide \((\text{V}_2\text{O}_5)\) for which strong absorption occurs at shorter visible wavelengths, i.e., where \( k_\lambda \) is not zero.

From the discussion in the present report of special techniques using extinction measurements in the infrared where \( k_\lambda \) is not zero, it is suggested that combinations of visible and infrared wavelengths can optimize desired parameters using polarization techniques. Since the size parameter \( mD_0/\lambda \) depends on wavelength, a change in the wavelength of the incident light will cause a shift in the position of the curve peaks in Figure 9 such that larger values of \( \lambda \) shift the peaks to the right.
Figure 9. Polarization Ratio $P(\theta)$ vs Scattering Angle $\theta$ for DOP.
Multi-wavelength observations have been used to show that particle size distributions obtained at different wavelengths are consistent, but when wavelength spacing is as great as $\lambda = 0.63$ and 10.5 $\mu$m additional work is needed to confirm this and also to determine whether the useful range of $\sigma_0$ can be extended beyond 0.3 in measurements using the polarization method.

The scattering ratio method for the determination of particle size distribution is a variation where the angle of observation is held constant and the polarization ratio is measured or calculated over a spectrum or over a number of discrete wavelengths.

The method was developed in the early 1960s and was perfected by using monodisperse polystyrene latex spheres and known combinations of single sphere sizes to give mixtures of known size distributions checked by electron microscopy.

Thus the scattering ratio method gives an alternative way of studying, e.g., the variation of $\sigma_0$ at some $D_H$ with $\lambda$ at some constant $\Theta$, although the same thing can be done by using the polarization method directly and plotting, e.g., Figure 9 with $\lambda$ as the abscissa at constant $\Theta$.

Turbidimetric particle size distribution methods invariably are similar to Wallach, et al. Nelson proposed a method by which calculated scattering cross sections from the Mie theory are compared to experimental extinction data using monodisperse polystyrene latex spheres in distilled water as a calibration means, and a computer program is used to null the error vs wavelength thus forcing a unique solution for the size distribution which is taken as bimodal to account for coagulation. The method also is said to yield particle volume concentration and the scattering cross section function, $Q_\lambda$. Extrapolation of the technique to tropospheric aerosols remains to be demonstrated.

4. SUMMARY AND CONCLUSIONS

This paper has concentrated on advances that have been made in the remote characterization of tropospheric aerosols by extinction and turbidimetric methods over the nearly two decades that have elapsed since the publication of Kerker's classic text. The technology has been updated by including results not previously published with improved techniques in polarization and angular scattering methods. Perhaps most importantly, the author has tried to support the argument that if all methods presently available in this technology are taken into account at this time, our ability to remotely characterize or quantitatively analyze atmospheric aerosols is further advanced than workers in the individual light measurement disciplines realize.

It is concluded that by combinations of well-established and recently-developed light scattering techniques, investigators can undertake promising new areas of research. Some examples include combinations of extinction and polarization measurements with computer programs to study isosbestic or Christiansen wavelengths, and optimization of desired parameters as functions of wavelength or particle size distribution.
LITERATURE CITED


APPENDIX A

LASER MONITORING OF MASS CONCENTRATIONS

OF MONODISPERSE TEST AEROSOLS

A1. INTRODUCTION

Aerosol sampling usually is considered to be a very complicated problem in aerosol physics. It may be desired, for example, to determine the efficiency, as a function of particle diameter \( D_p \) (\( \mu \text{m} \)), with which aerosol particles of various kinds are drawn into an aspirated sampling tube or head or into the human respiratory system.

The principal experimental difficulty is one of establishing and maintaining a test aerosol of known characteristics, including composition, mass or number concentration, and particle size distribution. The aerosol sampler is placed in the test aerosol and is operated at a known flow rate for a known period of time, after which a determination is made, usually by weighing a filter paper in the sampler, of the mass of the collected aerosol. This is ratioed to the particle mass contained in the sampled volume of the test aerosol to determine the sampler efficiency.

The problem of maintaining the test aerosol is not a trivial one. It must be sampled periodically to determine if (and how) it is changing with time as, for example, particles settle or agglomerate. This sampling disturbs and draws material from the test aerosol, and the subsequent analyses usually require at least several minutes to complete, during which period the characteristics of the test aerosol may continue to change.

This appendix describes a technique by which the test aerosol mass concentration is continuously sampled by a laser beam. This requires that the optical constants, particle size distribution, and other properties of the aerosol particles be known. The test aerosol materials discussed in this paper are liquids that are generated as spherical droplets, for which accurate (+ or - 10%) optical calculations can be made using the Mie theory.\(^2\)

For approximation purposes in optical calculations, it is often useful to treat an aerosol as if it consisted entirely of monodisperse particles of some equivalent diameter.\(^3\) But a better approach is to generate monodisperse droplet aerosols directly, as was done in the work reported here.
A2. THEORY

In the geometric optical scattering regime, particle diameters are much larger than the wavelength $\lambda$ (\mu m) of observation. That is, $D_{\mu} > \lambda$. Under these conditions, it is straightforward to show that:

$$\alpha_{\lambda} = \frac{3Q_{\lambda}/2D_{\mu}}{\rho}$$

where $\alpha$ (m$^2$g$^{-1}$) is the mass extinction coefficient of the spherical particles, $\rho$ (g cm$^{-3}$) is the particle density, and $Q_{\lambda}$ is the cross-section efficiency factor, which is very nearly constant and equal to 2.0 for the liquid aerosol droplets discussed in this paper,

$$\alpha_{\lambda} \sim \frac{3}{D_{\mu}\rho}$$

The mass extinction coefficient is used in the Beer-Lambert equation:

$$-\ln T_{\lambda} = \alpha_{\lambda}CL,$$

where $T_{\lambda}$ is the optical transmittance at laser wavelength $\lambda$, $C$ (g m$^{-3}$) is the aerosol mass concentration, and $L$ is the optical path length uniformly filled by the aerosol (m). $C$, the quantity sought for the test aerosol, is then:

$$C = -\frac{1}{\alpha_{\lambda}L} \ln T_{\lambda} \sim \frac{D_{\mu}\rho}{3L} \ln T_{\lambda}$$

Several liquids used to generate test aerosols in work described in this appendix are shown in the table.

<table>
<thead>
<tr>
<th>Table: Liquids Used to Generate Test Aerosols</th>
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<tbody>
<tr>
<td>Symbol</td>
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<tr>
<td>DMP</td>
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<tr>
<td>DEP</td>
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<tr>
<td>DHP</td>
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<tr>
<td>DOP</td>
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* The sodium D line is at the 0.589-\mu m wavelength, and the real index at the He-Ne laser wavelength of 0.63 \mu m is nearly identical to this value for transparent colorless liquids.
Mie calculations were performed for the He:Ne laser ($\lambda = 0.63 \mu m$), with results as shown in Figure A1, and it was confirmed that the approximation of Equation (A2) is quite precise, i.e., that the functional dependence of $a_{0.63}$, the extinction coefficient, on $D_D$ is very similar for all liquid aerosol materials considered, with only liquid densities $p$ having a predictable effect.

The laser wavelength (0.63 $\mu m$) lies very close to the sodium D line (0.589 $\mu m$), and the materials listed in Table A1 are transparent and colorless in the visible wavelengths (i.e., they have a real refractive index $n$ but a negligible absorption coefficient or imaginary index $k$). Thus it was very convenient to obtain the real indices of these materials using a standard laboratory refractometer at the sodium D line and to use these in Mie calculations for the 0.63-$\mu m$ wavelength. The error introduced by doing this is very small, as is indicated by Figure A2, which compares the $a_{\lambda}$ vs $D_D$ functions for DMP calculated at $\lambda = 0.589$ and 0.63 $\mu m$.

After preliminary testing, it was found convenient to use the liquid DOP (Figure A3) in most subsequent work. Since $p = 0.98$ g cm$^{-3}$ for DOP, the approximations in Equations (A2) and (A4) can be still further simplified:

$$a_{0.63} \sim 3/D_D,$$

and:

$$C \sim -D_D \ln T_{0.63}$$

(A5)

It can be seen from Figure A3 that the approximation of Equation (A5) is quite precise in the geometric regime ($D_D \gg \lambda$).

A3. EXPERIMENTAL PROCEDURE

A 1-m$^3$ test chamber was constructed and used as shown schematically in Figure 4. The He:Ne laser was optically aligned through pinholes in opposite chamber walls. The pinholes were utilized to prevent window effects. A pressure-equalization scheme between the chamber interior and room air prevented aerosol particles from flowing in either direction through the pinholes during testing.

Although Figure 4 shows an aerosol cloud coming from the generator, the chamber was stirred during testing to ensure uniform mixing. This was important because laser path length $L$ of the laser beam in the test chamber (which was 1 m long) must be filled uniformly with the test aerosol during measurements.

The aerosol generator was of the spinning disk design$^{29}$ to produce essentially monodisperse liquid droplets of test aerosol. The droplet size $D_D$ disseminated by the spinning disk generator is determined by the diameter and angular velocity of the disk, by the liquid flow rate, and to a lesser extent by other parameters including temperature.

APPENDIX A
Figure A1. Values of $\alpha_{\text{opt}} \cdot 10^3$ vs $D_n$ Calculated From the Mie Theory for Liquids Listed in Table A1.
Figure A2. Values of $\alpha_A$ vs $D_W$ calculated from the Hie Theory for Dimethyl Phthalate (DMP), using the Sodium D Line Refractive Index measured at its Wavelength of 0.589 $\mu$m (Dashed Curve) and assuming the same Index at the He:Ne Laser Wavelength $\lambda = 0.63$ $\mu$m (Solid Curve).

APPENDIX A
Figure A3. Values of $a_{0.63}$ vs $D_m$ Calculated From Mie Theory for Dioctyl Phthalate (DOP).

APPENDIX A
Figure A4. Schematic Representation of the Experimental Setup; the Chamber Was Stirred to Ensure Uniform Aerosol Mixing in the Laser Beam.
Aerosol droplet size was checked by microscopic examination of samples deposited on glass slides. The generator was easily capable of disseminating mass concentrations of test aerosols in the range $C = 0.1-1.0 \, \text{g m}^{-3}$ for these tests. Other monodisperse aerosol generators are commercially available, although some do not have the aerosol output capability of the unit used in these tests. Plastic microspheres also were used in some trials, for which the $\alpha_{0.63} D_P$ functions were not much different from those shown in Figure A1.

For testing, aerosol samplers were mounted completely inside the chamber with their inlets positioned perpendicular to and directly adjacent to the laser beam. This ensured that the laser beam and inlets intercepted identical samples. A refinement to this procedure involves using a second laser beam perpendicular to the first and nearly intersecting it, thus providing laser beam "cross hairs", which can be positioned directly in front of the sampler inlet tube under test.

**A4. RESULTS AND DISCUSSION**

The purpose of this appendix is to describe a technique by which the mass concentration of a sized test aerosol can be continuously sampled by a laser beam to improve experimental procedures. Only a brief discussion of typical test data for aspirated sampler efficiencies will be given here. Figure A5 presents some data for DOP droplets and illustrates the methodology discussed in this paper.

In Figure A5, the solid and dashed curves labeled Aspirated Sampler are typical of the efficiencies measured. For these aspirated samplers, the sampling efficiency falls to less than 10% in the DOP droplet size range $D_P = 30-40 \, \mu \text{m}$. The He:Ne laser can be considered 100% efficient in sampling the test aerosol, since laser attenuation depends entirely on extinction by the aerosol droplets ($\varepsilon_L = 100\%$).

Multiple scattering of the laser beam is not a problem, since the test aerosol concentration is always chosen to give high transmittances (0.7-1.0), thus conserving the aerosol material. The assumption that the actual mass concentration of the test aerosol, $C_{\text{actual}}$, is equal to that determined from the laser using Equation (A6), $C_{\text{laser}}$, is valid to within the error of the Mie calculations, which was found to be as good as plus or minus 10% in this application.

The equivalent mass concentration collected by the aspirated sampler, $C_{\text{sampled}}$, is determined by weighing the filter paper in the sampler before and after an aerosol trial and is reproducible within about plus or minus 10%. The percent efficiency of the sampler $\varepsilon_S$ is then $100 \times (C_{\text{sampled}}/C_{\text{actual}})$. When the test aerosol particle diameter is only

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* The Berglund-Liu monodisperse aerosol generator is marketed by Thermo-Systems, Inc. (TSI). P.O. Box 3394, St. Paul, Minnesota 55165.

** Microspheres of glass, polystyrene, and other materials are available from Duke Scientific Corp., 445 Sherman Avenue, Palo Alto, Calif. 94306.
DETERMINATION OF ASPIRATED SAMPLER EFFICIENCY USING LASER BEAM

Figure A5. Typical Sampling Efficiencies for COP Test Aerosols Are Shown by the Curves; the Methodology of Measurements Also Is Illustrated.
a few micrometers or less, the aspirated sampler efficiency approaches 100%, and these samplers can be used directly to confirm laser measurements of aerosol mass concentration as was reported previously for smoke aerosols. A6

A5. CONCLUSIONS

The discussion in this appendix has been limited to the determination of aspirated aerosol sampler efficiencies using laser reference techniques. These techniques, of course, are not limited to sampler efficiency measurements but can be used wherever it is desirable to continuously monitor the mass concentration of a test aerosol of spherical droplets, e.g., in studies of the human respiratory system.

By using a He:Ne laser in the visible wavelengths (\( \lambda = 0.63 \, \mu m \)), aerosol measurements are made in the geometric scattering regime \( (D_D, \lambda) \) using materials for which many simplifications and precise approximations are possible from the Mie theory. The method is fast, accurate, and eliminates the need for precalibration of standard samplers against which, for example, candidate samplers must be compared.

LITERATURE CITED


APPENDIX B

MASS EXTINCTION COEFFICIENTS ESTIMATED FOR
NONABSORBING SPHERICAL AEROSOL PARTICLES
IN THE GEOMETRIC SCATTERING REGIME

The Beer-Lambert equation for aerosols can be written

\[ \ln T_\lambda = \alpha_\lambda CL, \]

where \( T_\lambda \) is the optical transmittance at wavelength \( \lambda \) (\( \mu \text{m} \)), \( \alpha_\lambda \) is the mass extinction coefficient (\( \text{m}^2 \text{ g}^{-1} \)), \( C \) is the aerosol mass concentration (\( \text{g m}^{-3} \)), and \( L \) is the optical path length (\( \text{m} \)). It is straightforward to show that

\[ \alpha_\lambda = 3Q_\lambda/2D_\mu \rho, \]

where the Mie theory gives the value of \( Q_\lambda \) that depends on the particle diameter \( D_\mu \) (\( \mu \text{m} \)) compared to wavelength \( \lambda \), and \( \rho \) is the particle mass density (\( \text{g cm}^{-3} \)). In the geometric regime (\( D_\mu \gg \lambda \)), \( Q_\lambda \rightarrow 2.0 \) as \( D_\mu \) increases. Thus,

\[ \alpha_\lambda \rightarrow 3/D_\mu \rho. \quad (D_\mu \gg \lambda) \]

For example, Figure B1 shows values of \( \alpha_\lambda \) vs \( D_\mu \) at the Hg:Ne laser wavelength (\( \lambda = 0.63 \mu \text{m} \)) calculated for spherical droplets of water (dashed curve) and several phthalates (dibutyl, diethyl, dimethyl, dioctyl - solid curves); the latter are commonly used in aerosol chamber testing.

The figure shows that when \( D_\mu \gg 0.63 \mu \text{m} \), the Mie-calculated extinction coefficient curves merge into a tail whose function is given by Equation (B3) and whose thickness depends mainly on the differences in mass densities (\( \rho \)) of the liquids represented. The density differences are not obvious because of the logarithmic scale used for the ordinate of Figure B1.
Figure B1. He-Ne-Calculated Extinction Coefficient vs Spherical Droplet Diameter for the He:Ne Wavelength ($\lambda = 0.63 \, \mu m$); the Dashed Curve Is for Water Droplets, While the Solid Curves Are for Droplets of Phthalates (Dibutyl, Diethyl, Dimethyl and Dioctyl).

APPENDIX B
A great many liquid and solid substances exist for which, when they are dispersed as spherical particles at wavelengths where their absorption is negligible, remarkably good first approximations of $\alpha_\lambda$ can be made using Equation (B3). Hence this equation is useful for first calculations involving such aerosols when their optical properties are not known at some wavelength $\lambda$, but $D_\mu \gg \lambda$. A common example is that of water fog droplets in the visible wavelengths. Often, visible-wavelength observations can be combined with IR measurements to yield additional information about an aerosol, e.g., mean particle diameter.$^B_5$ Combining Equations (B1) and (B5) gives

$$-\ln T_\lambda = \frac{3CL}{D_\mu \rho}, \quad (D_\mu \gg \lambda) \quad (B4)$$

and it is straightforward to show that

$$-\ln T_\lambda = \frac{\pi (D_\mu^2 N L \times 10^{-6})}{2}, \quad (D_\mu \gg \lambda) \quad (B5)$$

where $N$ is the number of spherical particles per cubic centimeter. Equation (B5) can be extended to some rather practical applications. For example, Middleton$^B_6$ estimates meteorological range as that for which a high-contrast target is seen with 2% transmittance, i.e., $T_\lambda = 0.02$ and $-\ln T_\lambda = 3.912$, so that from Equation (B5)

$$N = \frac{(2.49 \times 10^6)}{D_\mu^2 L}. \quad (D_\mu \gg \lambda) \quad (B6)$$

As an example, a typical developing water fog is comprised of droplets having mean diameters in the 3-5 $\mu$m range. If an observer could barely see a high-contrast target at $L = 200$ m under such conditions, from Equation B6 the fog should contain approximately 500-1400 droplets cm$^{-3}$. Perhaps a more useful application would be to calculate mean values of $D_\mu$ from Equation (6) using observed ranges ($L$) and droplet populations ($N$) measured by independent means, as a water fog aged and dissipated.

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