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Development, validation, and potential enhancements to the second-generation operational aerosol product at the National Environmental Satellite, Data, and Information Service of the National Oceanic and Atmospheric Administration

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Abstract. A revised (phase 2) single-channel algorithm for aerosol optical thickness, τ_{SAT}^{A} , retrieval over oceans from radiances in channel 1 (0.63 μ m) of the advanced very high resolution radiometer (AVHRR) has been implemented at the National Oceanic and Atmospheric Administration's National Environmental Satellite Data and Information Service for the NOAA 14 satellite launched December 30, 1994. It is based on careful validation of its operational predecessor (phase 1 algorithm), implemented for NOAA 11 in 1989. Both algorithms scale the upward satellite radiances in cloud-free conditions to aerosol optical thickness using an updated radiative transfer model of the ocean and atmosphere. Application of the phase 2 algorithm to three matchup Sun-photometer and satellite data sets, one with NOAA 9 in 1988 and two with NOAA 11 in 1989 and 1991, respectively, show systematic error is less than 10%, with a random error of $\sigma_{\tau} \approx 0.04$. First results of τ_{SAT}^{A} retrievals from NOAA 14 using the phase 2 algorithm, and from checking its internal consistency, are presented. The potential two-channel (phase 3) algorithm for the retrieval of an aerosol size parameter, such as the Junge size distribution exponent, by adding either channel 2 (0.83 μ m) from the current AVHRR instrument, or a 1.6-µm channel to be available on the Tropical Rainfall Measurement Mission and the NOAA-KLM satellites by 1997 is under investigation. The possibility of using this additional information in the retrieval of a more accurate estimate of aerosol optical thickness is being explored.

1. Introduction

The upward radiance over dark surfaces, like the open ocean in the red and near-infrared spectrum, mainly varies with concentration and type of atmospheric aerosol particles, so that information about these particles can be derived from satellite data. Aerosols backscatter solar radiation in proportion to the aerosol optical thickness τ^{A} and the aerosol single scattering phase function $P^{A}(\chi)$, χ being scattering angle. The marine aerosol is almost purely scattering (albedo of single scattering $\omega \rightarrow 1$) when not mixed with continental sources (dust, smoke, soot) [Gordon and Morel, 1983; World Climate Program, 1983, 1986; D'Almeida et al., 1991]. The relationship between the satellite-detected radiance and τ^{A} is therefore illuminationobservation geometry specific. To retrieve τ^{A} , $P^{A}(\chi)$ must be known.

It was shown in some studies that the variability of $P^{\Lambda}(\chi)$ in backscatter is much less than that of τ^{Λ} . Griggs [1975] has experimentally shown that upward radiance was essentially proportional to τ^{Λ} , independent of aerosol type. Koepke and

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Paper number 96JD02132. 0148-0227/97/96JD-02132\$09.00 Quenzel [1979] carried out theoretical calculations for a wide range of aerosol models, representative of their natural variability, and have shown that variability of $P^{\Lambda}(\chi)$ in backscatter is well within $\approx\pm25\%$ (and can be lowered down to $\pm4\%$ if optimal scattering geometry is used), whereas τ^A varies more than an order of magnitude. The feasibility of τ^{Λ} retrieval from a single satellite reflectance channel using a "universal" aerosol model has been recently confirmed by Kaufman [1993], who has shown experimentally that the relationship between τ^{Λ} and atmospheric path radiance is close to universal. Such a onechannel procedure may be viewed as a normalization of the satellite radiances to a common observation-illumination geometry and their scaling to a range of aerosol optical depths, measured by customary Sun photometry. The advantage of this procedure is that it makes satellite measurements taken in different geographical regions and in different seasons uniform, allowing their direct comparison, and scales them to a well understandable physical quantity such as the aerosol optical depth.

This has been the rationale for the development of an aerosol remote sensing capability at the National Environmental Satellite, Data, and Information Service of the National Oceanic and Atmospheric Administration (NOAA/NESDIS), which commenced in 1977 [*Griggs*, 1983]. The algorithm has been specifically developed for use with channel 1 (0.63 μ m) of the advanced very high resolution radiometer (AVHRR) on-

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Figure 1. Dependence of reflected radiation at 0.63 μ m on aerosol optical thickness and ν (Junge particle size parameter) as computed from the Dave' code for the ocean/atmosphere model used in the phase 1 retrieval algorithm. Solar zenith angle is 60°, satellite zenith angle is 24°, and relative azimuth angle is 180°.

board NOAA polar orbiters, which provides global daily coverage on an operationally consistent long-term basis. Figure 1 illustrates the principle of the retrieval of τ_{SAT}^{A} from the AVHRR backscattered radiance. It shows that if the actual size of aerosol particles deviates from that used in the retrieval model ($\nu = 3.5$, see section 2 for details), than the τ_{SAT}^{A} retrievals would be in error. However, a ground truth experiment conducted in 1980–1981 with collocated Voltz-type Sunphotometer data sets from 10 island and coastal locations and one ship has shown that τ_{SAT}^{A} retrievals with $\nu = 3.5$ have little systematic (bias) error, with a random error of $\sigma_{\tau} \approx 0.05$.

After the phase 1 algorithm was implemented into NOAA operations, another ground truth campaign was conducted, with emphasis on collocating with higher-quality Sunphotometer data from research ships [Reddy et al., 1990; Kovotyev et al., 1993]. The results of this campaign have shown a sizeable systematic underestimation by the satellite retrievals [Ignatov et al., 1995a, b]. After taking into consideration the poor quality of the 1980-1981 Sun-photometer data, it was concluded that the retrieval model used in the phase 1 algorithm was incorrect and must be revised. The phase 1 algorithm and results of its validation are described in section 2. In its revision, extensive analysis of literature and matchup data has been performed (section 3). An improved secondgeneration (phase 2) algorithm has been formulated, validated (bias less than 10%; residual random error $\sigma_{\tau} \approx 0.04$), and applied to data of NOAA 14, launched December 30, 1994 (section 4). Both phase 1 and 2 algorithms use only one channel, so only one unknown can be retrieved. Even though the phase 2 algorithm corresponds to an "average" aerosol better than phase 1, it may still be prone to errors in retrieved τ_{SAT}^{A} because of nonuniversality of the aerosol model (resulting from seasonal and regional variability in the aerosol physical and chemical properties). Measurements in AVHRR channel 2 (0.83 μ m), or, with the launch of NOAA-K in 1996 and the Tropical Rainfall Measurement Mission (TRMM) in 1997, a third reflectance channel at 1.6 μ m, may be potentially used to retrieve particle size information, which may be further used to reduce errors in τ_{SAT}^{A} (section 5). The value of the current product is discussed in section 6, and concluding remarks are given in section 7.

2. Phase 1 Algorithm (July 1987 to September 1994)

2.1. Algorithm Description

The phase 1 algorithm for aerosol optical thickness $\tau^{\rm A}_{\rm SAT1}$ retrieval has been implemented into an already existing sea surface temperature system, which incorporates procedures useful for aerosol analysis such as cloud clearing and global field analysis [McClain et al., 1985]. To avoid the influence of specular reflection from the ocean, the solar side of the orbit and measurements whose viewing angle is less than 40° away from the specular ray have been excluded from the analysis. Experimental weekly production of global 8-day (1°)² composites of τ_{SAT1}^{A} from 2 × 2 arrays of 4-km global area coverage pixels only over oceans commenced in July 1987 [McClain, 1989; Rao et al., 1989; Stowe, 1991]. In July 1989 these global analyses began being archived onto magnetic tape at NOAA's National Climate Data Center, Asheville, North Carolina. In January 1990 they became an operational product at NOAA/ NESDIS.

The principle of the algorithm is shown in Figure 1. At the particular geometry, the $\nu = 3.5$ curve is used to retrieve τ_{SAT1}^{A} from cloud-free satellite radiances R at 0.63 μ m, reduced to dimensionless reflectance by normalization to solar flux, F_s , as $L = \pi(R/F_s)$, and corrected for the Sun-Earth distance. $\tau_{\rm SAT1}^{\rm A}$ is then scaled to $\lambda = 0.50 \ \mu m$, consistent with the retrieval model (multiplying by ratio of scattering coefficients (optical thicknesses) between 0.50 and 0.63 μ m, 1.348). In operation the relationship between τ_{SAT}^{A} and L is described by a four-dimensional look-up table (LUT), precalculated for different τ^{Λ} and view-Sun-azimuth angles using Dave's [1973] scalar radiative transfer model, assuming Elterman's vertical profile of aerosol concentration A(h); midlatitude ozone and water vapor profiles with integrated contents of 0.316 atm cm and 2.91 gm cm⁻², respectively. Other parameters to be specified as input to the LUT are Lambertian oceanic reflectance ρ^{s} ; the volume scattering and absorption coefficients, and the aerosol phase function, $P^{A}(\chi)$, derived from Mie calculations with a prescribed aerosol microphysical model. The oceanic albedo is set to $\rho^{s} = 1.5\%$. The aerosol model uses a modified Junge size distribution: dN/dr = 0 ($r < r_{min}$ and $r > r_{max}$); $dN/dr = A \ (r_{\min} \le r \le r_m); \ dN/dr = A \ (r/r_m)^{-(\nu+1)} (r_m \le r_m)$ $r \le r_{\text{max}}$), where $r_{\text{min}}, r_m, r_{\text{max}}$ are particle radii equal to 0.03, 0.1, and 10 μ m, respectively; size parameter $\nu = 3.5$ (equivalent Angstrom exponent $\alpha = 1.5$; a complex index of refraction n = 1.5-0.0i (albedo of single scattering, ω , equals 1); and A = 22510 cm⁻³ μ m⁻¹ is a normalization constant dependent on an assumed number of particles per unit volume [Dave, 1973]. This model was adopted from Griggs [1983]. The specified modeled LUT establishes a unique relationship between the upward radiance and τ_{SAT1}^{A} (e.g., Figure 1), which is further used to derive τ_{SAT1}^{Λ} globally.

2.2. Global Application of Phase 1 Algorithm to NOAA 11 Data

The phase 1 algorithm has been applied to data of NOAA 11, launched in September 1988. Satellite counts have been converted to radiances using prelaunch coefficients, with an empirical adjustment described by *Ignatov et al.* [1995a]. This product has provided the most complete description to date of the spatial and temporal variability of tropospheric and, with the eruption of Mount Pinatubo, stratospheric aerosols over oceans. Plate 1 shows the variation with time (weekly) from



Plate 1. Weekly 1° zonal time series of τ_{SAT1}^{A} at 0.5 μ m over oceans from NOAA 11 AVHRR.

June 29, 1989, to September 8, 1994, of the zonal average (1° latitude band) τ_{SAT1}^{A} . The most striking feature in Plate 1 is the high values in tropical optical thickness coincident with the eruption of Mount Pinatubo. The rapid increase and then gradual decrease in volcanic aerosol content of the tropical stratosphere has been captured, as well as the seasonal transport of this aerosol to high latitudes [*Stowe et al.*, 1992; *Long and Stowe*, 1994; *Grant et al.*, 1996]. Before the eruption and beginning about a year after the eruption, other seasonal latitudinal variations are evident but to a much smaller degree. These are associated with natural mechanisms which create and disburse aerosols in the troposphere. It is evident from Plate 1 that the northern hemisphere has more aerosol than the southern. This is due to the aerosol sources primarily being restricted to land area (wind-blown dust, fires, and industrial pollutants), of which there are more in the northern hemisphere.

The geographical separation between tropospheric aerosol sources can be seen in Plate 2, which shows the annual mean aerosol optical thickness as depicted by this product from the 2 years before the Mount Pinatubo eruption. It shows that the aerosol over the oceans is far from uniform, being separated into distinct regional areas of maximum optical thickness. The features adjacent to the continents are from wind transported aerosol sources originating on the land: mineral aerosol from the deserts of Africa, Saudi Arabia, and Asia [*Prospero and Carlson*, 1972]; smoke from biomass burning in central and southern Africa, Central and South America, and Indonesia [*Kaufman et al.*, 1990b; *Andreae et al.*, 1994]; and sulfate acrosols from fossil fuel combustion in the highly industrialized





regions of eastern United States, Europe, eastern Asia, and Japan [Husar and Wilson, 1993]. There is also a clear indication of longitudinal bands of aerosol over the southern and northern tropical oceans. These are probably the result of biogenically produced aerosols over the oceans [Charlson et al., 1987]. Of all the aerosol types present over the oceans, the mineral dust is the least similar to the assumed aerosol model and therefore is the most uncertain in terms of optical depth estimates from AVHRR.

2.3. Validation of the Phase 1 Algorithm

Sun-photometer data τ_{SP}^{A} collected during two oceanic cruises of the RV Akademik Vernadsky (hereafter AV-89 and AV-91, respectively), and one on the NOAA ship Mount Mitchell (MM-88; this latter corresponds to NOAA 9 data, which were calibrated using prelaunch calibration but without the empirical adjustment developed for NOAA 11, referenced in section 2.2) were used to quantify the accuracy of the τ_{SAT1}^{A} retrievals. The principles and results of selecting and processing the first two matchup data sets are described by Ignatov et al. [1995a, b]. The SP measurements taken within ± 2 hours of the satellite overpasses were selected. The 10 nearest satellite retrievals, within 300 km of the ship, were then searched. The 68 reliable matchups (20 obtained for AV-89, 38 for AV-91, and 10 for MM-88) cover large areas in the North Atlantic, in different years: 1988 and 1989 (before Mount Pinatubo) and 1991 (early post Mount Pinatubo), thus being representative of a variety of aerosols over the ocean. Averaging of τ_{SAT}^{A} (over

the 10 satellite retrievals closest to the ship) and $\tau_{\rm SP}^{\rm A}$ (averaged within $\approx \pm 2$ hours of satellite overpass) was applied to suppress noise resulting from natural variability and measurement errors in both data sources. Comparing the averaged characteristics is also consistent with the gridded NOAA operational product which provides an objective analysis of τ_{SAT1}^A fields over a $(1^{\circ})^2 \times 1$ week space/time box. The accuracy of ${}^{\Lambda}\tau_{SP}$ is ~0.02-0.03 [Ignatov et al., 1995a, b; Reddy et al., 1990].

Figure 2 shows the results of validation of τ_{SAT1}^{A} retrievals from NOAA 9 and NOAA 11. The regression is



Figure 2. Regression of τ_{SAT1}^{A} against Sun-photometer aerosol optical thickness at 0.5-µm wavelength.



Plate 3. Global distribution of τ_{SAT2}^{A} at 0.63 μ m from NOAA 14 (top) for July 1995 with pre-launch calibration and (bottom) for August 1995 with postlaunch calibration.

$$\tau_{\rm SAT1}^{\rm A} = (0.64 \pm 0.05)\tau_{\rm SP}^{\rm A} - 0.02 \tag{1}$$

with only a slightly nonzero bias. The random error of the regression is about $\sigma_{\tau} \approx 0.045$, with a correlation coefficient of

R = 0.87. Thus, to a first approximation, the maps shown in section 2.2 should be scaled by the formula (1.56 $\tau_{SATI}^{\Lambda} + 0.03$) to more correctly represent the optical thickness at 0.50 μ m observed by Sun photometry.

3. Revision of the Phase 1 Algorithm: Guidance to Phase 2

The cause of this bias was the subject of an intense investigation [Ignatov et al., 1995a]. After removal of the scaling of τ_{SAT}^{A} from 0.63 to 0.50 μ m and recalibration of the satellite radiances with coefficients determined from postlaunch analysis and adjusted for the dates of the matchup data sets [Rao and Chen, 1995], the inadequacy of the retrieval model is the only reason of discrepancy between τ_{SAT}^{A} and τ_{SP}^{A} . The diagnostic analysis was aided by use of the linearized single scattering approximation of the radiative transfer equation [Viollier et al., 1980; Gordon and Morel, 1983], given by the following equation for τ^{A} retrieval:

$$\tau_{\rm SAT}^{\rm A} = \frac{\rho - \rho^{\rm R} - \rho^{\rm S} T}{\omega P^{\rm A}} \, 4\,\mu_{\rm s}\mu_{\nu} \tag{2}$$

where ρ is an apparent reflectance of the ocean-atmosphere system, defined by normalization of L to the solar flux and overhead illumination geometry; ρ^{R} is the Rayleigh scattering contribution to the total signal ρ ; ρ^{S} is the diffuse surface reflectance; T is the total atmospheric transmittance; P^{A} and ω are the aerosol phase function and albedo of single scattering; $\mu_{\nu} = \cos \Theta_{\nu}; \ \mu_{s} = \cos \Theta_{s}; \ \Theta_{\nu} \ \text{and} \ \Theta_{s} \ \text{are the view and solar}$ zenith angles. Equation (1) shows that the errors in τ_{SAT}^{A} may result from incorrect ρ^{S} , ω , or P^{A} since all other terms are reasonably well known. Note that ρ^{s} participates in (2) as an additive term, and (ωP^A) as a multiplicative one. This suggests that the negative bias in the satellite retrievals (removed in operational practice by setting these to zero; note the large number of zero values in Figure 2) comes from having overestimated the oceanic reflectance, and the depressed slope $(d\tau_{SAT}^A/d\tau_{SP}^A < 1)$ from an incorrect aerosol model (ω and/or P^{Λ}).

Analysis of the literature by *Ignatov et al.* [1995a] suggests that $\rho^{\rm S} \approx 0.2\%$ rather than 1.5% for deep clean ocean. Correcting this in the retrieval model and taking into account specular reflection of diffuse downward scattered radiation in a single scattering approximation [*Viollier et al.*, 1980; *Gordon and Morel*, 1983] removed the negative bias, but a multiplicative underestimation in $\tau_{\rm SAT}^{\rm A}$ remained. This result strongly indicated that the $P_{3.5}^{\rm A}(\chi)$ used in the operational retrievals is not representative of the average Atlantic aerosol.

Ignatov et al. [1995a, b] attempted to reconcile τ_{SAT}^{A} and τ_{SP}^{A} by inferring a microphysical model from the wavelength dependence of the τ_{SP}^{A} . This gave an Angstrom exponent of about $\alpha = 0.6$, which led to setting $\nu \approx 2.5$ ($\nu = \alpha + 2$) in the Junge size distribution. However, using this model removed the systematic discrepancy between τ_{SAT}^{A} and τ_{SP}^{A} only when it was supplemented by adjusting the imaginary part of the aerosol refractive index to 0.01. Correlation statistics indicate that this algorithm estimates τ_{SAT}^A with little systematic bias or offset: $\tau_{\text{SAT}}^{\text{A}} \approx -0.004 + (0.98 \pm 0.08) \tau_{\text{SP}}^{\text{A}}$. The correlation coefficient for the 68 cases is R = 0.92, with a random error of $\sigma_{\tau} \approx$ 0.046. Additional analyses of Mie calculations have shown that the imaginary part influences the results of $\tau^{\mathbf{A}}_{\mathbf{SAT}}$ retrieval mainly through decreasing the phase function in the backscatter direction (about 30%) rather than through absorption itself (only 10%, consistent with the corresponding albedo of single scattering, $\omega \approx 0.9$).

The value of $\omega \approx 0.9$ was thought of as unrealistic, and A. Ignatov (submitted manuscript, 1996) tried to estimate an empirical phase function, $P_{\text{emp},i}^{\Lambda}(\chi_i)$, at each matchup point *i* for



Figure 3. Two model phase functions: $P_{3,5}^A$ (Junge, $\nu = 3.5$) (dashed) and P_{LN}^A derived from Mie calculations with lognormal size distribution ($r_m = 0.1 \ \mu m$; $\sigma = 2.03 \ \mu m$; n = 1.40-0.0i] (dotted), and results of the empirical phase function, P_{emp}^A , estimation for AV-89 (crosses) and AV-91 (open circles). All phase functions are normalized to 4π .

the particular scattering angle χ_i , having the τ_{SP}^A known from the Sun-photometer measurements. The retrieved values of $P_{emp,i}^A(\chi_i)$ are plotted versus scattering angle in Figure 3, each point corresponding to one matchup. All the experimental points fall well below $P_{3,5}^A$, which explains the systematic underestimation in the operational retrievals (Figure 2 and (2)). One has therefore to find an aerosol phase function which describes the experimental points in Figure 3.

According to D'Almeida et al. [1991], clean maritime aerosol is composed mostly of two components: sea salt (in which the nucleation mode dominates) and sulfate. For a typical relative humidity over the oceans of 70-80% the lognormal distributions of each are close together with $r_m \approx 0.1 \ \mu m$, and $\sigma = 2.03 \ \mu m$. Complex indices of refraction for both are also close, about 1.4-0.0*i*. As a result, the bimodal size distribution of the marine aerosol can be effectively represented by a monomodal lognormal function. Its phase function, shown in Figure 3, describes the experimental points quite well. Figure 4 shows τ_{SAT}^{A} retrieved with the described lognormal model versus $^{A}\tau_{SP}$. The regression gives

$$\tau_{\rm SAT}^{\rm A} \approx 0.01 + (0.91 \pm 0.04) \tau_{\rm SP}^{\rm A} \tag{3}$$

with R = 0.93 and an error of $\sigma_{\tau} \approx 0.04$. This error is inherent to the present single-channel algorithm, and can possibly be reduced with future multispectral algorithms.

Note that the A. Ignatov (submitted manuscript, 1996) study includes only matchups with large aerosol loadings ($\tau_{SP}^A \ge 0.1$), which are not typical for clean maritime conditions [Smirnov et al., 1995]. As a result, the derived empirical phase function may not be representative of clean maritime aerosol, and one needs to be cautious in the geophysical and microphysical interpretation of the derived phase function. Also, application of the derived phase function in the case of small aerosol loadings may be questionable. However, for practical retrievals from AVHRR, error in P^A is of more concern for large aerosol loadings, since absolute systematic error $\delta \tau_{SAT}^{A}$ is proportional to $\tau_{\text{SAT}}^{\text{A}}$ for the same relative error in P^{A} (see equation (2)). Note that reducing the absolute uncertainty $\delta \tau_{SAT}^{A}$ is important for climate modeling [e.g., Charlson et al., 1992]. To that end, the lognormal phase function is a reasonable candidate for replacing, the $P_{3,5}^{A}(\chi)$ used in the phase 1 retrieval algorithm.



Figure 4. Regression of τ_{SAT2}° against same Sun-photometer data set as in Figure 4 but at 0.63 μ m.

The phase 2 model's apparent contradiction to other literature, which recommends using a two- or three-mode lognormal function [World Climate Program, 1983, 1986; Shettle and Fenn, 1979; Gathman, 1983; Gras, 1991; Hoppel et al., 1994], requires further investigation but is of little concern for practical use, since the P_{LN}^{A} is close to the empirically derived one. The premise of that model is its simplicity, in particular fewer numbers of descriptive parameters (each mode requires values of modal radius, standard deviation, and complex index of refraction). This model has been adopted for the operation of NOAA 14, launched in December 1994. Its summary and some preliminary attempts at validation are described in the next section.

4. Phase 2 Algorithm (March 1995 Onward)

4.1. Algorithm Summary

The second-generation (phase 2) acrosol retrieval algorithm has many features common with phase 1 described in section 2. The difference between the two comes from the fact that the phase 1 algorithm has been carefully validated, and its parameters have been revised based on experimental data, and careful analysis of the recent literature. The Lambertian surface reflectance in the second generation product is lowered 7.5 times, from 1.5% down to 0.2%. The bidirectional Fresnel reflectance from the flat ocean surface is approximately taken into account by introducing a diffuse glint correction to the phase function, derived from the single scattering approximation [Viollier et al., 1980; Gordon and Morel, 1983]. Its application to the phase 2 algorithm is described in detail by Ignatov et al. [1995a]. We understand restrictions of this procedure, especially at high aerosol loadings and in the vicinity of glint, and are working now on more appropriate quantification of this effect. The aerosol phase function used in the phase 2 algorithm, was derived assuming a lognormal size distribution with $r_{\rm m} = 0.1 \ \mu {\rm m}$, $\sigma = 2.03 \ \mu {\rm m}$; n = 1.4-0i.

No scaling is made anymore from the wavelength of measurement 0.63 μ m to the wavelength of 0.50 μ m, recommended by the World Meteorological Organization as a reference. The reason is that this scaling factor is strongly atmospheric model dependent and may result in a systematic error in the retrieved τ_{SAT}^{Λ} . For those wanting to convert, the scaling factor for the first-generation model is 1.348 and for the second-generation is 1.164.

4.2. Testing Phase 2 Algorithm With NOAA 11 Pathfinder Data (September 1989)

Expected differences in the performance of the two algorithms has been verified globally with data from the AVHRR Atmosphere Pathfinder Program. This is a NASA/NOAA joint program to reprocess NOAA satellite data with community consensus calibration and remote sensing algorithms into a long-term product data set for weather and climate change research [University Corporation for Atmospheric Research (UCAR), 1994]. The AVHRR atmosphere pathfinder data set is being produced at NOAA/NESDIS. Phase A1 of this program uses the CLAVR-1 (clouds from AVHRR phase 1) cloud screening algorithm [Stowe et al., 1991], to produce among some 71 parameters, cloud-free and cloudy radiances in all five channels averaged in 110 km equal area regions over the globe. Phase A2 uses these area averaged radiances to derive aerosol optical thickness over oceans and radiation budget parameters at the top of the atmosphere. Initially, an 18-month "benchmark" period from April 1987 to November 1988 (NOAA 9) and September 1989 (NOAA 11) is being processed. For this initial run, τ_{SAT}^{A} has been computed with both phase 1 and 2 aerosol algorithms for validation purposes and for continuity with the over 5 years of NOAA 11 data. Figure 5 shows the result of plotting $\tau_{\text{SAT1}}^{\Lambda}$ versus $\tau_{\text{SAT2}}^{\Lambda}$ (converted to 0.50 μ m for consistency) for September 6, 1989, NOAA 11 data for all 110-km grid cells with cloud-free radiances. The regression line has the form

$$\tau_{\rm SAT1}^{\rm A} = (0.69 \pm 0.003)\tau_{\rm SAT2}^{\rm A} - 0.03 \tag{4}$$

with R = 0.97 and $\sigma_{\tau} = 0.023$. The relationship between $\tau_{\text{SATI}}^{\Lambda}$ and $\tau_{\text{SP}}^{\Lambda}$ (cf. equation (1)) produced a slope and intercept which agrees to those in (3) within the error of estimation. Thus it can be concluded that when globally applied, the phase 2 algorithm yields a result that is similar to what would have been observed had we performed validation with Sun photometers.

4.3. Application of the Phase 2 Algorithm to NOAA 14 Data

After the failure of NOAA 11 AVHRR in September 1994, operational production of τ_{SAT2}^{A} did not commence until March 1995 using NOAA 14 data. As an example of these initial results, Plate 3 shows a global image of τ_{SAT2}^{A} for the month of July 1995. It reproduces many of the features present



Figure 5. Regression of τ_{SAT1}^{c} against τ_{SAT2}^{c} at 0.5 μ m globally, using the AVHRR atmosphere pathfinder processing system for September 6, 1989.



Figure 6. Effect of prelaunch calibration error on τ_{SAT2}^{A} .

in the NOAA 11 maps (cf. Plate 2), but the optical depths appear to be elevated by more than the expected factor of 1.34 (a ratio of underestimation of 1.56 from (1) and conversion factor from 0.63 to 0.5 μ m of 1.164). Postlaunch investigations into the calibration of NOAA 14 AVHRR channels 1 and 2 (N. Rao, personal communication, 1995), have confirmed revision of prelaunch calibration by R. Mitchell (In-flight characteristics of NOAA AVHRR channels 1 and 2, submitted to IEEE Transactions on Geoscience and Remote Sensing, 1996) and have led to the conclusion that the prelaunch values of calibration coefficients were in error. Rao's revision of these terms was implemented into operations the end of July. The effect on the NOAA 14 aerosol product is shown also in Plate 3, where much more reasonable values of τ_{SAT2}^{A} appear to be distributed over the globe for August 1995, having about the right degree of enhancement compared with Plate 2. Figure 6 shows quantitatively the effect of applying Rao's postlaunch calibration coefficients to a sample of NOAA 14 data. This shows that τ_{SAT2}^{A} was being overestimated before August 1995 by about 0.08, on average.

The Sun-photometer method of validation is nearing a reality for NOAA 14. NESDIS has begun collecting τ_{SAT2}^{A} observations in real time at seven coastal locations where well calibrated and maintained Sun-photometer observations are available, but none of these data sets have been processed as yet [Holben et al., 1996].

5. Possible Enhancements to the AVHRR Product

5.1. Multispectral Retrieval Algorithms

A third-generation (phase 3) algorithm is currently under development. It uses two channels from the current AVHRR. The algorithm will be used to estimate an additional parameter: the Angstrom wavelength exponent α traditionally measured in Sun photometry. This could be used to retrieve an aerosol size parameter, which in turn could possibly be used to adjust the size distribution (scattering phase function) used in the single-channel retrieval algorithm [e.g., *Durkee et al.*, 1991]. This could potentially remove residual errors resulting from departures of the actual aerosol size distribution from the one used in the single-channel retrieval model.

One example of such an algorithm was developed several years ago [*Ahmad et al.*, 1989]. This approach relied upon regression relationships from theoretical radiative transfer

model calculations which relate the ratio of aerosol path radiance between two channels to the exponent of the Junge power law model. The retrieved Junge exponent ($\nu = \alpha + 2$) determines the aerosol particle size distribution. It is then used to retrieve the optical thickness from one of the channels, again through the use of theoretical regression relationships, but specifically for that particular Junge exponent (aerosol size distribution).

A graphical illustration of such an algorithm is shown in Figure 7a. On the ordinate normalized radiance (defined in section 2.1) is plotted at the effective wavelength (average wavelength weighted by product of channel response function and solar spectral irradiance) of channel 2 (0.83 µm) of AVHRR, and on the abscissa the same quantity is plotted but for channel 1. Solar and satellite geometries are identical to those used in Figure 1. This has been done for three atmospheric models (tropical, midlatitude, and Arctic) to represent the possible range of water vapor content which affects the radiance in channel 2 due to its filter encompassing several strong bands of water vapor absorption. To each of these three atmospheric models is added the ocean surface reflection model and Junge aerosol size distribution dependent scattering and absorption models used in our initial efforts to improve upon the phase 1 single-channel algorithm [Ignatov et al., 1995a]. Three values of the Junge exponent have been used in this example: $\nu = 3.5$ (effective radius, third moment divided by second moment of size distribution, $r_{\rm eff} = 0.21 \ \mu m$), $\nu = 2.5$ $(r_{\rm eff} = 0.79 \ \mu {\rm m})$ and $\nu = 1.5 \ (r_{\rm eff} = 3.64 \ \mu {\rm m})$. The total number of particles in each model atmosphere was adjusted to yield the same aerosol optical thickness at $0.5 \ \mu m$ (heavy dashed line in Figure 7 shown only for midlatitude model). The result is a set of three families of curves, one for each Junge exponent value, relating the change in radiance for the two channels as optical thickness changes. Making up each of these are three curves for each of the three different atmospheric models, showing the effect of variable water vapor on the radiance in channel 2.

Given an AVHRR observation over a clear ocean at the viewing and illumination angles of Figure 7 (it should fall somewhere in the space bounded by the outermost curves), the corresponding values of ν and τ_{SAT3}^A can then be read. Automating this procedure is relatively trivial, once a similar set of figures (i.e., look-up tables) have been computed for the range of geometries likely to be observed. The accuracy of the retrieved optical thickness and Junge parameter is dependent not only on the correctness of the assumed model parameters but also on measurement error. As is evident from Figure 10a, measurement noise will introduce larger relative error in the derived parameters as the optical thickness of the aerosol approaches zero (curves converge at $\tau_{SAT3}^{A} = 0$, where the ability to separate the effect of different aerosol models vanishes). Quantifying the magnitude of these errors and their sensitivity to independent variables is the subject of current research (e.g., A. Ignatov, Accuracy of aerosol Angstrom exponent derived from two-channel satellite measurement over the oceans, submitted to Applied Optics, 1996).

One source of error, the variable amount of water vapor, can be reduced in two ways: either by providing independent knowledge of the water vapor amount and vertical profile relative to the aerosol profile, not presently available from satellite remote sensing technology, or by using a pair of channels weakly affected by water vapor absorption. Such a pair of channels (0.63 and 1.61 μ m) will become available from



Figure 7. Illustration of the principle of τ_{SAT3}^{A} and ν retrieval from two-channel reflectance measurements at (a) 0.63 and 0.83 μ m and (b) 0.63 and 1.61 μ m (the heavy dashed lines are contours of τ_{SAT3}^{A} at 0.5 μ m, shown only for the midlatitude model).

AVHRR with the NOAA-KLM series of satellites, scheduled to commence in 1996. However, because of data storage constraints on-board the spacecraft, only the afternoon satellite, NOAA-L, will provide global coverage at 1.61 μ m in daytime. To show the potential of retrieving improved aerosol information from these two channels, Figure 7b has been constructed. Not only is the effect of variable water vapor greatly reduced, but the separation of the three families of curves has been increased. This means that for the same relative measurement error, one should be able to obtain more accurate aerosol information at lower values of optical thickness than with the current two AVHRR reflectance channels. A similar pair of channels are available from the Thematic Mapper on the Landsat series of satellites, from the along track scanning radiometer (ATSR) instrument on the ERS-2 satellite, and from the visible infrared scanner (VIRS) instrument to be launched on the TRMM satellite in 1997.

5.2. Other Potential Enhancements

With regard to other aerosol remote sensing issues not yet addressed in our program, it is highly desirable to make retrievals over surfaces with relatively high reflectivity, such as in the regions of oceanic specular reflection or for most land surfaces. A climatology of clear-sky reflectance and its variance could be used as input to radiative transfer model LUTs for the extraction of optical thickness and particle size parameters in these areas, although the sensitivity of reflected radiation to the presence of aerosols is diminished relative to that over dark ocean surfaces [Kaufman et al., 1990a]. Also, use of infrared retrieval techniques, as other investigators have shown [Legrand et al., 1989], can retrieve aerosol optical thickness information over certain geographical areas, such as deserts. Eventually, with the launch of the ocean color temperature scanner (OCTS) and polarization and directionality of Earth reflectances (POLDER) instruments on the Japanese ADEOS satellite and SeaWifs instrument on a commercial satellite in 1996, and the launch of the EOS polar platform instruments (moderate-resolution imaging spectrometer (MODIS), multiangle imaging spectrometer (MISR), and Earth observing scanning polarimeter (EOSP)) between 1998 and 2003, many other spectral channels as well as linear polarization parameters will become available for the description of the atmospheric aerosol content.

Also, even at the single-channel algorithm level, further improvements could be made by more accurately describing the diffuse glint and introducing the direct glint, both of which will require information about wind speed. Using a microphysical aerosol model that varies by geographical region and provides a phase function that closely matches empirically derived values for each region separately may also improve accuracy. Establishment of ground truth sites with in situ as well as remote sensing data would greatly assist the validation of multispectral techniques.

6. Discussion

The phase 2 retrievals show reasonable agreement with Sunphotometer data on the average. Residual scatter indicates that the phase function for real maritime aerosol varies and can be assumed "universal" only within the reported uncertainty limits. For single-channel retrievals the only alternative to a "universal" empirical phase function would involve its parameterization versus τ^{Λ} : $P^{\Lambda} = f(\tau^{\Lambda})$, which allows one to keep one unknown under retrieval τ^{Λ} . The analysis of A. Ignatov (submitted manuscript, 1996) did not confirm this hypothesis, however. Also, oceanic diffuse reflectance is not a universal constant, $\rho^{S} = 0.2\%$. It may be much higher over turbid coastal waters, and is bidirectional.

We believe that a more complicated two-channel procedure should be introduced only after the potential of the onechannel algorithm is fully realized, its advantages and deficiencies understood, and the necessity and sufficiency of additional measurements convincingly proved. To that end, using the microphysical model which closely reproduces the empirically derived phase function is the logical next step, providing a bridgehead between the present and future generation of aerosol retrieval algorithms. Additionally, the spectral position of AVHRR channel 1 is not far from the optimal for aerosol retrieval, recommended by *Koepke and Quenzel* [1981]. So far, the rms accuracy of the Phase 2 algorithm is $\sigma_{\tau} \approx 0.04$. This estimate is consistent with results of *Kaufman* [1993], who shows that using a universal relationship between aerosol optical depth and path radiance leads to a maximum error of 0.10 (about $3\sigma_{\tau}$, assuming a Gaussian distribution of error).

It is recognized that the AVHRR instrument is not the optimum instrument for making aerosol measurements. It has no onboard calibration for its two (a third will be added in 1997) reflectance measuring channels. With the development of spaceborne multispectral imaging spectrometers like MO-DIS [King et al., 1992], multiangular viewing radiometers like MISR [Martonchik and Diner, 1992], and ATSR [Mutlow et al., 1994], and polarimetric imaging radiometers like POLDER [Deuze et al., 1993], remote sensing of aerosol physical and chemical properties should be improved beyond AVHRR's capabilities. Nonetheless, the work that has been done with AVHRR has clearly demonstrated that aerosol remote sensing can be done from space, and that the resulting products over global oceans can provide valuable information on spatial and temporal variability of this important climate forcing parameter [Joussaume and Sadourny, 1989; Long and Stowe, 1994; Lacis and Mishechenko, 1995]. Efforts are now underway to reprocess the entire AVHRR record from the NOAA afternoon satellites back to 1981 [UCAR, 1994]. Such a climatology. having accounted for calibration drift and the same aerosol retrieval algorithm throughout, will provide a unique data set from which a greater knowledge of the Earth's aerosol characteristics will be gleaned, and against which the value of the future instruments can be measured.

7. Conclusion

The second generation single-channel aerosol retrieval algorithm has been described. It has been implemented into the operational processing system for the NOAA 14 satellite, launched on December 30, 1994. The algorithm is based on analysis of three matchup Sun-photometer-NOAA satellite data sets, which revealed errors in the operational NOAA/ NESDIS τ_{SAT1}^{A} retrievals. A fundamental revision of the oceanic reflectance model was undertaken from analysis of the scientific literature. A new empirical aerosol phase function has been derived from the matchup data. A microphysical aerosol model, which matches the empirical phase function, has been chosen for the retrieval of τ_{SAT}^A . These adjustments to the first generation NOAA/NESDIS aerosol product bring, to a first approximation, the Sun-photometer and satellite data into better than 10% agreement. It also has been shown that when applied to a global analysis with AVHRR atmosphere pathfinder data sets, the phase 2 algorithm provides $\tau^{\Lambda}_{S\Lambda T2}$ which deviates from $\tau^{\rm A}_{\rm SAT1}$ in the same way that $\tau^{\rm A}_{\rm SP}$ deviated from τ_{SAT1}^{A} . Examples of the NOAA 14 aerosol global maps have been presented, which before August 1995, are biased high by AVHRR calibration errors.

Since the τ_{SAT}^{Λ} are retrieval model dependent, we call the satellite product a "radiatively equivalent" aerosol optical thickness. The phase 2 algorithm provides unbiased, in the mean statistical sense, aerosol retrievals over oceans. It can be viewed as a procedure for removing the strong angular dependencies from upward radiances (normalizing them to a common geometry), with scaling to aerosol optical thickness measured by Sun photometry. Any remaining noise in the satellite

retrievals ($\sigma_r \approx 0.04$, see (3)) follows from variability of the aerosol phase function, aerosol absorption, and oceanic reflectance about their average values, typical for the oceans. Some of this variability may be potentially accounted for by further adding other spectral or angular measurements to the retrieval algorithms; for those, the present NOAA/NESDIS aerosol product provides a reference level from which improvements can be measured. An example of such an algorithm, with the improvements possible with the 1.61- μ m channel on the NOAA/KLM series of AVHRR instruments, was presented. Further investigations are needed, together with intensive field measurements, to establish the most globally robust multichannel aerosol retrieval algorithm.

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