Fast and simple model for atmospheric radiative transfer

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Abstract. Radiative transfer models (RTMs) are of utmost importance for quantitative remote sensing, especially for compensating atmospheric perturbation. A persistent trade-off exists between approaches that prefer accuracy at the cost of computational complexity, versus those favouring simplicity at the cost of reduced accuracy. We propose an approach in the latter category, using analytical equations, parameterizations and a correction factor to efficiently estimate the effect of molecular multiple scattering. We discuss the approximations together with an analysis of the resulting performance and accuracy. The proposed Simple Model for Atmospheric Radiative Transfer (SMART) decreases the calculation time by a factor of more than 25 in comparison to the benchmark RTM 6S on the same infrastructure. The relative difference between SMART and 6S is about 5% for spaceborne and about 10% for airborne computations of the atmospheric reflectance function. The combination of a large solar zenith angle (SZA) with high aerosol optical depth (AOD) at low wavelengths lead to relative differences of up to 15%. SMART can be used to simulate the hemispherical conical reflectance factor (HCRF) for spaceborne and airborne sensors, as well as for the retrieval of columnar AOD.

1 Introduction

The terrestrial atmosphere attenuates the propagation of the solar radiation down to the Earth’s surface and back up to a sensor. The scattering and absorption processes involved disturb the retrieval of quantitative information on surface properties. Radiative transfer models (RTMs) and their inversions are commonly used to correct for such effects on the propagation of light. Well-known RTMs are 6S (Second Simulation of a Satellite Signal in the Solar Spectrum) (Vermote et al., 1997), SCATRAN (Rozanov et al., 2005), SHARM (Muldashev et al., 1999; Lyapustin, 2005), RT3 (Evans and Stephens, 1991), RTMOM (Gov- aerts, 2006), RAY (Zege and Chaikovskaya, 1996), STAR (Ruggaber et al., 1994) and Pstar2 (Nakajima and Tanaka, 1986; Ota et al., 2010), as well as DISORT (Stamnes et al., 1988), which is used in MODTRAN (Berk et al., 1989), STREAMER (Key and Schweiger, 1998) and SBDART (Ricchiazzi et al., 1998). These accurate but complex RTMs are frequently run in a forward mode, generating look-up tables (LUTs), which are later used during the inversion process for atmospheric compensation (Gao et al., 2009) or aerosol retrieval (Kokhanovsky, 2008; Kokhanovsky and Leeuw, 2009; Kokhanovsky et al., 2010), for instance. There are also a series of highly accurate, but computationally intensive Monte Carlo photon transport codes available. However, the best accuracy may not be always desirable for a RTM. Approximative equations have been developed before computers were widely available (Hammad and Chapman, 1939; Sobolev, 1972). With regard to the growing size and frequency of remote sensing datasets, approximative and computationally fast RTMs are becoming relevant again (Kokhanovsky, 2006; Katsev et al., 2010; Carrer et al., 2010). In particular, RTMs of the vegetation canopy and further algorithms that exploit data from imaging spectroscopy instruments (Itten et al., 2008) often rely on fast atmospheric RTM calculations.

In this context, we propose the fast Simple Model for Atmospheric Radiative Transfer (SMART). It is based on approximative analytical equations and parameterizations, which represent an favourable balance between speed and accuracy. We consider minimised complexity and computational speed as important assets for downstream applications and define an acceptable uncertainty range of up to 5–10% for the modelled reflectance factor at the sensor level, under typical mid-latitude remote sensing conditions. SMART can
therefore be used as a physical model, maintaining a cause-
and-effect relationship in atmospheric radiative transfer. In
stead of depending on the classic LUT approach, it permits
parameter retrieval in near-real-time. This enables the rapid
assessment of regional data requiring exhaustive correction,
such as imaging spectrometer data. Furthermore, it supports
the straightforward inversion of aerosol optical depth (AOD;
$\tau_{\text{aer}}$) by implementing radiative transfer equations as a func-
tion of $\tau_{\text{aer}}$. The theoretical feasibility for the retrieval of
aerosols in terms of the sensor performance was shown in
Seidel et al. (2008) for the APEX instrument (Itten et al.,
2008).

In this paper, we describe the two-layer atmospheric model
with the implementation of approximative radiative transfer
equations in both layers and at the Earth’s surface. We then
assess the accuracy and performance of SMART in com-
parison with 6S.

2 SMART – a simple model for atmospheric
radiative transfer

A remote sensing instrument measures the spectral radiance as a function of the spectral atmospheric
properties and the illumination/observation geometry
$R_{\lambda}(t_{\text{s}}, P_{\lambda}(\Theta), \omega_{\text{s}}; \mu_{0}, \mu, \phi - \phi_{0})$, where $t_{\text{s}}$ is the optical
depth, $P_{\lambda}(\Theta)$ is the phase function at the scattering angle $\Theta$, \omega_{s} is the single scattering albedo, $\mu_{0} = \cos \theta_{0}$, $\mu = \cos \theta$, $\theta_{0}$ and $\theta$ represent
the solar and viewing zenith angles (SZA, VZA), $\phi - \phi_{0}$ is the relative azimuth between viewing $\phi$ and
directional $\phi_{0}$. However, from a modelling perspective,
it is more convenient to use a dimensionless reflectance
function. The relationship between radiance and reflectance
is given by:

$$ R_{\lambda} = \frac{\pi L_{\lambda}}{\mu_{0} F_{0,\lambda}}, $$

where $F_{0,\lambda}$ is the spectral solar flux or irradiance on a unit
area perpendicular to the beam. For readability, we omit
the arguments. The subscripted wavelength denotes spectral
dependence.

SMART assumes a plane-parallel, two-layer atmosphere.
We will use the superscript I to denote the upper layer, super-
script II for the lower layer. While the lower layer contains
aerosol particles and molecules, the upper layer contains only
molecules. The surface elevation, the transition altitude of the
two layers, as well as the top-of-atmosphere (TOA) altitude
can be chosen freely. The planetary boundary layer (PBL) height
is a good estimate for the vertical extent of the lower
layer. The sensor altitude can be set to any altitude within
the atmosphere or to the TOA. Altitudes are related to
air pressure $p$ according to the hydrostatic equation. This
1-D coordinate system is used in Eqs. (3) and (25) to
determine $t_{\text{s}}$ and to scale the atmospheric reflectance and trans-
mitance function corresponding to a specific altitude within
atmosphere.

SMART accepts any combination of $\tau_{\lambda}$, $\theta_{0}$, $\theta$ and
$\lambda$. The current implementation executes on the 2-D ar-
ray $[\lambda, \tau_{550nm}]$, where $\lambda \in [400nm, 800nm]$ and $\tau_{550nm} \in
[0.0, 0.5]$. The spectral dependence of the AOD is approx-
imated by:

$$ \tau_{\lambda} = \tau_{550nm} \left( \frac{\lambda}{550nm} \right)^{-a}, $$

according to Ångström’s law (Ångström, 1929). Aerosol
optical properties, such as the asymmetry factor $S_{1,\lambda}$, $\omega_{1,\lambda}$
and the Ångström parameter $\alpha$ are taken from d’Almeida
et al. (1991) for the following aerosol models: clean-
continental, average-continental, urban, clean-maritime,
maritime-polluted and maritime-mineral.

2.1 Radiative transfer in layer I

By definition, the layer I contains no aerosols and the to-
total optical depth is therefore given by the molecular optical
depth $r_{\lambda}^{\text{mle}} = r_{550nm}^{\text{mle}} (1 - h_{\text{PBL}})$, where

$$ h_{\text{PBL}} = \frac{\rho_{\text{SFC}} - \rho_{\text{TOA}}}{\rho_{\text{SFC}} - \rho_{\text{TOA}}} $$

is the relative height of the PBL within the atmosphere. It
ranges from 0 at the surface (SFC) to 1 at TOA. Values for
$r_{550nm}^{\text{mle}}$ are computed using semi-empirical equations from Bod-
haine et al. (1999).

The downward total transmittance $T_{\lambda}^{\text{dfs}}$ is the sum of the
downward direct transmittance $T_{\lambda}^{\text{dfs}}$ and the downward dif-
fuse transmittance $T_{\lambda}^{\text{dfs}}$:

$$ T_{\lambda}^{\text{dfs}} = T_{\lambda}^{\text{dfs}} + T_{\lambda}^{\text{dfs}} = e^{-v_{0}} + T_{\lambda}^{\text{dfs}} e^{-u_{0} - v_{0} r_{\lambda}^{\text{dfs}} - w_{0} (r_{\lambda}^{\text{dfs}})^{2}}. $$

$T_{\lambda}^{\text{dfs}}$ is approximated by using a fast and accurate pa-
parameterization suggested by Kokhanovsky et al. (2005) for
$\omega_{e} = 1$, where

$$ u_{0} = \sum_{m=0}^{3} h_{m} \mu_{0}^{m}, $$

$$ v_{0} = p_{0} + p_{1} \mu_{0}, $$

$$ w_{0} = q_{0} + q_{1} \mu_{0}. $$

The constants $p_{0}, q_{0}, p_{1}, q_{1}, p_{2}, q_{2}$ and $h_{m}$ are parameterized
using polynomial expansions with respect to $g_{\lambda}$, e.g.

$$ p_{0} = \sum_{s=0}^{3} p_{0,s} g_{\lambda}. $$

$p_{0,s}$ and all other expansion coefficients are given in
Kokhanovsky et al. (2005). The upward transmittance $T_{\lambda}^{\text{dfs}}$
is defined according to Eqs. (4) to (8) by substituting $\mu_{0}, u_{0}, v_{0}, w_{0}$
for $\mu, u, v, w$, respectively.
The transmitted light is scattered in all directions. The ratio of scattering to total light extinction $\omega_\lambda$ and the angular distribution of the scattered light $P_\lambda(\Theta)$ are used to describe the scattering process. To simplify the approach, the total intrinsic atmospheric scattering function can be decomposed into the single scattering approximation (SSA) and multiple scattering (MS). The first order atmospheric reflectance function $R_\lambda^{1,\text{SSA}}$ can be expressed using the analytical equation as given in van de Hulst (1948); Sobolev (1972); Hansen and Travis (1974); Kokhanovsky (2006):

$$ R_\lambda^{1,\text{SSA}} = \frac{\omega_\lambda^{\text{mlc}} P_\lambda^{\text{mlc}}(\Theta)}{4(\mu_0 + \mu)} \left(1 - e^{-m\tau}\right), \quad (9) $$

where the molecular single scattering albedo $\omega_\lambda^{\text{mlc}} = 1$ and the molecular (Rayleigh) scattering phase function for reflected, unpolarised solar radiation is given by:

$$ P_\lambda^{\text{mlc}}(\Theta) = \frac{3}{4} \left(1 + \cos^2 \Theta\right), \quad (10) $$

with the scattering angle

$$ \Theta = \arccos \left[-\mu_0 \mu + \cos(\phi - \phi_0) \sqrt{(1 - \mu_0)(1 - \mu)}\right], \quad (11) $$

and the geometrical air mass factor $m = (\mu_0^{-1} + \mu^{-1})$. $P_\lambda^{\text{mlc}}(\Theta)$ is plotted in Fig. 1.

Standard RTMs spend most of their computational time calculating multiple scattering with iterative integration procedures. In the case of layer I, we therefore suggest a generic correction factor $f_{\mu_0}^{\text{corr}}$ to approximate Rayleigh multiple scattering. We derive one $f_{\mu_0}^{\text{corr}}$ per SZA as a function of $\lambda$ and $\tau$ from accurate MODTRAN/DISORT calculations, however without polarisation. The correction factor is defined as the ratio between the total reflectance and the SSA at sensor level:

$$ f_{\mu_0}^{\text{corr}}(\lambda, \tau) = \frac{R_\lambda^{\text{Sensor,MODTRAN}}}{R_\lambda^{\text{Sensor,SSA,MODTRAN}}} . \quad (12) $$

The total reflectance function of layer I is then given by Eqs. (9) and (12):

$$ R_\lambda^I = R_\lambda^{1,\text{mlc}} + \frac{\omega_\lambda^{\text{aer}} P_\lambda^{\text{aer}}(\Theta)}{4(\mu_0 + \mu)} \left(1 - e^{-m\tau}\right) f_{\mu_0}^{\text{corr}} . \quad (13) $$

### 2.2 Radiative transfer in layer II

The down- and upward total transmittances $T_{\lambda}^{\text{II} \downarrow}$, $T_{\lambda}^{\text{II} \uparrow}$ in layer II are calculated according to Eq. (4) by using $g_\lambda^{\text{aer}}$ and substituting $\tau_1^{\text{aer}}$ to the total spectral optical depth of layer II $\tau_\lambda^{\text{II}} = \tau_\lambda^{\text{aer}} + \tau_\lambda^{\text{mcl}} h^{\text{PBL}}$.

The atmospheric reflectance function of layer II is simplified by the decomposition into molecular and aerosol parts. As a consequence, the aerosol-molecule scattering interactions are neglected. The related error is examined in Sect. 3.3. The molecular reflectance function $R_\lambda^{1,\text{mlc}}$ is derived directly from Eq. (13), where $\tau_1^I$ is changed to $\tau_\lambda^{\text{mcl}} h^{\text{PBL}}$. Thus, the total reflectance function of layer II is given by:

$$ R_\lambda^I = R_\lambda^{1,\text{mlc}} + \frac{\omega_\lambda^{\text{aer}} P_\lambda^{\text{aer}}(\Theta)}{4(\mu_0 + \mu)} \left(1 - e^{-m\tau}\right) + R_\lambda^{\text{aer,MS}} . \quad (14) $$

The aerosol scattering phase function $P_\lambda^{\text{aer}}(\Theta)$ is defined by the approximate Henyey-Greenstein (HG) phase function (Henyey and Greenstein, 1941), which depends on the aerosol asymmetry factor $g_\lambda^{\text{aer}}$ and the scattering angle $\Theta$:

$$ P_\lambda^{\text{aer}}(\Theta) = \frac{1 - (g_\lambda^{\text{aer}})^2}{\left[1 + (g_\lambda^{\text{aer}})^2 - 2g_\lambda^{\text{aer}} \cos \Theta\right]^{2/3}} . \quad (15) $$

This HG phase function is plotted in Fig. 1 with $g_{\lambda}^{\text{aer}} = 0.63$ for a dry water soluble aerosol according to d’Almeida et al. (1991). The exact phase function derived from the Lorenz-Mie theory is superimposed to illustrate the imperfection of the HG approximation in the forward scattering domain for $\Theta > 150^\circ$. This influence on the accuracy of SMART is discussed in the second half of Sect. 3.2.

The second order (or secondary) scattering is calculated according to the Successive Orders of Scattering (SOS)
the method described by Hansen and Travis (1974):

\[ \frac{R_{\text{aer,MS}}(\mu, \mu_0, \phi - \phi_0)}{4\pi} = \frac{r_{\text{aer,MS}}}{4\pi} \]  

(16)

\[ \cdot \int_0^1 \int_0^1 \left[ \frac{1}{\mu} P_{t}^{\text{aer}}(\mu, \mu', \phi - \phi') R_{\text{SSA}}^T(\mu, \mu_0, \phi - \phi_0) \ight. \]

\[ \left. + \frac{1}{\mu_0} R_{\text{SSA}}(\mu, \mu', \phi - \phi') P_{t}^{\text{aer}}(\mu, \mu_0, \phi - \phi_0) \right] \cdot \frac{e^{-\frac{\mu r_{\text{aer}}}{\mu_0}}}{\mu} \cdot T_{\text{SSA}}(\mu, \mu_0, \phi - \phi_0) \cdot \left( e^{\frac{\mu r_{\text{aer}}}{\mu_0}} - e^{\frac{\mu r_{\text{aer}}}{\mu}} \right) d\mu' d\phi'. \]

However, for our accuracy requirements and under favourable remote sensing conditions, second order scatterings can be met by SMART. We compare SMART with other non-angular arguments are omitted for the sake of readability. \( P_{t}^{\text{aer}} \) and \( P_{t}^{\text{aer}} \) denote the aerosol HG phase function (Eq. (15)) using the scattering angle \( \Theta_t \) in case of reflectance (Eq. (11)) and the scattering angle \( \Theta_t = \arccos \left[ \mu_0 \mu + \cos(\phi - \phi_0) \sqrt{(1 - \mu_0^2)(1 - \mu^2)} \right] \) in case of transmittance. The single scattering transmittance \( T_{\text{SSA}} \) is given in van de Hulst (1948); Sobolev (1972); Hansen and Travis (1974); Kokhanovsky (2006):

\[ T_{\text{SSA}}^{\lambda} = \frac{\omega_{\text{aer}} P_{t}^{\text{aer}}(\Theta_t)}{4(\mu_0 - \mu)} \exp \left( -\frac{\mu r_{\text{aer}}}{\mu_0} \right) . \]

(17)

In case of \( \mu_0 = \mu \), we modify Eq. (18) to avoid indeterminacy with l'Hôpital's (Bernoulli's) rule:

\[ T_{\text{SSA}}^{\lambda} = \frac{\omega_{\text{aer}} P_{t}^{\text{aer}}(\Theta_t)}{4\mu^2} \exp \left( -\frac{\mu r_{\text{aer}}}{\mu} \right) . \]

(19)

We use a numerical approximation to calculate the integrals of Eq. (16). This is by far the most computationally intensive step in SMART. Therefore, we currently neglect scattering orders higher than two. A third order term could be added to Eq. (16) as given by Hansen and Travis (1974). However, for our accuracy requirements and under favourable remote sensing conditions, second order scattering is sufficient. More details are given in the first half of Sect. 3.2.

If fast computation is more important than accuracy, \( R_{\text{aer,MS}} \) can be substituted by \( \cdot \frac{f_{\text{corr}}}{\mu_0} (\lambda, \tau_{\text{aer}}^{\text{550nm}}) \) in analogy to Eq. (12). The expense is roughly 20% in decreased accuracy.

2.3 Radiative transfer at the surface

The modelling of optical processes at the surface can be elaborate due to adjacency and directional effects. Here we assume the simple case with isotropically reflected light on a homogeneous surface according to Lambert’s law (Ångström, 1925; Chandrasekhar, 1960; Sobolev, 1972):

\[ R_{\lambda}^{\text{SFC}} = \frac{a_{\lambda}}{1 - s_{\lambda} a_{\lambda}}. \]

(20)

where \( a_{\lambda} \) is the surface albedo and \( s_{\lambda} \) is the spherical albedo to account for multiple interaction between surface and atmosphere. We use the parameterization suggested by Kokhanovsky et al. (2005) for \( s_{\lambda} \), where:

\[ s_{\lambda} = \frac{T_{\lambda}^\Pi}{\mu^2} \left( a e^{-\frac{\mu}{4}} + b e^{-\frac{\mu}{2}} + c \right) . \]

(21)

The constants \( a, \alpha, b, \beta \) and \( c \) are parameterized according to Eq. (8). The corresponding expansion coefficients are given in Kokhanovsky et al. (2005). The resulting \( R_{\lambda}^{\text{SFC}} \) is also known as the hemispherical conical reflectance factor (HCRF) according to Schepman-Strub et al. (2006).

2.4 At-sensor reflectance function

Finally, we put the above equations together along the optical path to resolve the reflectance function \( R_{\lambda} \). Multiple retroreflections between layers I and II are neglected. A sensor at TOA or within levels I or II is simulated as follows:

\[ R_{\lambda}^{\text{S,TOA}} = R_{\lambda}^I + T_{\lambda}^{\Pi} \left[ R_{\lambda}^{\Pi} + R_{\lambda}^{\text{SFC}} T_{\lambda}^{\Pi} \right] T_{\lambda}^{\Pi}, \]

(22)

\[ R_{\lambda}^{\text{S,I}} = R_{\lambda}^I s_h^I + T_{\lambda}^{\Pi} \left[ R_{\lambda}^{\Pi} + R_{\lambda}^{\text{SFC}} T_{\lambda}^{\Pi} \right] \left( 1 - s_h^I + s_h^I T_{\lambda}^{\Pi} \right). \]

(23)

\[ R_{\lambda}^{\text{S,II}} = T_{\lambda}^{\Pi} \left[ R_{\lambda}^{\Pi} s_h^I + T_{\lambda}^{\Pi} R_{\lambda}^{\text{SFC}} \left( 1 - s_h^I + s_h^I T_{\lambda}^{\Pi} \right) \right]. \]

(24)

where \( T_{\lambda}^{\Pi} = T_{\lambda}^{\Pi} \), \( s_h^I = \frac{p_{\text{PBL}} - p_{\text{Sensor}}}{p_{\text{TOA}} - p_{\text{PBL}}} \) and \( s_h^I = \frac{p_{\text{SFC}} - p_{\text{Sensor}}}{p_{\text{SFC}} - p_{\text{PBL}}} \).

(25)

These scaling factors are used to account for the relative height of the sensor within the corresponding layer. \( s_h^I \) ranges from 1 at TOA to 0 at the PBL, while \( s_h^I \) varies from 1 at the PBL to 0 at the Earth’s surface (SFC).

3 Accuracy assessment

For typical airborne remote sensing conditions in the mid-latitudes we choose the representative uncertainty of imaging spectroscopy data of approximately 5% (Itten et al., 2008) as the accuracy requirement for SMART. Less typical conditions are analysed as well; in these cases we will accept larger errors. The definition of the conditions is given in Table 1. The AOD range was chosen according to the findings of Ruckstuhl et al. (2008), the wavelength range selected with regard to the optimal sensor performance (Seidel et al., 2008), while also avoiding strong water vapour absorption. We assume a black surface at the sea level (\( a_{\lambda} = 0 \)) to focus on the atmospheric part of SMART. Furthermore, we solely use the nadir viewing direction (\( \mu = 1 \)), which is approximated by small field-of-view sensors (FOV <30°).

This section evaluates if the prior accuracy requirements can be met by SMART. We compare SMART with
Table 1. Definition of the conditions and the related accuracy requirements for SMART. The limited conditions refer to typical airborne remote sensing needs in the mid-latitudes, which SMART was developed for. The analysed conditions refer to the accuracy assessment.

<table>
<thead>
<tr>
<th>remote sensing conditions</th>
<th>limited</th>
<th>analysed</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \omega_{\text{aer}}^{\text{550nm}} )</td>
<td>0–0.5</td>
<td>0–0.5</td>
</tr>
<tr>
<td>solar zenith angle, degrees</td>
<td>20–60</td>
<td>nadir–70</td>
</tr>
<tr>
<td>viewing zenith angle</td>
<td>nadir</td>
<td>nadir</td>
</tr>
<tr>
<td>wavelength, nm</td>
<td>500–700</td>
<td>400–800</td>
</tr>
<tr>
<td>surface albedo</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>accuracy requirement, %</td>
<td>5</td>
<td>15</td>
</tr>
</tbody>
</table>

In the following, the accuracy of SMART is investigated for specific approximation uncertainties, as well as for the overall accuracy. As an indicator of the accuracy, we calculate the relative difference or percent error of the reflectance function to the benchmark 6S:

\[
\delta R \cdot 100 = \frac{R_{\text{SMART}} - R_{6S}^{\text{S}}}{R_{6S}^{\text{S}}} \cdot 100. \tag{26}
\]

3.1 Rayleigh scattering approximation and polarisation

The total Rayleigh scattering is \( R_{\lambda}^{\text{nlc}} = R_{\lambda}^{\text{nlc}1} + R_{\lambda}^{\text{nlcII}} \) as given by Eqs. (13) and (14). The associated approximations include the Rayleigh scattering phase function (Eq. 10), the multiple scattering correction factor from MODTRAN (Eq. 12) and the neglected polarisation due to the scalar equations. The percent error is a distinct function of the wavelength and SZA, induced mainly by polarisation. Figure 2a shows that it grows towards shorter wavelengths and larger

![Graph](image)

Fig. 2. Percent error due to Rayleigh scattering and polarisation with respect to wavelength and solar zenith angle (SZA) at top-of-atmosphere.

SZA. It is known that the scalar approximation can introduce uncertainties of up to 10% in the blue spectral region (van de Hulst, 1980; Mishchenko et al., 1994). The SZA dependency of this uncertainty is shown in Fig. 2b. At 550 nm, the Rayleigh scattering uncertainty in the typical SZA range from 20–50° is below 3%.

3.2 Aerosol scattering approximation

The main approximations for the aerosol scattering are the double scattering (Eq. 16) and the HG phase function (Eq. 15). Initially, we use the exactly same phase function as in 6S in order to study the error induced only by the neglected higher orders of scattering. This phase function for dry water soluble aerosols was derived from the Lorenz-Mie scattering theory. Subsequently, we compare the combined
The percent error introduced by the double scattering approximation is plotted in Fig. 3. It is almost constant over the spectra due to the higher reflectance at shorter wavelengths (see Fig. 3a). It is obvious that the reflectance function $R_S$ is increasingly underestimated by SMART for larger AOD due to the neglected third and higher orders of aerosol scattering (see Fig. 3b). Figure 3c shows that larger SZA leads to an underestimation of the atmospheric reflectance for the same reason.

To study the accuracy of the total aerosol scattering $R_S^a$ as part of Eq. (14), we include the approximative HG phase function in SMART. 6S still uses the same Mie phase function as before. The input parameter for the HG phase function $g_{550}$ corresponds to the same dry water soluble aerosol, which is used in 6S. The exact Mie and the approximative HG phase function are shown in Fig. 1 for the same aerosol. The latter provides a reasonable approximation for scattering angles around 130°, which corresponds to a 50° SZA for nadir observations. The resulting combination of the aerosol double scattering error with the HG approximation error is examined in Fig. 4. It suggests that the use of the HG approximation does not introduce large percent errors within the range of typical SZA, as defined in Table 1. Given a range of 20–45° SZA, SMART is quite accurate at all investigated wavelengths and AOD values.

By comparing Figs. 3a with 4a and Figs. 3b with 4b, it can be seen that the HG approximation reverses some of the errors due to the aerosol double scattering approximation. The HG phase function for dry water soluble aerosols tends to overestimate of the aerosol scattering, which finally leads to a less distinct underestimation due to the neglected third and higher orders of aerosol scattering.

### 3.3 Coupling of Rayleigh and aerosol scattering

The current version of SMART does not yet account for the scattering interaction between molecules and aerosols. We analyse this effect by comparing 6S computations with the coupling switched on and off. The relative error related to this specific approximation is shown in Fig. 5. It remains within about 3%, reaching a maximum at large SZA (see Fig. 5c) and short wavelengths (see Fig. 5a). With errors of less than 2%, small SZAs are almost not influenced by the coupling and there is no distinct dependency on AOD noticeable (see Fig. 5b).

### 3.4 Overall accuracy

Previous Sects. 3.1–3.3 demonstrated that the approximations in SMART are adequate. Most of them are within the desired accuracy range of ±5% for the limited remote sensing conditions as defined in Table 1. Errors of up to ±15% are found for large SZA, however, they are mainly related to SMART’s simple two-layer atmospheric structure.

In the following, we examine the overall accuracy of SMART by comparing it according to Eq. (26) with independent computations of 6S. The computations of SMART are performed by Eq. (22) for a TOA sensor altitude at 80 km and by Eq. (23) for an airborne sensor altitude at 5500 m a.s.l. The percent error due to the excluded coupling between molecules and aerosols is inherent in the results of this subsection.

Figure 6 shows the result of two independent calculations using SMART (solid line) and 6S (dashed line) with respect to $\lambda$ and $\tau_{550}$. The qualitative agreement between the two models is evident. A quantitative perspective by statistical means of the overall accuracy is provided in Table 3, where

$$\text{RMSE} = \sqrt{\frac{1}{N} \sum (R_{\text{SMART}} - R_{6S})^2},$$

is the root mean square error and

$$\text{NRMSE} = \frac{\text{RMSE} \cdot 100}{\max (R_{\text{SMART}}) - \min (R_{\text{SMART}})},$$

is the normalised RMSE. The statistics are derived from all combinations of input parameters defined in Tables 1 and 2 within the limited conditions. The resulting correlation between SMART and 6S is almost perfect. The RMSE is approximately 0.16 reflectance values and the NRMSE is between 1.8% and 3.5%. The differences are smaller at TOA in comparison to those at 5500 m.
3.2 Aerosol scattering approximation

The main approximations for the aerosol scattering are the aerosol double scattering approximation. This phase function for dry water soluble aerosols tends to HG phase function for the same aerosol. The latter provides a reasonable approximation with 6S.

The percent error introduced by the double scattering approximation is shown in Fig. 3. It remains within the range of typical SZA, as defined in Table 1.

Percent error of the SMART reflectance function due to aerosol scattering with respect to wavelength, aerosol optical depth (AOD) and solar zenith angle (SZA) at top-of-atmosphere. SMART and 6S use the same phase function from Lorenz-Mie theory.

Fig. 3. Percent error of the SMART reflectance function due to aerosol scattering with respect to wavelength, aerosol optical depth (AOD) and solar zenith angle (SZA) at top-of-atmosphere. SMART and 6S use the same phase function from Lorenz-Mie theory.

Fig. 4. Percent error of the SMART reflectance function due to aerosol scattering with respect to wavelength, aerosol optical depth (AOD) and solar zenith angle (SZA) at top-of-atmosphere. SMART uses the HG phase function, 6S the phase function from Lorenz-Mie theory.

www.atmos-meas-tech.net/3/1129/2010/
The percent error due to the excluded coupling between backscattering and forward scattering is almost 0% for different scenarios.

In the following, we analyse the overall accuracy of Eq. (22) by Eq. (26) in more details with respect to wavelength, SZA and AOD. SMART computes very similar results compared to 6S at TOA with an SZA between 30° and 40°. This conclusion can be drawn from the combination of Figs. 2b, 4c and 5c, as well as from the total percent error in Fig. 7a. The overall percent error does not exceed ±5% at any investigated wavelength or AOD. At the large SZA of 60°, SMART overestimates $R_{\text{S,TOA}}$ by more than 10% at short wavelengths. Nevertheless, the overall accuracy is still well within the acceptable range of 10% at any wavelength larger than 450 nm (see Fig. 7b). At 550 nm, only the combination of very small SZA with a strong AOD or a high SZA with low AOD leads to a percent error just outside of the desired 5% margin (see Fig. 7c). In the blue part of the spectrum, high or low SZA lead to significant percent errors in a pure Rayleigh scattering atmosphere (see Fig. 7e). The same is true in an atmosphere containing aerosols, where the aerosols introduce additional percent errors in the red part of the spectrum for small SZA (see Fig. 7d and f).

Since SMART is also intended for the use with airborne remote sensing data, we additionally analyse the overall accuracy of Eq. (23) by Eq. (26). We place the sensor at 5500 m above the assumed black surface at sea level. The airborne scenario is more sensitive to the approximative two-layer setup in SMART. The 26 atmospheric layers in 6S can better account for the vertically inhomogeneous atmosphere. In fact, the percent error is slightly larger in the airborne case in comparison with the TOA case. The error distribution in the contour plots of Fig. 8a–f show that SMART underestimates the reflectance factors at 5500 m. Nevertheless, the hypothetical pure Rayleigh atmosphere still performs well,

![Figure 6](image-url)
Fig. 7. Overall accuracy with a sensor at top-of-atmosphere.

(a) $\delta R_{S,TOA}^{S,AER}(\lambda, \tau_{550 \text{ nm}}) \cdot 100$ at $30^\circ$ SZA

(b) $\delta R_{S,TOA}^{S,AER}(\lambda, \tau_{550 \text{ nm}}) \cdot 100$ at $60^\circ$ SZA

(c) $\delta R_{550 \text{ nm}}^{S,TOA}(SZA, \tau_{550 \text{ nm}}) \cdot 100$ at $\lambda = 550\text{nm}$

(d) $\delta R_{S,TOA}^{S,AER}(\lambda, SZA) \cdot 100$ at $\tau_{550 \text{ nm}} = 0.0$

(e) $\delta R_{S,TOA}^{S,AER}(\lambda, SZA) \cdot 100$ at $\tau_{550 \text{ nm}} = 0.2$

(f) $\delta R_{S,TOA}^{S,AER}(\lambda, SZA) \cdot 100$ at $\tau_{550 \text{ nm}} = 0.5$
Table 3. Quantitative comparison between SMART and 6S by statistical means for the limited conditions as defined in Table 1. SMART uses the HG phase function; 6S used the phase function from Mie calculations. \( R^2 \) denotes the squared correlation coefficient, RMSE the root mean square error and NRMSE the normalised RMSE.

<table>
<thead>
<tr>
<th>sensor altitude</th>
<th>( R^2 )</th>
<th>RMSE</th>
<th>NRMSE</th>
</tr>
</thead>
<tbody>
<tr>
<td>TOA</td>
<td>0.998</td>
<td>0.157</td>
<td>1.77%</td>
</tr>
<tr>
<td>5500 m</td>
<td>0.998</td>
<td>0.167</td>
<td>3.52%</td>
</tr>
</tbody>
</table>

with a maximum percent error of 6% (see Fig. 8a, b and c). The aerosols worsen the underestimation in the lower half of the visible spectrum, especially at very small and very large SZAs. At 550 nm and a 30° SZA, the percent error is 6% or less for an AOD up to 0.5. With the same constellation but an extreme SZA, the percent errors reach about 10% (see Fig. 8c, e and f). The largest offset between SMART and 6S is found at 60° SZA, 400 nm and an AOD of 0.5 with 18% relative difference. However, it should be noted that absolute difference \( R^S_{\text{SMART}} - R^S_{\text{6S}} \) is in fact smaller in the airborne case compared to the TOA case (not shown). Nonetheless, the relative error given by Eq. (26) is larger due to the smaller \( R^S_{\text{6S}} \) in the denominator.

4 Performance assessment

SMART is designed to optimally balance the opposing needs for accuracy and computational speed; the speed decreases with increasing model complexity and accuracy. We use the 6S vector version 1.1 (Vermote et al., 1997) as a benchmark RTM (same as in Sect. 3) to assess the performance of SMART. 6S is compiled with GNU Fortran and SMART is implemented in IDL. Both run on the same CPU infrastructure.

SMART needs only approximately 0.05 s for the calculation of one reflectance factor value. The more complex 6S needs about 1.4 s under identical conditions. Consequently, SMART computes more than 25 times faster. If \( R^S_{\text{aer,MS}} \) (Eq. 16) is substituted by a simple correction factor \( f^\text{corr}_a (\lambda, \tau) \) for aerosol multiple scattering (similar to Eq. 12), SMART runs 220 times faster than by numerically solving Eq. (16) in the presented configuration.

5 Summary and conclusions

We introduced SMART, as well as its approximative radiative transfer equations and parameterizations. Results of the atmospheric at-sensor reflectance function computed by SMART were compared with benchmark results from 6S for accuracy and performance. The overall percent error was examined and discussed, as were the individual errors resulting from Rayleigh scattering, aerosol scattering and molecule-aerosol interactions. The aerosol scattering was compared to 6S with and without the effect of the HG phase function approximation.

We found that SMART fulfils its design principle: it is fast and simple, yet accurate enough for a range of applications. One example may include the assessment of atmospheric effects when inspecting the quality of airborne or spaceborne data against ground truth measurements in near-real-time. The generation of atmospheric input parameters for vegetation canopy RTM inversion schemes, could be another application. SMART computes more than 20 reflectance results per second on a current customary desktop computer. This is more than 25 times faster than the benchmark RTM. The overall percent error under typical mid-latitude remote sensing conditions was found to be about 5% for the spaceborne and 5% to 10% for the airborne case. Large AOD or SZA values lead to larger percent errors of up to 15%. In general, the included approximations are sensitive to the strong scattering in the blue spectral region, which leads to larger percent errors. Together with the effect of polarisation, the total percent error of SMART exceeds the desired accuracy goal of 5% only in the blue region. It is therefore suggested that SMART be used preferably in the spectral range between roughly 500 nm and 680 nm, avoiding the blue and strong absorption bands. However, the neglected ozone absorption in this spectral interval leads to a small overestimation of up to 0.007 reflectance units at large SZA and 600 nm. It is also recommended to use SMART for computations with a sensor above the PBL to avoid uncertainties in the vertical distribution of the aerosols.

SMART can be improved by implementing other phase functions instead of the HG approximation, including those derived from Lorenz-Mie theory, geometrical optics (ray-tracing), and T-matrix approaches (Liou and Hansen, 1971; Mishchenko et al., 2002). Further refinements may include the coupling between molecules and aerosols, as well as the implementation of freely mixable aerosol components and hygroscopic growth (Hess et al., 1998). To account for polarisation, the scalar equations can be extended to the vector notation. Furthermore, a similar approach as used for the Rayleigh multiple scattering in this study (Eq. 12) may perhaps be used to perform a rough polarisation correction. Other issues for further developments may include additional atmospheric layers, gaseous absorption (foremost ozone), adjacency effects and the treatment of a directional, non-Lambertian surface.

A recent inter-comparison study for classic RTMs such as 6S, RT3, MODTRAN and SHARM, found discrepancies of \( \delta R \leq 5\% \) at TOA (Kotchenova et al., 2008). Even larger errors were found when polarisation was neglected or the HG phase function was used. SMART does not yet account for polarisation and uses the HG approximation by default, however...
Fig. 8. Overall accuracy with a sensor at 5500 m.

(a) \( \delta R_{S, 5500 \text{ m}}(\lambda, \tau_{aer}^{550 \text{ nm}}) \cdot 100 \) at 30° SZA

(b) \( \delta R_{S, 5500 \text{ m}}(\lambda, \tau_{aer}^{550 \text{ nm}}) \cdot 100 \) at 60° SZA

(c) \( \delta R_{S, 550 \text{ nm}}(\lambda, \tau_{aer}^{550 \text{ nm}}) \cdot 100 \) at \( \tau_{aer}^{550 \text{ nm}} = 0.2 \)

(d) \( \delta R_{S, 5500 \text{ m}}(\lambda, \tau_{aer}^{550 \text{ nm}}) \cdot 100 \) at \( \tau_{aer}^{550 \text{ nm}} = 0 \)

(e) \( \delta R_{S, 5500 \text{ m}}(\lambda, \tau_{aer}^{550 \text{ nm}}) \cdot 100 \) at \( \tau_{aer}^{550 \text{ nm}} = 0.5 \)

(f) \( \delta R_{S, 5500 \text{ m}}(\lambda, \tau_{aer}^{550 \text{ nm}}) \cdot 100 \) at \( \tau_{aer}^{550 \text{ nm}} = 0 \).
References


