Atmospheric / Topographic Correction for Airborne Imagery


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DLR-IB 565-02/15
The cover image shows Sequence of ATCOR/BREFCOR process for a mosaic of five image lines of CASI imagery. Upper left: original image, middle: elevation data (ranging from 500 to 1200 m), right: ATCOR standard correction using the given DEM, lower left: BCI image (ranging from -0.5 to 0.8), middle: ANIF factor (ranging from 0.9 to 1.1, approx), lower right: BREFCOR corrected image.

An improved BRDF correction algorithm (BREFCOR) has been introduced in the ATCOR-2015 release.
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Chapter 1

Introduction

The objective of any radiometric correction of airborne and spaceborne imagery of optical sensors is the extraction of physical earth surface parameters such as spectral albedo, directional reflectance quantities, emissivity, and temperature. To achieve this goal the influence of the atmosphere, solar illumination, sensor viewing geometry, and terrain information have to be taken into account. Although a lot of information from airborne and satellite imagery can be extracted without radiometric correction, the physically based approach offers advantages, especially when dealing with multitemporal data and when a comparison of different sensors is required. In addition, the full potential of imaging spectrometers can only be exploited with this approach.

Although physical models can be quite successful to eliminate atmospheric and topographic effects they inherently rely on an accurate spectral and radiometric sensor calibration and on the accuracy and appropriate spatial resolution of a digital elevation model (DEM) in rugged terrain. In addition, many surfaces have a bidirectional reflectance behavior, i.e., the reflectance depends on the illumination and viewing geometry. The usual assumption of an isotropic or Lambertian reflectance law is appropriate for small field-of-view (FOV < 30°, scan angle < ±15°) sensors if viewing does not take place in the solar principal plane. However, for large FOV sensors and / or data recording close to the principal plane the anisotropic reflectance behavior of natural surfaces causes brightness gradients in the image. These effects can be removed with an empirical method that normalizes the data to nadir reflectance values. In addition, for rugged terrain areas illuminated under low local solar elevation angles, these effects also play a role and can be taken care of with an empirical method included in the ATCOR package.

The ATCOR software was developed to cover about 80% of the typical cases with a reasonable amount of coding. It is difficult if not impossible to achieve satisfactory results for all possible cases. Special features of ATCOR are the consideration of topographic effects and the capability to process thermal band imagery.

There are two ATCOR models available, one for satellite imagery, the other one for airborne imagery ([72], [73]). An integral part of all ATCOR versions is a large database containing the results of radiative transfer calculations based on the MODTRAN®5 code (Berk et al. 1998, 2008). While ATCOR uses the AFRL MODTRAN® code to calculate the database of atmospheric look-up tables (LUT), the correctness of the LUTs is the responsibility of ATCOR.

Historical note: For historical reasons, the satellite codes are called ATCOR-2 (flat terrain, two geometric degrees-of-freedom DOF [60]) and ATCOR-3 (three DOF’s, mountainous terrain [63]). They support all operationally available small to medium FOV optical and thermal satellite sen-
sors with a sensor-specific atmospheric database. The scan angle dependence of the atmospheric correction functions within a scene is neglected here.

The airborne version is called ATCOR-4, to indicate the four geometric DOF’s $x$, $y$, $z$, and scan angle $\theta$. It includes the scan angle dependence of the atmospheric correction functions, a necessary feature, because most airborne sensors have a large FOV up to $60^\circ-90^\circ$. While satellite sensors always operate outside the atmosphere, airborne instruments can operate in altitudes of a few hundred meters up to 20 km. So the atmospheric database has to cover a range of altitudes. Since there is no standard set of airborne instruments and the spectral / radiometric performance might change from year to year due to sensor hardware modifications, a monochromatic atmospheric database was compiled based on the MODTRAN®5 radiative transfer code. This database has to be resampled for each user-defined sensor.

Organization of the manual:

Chapters 2 and 3 contain a short description of the basic concepts of atmospheric correction which will be useful for newcomers. Chapter 2 discusses the solar spectral region, while chapter 3 treats the thermal region. Chapter 4 presents the workflow in ATCOR, and chapter 5 contains a detailed description of all graphical user interface panels.

It is followed by chapters on batch processing, value added products available with ATCOR, sensor simulation, internal reference, and finally a comprehensive chapter on the theoretical background of atmospheric correction.

Information on the IDL version of ATCOR can be found on the internet: http://www.rese.ch. What is new in the 2016 version:

- The ATCOR GUI supports the input of a negative visibility. This means the visibility is not changed during processing (even if it causes negative surface reflectance pixels). This option was previously only accepted if the visibility parameter was set in the ’.inn’ file.

- An external aerosol optical thickness map (AOT550nm) is supported. IF the file name of the input scene is ‘scene_xxx.bsq’, then the external AOT map should be named ’scene_xxx_aot550.bsq’, and it supersedes the internal visibility/AOT calculation.

- The pre-classification map is improved.

- Optional scene-dependent scaling of path radiance in the blue-green spectral region for constant visibility.
Chapter 2

Basic Concepts in the Solar Region

Standard books on optical remote sensing contain an extensive presentation on sensors, spectral signatures, and atmospheric effects where the interested reader is referred to (Slater 1980 [90], Asrar 1989 [4], Schowengert 1997 [87]).

This chapter describes the basic concept of atmospheric correction. Only a few simple equations (2.1-2.25) are required to understand the key issues. We start with the radiation components and the relationship between the at-sensor radiance and the digital number or grey level of a pixel. Then we are already able to draw some important conclusions about the radiometric calibration. We continue with some remarks on how to select atmospheric parameters. Next is a short discussion about the thermal spectral region. The remaining sections present the topics of BRDF correction, spectral / radiometric calibration, and de-shadowing. For a discussion of the haze removal method the reader is referred to chapter 10.5.3.

Two often used parameters for the description of the atmosphere are 'visibility' and 'optical thickness'.

Visibility and optical thickness

The visibility (horizontal meteorological range) is approximately the maximum horizontal distance a human eye can recognize a dark object against a bright sky. The exact definition is given by the Koschmieder equation:

\[ VIS = \frac{1}{\beta} \ln \frac{1}{0.02} = \frac{3.912}{\beta} \]  \hspace{1cm} (2.1)

where \( \beta \) is the extinction coefficient (unit \( km^{-1} \)) at 550 nm. The term 0.02 in this equation is an arbitrarily defined contrast threshold. Another often used concept is the optical thickness of the atmosphere (\( \delta \)) which is the product of the extinction coefficient and the path length \( x \) (e.g., from sea level to space in a vertical path):

\[ \delta = \beta x \]  \hspace{1cm} (2.2)

The optical thickness is a pure number. In most cases, it is evaluated for the wavelength 550 nm. Generally, there is no unique relationship between the (horizontal) visibility and the (vertical) total optical thickness of the atmosphere. However, with the MODTRAN® radiative transfer code a certain relationship has been defined between these two quantities for clear sky conditions as shown in Fig. 2.1 (left) for a path from sea level to space. The optical thickness can be defined separately for the different atmospheric constituents (molecules, aerosols), so there is an optical thickness due
to molecular (Rayleigh) and aerosol scattering, and due to molecular absorption (e.g., water water, ozone etc.). The total optical thickness is the sum of the thicknesses of all individual contributors:

$$\delta = \delta(\text{molecular scattering}) + \delta(\text{aerosol}) + \delta(\text{molecular absorption})$$ (2.3)

The MODTRAN® visibility parameter scales the aerosol content in the boundary layer (0 - 2 km altitude). For visibilities greater than 100 km the total optical thickness asymptotically approaches a value of about 0.17 which (at 550 nm) is the sum of the molecular thickness ($\delta = 0.0973$) plus ozone thickness ($\delta = 0.03$) plus a very small amount due to trace gases, plus the contribution of residual aerosols in the higher atmosphere (2 - 100 km) with $\delta = 0.04$. The minimum optical thickness or maximum visibility is reached if the air does not contain aerosol particles (so called "Rayleigh limit") which corresponds to a visibility of 336 km at sea level and no aerosols in the boundary layer and higher atmosphere. In this case the total optical thickness (molecular and ozone) is about $\delta = 0.13$. Since the optical thickness due to molecular scattering (nitrogen and oxygen) only depends on pressure level it can be calculated accurately for a known ground elevation. The ozone contribution to the optical thickness usually is small at 550 nm and a climatologic/geographic average can be taken. This leaves the aerosol contribution as the most important component which varies strongly in space and time. Therefore, the aerosol optical thickness (AOT) at 550 nm is often used to characterize the atmosphere instead of the visibility.

The atmospheric (direct or beam) transmittance for a vertical path through the atmosphere can be calculated as:

$$\tau = e^{-\delta}$$ (2.4)

Fig. 2.1 (right) shows an example of the atmospheric transmittance from 0.4 to 2.5 $\mu$m. The spectral regions with relatively high transmittance are called "atmospheric window" regions. In absorbing regions the name of the molecule responsible for the attenuation of radiation is included.
**Apparent reflectance**

The apparent reflectance describes the combined earth/atmosphere behavior with respect to the reflected solar radiation:

\[
\rho(\text{apparent}) = \frac{\pi d^2 L}{E \cos \theta_s}
\]  

(2.5)

where \(d\) is the earth-sun distance in astronomical units, \(L = c_0 + c_1 \, DN\) is the at-sensor radiance, \(c_0, c_1, DN\), are the radiometric calibration offset, gain, and digital number, respectively. \(E\) and \(\theta_s\) are the extraterrestrial solar irradiance and solar zenith angle, respectively. For airborne imagery the use of the downwelling solar flux \(E_d\) at the aircraft altitude would be a more accurate description, but \(E_d\) is not available in the code. Therefore, the extraterrestrial \(E\) is employed which is a useful approximation. For high flight altitudes above 4 km the difference between \(E\) and \(E_d\) is small.

For imagery of satellite sensors the apparent reflectance is also named top-of-atmosphere (TOA) reflectance.

### 2.1 Radiation components

We start with a discussion of the radiation components in the solar region, i.e., the wavelength spectrum from 0.35 - 2.5 \(\mu\)m. Figure 2.2 shows a schematic sketch of the total radiation signal at the sensor. It consists of three components:

1. path radiance \((L_1)\), i.e., photons scattered into the sensor’s instantaneous field-of-view, without having ground contact.

2. reflected radiation \((L_2)\) from a certain pixel: the direct and diffuse solar radiation incident on the pixel is reflected from the surface. A certain fraction is transmitted to the sensor. The sum of direct and diffuse flux on the ground is called global flux.

3. reflected radiation from the neighborhood \((L_3)\), scattered by the air volume into the current instantaneous direction, the adjacency radiance. As detailed in [69] the adjacency radiation \(L_3\) consists of two components (atmospheric backscattering and volume scattering) which are combined into one component in Fig. 2.2 to obtain a compact description.

Only radiation component 2 contains information from the currently viewed pixel. The task of atmospheric correction is the calculation and removal of components 1 and 3, and the retrieval of the ground reflectance from component 2.

So the total radiance signal \(L\) can be written as:

\[
L = L_{\text{path}} + L_{\text{reflected}} + L_{\text{adj}} (= L_1 + L_2 + L_3)
\]  

(2.6)

The path radiance decreases with wavelength. It is usually very small for wavelengths greater than 800 nm. The adjacency radiation depends on the reflectance or brightness difference between the currently considered pixel and the large-scale (0.5-1 km) neighborhood. The influence of the adjacency effect also decreases with wavelength and is very small for spectral bands beyond 1.5 \(\mu\)m [69].

For each spectral band of a sensor a linear equation describes the relationship between the recorded brightness or digital number \(DN\) and the at-sensor radiance (Fig. 2.2):

\[
L = c_0 + c_1 \, DN
\]  

(2.7)
CHAPTER 2. BASIC CONCEPTS IN THE SOLAR REGION

Figure 2.2: Schematic sketch of solar radiation components in flat terrain. $L_1$ : path radiance, $L_2$ : reflected radiance, $L_3$ : adjacency radiation.

The $c_0$ and $c_1$ are called radiometric calibration coefficients. The radiance unit in ATCOR is $mWcm^{-2}sr^{-1}µm^{-1}$. For instruments with an adjustable gain setting $g$ the corresponding equation is:

$$L = c_0 + \frac{c_1}{g} * DN$$

During the following discussion we will always use eq. (2.7). Disregarding the adjacency component we can simplify eq. (2.6)

$$L = L_{path} + L_{reflected} = L_{path} + \tau \rho E_g/\pi = c_0 + c_1 DN$$

where $\tau$, $\rho$, and $E_g$ are the ground-to-sensor atmospheric transmittance, surface reflectance, and global flux on the ground, respectively. Solving for the surface reflectance we obtain:

$$\rho = \frac{\pi \{d^2(c_0 + c_1 DN) - L_{path}\}}{\tau E_g}$$

The factor $d^2$ takes into account the sun-to-earth distance (d is in astronomical units), because the LUT’s for path radiance and global flux are calculated for d=1 in ATCOR. Equation (2.9) is a key formula to atmospheric correction. A number of important conclusions can now be drawn:

- An accurate radiometric calibration is required, i.e., a knowledge of $c_0$, $c_1$ in each spectral band.

- An accurate estimate of the main atmospheric parameters (aerosol type, visibility or optical thickness, and water vapor) is necessary, because these influence the values of path radiance, transmittance, and global flux.

- If the visibility is assumed too low (optical thickness too high) the path radiance becomes high, and this may cause a physically unreasonable negative surface reflectance. Therefore, dark surfaces of low reflectance, and correspondingly low radiance $c_0 + c_1 DN$, are especially sensitive in this respect. They can be used to estimate the visibility or at least a lower
CHAPTER 2. BASIC CONCEPTS IN THE SOLAR REGION

bound. If the reflectance of dark areas is known the visibility can actually be calculated. The interested reader may move to chapter 10.4.2, but this is not necessary to understand the remaining part of the chapter.

- If the main atmospheric parameters (aerosol type or scattering behavior, visibility or optical thickness, and water vapor column) and the reflectance of two reference surfaces are measured, the quantities \( L_{\text{path}} \), \( \tau \), \( \rho \), and \( E_g \) are known. So, an "inflight calibration" can be performed to determine or update the knowledge of the two unknown calibration coefficients \( c_0(k) \), \( c_1(k) \) for each spectral band \( k \), see section 2.4.

Selection of atmospheric parameters

The optical properties of some air constituents are accurately known, e.g., the molecular or Rayleigh scattering caused by nitrogen and oxygen molecules. Since the mixing ratio of nitrogen and oxygen is constant the contribution can be calculated as soon as the pressure level (or ground elevation) is specified. Other constituents vary slowly in time, e.g., the \( \text{CO}_2 \) concentration. ATCOR calculations were performed for a concentration of 400 ppmv. Ozone may also vary in space and time. Since ozone usually has only a small influence ATCOR employs a fixed value of 331 DU (Dobson units, corresponding to the former unit 0.331 atm-cm, for a ground at sea level) representing average conditions. The three most important atmospheric parameters that vary in space and time are the aerosol type, the visibility or optical thickness, and the water vapor. We will mainly work with the term visibility (or meteorological range), because the radiative transfer calculations were performed with the MODTRAN®5 code (Berk et al., 1998, 2008), and visibility is an intuitive input parameter in MODTRAN®, although the aerosol optical thickness can be used as well. ATCOR employs a database of LUTs calculated with MODTRAN®5.

Aerosol type

The aerosol type includes the absorption and scattering properties of the particles, and the wavelength dependence of the optical properties. ATCOR supports four basic aerosol types: rural, urban, maritime, and desert. The aerosol type can be calculated from the image data provided that the scene contains vegetated areas. Alternatively, the user can make a decision, usually based on the geographic location. As an example, in areas close to the sea the maritime aerosol would be a logical choice if the wind was coming from the sea. If the wind direction was toward the sea and the air mass is of continental origin the rural, urban, or desert aerosol would make sense, depending on the geographical location. If in doubt, the rural (continental) aerosol is generally a good choice.

The aerosol type also determines the wavelength behavior of the path radiance. Of course, nature can produce any transitions or mixtures of these basic four types. However, ATCOR is able to adapt the wavelength course of the path radiance to the current situation provided spectral bands exist in the blue-to-red-region and the scene contains reference areas of known reflectance behavior. The interested reader may read chapter 10.4.2 for details.

Visibility estimation

Two options are available in ATCOR:

- An interactive estimation in the SPECTRA module (compare chapter 5). The spectra of different targets in the scene can be displayed as a function of visibility. A comparison with reference spectra from libraries determines the visibility. In addition, dark targets like vegetation in the blue-to-red spectrum or water in the red-to-NIR can be used to estimate the visibility.
• An automatic calculation of the visibility can be performed if the scene contains dark reference
pixels. The interested reader is referred to chapter 10.4.2 for details.

Water vapor column

The water vapor content can be automatically computed if the sensor has spectral bands in water
vapor regions (e.g., 920-960 nm). The approach is based on the differential absorption method
and employs bands in absorption regions and window regions to measure the absorption depth,
see chapter 10.4.3. Otherwise, if a sensor does not possess spectral bands in water vapor regions,
e.g. Landsat TM or SPOT, an estimate of the water vapor column based on the season (summer
/ winter) is usually sufficient. Typical ranges of water vapor columns are (sea-level-to space):
tropical conditions: $wv=3-5$ cm (or g cm$^{-2}$)
midlatitude summer: $wv= 2-3$ cm
dry summer, spring, fall: $wv=1-1.5$ cm
dry desert or winter: $wv=0.3-0.8$ cm

2.2 Spectral calibration

This section can be skipped if data processing is only performed for imagery of broad-band sensors.
Sensor calibration problems may pertain to spectral properties, i.e., the channel center positions
and / or bandwidths might have changed compared to laboratory measurements, or the radiometric
properties, i.e., the offset ($c_0$) and slope ($c_1$) coefficients, relating the digital number (DN) to the
at-sensor radiance $L = c_0 + c_1 \times DN$. Any spectral mis-calibration can usually only be detected
from narrow-band hyperspectral imagery as discussed in this section. For multispectral imagery,
spectral calibration problems are difficult or impossible to detect, and an update is generally only
performed with respect to the radiometric calibration coefficients, see chapter 2.4.

Surface reflectance spectra retrieved from narrow-band hyperspectral imagery often contain spikes
and dips in spectral absorption regions of atmospheric gases (e.g., oxygen absorption around 760
nm, water vapor absorption around 940 nm). These effects are most likely caused by a spectral
mis-calibration. In this case, an appropriate shift of the center wavelengths of the channels will
remove the spikes. This is performed by an optimization procedure that minimizes the deviation
between the surface reflectance spectrum and the corresponding smoothed spectrum. The merit
function to be minimized is

$$\chi^2(\delta) = \sum_{i=1}^{n} \left( \rho_{i}^{surf}(\delta) - \rho_{i}^{smooth} \right)^2$$  \hspace{1cm} (2.11)

where $\rho_{i}^{surf}(\delta)$ is the surface reflectance in channel $i$ calculated for a spectral shift $\delta$, $\rho_{i}^{smooth}$ is the
smoothed (low pass filtered) reflectance, and $n$ is the number of bands in each spectrometer of a
hyperspectral instrument. So the spectral shift is calculated independently for each spectrometer.
In the currently implemented version, the channel bandwidth is not changed and the laboratory
values are assumed valid. More details of the method are described in [30]. A spectral re-calibration
should precede any re-calibration of the radiometric calibration coefficients; see section 5.8.4 for
details about this routine.

Figure 2.3 shows a comparison of the results of the spectral re-calibration for a soil and a vegetation
target retrieved from an AVIRIS scene (16 Sept. 2000, Los Angeles area). The flight altitude was
20 km above sea level (asl), heading west, ground elevation 0.1 km asl, the solar zenith and azimuth angles were 41.2° and 135.8°. Only part of the spectrum is shown for a better visual comparison of the results based on the original spectral calibration (thin line) and the new calibration (thick line). The spectral shift values calculated for the 4 individual spectrometers of AVIRIS are 0.1, -1.11, -0.88, and -0.21 nm, respectively.

**Figure 2.3:** Wavelength shifts for an AVIRIS scene.

### 2.3 Wavelength and refractive index

As the wavelength of electromagnetic radiation depends on the refractive index of the medium, this effect has to be calculated for airborne sensors if a high accuracy is needed, especially for hyperspectral instruments. The spectral channel filter functions are usually measured in the laboratory. So the measured wavelength depends on the refractive index $n_{\text{lab}}$ or pressure $p_{\text{lab}}$ at the elevation $h_{\text{lab}}$, during lab measurement. If $\lambda_0$ denotes the wavelength in vacuum, i.e. $n_{\text{vac}} = 1$, the sensor wavelength during a lab measurement is:

$$\lambda_{\text{sen}}(p_{\text{lab}}, h_{\text{lab}}) = \frac{\lambda_0}{n_{\text{lab}}} \quad (2.12)$$

We assume a typical scale height $H = 8$ km for the height dependence of pressure and air density, i.e.

$$p(h) = p_0 \exp\left(-\frac{h}{H}\right) \quad (2.13)$$

For a standard atmosphere (mid-latitude summer) we have $p_0 = 1013$ mbar (hPa).

For a spaceborne sensor the lab measurement is performed in a vacuum chamber, therefore $n_{\text{lab}}$ is close to 1 and $\lambda_{\text{sen}} = \lambda_0$. The MODTRAN radiative transfer calculations for the ATCOR look-up tables (LUTs) are performed on the basis of wavenumber $w$ ($cm^{-1}$) which is converted into wavelength $\lambda$ ($\mu m$) using

$$\lambda = \frac{10000}{w \ n(h)} \quad (2.14)$$
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For a spaceborne sensor we have \( n = 1 \), but for airborne sensors we have to account for the refractive index \( n(h) \) in two respects:

- The MODTRAN wavenumber has to be converted into a wavelength \( \lambda \) using eq. (2.14) taking care of the refractive index for the corresponding flight altitude \( h \) (or pressure level \( p \)). Eq. (2.13) is used to convert the flight altitude into the corresponding pressure. Switching to wavelength is required, because the high-resolution spectral database of atmospheric LUTs has to be convolved with the channel filter functions delivered as wavelength data.

- The lab measured wavelength of the channel filter functions (spectral response files) also has to be adapted to the refractive index at the flight altitude.

We use the equation:

\[
\phi(h) = 1 + 0.000293 \exp\left(-\frac{h}{H}\right)
\]  

(2.15)

Therefore, the MODTRAN wavelength conversion is

\[
\lambda_{MOD} = \frac{10000}{w n(h)} = \frac{\lambda_0}{n(h)}
\]  

(2.16)

The lab wavelength conversion for pressure \( p_{lab} \) and height \( h_{lab} \) is

\[
\lambda_{sen}(h_{lab}) = \frac{\lambda_0}{n(h_{lab})}
\]  

(2.17)

Eq. (2.13) is used to calculate the pressure \( p \) for a given flight altitude \( h \) and vice versa. Using the parameter \( h \) to indicate the pressure-dependence of the refractive index \( n(h) \) we get

\[
\lambda_{sen}(h) = \frac{\lambda_0}{n(h)} = \frac{\lambda_{sen}(h_{lab}) n(h_{lab})}{n(h)}
\]  

(2.18)

The wavelength change or shift is calculated as :

\[
\Delta_{MOD} = \lambda_{MOD}(h) - \lambda_0
\]  

(2.19)

\[
\Delta_{sen} = \lambda_{sen}(h) - \lambda_{sen}(h_{lab})
\]  

(2.20)

Figure 2.4 (top) shows the calculated wavelength shifts for MODTRAN required for 3 flight altitudes (1, 4.2, 100 km). The 4.2 km corresponds to a pressure level of 600 (hPa, mbar). Note: If the sensor is contained in a pressurized chamber at \( p=600 \) hPa, this pressure level has to be used for the calculation of the sensor wavelength shift independent of the actual flight altitude. This means the MODTRAN wavelength shift also has to be adapted to this pressure level, i.e. using the corresponding virtual flight altitude of 4.2 km.

Figure 2.4 (middle, left and right) show the lab wavelength shifts for the 2 cases of \( p_{lab} = 1013 \) hPa and \( p_{lab} = 940 \) hPa to study the influence of lab measurements at sea level (1013 hPa) and at a higher elevation (598 m above sea level).

The new MODTRAN wavelengths have a negative shift, indicating they are smaller than the original \( \lambda_0 \) (shifted left to shorter wavelengths), whereas the lab wavelengths have a positive shift, i.e. they are shifted to the right to longer wavelengths. Therefore, the combined shift (MODTRAN and lab) is not the sum but the difference of these shifts.
The combined total shift is shown in the bottom two plots: the left one represents the case with $p_{lab} = 1013$ hPa, the right one with $p_{lab} = 940$ hPa. Both cases are very similar with a slightly higher shift for the 1013 hPa case.

The total shift plots show that some compensation effects exist, e.g. for $h=100$ km the MODTRAN wavelength shift is 0 and the lab shift is largest. The opposite trend is observed for $h=1$ km, where the MODTRAN shift is largest and the lab wavelength shift is small. Therefore, the three altitude cases coincide on one line. The total shift increases with wavelength and is largest in the thermal spectral region.

There is a slight dependence of the results on the assumed scale height $H=8$ km and the sea level pressure $p_0=1013$ hPa. The scale height actually depends on the temperature and humidity profile, the average mass of atmospheric particles, and location (because of the acceleration of gravity). It can be approximately calculated with the equation of hydrostatic equilibrium using the ideal gas law, see any textbook on atmospheric physics.

As ATCOR is used by customers all over the world and the specific atmospheric state is usually not known, a typical standard scale height of $H=8$ km is assumed in ATCOR. For typical summer and winter conditions, the scale height varies between 8.0 and 8.5 km. The wavelength difference due to the air refractive index for $H=8.0$ km versus $H=8.5$ km is smaller than 0.06 nm in the wavelength region 0.4 to 10 $\mu$m. Therefore, the use of $H=8.0$ km is sufficient for practical purposes.

Some examples:

- The AVIRIS NG (Next Generation) spectrometer is operated under near vacuum (10 Torr, 13.3 mbar) conditions.
- The APEX spectrometer has a pressure regulation unit keeping the optical subunit at 200 mbar above ambient flight altitude pressure [35].
- Most airborne spectrometers operate under ambient flight altitude pressure.

The next table shows the default contents of file "pressure.dat". The file is created for each sensor in the sensor-specific folder during the first run of the RESLUT (resampling) module if no "pressure.dat" exists. The user should edit the first line of the file if necessary.

<table>
<thead>
<tr>
<th>1013.0</th>
<th>R0.0</th>
<th>lab pressure, instrument pressure (mbar, hPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>instrument pressure is relative or absolute</td>
</tr>
<tr>
<td></td>
<td></td>
<td>$R=r=$relative pressure above ambient flight altitude</td>
</tr>
<tr>
<td></td>
<td></td>
<td>$A=a=$absolute pressure</td>
</tr>
</tbody>
</table>

*Table 2.1: Default file "pressure.dat" to be edited if necessary.*

## 2.4 Inflight radiometric calibration

Inflight radiometric calibration experiments are performed to check the validity of the laboratory calibration. For spaceborne instruments processes like aging of optical components or outgassing during the initial few weeks or months after launch often necessitate an updated calibration. This approach is also employed for airborne sensors because the aircraft environment is different from the...
laboratory and this may have an impact on the sensor performance. The following presentation only
discusses the radiometric calibration and assumes that the spectral calibration does not change,
i.e., the center wavelength and spectral response curve of each channel are valid as obtained in the
laboratory, or it was already updated as discussed in chapter 2.2.

The radiometric calibration uses measured atmospheric parameters (visibility or optical thickness
from sun photometer, water vapor content from sun photometer or radiosonde) and ground re-

fectance measurements to calculate the calibration coefficients \( c_0 \), \( c_1 \) of equation (2.7) for each
band. For details, the interested reader is referred to the literature (Slater et al., 1987, Santer et
al. 1992, Richter 1997). Depending of the number of ground targets we distinguish three cases: a
single target, two targets, and more than two targets.

**Calibration with a single target**
In the simplest case, when the offset is zero (\( c_0 = 0 \)), a single target is sufficient to determine the
calibration coefficient \( c_1 \):

\[
L_1 = c_1 D N_1^* = L_{\text{path}} + \tau \rho_1 E_g / \pi
\]

\( L_{\text{path}} \), \( \tau \), and \( E_g \) are taken from the appropriate LUT’s of the atmospheric database, \( \rho_1 \) is the
measured ground reflectance of target 1, and the channel or band index is omitted for brevity.

\( D N_1^* \) is the digital number of the target, averaged over the target area and already corrected for
the adjacency effect. Solving for \( c_1 \) yields:

\[
c_1 = \frac{L_1 - L_{\text{path}}}{D N_1^*} = \frac{L_{\text{path}} + \tau \rho_1 E_g / \pi}{D N_1^*}
\]

Remark: a bright target should be used here, because for a dark target any error in the ground
reflectance data will have a large impact on the accuracy of \( c_1 \).

**Calibration with two targets**
In case of two targets a bright and a dark one should be selected to get a reliable calibration. Using
the indices 1 and 2 for the two targets we have to solve the equations:

\[
L_1 = c_0 + c_1 * D N_1^* \quad L_2 = c_0 + c_1 * D N_2^*
\]

This can be performed with the \( c_0 & c_1 \) option of ATCOR’s calibration module, see chapter 5. The
result is:

\[
c_1 = \frac{L_1 - L_2}{D N_1^* - D N_2^*} \quad c_0 = \frac{L_1 - c_1 * D N_1^*}{L_2 - c_1 * D N_2^*}
\]

Equation (2.24) shows that \( D N_1^* \) must be different from \( D N_2^* \) to get a valid solution, i.e., the two
targets must have different surface reflectances in each band. If the denominator of eq. (2.24) is
zero ATCOR will put in a 1 and continue. In that case the calibration is not valid for this band.
The requirement of a dark and a bright target in all channels cannot always be met.

**Calibration with \( n > 2 \) targets**
In cases where \( n > 2 \) targets are available the calibration coefficients can be calculated with a least
squares fit applied to a linear regression equation, see figure 2.5. This is done by the "cal_regress"
program of ATCOR. It employs the ",rdn" files obtained during the single-target calibration (the
"c1 option" of ATCOR’s calibration module. See section 5.8.5 for details about how to use this routine.

Note: If several calibration targets are employed, care should be taken to select targets without spectral intersections, since calibration values at intersection bands are not reliable. If intersections of spectra cannot be avoided, a larger number of spectra should be used, if possible, to increase the reliability of the calibration.

2.5 De-shadowing

Remotely sensed optical imagery of the Earth’s surface is often contaminated with cloud and cloud shadow areas. Surface information under cloud covered regions cannot be retrieved with optical sensors, because the signal contains no radiation component being reflected from the ground. In shadow areas, however, the ground-reflected solar radiance is always a small non-zero signal, because the total radiation signal at the sensor contains a direct (beam) and a diffuse (reflected skylight) component. Even if the direct solar beam is completely blocked in shadow regions, the reflected diffuse flux will remain, see Fig. 2.6. Therefore, an estimate of the fraction of direct solar irradiance for a fully or partially shadowed pixel can be the basis of a compensation process called de-shadowing or shadow removal. The method can be applied to shadow areas cast by clouds or buildings.

Figure 2.7 shows an example of removing cloud shadows from HyMap imagery. It is a sub-scene of a Munich flight line acquired 25 May 2007, with a flight altitude of 2 km above ground level. Occasional clouds appeared at altitudes higher than the aircraft cruising altitude. After shadow removal many details can be seen that are hidden in the uncorrected scene. The bottom part shows the shadow map, scaled between 0 and 1000. The darker the area the lower the fractional direct solar illumination, i.e. the higher the amount of shadow.

The proposed de-shadowing technique works for multispectral and hyperspectral imagery over land acquired by satellite / airborne sensors. The method requires a channel in the visible and at least one spectral band in the near-infrared (0.8-1 $\mu$m) region, but performs much better if bands in the short-wave infrared region (around 1.6 and 2.2 $\mu$m) are available as well. A fully automatic shadow removal algorithm has been implemented. However, the method involves some scene-dependent thresholds that might be optimized during an interactive session. In addition, if shadow areas are concentrated in a certain part of the scene, say in the lower right quarter, the performance of the algorithm improves by working on the subset only.

The de-shadowing method employs masks for cloud and water. These areas are identified with spectral criteria and thresholds. Default values are included in a file in the ATCOR path, called "preferences/preference_parameters.dat". As an example, it includes a threshold for the reflectance of water in the NIR region, $\rho=5\%$. So, a reduction of this threshold will reduce the number of pixels in the water mask. A difficult problem is the distinction of water and shadow areas. If water bodies are erroneously included in the shadow mask, the resulting surface reflectance values will be too high.

Details about the processing panels can be found in section 5.4.9.

2.6 BRDF correction

The reflectance of many surface covers depends on the viewing and solar illumination geometry. This behavior is described by the bidirectional reflectance distribution function (BRDF). It can
clearly be observed in scenes where the view and / or sun angles vary over a large angular range. Most across-track brightness gradients that appear after atmospheric correction are caused by BRDF effects, because the sensor’s view angle varies over a large range. In extreme cases when scanning in the solar principal plane, the brightness is particularly high in the hot spot angular region where retroreflection occurs, see Figure 2.8, left image, left part. The opposite scan angles (with respect to the central nadir region) show lower brightness values.

A simple method, called nadir normalization or across-track illumination correction, calculates the brightness as a function of scan angle, and multiplies each pixel with the reciprocal function (compare Section 10.6.1).

The BRDF effect can be especially strong in rugged terrain with slopes facing the sun and others oriented away from the sun. In areas with steep slopes the local solar zenith angle $\beta$ may vary from $0^\circ$ to $90^\circ$, representing geometries with maximum solar irradiance to zero direct irradiance, i.e., shadow. The angle $\beta$ is the angle between the surface normal of a DEM pixel and the solar zenith angle of the scene. In mountainous terrain there is no simple method to eliminate BRDF effects. The usual assumption of an isotropic (Lambertian) reflectance behavior often causes an overcorrection of faintly illuminated areas where local solar zenith angles $\beta$ range from $60^\circ$- $90^\circ$. These areas appear very bright, see Figure 2.9, left part.

To avoid a misclassification of these bright areas the reflectance values have to be reduced (Fig. 2.9, center part). In ATCOR empirical geometry-dependent functions are used for this purpose. In the simplest cases, the empirical BRDF correction employs only the local solar zenith angle $\beta$ and a threshold $\beta_T$ to reduce the overcorrected surface reflectance $\rho_L$ with a factor, depending on the incidence angle. For details the interested reader is referred to section 10.6.2.

A more sophisticated method available in ATCOR is the BRDF effects correction (BREFCOR) method. It uses both the surface cover type characterization and the per-pixel observation angle to find an appropriate anisotropy factor for correction. The method follows a novel scheme based on a fuzzy surface characterization and uses semi-empirical BRDF models for the correction. The process follows the below steps:

1. perform a fuzzy BRDF-Cover-Index (BCI) image characterization
2. calibrate the BRDF-model using a number of scenes of the same area and time of the year
3. calculate the anisotropy index for each spectral band using the calibrated model and the BCI
4. correct the image using the anisotropy index

Further details about this methods can be found in section 10.6.3.
Figure 2.4: MODTRAN and lab wavelength shifts (see discussion in the text).
CHAPTER 2. BASIC CONCEPTS IN THE SOLAR REGION

Figure 2.5: Radiometric calibration with multiple targets using linear regression.

Figure 2.6: Sketch of a cloud shadow geometry.
Figure 2.7: De-shadowing of a HyMap sub-scene of Munich. Color coding: RGB = channels 860/646/543 nm. Top left: original, right: de-shadowed image, bottom: shadow map.

Figure 2.8: Nadir normalization of an image with hot-spot geometry. Left: reflectance image without BRDF correction. Right: after empirical BRDF correction.
Figure 2.9: BRDF correction in rugged terrain imagery. Left: image without BRDF correction. Center: after BRDF correction with threshold angle $\beta_T = 65^\circ$. Right: illumination map $= \cos\beta$.

Figure 2.10: Effect of BRDF correction in an image mosaic (ADS image, ©swisstopo)
Chapter 3

Basic Concepts in the Thermal Region

Fig. 3.1 (left) presents an overview of the atmospheric transmittance in the 2.5 - 14 \( \mu m \) region. The main absorbers are water vapor and \( CO_2 \) which totally absorb in some parts of the spectrum. In the thermal region (8 - 14 \( \mu m \)) the atmospheric transmittance is mainly influenced by the water vapor column, ozone (around 9.6 \( \mu m \)) and \( CO_2 \) (at 14 \( \mu m \)). Fig. 3.1 (right) shows the transmittance for three levels of water vapor columns \( w=0.4, 1.0, 2.9 \) cm, representing dry, medium, and humid conditions. The aerosol influence still exists, but is strongly reduced compared to the solar spectral region because of the much longer wavelength. So an accurate estimate of the water vapor column is required in this part of the spectrum to be able to retrieve the surface properties, i.e., spectral emissivity and surface temperature.

![Figure 3.1: Atmospheric transmittance in the thermal region.](image)

Similar to the solar region, there are three radiation components: thermal path radiance \( (L_1) \), i.e., photons emitted by the atmospheric layers, emitted surface radiance \( (L_2) \), and reflected radiance \( (L_3) \).

In the thermal spectral region from 8 - 14 \( \mu m \) the radiance signal can be written as

\[
L = L_{path} + \tau \epsilon L_{BB}(T) + \tau(1 - \epsilon)F/\pi
\]  

(3.1)
where $L_{\text{path}}$ is the thermal path radiance, i.e., emitted and scattered radiance of different layers of the air volume between ground and sensor, $\tau$ is the atmospheric ground-to-sensor transmittance, $\epsilon$ is the surface emissivity ranging between 0 and 1, $L_{BB}(T)$ is Planck’s blackbody radiance of a surface at temperature $T$, and $F$ is the thermal downwelling flux of the atmosphere, see Fig. 3.2.

So the total signal consists of path radiance, emitted surface radiance, and reflected atmospheric radiation. The adjacency radiation, i.e., scattered radiation from the neighborhood of a pixel, can be neglected because the scattering efficiency decreases strongly with wavelength.

For most natural surfaces the emissivity in the 8-12 $\mu$m spectral region ranges between 0.95 and 0.99. Therefore, the reflected downwelling atmospheric flux contributes only a small fraction to the signal. Neglecting this component for the simplified discussion of this chapter we can write

$$L_{BB}(T) = \frac{L - L_{\text{path}}}{\tau \epsilon} = \frac{c_0 + c_1DN - L_{\text{path}}}{\tau \epsilon}$$  \hspace{1cm} (3.2)

In the thermal region the aerosol type plays a negligible role because of the long wavelength, and atmospheric water vapor is the dominating parameter. So the water vapor, and to a smaller degree the visibility, determine the values of $L_{\text{path}}$ and $\tau$. In case of coregistered bands in the solar and thermal spectrum the water vapor and visibility calculation may be performed with the solar channels. In addition, if the surface emissivity is known, the temperature $T$ can be computed from eq. (3.2) using Planck’s law.

For simplicity a constant emissivity $\epsilon = 1.0$ or $\epsilon = 0.98$ is often used and the corresponding temperature is called brightness temperature. The kinetic surface temperature differs from the brightness temperature if the surface emissivity does not match the assumed emissivity. With the assumption $\epsilon = 1.0$ the kinetic temperature is always higher than the brightness temperature. As a rule of thumb an emissivity error of 0.01 (one per cent) yields a surface temperature error of 0.5K.

For rugged terrain imagery no slope/aspect correction is performed for thermal bands, only the elevation-dependence of the atmospheric parameters is taken into account.
CHAPTER 3. BASIC CONCEPTS IN THE THERMAL REGION

3.1 Thermal spectral calibration

The spectral calibration in the thermal region using atmospheric absorption features can be conducted in a similar way as for the solar region. A spectral mis-calibration will cause spikes and dips in the surface emissivity spectrum. An appropriate shift of the center wavelengths of the channels will remove these artifacts. This is performed by an optimization procedure that minimizes the deviation between the surface emissivity spectrum and the corresponding smoothed spectrum. However, in the thermal region one also has to account for the unknown surface temperature. Therefore, the merit function also has to be evaluated for a range of surface temperatures \( T_k \) and the calculated emissivity depends on the assumed temperature:

\[
\varepsilon(i, T_k) = \frac{L(i) - L_p(i) - F(i)/(\pi)}{L_{bb}(i, T_k) \cdot \tau(i) - F(i)/(\pi)}
\]  (3.3)

Here the index \( i \) indicates the channel, \( L \) is the measured at-sensor radiance, \( L_p \) the path radiance, \( L_{bb} \) the blackbody radiance, and \( F \) the downwelling thermal flux multiplied with the ground-to-sensor transmittance \( \tau(i) \). The merit function to be minimized as a function of the wavelength shift \( \delta \) is:

\[
\chi^2(\delta) = \sum_{k=1}^{m} \sum_{i=1}^{n} (\varepsilon(i, T_k, \delta) - \bar{\varepsilon}(i, T_k, \delta))^2 \rightarrow Min
\]  (3.4)

The moving average of the emissivity is performed over 5 channels. In the present version, only channels in the 8.5 - 13.5 \( \mu m \) region are taken into account to avoid strong atmospheric absorption regions. Since MODTRAN look-up tables are used, the resulting wavelength shift also depends on the accuracy of these LUTs. The default temperature range is 280 - 310 K, but the user can specify it with the keyword \( \text{trange} \), e.g. \( \text{trange}=[270,320] \). The temperature increment is fixed at 1 K.

Input to the spectral calibration is the thermal scene (ENVI band sequential format) in the original geometry (i.e. not geocoded). The program will select 10 pixels from 10 image lines in the image center (nadir), calculate the wavelength shift, and the mean and standard deviation. Additionally, there is an optional keyword \( \text{box} \) where the averaging over a specified box of pixels can be specified to reduce the influence of noise. The default is \( \text{box}=1 \) (no pixel averaging, \( \text{box}=3 \) performs an averaging over 3 x 3 pixels). A wavelength shift < \( \text{FWHM}/30 \) will have a negligible effect and usually does not require an update of the sensor response functions and sensor-specific atmospheric LUTs.

When starting the spectral calibration program (either in the GUI or batch mode, see chapters 5 and 6, respectively) for an image named 'scene.bsq' the corresponding 'scene.inn' must already be available, because the sensor name and atmospheric LUTs are taken from this file. Note: this 'scene.inn' file contains the (sea level) water vapor column in its name, e.g. the string 'wv10' in 'h02000_wv10.tem'. The 'wv10' might not be the correct water vapor column, and for instance the 'wv04' or 'wv29' could be more realistic. However, this mainly influences the depth of the atmospheric absorption spectrum, it has a small influence on the wavelength shift calculated during the spectral calibration.

The spectral sampling distance SSD of the high-resolution thermal database (*.bt7 files) is SSD = 0.4 cm\(^{-1}\) in the 7 - 10 \( \mu m \) region, and SSD = 0.3 cm\(^{-1}\) in the 10 - 14.9 \( \mu m \) region. The full width at half max (FWHM) is always twice the sampling distance. This means we have a variable SSD
in wavelength, about 2 - 4 nm below 10 \( \mu m \), and 3 - 5 nm in the 10 - 13 \( \mu m \) part of the spectrum. This is adequate for the processing of thermal band imagery with bandwidths greater than 25 nm.
Chapter 4

Workflow

This chapter familiarizes the user with ATCOR-4’s workflow and with the program’s basic functionality using the graphical User interface. A detailed description of all modules and user interface panels is given in the subsequent chapter 5.

ATCOR may also be used in batch mode for most of its functions. A description of the batch mode can be found in chapter 6.

4.1 Menus Overview

To start ATCOR-4, double click the file 'atcor4.sav'. It will be opened through IDL or the IDL virtual machine. Alternatively, type atcor4 on the IDL command line after having added the atcor4 directory to the IDL search path. The graphical user interface of Fig. 4.1 will pop up. A large number of processing modules is available from this level as described in chapter 5. Most of them can be used without reading a detailed manual description because they contain explanations in the panels themselves. However, the next section guides the ATCOR newcomer during the atmospheric correction of a sample scene. The functions in the "File" menu allow the display of an image file, the on-screen display of calibration files, sensor response curves etc, see Fig. 4.2. More details about this menu are given in chapter 5.1.

![Figure 4.1: Top level graphical interface of ATCOR.](image_url)

The "Sensor" menu of Fig. 4.1 contains routines to create spectral filter curves (rectangular, Gaussian, etc) from a 3-column ASCII file (band number, center wavelength, bandwidth, one line per channel) provided by the user, calculates atmospheric look-up tables (LUTs) for new sensors, and computes the radiance/temperature functions for thermal bands; see Fig. 4.3 and chapter 5.2.

The "Topographic" menu contains programs for the calculation of slope/aspect images from a digital elevation model, the skyview factor, and topographic shadow. Furthermore, it supports the import and smoothing of DEMs and its related layers; see chapter 5.3.
The menu "ATCOR" gives access to the ATCOR-4 core processes for atmospheric correction in flat and rugged terrain. It also allows the tiled processing. It is further described in chapter 4.2 below and in chapter 5.4.

The "BRDF" menu provides access to the BREFCOR BRDF effects correction method and to the nadir normalization for wide field-of-view imagery; see chapters 5.5 and 5.5.2.

The "Filter" menu provides spectral filtering of single spectra (reflectance, emissivity, radiance) provided as ASCII files, spectral filtering of image cubes, and spectral polishing; see chapter 5.6.

The "Simulation" menu provides programs for the simulation of at-sensor radiance scenes based on surface reflectance (or emissivity and temperature) images; see chapter 5.7.

The "Tools" menu contains a collection of useful routines such as the calculation of the solar zenith and azimuth angles, spectral classification, nadir normalization for wide field-of-view imagery, spectral calibration, conversion of the monochromatic atmospheric database from one to another solar irradiance spectrum, scan angle file creation, and more; see chapter 5.8.
CHAPTER 4. WORKFLOW

Finally, the "Help" menu allows browsing of the ATCOR user manual, provides a link to web resources, and displays license and credits information, and serves to update your software; see chapter 5.9.

Figure 4.4: Topographic modules.
CHAPTER 4. WORKFLOW

4.2 First steps with ATCOR-4

The 'ATCOR' menu of Fig. 4.5 displays the choices 'ATCOR4f: flat terrain' and 'ATCOR4r: rugged terrain', compare Fig. 4.5. The last button starts the ATCOR processing in the image tiling mode, i.e., the image is divided into sub-images in x and y direction as specified by the user. This mode is intended for large scenes, compare section 5.4.11, and the '.inn' file with the processing parameters must already exist.

Let us start with a scene from a flat terrain area where no digital elevation model (DEM) is needed. Then the panel of Fig. 4.6 will pop up. First, the 'INPUT IMAGE FILE' has to be selected. ATCOR requires the band sequential format (BSQ) for the image data with an ENVI header. Next the acquisition date of the image has to be updated with the corresponding button. We work from top to bottom to specify the required information. The scan angle file is only required if the image geometry does not correspond to the original geometry as specified in the 'sensor*.dat' file which contains the number of pixels per line and the sensor field-of-view (FOV), see chapter 4.6. The scale factor defines the multiplication factor for surface reflectance (range 0 - 100%) in the output file. A scale factor of 1 yields the output as float data (4 bytes per pixel). However, a scale factor of 100 is recommended, so a surface reflectance value of say 20.56% is coded as 2056 and is stored as a 2 byte integer which means the file size is only half of the float file size with no significant loss of information.

If the input file name is "image.bsq" then the default output file name for the atmospherically corrected image is "image_atm.bsq". The user may modify the output name, but it is recommended to keep the "_atm.bsq" qualifier to facilitate the use of subsequent programs. Then the flight and solar geometry have to be specified as well as the sensor and the calibration file. The atmospheric file contains the look-up table (LUT) results of the radiative transfer calculations, separately for the solar and thermal region.

For a new user-specified sensor these LUTs have to be calculated once prior to the first call of ATCOR. This is done with the module RESLUT available under the 'Sensor' button of Fig. 4.1.

It is recommended to check the quality of the atmospheric correction before processing the image data. For that purpose, the SPECTRA module should be used where the surface reflectance of small user-defined boxes can be evaluated and compared with library spectra, compare chapter 5.5.2. In case of calibration problems, the spectral calibration module, available from the 'Tools' button of Fig. 4.1, and the radiometric (inflight) calibration may be employed before finally processing the image data. //

The 'AEROSOL TYPE' button provides an estimate for the recommended aerosol type (e.g. rural, maritime, urban) derived from the scene. This module also provides a visibility value for each aerosol type based on reference pixels (dark vegetation) in the scene. The 'VISIB. ESTIMATE'
button provides a visibility value for the selected aerosol type by checking dark scene pixels in the red band (vegetation, water) and NIR band (water). It is assumed that the lowest reflectance in the red band is 0.01 (1 percent) and 0.0 in the NIR band. Therefore, the obtained visibility value usually can be considered as a lower bound. The higher visibility value (of 'AEROSOL TYPE' and 'VISIB. ESTIMATE') is recommended as a start visibility for the SPECTRA module.

The 'Inflight Calibration' routine is described in chapter 2.4 and chapter 5.4.8.

The 'WATER VAPOR' button can be used to test the appropriate band combinations for the retrieval of a water vapor map without a calculation of the surface reflectance cube. Fig. 4.7 shows the panel with the image processing options. Some options may not be accessible, they are blocked if the required spectral bands are missing.

In case of a rugged terrain the ATCOR4r button has to be selected (Fig. 4.1, "ATCOR"). This panel is similar to Fig. 4.6, but an additional panel for the specification of the DEM files will appear (Fig. 4.8). The user has to provide the DEM file matched to the size of the input image. The slope and aspect files can be calculated from the corresponding module under "Topographic" (Fig. 4.1). These two files may need a special treatment as discussed in chapter 5.5.2. Therefore, they are not automatically created from the elevation file. The "skyview" file and cast shadow file are optional, only required in extremely steep terrain. The "skyview" calculation can also be found under the "Topographic" label of Fig. 4.1.

Depending on the selected image processing option some additional panels may pop up. Most of them are listed in chapter 5.4.10. They are mostly self-explaining and are not discussed here. They also contain default settings which can be used in most cases. When the main panel (Fig.
Figure 4.7: Image processing options. Right panel appears if a cirrus band exists.

Figure 4.8: Panel for DEM files.

4.6 is left and the SPECTRA or "IMAGE PROCESSING" sections are entered, all information is written to a processing initialization (.inn) file, e.g., "image.inn". When reloading the input file this information is read from the .inn file, so a new specification of all processing parameters is not necessary. Therefore, this .inn file can also be used for a batch processing, see chapter 6.

4.3 Survey of processing steps

Figure 4.9 shows the typical workflow of atmospheric correction. A detailed description of the corresponding graphical user interface for each module is given in chapter 5. First, the image is loaded with possibly some additional information (DEM files). Then the sensor has to be defined, the radiometric calibration file, and a basic atmosphere/aerosol type combination, e.g. a summer atmosphere with a rural aerosol. It is recommended to check the validity of the calibration and to estimate the visibility (and perhaps the atmospheric water vapor column wv) before processing the image cube. The SPECTRA module can be employed for this purpose, see chapter 5.4.5. Reflectance spectra of scene targets can be displayed as a function of visibility and water vapor.
and compared with field or library spectra. If calibration problems exist in a few channels a copy of the calibration file can be edited in these channels to match the reference spectrum. If there are problems in many channels the inflight radiometric calibration module should be used to generate a calibration file as discussed in chapters 2.2, 2.4, and 5.4.8.//

Figure 4.9: Typical workflow of atmospheric correction.

When the calibration file is OK the user can continue with the image processing. Depending on the available sensor channels there are options to process the imagery with constant or variable visibility and atmospheric water vapor. For large FOV sensors an option is available to correct for across-track illumination (BRDF) effects. This is especially useful if the image recording took place in the solar principal plane. In addition, a spectral polishing can be performed for the atmospherically and / or BRDF corrected data as indicated by the dotted lines of figure 4.9. The polishing requires hyperspectral imagery. Finally, a classification may be performed.

Figure 4.10 shows the input / output image files associated with ATCOR processing. On the left part the flat terrain case is treated, on the right part the rugged terrain case. If the image has the original scan geometry, i.e., the number of pixels per line corresponds to the number specified in the "sensor.dat" file, then the scan angle assignment for each pixel is calculated internally employing the total FOV and pixels per line. Otherwise, a scan angle file (created with PARGE) has to be provided.

In mountainous terrain, the DEM, DEM slope and aspect files are required. Optional input are the skyview file and the shadow map, the latter can also be calculated on-the-fly. The slope and aspect files can be calculated from ATCOR’s interactive menu, or run as a batch job (slopasp_batch see chapter 5). The skyview file has to be computed with the skyview program, see chapter 5.3.4.
### 4.4 Directory structure of ATCOR-4

Figure 4.11 shows the directory structure of the airborne version of ATCOR. There are a number of sub-directories with the following content: The 'bin' directory holds the ATCOR4 program with all modules as listed in chapter 5. The 'sensor' directory holds all supported airborne sensors in sensor-specific sub-directories.

As a possible convention, the last 2 digits indicate the year of calibration, e.g. hymap04 means the spectral and radiometric calibration of HyMap conducted in 2004. The "atm_database" contains the files of the monochromatic ATCOR database, see chapter 9.1. The "atm_lib" contains the results of the atmospheric database after resampling with the sensor-specific spectral response curves. The "spec_lib" is an optional subdirectory where the user can put field measurements of surface reflectance spectra resampled for the appropriate sensor. This is useful for inflight calibration or comparison of scene spectra with ground spectra. Finally, the "demo_data" contains some demo imagery to be able to run ATCOR-4 immediately.

### 4.5 Convention for file names

Although file names are arbitrary it is useful to agree on some conventions to facilitate the search of files, especially concerning the extensions of file names.

Input images to ATCOR must have the band sequential format (BSQ), therefore it is recommended to employ the '.bsq' as an extension, e.g. 'image1.bsq'. Then in this example the default output file name of ATCOR will be 'image1.atm.bsq' and a log report of the processing is available in 'image1.atm.log'.

Once an image is processed with ATCOR all input parameters are saved in a file '*.inn' that is automatically loaded should the image be processed again.

The recommended extension for the radiometric calibration files is '.cal'. Other extensions are '*.ele.bsq' for the digital elevation file, '*.slp.bsq' for the DEM slope file, '*.asp.bsq' for the DEM aspect, '*.sky.bsq' for the sky view file, '*.ilu.bsq' for the solar illumination file in rugged terrain, and '*.cla.bsq' for the classification map of SPECL. The interface to PARGE (Schläpfer and Richter 2002 [66]) uses a scan angle file '.sca.bsq', where band 1 provides the scan view angle for each pixel.
and band 2 the absolute scan azimuth angle. The visibility index map is named ‘*_visindex.bsq’, the aerosol optical thickness is ‘*_aot.bsq’, the cloud/building shadow map is named ‘*_fshd.bsq’, and the atmospheric water vapor map ‘*_wv.bsq’.

**Thermal band imagery**

In case of thermal band imagery the surface temperature and emissivity are stored in separate files. Surface temperature is appended to the reflectance cube, i.e. is included as the last channel in the ‘*_atm.bsq’ file, e.g., ‘image1_atm.bsq’.

The surface temperature calculation is based on an assumption for the emissivity in one spectral band. Three options have been implemented:

- a constant scene emissivity of $\epsilon = 0.98$ in the channel used for the surface temperature calculation.

- a map of 3 emissivity classes (vegetation, soil, others) depending on surface cover. The surface cover class is calculated on the fly based on the vegetation index and red/NIR reflectance values for each pixel. Typical average emissivity values were assumed (vegetation $\epsilon = 0.97$; soil, asphalt, concrete $\epsilon = 0.96$; others $\epsilon = 0.98$). The file name of the 3-class emissivity map is ‘image1_atm_emis3.bsq’, the ‘3’ indicating the 3 classes, if the file name of the scene is ‘image1.bsq’.

- a detailed map of up to 15 emissivity classes, compare table 10.1. The user has full control over the assignment of an emissivity value for each class, i.e. the file ‘emissivity.dat’ can be edited by the user. This emissivity assignment pertains to the channel for which the surface temperature is calculated, compare the sensor definition file in chapter 4.6. The emissivity
map for each scene can be calculated with program SPECL prior to the processing of the thermal bands, but after processing of the coregistered reflective bands.

SPECL classifies the surface reflectance spectrum of each pixel by comparing it to a set of template surface cover spectra and assigns the class for which the best match is found. If no sufficient match is found, the pixel is assigned to 'not classified'. The file name is "image1_atm_cla.emi.bsq". It is generated simultaneously with the classification file "image1_atm_cla.bsq", see chapter 5.8.2.

One of the first two options is usually selected for sensors with a single thermal band. The last option can be selected in case of multispectral thermal bands. After the surface temperature has been calculated based on the provided emissivity map in one channel, the remaining emissivity channels are computed and put into the file "image1_atm_emiss.bsq". The multi-band emissivity file is coded with 16 bits per pixel scaled with the factor 1000.

4.6 Definition of a new sensor

A few steps have to be taken to include a new airborne sensor in ATCOR, compare Fig. 5.14. These are:

- A new sensor subdirectory in the "/atcor/sensor/" folder has to be created. This is easiest done using the routine Define Sensor Parameters as described in chapter 5.2.1. Please make sure this name does not agree with an existing multispectral sensor name in the "../atcor/cal/" folder. This process may also be done manually by copying and adapting an existing user-defined sensor or one of the samples provided.

- A sensor-definition file must be specified. Just copy any of the existing files, e.g., "sensor_hymap2003.dat", and modify the appropriate lines, see the next table (or use the function above to make the changes)

- A wavelength file ("*.wvl") has to be specified. It is a simple ASCII file with three columns (band number, center wavelength, and bandwidth), compare Fig. 5.14. Center wavelength and bandwidth may be given in the nm or µm unit. The first line may contain an optional header with text. This wavelength file will be used to create the spectral response function for each band as a numerical table (the 'band*.rsp' files), compare Fig. 5.14.

Eight analytical filter shapes can be selected from the top level graphical interface (Fig. 4.1 when selecting the menu ‘Sensor’, then choose 'Create Channel Filter Files' button. Then the menu of Fig. 4.12 will pop up, and one of these 8 filter shapes can be selected. Filter numbers 1 to 4 are of the Butterworth type, the slow drop-off for the Butterworth order 1 is truncated at the 0.05 response and set to zero. The filter type 9 (parameter filter_type in Table 4.3) is reserved for arbitrary user-specified channel filter functions.

- A calibration file has to be provided (e.g., "hymap2003.cal") in the new sensor sub-directory.

- The RESLUT (resample atmospheric LUTs) program has to be run to generate the atmospheric LUTs for the new sensor employing the monochromatic atmospheric database in "../atcor4/atm_database/". These resampled "*.atm" files will automatically be placed in a sub-directory of './atcor4/atm_lib/' with the name of the selected sensor. RESLUT will also create the resampled spectrum of the extraterrestrial solar irradiance in the appropriate
".../sensor/hymap2003/" folder, see chapter 9.2, e.g., "e0_solar_hymap2003.spc". In addition, the first use of RESLUT will create the "pressure.dat" file in the sensor-specific folder.

Now you may start processing imagery of the just defined sensor. If the sensor has thermal spectral bands the program RESLUT will automatically calculate the coefficients of the temperature / radiance relationship. They will be stored in a "sensor*.bbfit" file in the appropriate sensor subdirectory. RESLUT will also create the resampled spectrum of the extraterrestrial solar irradiance, e.g. "e0_solar_hymap04.spc".

Two examples of the file "sensor_example.dat" are given below. The first table presents a sensor without thermal bands, and without gain settings. The second table defines a sensor with 79 bands (DAIS-7915), having 6 thermal channels. The mid-IR bands have to be specified separately, no atmospheric correction is performed for these bands. The input channels are included at the same position as output channels. In the ENVI header, they are labeled "DN".

| 60.0  512 across-track FOV [degree], pixels per line | 1  128 first, last reflective band (0.35-2.55 \(\mu m\)) |
| 0  0 first, last mid IR band (2.6 -7.0 \(\mu m\)) | 0  0 first, last thermal band (7.0 -14 \(\mu m\)) |
| 0  no tilt in flight direction | 0  required dummy |

Table 4.1: Sensor definition file: no thermal bands.

Line 5 of the sensor definition file is retained for compatibility with ATCOR-4 versions below 3.0. The tilt parameter is always zero, i.e., tilt sensors are not supported in the atmospheric database. In case of tilt sensors the old ATCOR-4 (version below 3.0) must be used which requires a MODTRAN® 5 license to calculate the atmospheric LUTs, because no database is available for tilt sensors. Line 6 is a required dummy to be compatible with previous versions.
Table 4.2: Sensor definition file: instrument with thermal bands.

<p>| | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>60</td>
<td>0</td>
<td>across-track FOV [degree], pixels per line</td>
</tr>
<tr>
<td>72</td>
<td>1</td>
<td>first, last reflective band (0.35-2.55 μm)</td>
</tr>
<tr>
<td>73</td>
<td>3</td>
<td>first, last mid IR band (2.6 -7.0 μm)</td>
</tr>
<tr>
<td>74</td>
<td>9</td>
<td>first, last thermal band (7.0 -14 μm)</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>no tilt in flight direction</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>required dummy</td>
</tr>
<tr>
<td>77</td>
<td>7</td>
<td>temperature band (itemp_band=77)</td>
</tr>
</tbody>
</table>
4.7 Spectral smile sensors

Imaging systems can employ different techniques to record a scene: the whiskbroom design uses a rotating or oscillating mirror to collect an image line across-track direction (with one or a few detector elements per spectral band). The forward direction is provided by the motion of the platform. Secondly, a pushbroom linear array can perform the same task without moving optical elements, but the number of array lines (each recording a certain spectral channel) in the focal plane is limited. The third imaging technique employs an area detector array where one direction collects the spatial information (across-track) and the orthogonal direction covers the spectral dimension. The advantage of the last two techniques is a longer pixel dwell time and a potentially improved signal-to-noise ratio (SNR). The drawback is a substantial increase in the spectral and radiometric characterization, i.e., a change of the channel center wavelength across the columns of the array (spectral ”smile”), spatial misregistration (“keystone”), and detector non-uniformity problems [52], [19], [74].

Typical representatives of the whiskbroom type are Landsat TM/ ETM, HyMap, AVIRIS, and Daedalus. These instruments almost show no spectral smile, i.e., the channel center position and bandwidth do not depend on column pixel location.

Spaceborne hyperspectral instruments showing the ”smile” effect are Hyperion and CHRIS/Proba, airborne instruments are for example CASI-1500 and APEX.

This section describes the ATCOR input files required for smile sensors. There are only two changes compared to the ”non-smile” instruments:

- The sensor definition file, e.g., ”sensor_casi1500.dat”, needs one more line (see Table 4.3) containing the parameters ismile (=1 if smile sensor, otherwise 0) and filter_type (a number between 1 and 9 for the type of channel filter function) compare section 4.6 and Fig. 4.12. The filter types 1 to 8 are analytical functions, filter type 9 is reserved for arbitrary user-defined channel filter functions (the ’band*.rsp’ files). Center wavelength and bandwidth for each channel are defined in the wavelength file (*.wvl) pertaining to the center pixel=column of the detector array.

- For each spectral channel j the channel center wavelength \( \lambda_c(j) \) depends on the image column or pixel position x. The absolute value of \( \lambda_c(j) \) is specified in the wavelength file used to generate the spectral channel response functions, and it is also included in the sensor-specific solar irradiance file (e.g., ”e0_solar_casi1500.spc”). If n is the number of image columns, the change \( \Delta(x, j) \) of the center wavelength \( \lambda_c(j) \) with the pixel position x can be described as a 4th order polynomial (using the nm unit):

\[
\Delta(x, j)[\text{nm}] = a_0(j) + a_1(j) \cdot x + a_2(j) \cdot x^2 + a_3(j) \cdot x^3 + a_4(j) \cdot x^4
\]

\[
\lambda_c(x, j) = \lambda_c(j) + \Delta(x, j)
\]

The first left-hand image pixel is x=0, the last right-hand image pixel is x=(n-1). The coefficients \( a_i(j) \) have to be stored in an ASCII file, corresponding to the band sequence. The first column must contain the wavelength or band number, followed by the five channel-dependent coefficients (beginning with \( a_0 \) and ending with \( a_4 \)), one line per channel. The fixed file name is smile_poly_ord4.dat and it has to be located in the corresponding sensor subdirectory. In the ideal case these coefficients should be derived from laboratory measurements.
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Since an accurate description is only required for channels in atmospheric absorption regions, the 5 coefficients can be set to zero for the remaining regions, but they must be provided for each channel. If all 5 coefficients are set to zero for a certain channel, this channel is processed in the "non-smile" mode which will expedite the processing.

- Optionally, the spectral bandwidth (FWHM) might also depend on the image column. Again, a 4th-order polynomial is used to describe the bandwidth change depending on column position x and channel j:

$$\Delta_{1}(x, j)[nm] = b_0(j) + b_1(j) \cdot x + b_2(j) \cdot x^2 + b_3(j) \cdot x^3 + b_4(j) \cdot x^4$$ (4.3)

$$FWHM(x, j) = FWHM(j) + \Delta_{1}(x, j)$$ (4.4)

The fixed file name is smile_poly_ord4_fwhm.dat and it has to be located in the corresponding sensor sub-directory. It is an ASCII file with 6 columns per channel, the first column is the channel number or wavelength, the remaining columns contain the polynomial coefficients starting with $b_0$.

| 30.0 | 512 | across-track FOV [degree], pixels per line |
| 1   | 128 | first, last reflective band (0.35-2.55 µm) |
| 0   | 0   | first, last mid IR band (2.6-7.0 µm) |
| 0   | 0   | first, last thermal band (7.0-14 µm) |
| 0   |     | no tilt in flight direction |
| 0   |     | required dummy |
| 1   | 5   | 1=smile sensor, 5 = Gaussian spectral channel filter |

Table 4.3: Sensor definition file: smile sensor without thermal bands.

Imagery from smile sensors must be processed in the raw geometry (IGM Image Geometry Map) to preserve the original image columns. During the surface reflectance retrieval the atmospheric/topographic correction is performed on a per column basis, i.e., to each image column its appropriate center wavelength/bandwidth is associated.

The per-column processing typically implies a factor of 8 increase in processing time. The following steps are to be performed:

1. Define a sensor (*wvl, *cal, *rsp files, RESLUT) using the original set of wavelengths (pre-launch values, as provided with the data from the data provider),

2. Run the smile detection tool (compare Section 5.8.3, using the sensor defined in 1 and appropriate absorption features, to derive the polynomial coefficients smile_poly_ord4.dat for smile correction in step 3; alternatively: enter the smile polynomial factors from laboratory calibration. Note: if two detectors are in the sensor system, this should be done separately for VNIR and SWIR, option 'repeat values', resolution 0.02 nm. Combine the two files for VNIR and SWIR manually into one file afterwards.

3. Using the same sensor as above, run the atmospheric correction with the smile correction option switched 'ON'; (after putting the new file smile_poly_ord4.dat into the sensor definition directory),
4. Apply the spectral polishing routine (see Section 5.6.3 and 5.6.4, and
5. Run the Spectral Smile Interpolation module (see Section 5.6.7 on the atmospherically corrected image.

4.8 Haze, cloud, water map

Although the surface reflectance cube (and temperature / emissivity for thermal channels) is the main result of the atmospheric correction, some additional products are often requested. One of these products is a map of the haze, cloud, water and land pixels of a scene. This map not only delivers a basic scene classification, but it may also contain information about potential processing problems. For example, if turbid water pixels are not included in the water mask, the haze mask may also be not appropriate and consequently, results of the haze removal over land might be of poor quality. Such a pre-classification as part of the atmospheric correction has a long history [22, 40, 59, 60, 61, 41, 49]. It is also employed as part of NASA’s automatic processing chain for MODIS [1] using the classes land, water, snow/ice, cloud, shadow, thin cirrus, sun glint, etc.

Therefore, the calculated haze/cloud/water map is a useful optional output of ATCOR. It is enabled by setting the parameter ihcw = 1 in the preference_parameters.dat file (see chapter 9.4). If the file name of the imagery is "image.bsq" the corresponding map is named "image_out_hcw.bsq". It is a 1-channel false-color coded ENVI file. In principle, if a certain mask of "image_out_hcw.bsq" (say haze pixels) contains artifacts, it may be edited, and if the edited file is named "image_hcw.bsq" it will automatically be used for the ATCOR processing. This means ATCOR can repeat the processing with an improved (edited) haze mask.

The file "image_hcw.bsq" can also be provided by an external (ATCOR-independent) source. In any case, if this files exists, ATCOR will skip its internal calculations of these masks and use the pre-calculated map.

![Diagram](image)

**Figure 4.13:** Optional haze/cloud/water output file.

The haze/cloud/water file contains the following classes (see Table 4.4):

- land
- water
- boundary layer haze (two classes: thin-to-medium haze, and medium-to-thick haze)
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<table>
<thead>
<tr>
<th>label</th>
<th>definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>geocoded background</td>
</tr>
<tr>
<td>1</td>
<td>shadow</td>
</tr>
<tr>
<td>2</td>
<td>thin cirrus (water)</td>
</tr>
<tr>
<td>3</td>
<td>medium cirrus (water)</td>
</tr>
<tr>
<td>4</td>
<td>thick cirrus (water)</td>
</tr>
<tr>
<td>5</td>
<td>land</td>
</tr>
<tr>
<td>6</td>
<td>saturated</td>
</tr>
<tr>
<td>7</td>
<td>snow</td>
</tr>
<tr>
<td>8</td>
<td>thin cirrus (land)</td>
</tr>
<tr>
<td>9</td>
<td>medium cirrus (land)</td>
</tr>
<tr>
<td>10</td>
<td>thick cirrus (land)</td>
</tr>
<tr>
<td>11</td>
<td>thin-medium haze/land</td>
</tr>
<tr>
<td>12</td>
<td>medium-thick haze/land</td>
</tr>
<tr>
<td>13</td>
<td>thin-medium haze/water</td>
</tr>
<tr>
<td>14</td>
<td>medium-thick haze/water</td>
</tr>
<tr>
<td>15</td>
<td>cloud/land</td>
</tr>
<tr>
<td>16</td>
<td>cloud/water</td>
</tr>
<tr>
<td>17</td>
<td>water</td>
</tr>
<tr>
<td>18</td>
<td>cirrus cloud</td>
</tr>
<tr>
<td>19</td>
<td>cirrus cloud thick</td>
</tr>
</tbody>
</table>

Table 4.4: Class label definition of "hcw" file.

- cirrus (three classes for thin, medium, thick; and cirrus cloud, thick cirrus cloud), provided a narrow channel around 1.38 $\mu$m exists.
- cloud over land, cloud over water
- snow (requires a 1.6 $\mu$m channel)
- saturated pixels using the criterion $T > 0.9 \times DN_{\text{max}}$ where $T$ is a threshold set at 0.9 times the maximum digital number. This criterion is only used for 8 bit and 16 bit (signed or unsigned) data, no threshold is defined for 32 bit (integer or float) data. As an example, for an 8 bit/pixel data encoding we obtain $T = 0.9 \times 255 = 230$, and pixels with a grey value greater than 230 are flagged as (truly or potentially) saturated.

Although the saturation nominally starts at $DN_{\text{max}}$ some sensors already show a non-linear behavior around 0.9 $\times$ $DN_{\text{max}}$, so the factor 0.9 is a precaution to be on the safe side.

This saturation check is performed for two channels in the visible region (blue band around 470-500 nm, and a green band around 550 nm). For multispectral sensors, the blue band is usually the critical one concerning saturation. For hyperspectral sensors with many blue bands the one closest to 450 nm is taken. Although the haze/cloud/water file contains saturated pixels based on two visible bands, the percentage of saturated pixels for all bands will be given in the corresponding ".log" file. However, the check of the blue and green channel normally captures all saturated pixels.

The thresholds for the cirrus class definition are described in chapter 10.2.
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Note: cloud (or building) shadow pixels are not included here, they are stored separately (file image_fshd.bsq”).

4.9 Processing of multiband thermal data

Several options have been implemented to process multiband thermal data, see chapter 10.1.5 for details. Apart from the final products (surface temperature and emissivity) intermediate products are available such as surface radiance, at-sensor blackbody temperature, and surface blackbody temperature. The intermediate products might be useful to trace back spectral or radiometric problems. If image.bsq denotes the file name of the input image then the following products are available:

- image_atm_emis3.bsq: 3 or 4 emissivity classes obtained from an on-the-fly (in memory) pre-classification (vegetation, soil, sand, water). The pre-classification requires daytime data acquisition and spectral bands in the solar region. This file has one channel with the emissivity values for the specified thermal band, or in case of ANEM the pixel-dependent values assign the maximum emissivity of all available thermal bands.

- image_atm_emiss.bsq contains the spectral emissivity map for all thermal channels.

- image_atm_emiss_lp3.bsq is the same emissivity map but filtered with a 3-channel low pass filter to smooth spectral noise features (requires at least 10 thermal bands).

- image_atm_emiss_lp5.bsq is the same emissivity map but filtered with a 5-channel low pass filter to smooth spectral noise features (requires at least 30 thermal bands).

- image_atm_isac_emiss.bsq: emissivity cube for the ISAC algorithm.

- image_at_sensor_channel_tmax.bsq: map of channel numbers with maximum at-sensor temperature.

- image_at_surface_channel_tmax.bsq: map of channel numbers with maximum surface temperature.

- image_at_sensor_tbb.bsq: at-sensor brightness temperature cube

- image_at_surface_tbb.bsq: at-surface brightness temperature cube.

The last channel of image_atm.bsq contains the surface temperature map evaluated with the appropriate emissivity, the preceding thermal channels in this file contain the surface radiance.

In case of the ISAC algorithm an additional file image_isac_lpath_trans.dat contains the spectral path radiance and transmittance estimates for the scene. Fig. 4.14 shows an example of these spectra derived from a SEBASS scene.

Fig. 4.15 presents the at-sensor/at-surface radiance and brightness temperatures. The at-sensor products clearly show the atmospheric absorption features which are removed in the at-surface quantities, apart from small residual effects. The bottom graphic presents the corresponding surface emissivity spectrum.
Figure 4.14: Path radiance and transmittance of a SEBASS scene derived from the ISAC method.

Figure 4.15: Comparison of radiance and temperature at sensor and at surface level.
4.10 External water vapor map

Sometimes, it is convenient to use an external water vapor map, even if this map could be derived from the scene. If the scene is named "scene.bsq" then the external map should be named "scene_wv.bsq" and it must have the same number of image lines and pixels per line as the scene. If this file is available in the same directory as the scene, it will be automatically used during the processing and the usage is also documented in the log file ("scene_atm.log"). Note: this feature is only supported for instruments that are able to retrieve the water vapor column with the intrinsic channels, because the prerequisite is the availability of the corresponding LUTs.

4.11 External AOT 550 nm map

Although ATCOR calculates the aerosol optical thickness map (at 550 nm) if the variable aerosol option is set (parameter npref=1 in the '*.inn' file), the import of an external AOT(550nm) map is also supported. This file should be located in the same folder as the scene. If the file name of the image is 'scene.bsq' then the corresponding AOT map should be named 'scene_aot550.bsq'. The size of this map should match the scene size (i.e. same number of columns and lines as the scene). If this map is stored as float data, then it contains the original AOT(550nm) values (i.e. without scale factor). This map can also be provided as 16-bit signed or unsigned integer with a mandatory scale factor of 1,000.

4.12 Filter for HySpex

HySpex is a suite of hyperspectral cameras developed and manufactured by Norsk Elektro Optikk, Norway (www.neo.no). There is a navigation file for each acquired airborne hyperspectral cube in the following format: an ASCII file with 8 columns separated by blanks, consisting of:

- line, longitude (deg), latitude (deg), ALT (meters), roll (deg), pitch (deg), heading (deg), UTCx

ALT is the flight altitude above sea level, and UTCx is the UTC time in seconds since midnight. The acquisition date is specified in the ENVI header. This information enables the automatic calculation of the solar zenith and azimuth angles evaluated for the center scan line of the scene. For this purpose, the nav ".txt" file must be in the same folder as the scene, named "scene.txt" if the scene name is "scene.bsq" or "scene.img".

In addition, the ENVI header of each HySpex scene contains the radiometric scaling factor in the 'description' tag, and it is used to create the radiometric calibration file "scene.cal" with the gain c1 = 100 / scaling (unit mWcm\(^{-2}\)sr\(^{-1}\)μm\(^{-1}\)). It is the same factor for all bands, and the offset c0 is zero.

4.13 Airborne FODIS instrument

Some airborne hyperspectral sensors are optionally equipped with an add-on instrument that can measure the downwelling hemispherical solar flux at the aircraft altitude. This add-on instrument is usually named FODIS (Fiber Optic Downwelling Irradiance Sensor). FODIS data could be useful for atmospheric correction, especially during adverse weather conditions. However, currently this information is seldom used for three reasons: atmospheric correction software usually does not
support processing of FODIS data, the data must be corrected for the roll, pitch, yaw movements of the aircraft, and FODIS calibration can be a problem.

ATCOR now offers a tool to use FODIS data during the atmospheric correction. The general restrictions are:

- It can only be used in flat terrain processing, because the direct and diffuse solar fluxes (required for a rugged terrain processing) are no more available, as FODIS measures only the total downwelling solar flux at the aircraft level.

- FODIS data contain no information on the atmosphere below the flight level, so the precalculated atmospheric LUTs containing path radiance, direct and diffuse transmittances have to be used, only the global solar flux on the ground \( E_g \) is re-calculated based on the FODIS measurements.

- The hyperspectral scene must be processed in the original geometry, because FODIS measures one spectrum per original scan line. Geocoding can be performed later if required.

Without FODIS data the global flux at the ground is calculated as the sum of the direct and diffuse flux:

\[
E_g = E_0 \tau_s \cos \theta_s + E_{\text{dif}} \tag{4.5}
\]

where \( E_0 \) is extraterrestrial solar irradiance, \( \tau_s \) = sun - ground (beam) transmittance, \( \theta_s \) = solar zenith angle, and \( E_{\text{dif}} \) the diffuse flux on the ground. With FODIS data eq. (4.5) is replaced by:

\[
E_g = E_f (\tau_{\text{dir}} + \tau_{\text{dif}}) \tag{4.6}
\]

where \( E_f \) is the geometrically corrected spectral FODIS flux per scan line, and \( \tau_{\text{dir}}, \tau_{\text{dif}} \) are the direct and diffuse transmittances. \( E_f \) is calculated from the originally measured flux \( E_f^{(0)} \) as

\[
E_f = E_f^{(0)} \cos \theta_s / \cos \beta \tag{4.7}
\]

where \( \theta_s \) is the solar zenith angle and \( \beta \) is the incident solar angle (between FODIS normal and incident solar beam):

\[
\cos \beta = \cos \theta_s \cos \theta_f + \sin \theta_s \sin \theta_f \cos (\phi_s - \phi_f) \tag{4.8}
\]

The FODIS slope and azimuth angles \( \theta_f, \phi_f \) are computed from the FODIS attitude angles (roll: \( \theta_{\text{roll}} \), pitch: \( \theta_{\text{pitch}} \), yaw or heading: \( \theta_{\text{yaw}} \) ):

\[
\begin{align*}
\theta_f &= \sqrt{\theta_{\text{roll}}^2 + \theta_{\text{pitch}}^2} \tag{4.9} \\
\phi_f &= (\theta_{\text{yaw}} + 180 + \arctan(\theta_{\text{roll}}/\theta_{\text{pitch}})) \mod 360^\circ \tag{4.10}
\end{align*}
\]

The last equation holds for the standard case of the aircraft nose up and the sign conventions are:

- roll angle is positive for the aircraft right wing up
- pitch angle is positive for the aircraft nose up
- yaw or heading ranges between 0° and 360° (90° = east)
The panel "TOOLS" contains a graphical user interface to process FODIS data in the Specim CaliGeo and NERC (Natural Environment Research Council, UK) formats. These are the currently supported formats and more will be added on demand.

Input to the GUI is the image file (example: "scene.bsq"), sensor, and FODIS format, see Fig. 4.16. The FODIS nav file(s) and irradiance file are included automatically if the mandatory file name conventions are kept, as described below. The output (example: "scene_fodis.slb") is the geometrically corrected FODIS at-sensor flux in the ENVI spectral library format, stored as one reference spectrum per image line (unit \(mWc m^{-2} \mu m^{-1}\)). The data format is specified in the ENVI header: parameter 'lines' contains the number of image lines, 'samples' contains the number of bands, parameter 'bands = 1', and 'data type = 4', i.e. binary float data.

If the "scene_fodis.slb" file exists in the image folder, ATCOR(flat terrain) will automatically use this file during atmospheric correction, and this will be documented in the corresponding "scene_atm.log" file.

Mandatory name conventions apply for the FODIS processing:

- **Specim CaliGeo format**: if the scene file name is "scene.bsq", then the navigation data must be named "scene_nav.txt", and the FODIS data (coming in the binary ENVI BIP format) "scene_fodis0.bip". These files have to be in the same folder as the scene. The angles in the nav file use the degree unit. The ASCII file '*nav.txt' contains 9 columns separated with blanks:
  - scan line number, time, x, y coordinates, flight altitude, heading, roll, pitch, and aircraft speed.
  - Only data in columns 6 to 8 (heading, roll, pitch) is used. Heading is in the (-180,180) degrees interval and will be converted into the (0,360) interval.
  - The FODIS flux measurements in '*fodis0.bip' are float data, and the ENVI header specifies the parameters 'samples = 1', 'lines = n' (where n is the number of image lines), 'bands = m', (where m is the number of bands), and 'data type = 4', i.e. float. All other ENVI header information is not used. The standard spectral flux unit is \((mWcm^{-2}nm^{-1})\), requiring a conversion factor of 0.001 for the unit \((mWcm^{-2}\mu m^{-1})\) which is used by ATCOR. However, the FODIS GUI (see Fig. 4.16) provides a flexible widget for re-scaling if necessary.

- **NERC format**: again, if the scene file name is "scene.bsq", then the roll, pitch, yaw files must be named "scene_COroll.txt", "scene_COpitch.txt", "scene_COhead.txt" and the FODIS measurement file has to be "scene_FODIS.txt". These files have to be in the same folder as the scene.
  - The angles in the roll, pitch, yaw files are in degrees times a scale factor 1,000. The COroll, COpitch, COhead.txt ASCII files contain one value per scan line, representing the corresponding angle times 1,000 stored as integer. The ASCII file "scene_FODIS.txt" contains the measured flux spectra with n columns (n = number of scan lines of scene) and m lines (m = number of bands). All flux values are in the unit \((mWcm^{-2}nm^{-1})\) stored as integer in the FORTRAN "I8" format.

**Note**: The capability of water vapor retrieval is switched off in case of FODIS processing, as it relies on the sun-ground-sensor radiation components and FODIS allows only the calculation of the sensor-ground-sensor radiation components. So the water vapor of the selected '.atm' LUT is taken. However, there is the possibility of providing an external water vapor map (see chapter 4.10), which can be calculated for the same scene in the non-FODIS mode (e.g. by moving the file "scene_fodis.slb" to another folder or renaming it temporarily in the same folder).
4.14 External float illumination file and de-shadowing

If the scene is processed with a DEM the additional files of slope, aspect (and optionally skyview and cast shadow) are used to calculate the solar illumination map (cosine of the local solar zenith angle), which is stored as byte data (range 0-100, scale factor 100). If the file name of the input scene is `scene.bsq` then the corresponding illumination file is named `scene_ilu.bsq`.

However, there is an option to process an external illumination map and skip the internal calculation. In this case the requested condition is a float-coded file (range 0 - 1) with the nomenclature `scene_raw_ilu.bsq` (first priority) or `scene_ilu.bsq` (second priority) to distinguish it from an internal calculation (if an internal calculation was performed previously). In case of a rugged terrain, the illumination-based de-shadowing algorithm stores results in a multi-layer file, and the first layer is the float-coded illumination.

Two de-shadowing methods are implemented in ATCOR: the matched filter approach (see chapter 10.5.6) and the illumination-based approach (see chapter 10.1.2). The latter delivers a float-coded illumination file (range 0 - 1) containing the fractional shadow.

If a float-coded illumination file exists in the folder of the input scene, then it is used for de-shadowing, also for the flat terrain case (without DEM files). Therefore, it has always the first priority. If the matched filter de-shadowing is intended, then the float-coded illumination file has to be renamed or temporarily removed from this folder.

Internally, the range 0 - 1 of the float illumination is then re-scaled to 0 - 1000 to be compatible with the matched filter de-shadowing case (only for the purpose of de-shadowing, not for the correction of terrain effects).

4.15 BRDF Correction

For the BRDF correction, including the BREFCOR approach, the following workflow is recommended:
CHAPTER 4. WORKFLOW

1. Perform the atmospheric compensation using ATCOR-4. The indigence BRDF needs to be corrected in this step by selecting the BRDF option (for terrain correction). The observer BRDF correction is a step done subsequently to the atmospheric correction.

2. Apply spectral polishing routines (if required).

3. Prepare the inputs for the BREFCOR model calibration; i.e. check the scan angle file, specifically the second layer should be contained therein as absolute azimuth.

4. Select calibration data set: a number of 2-4 image scenes from a whole campaign should be used for the calibration. These scenes should have good statistical distribution of the objects of interest in across track direction and shouldn’t contain too many clouds and cast shadows.

5. Start the model calibration with standard 5 levels on a 3-band image (RGB or NRG) first and select ‘write ANIF outputs’. Check the BCI side output image and compare to the selected bci level limits.

6. If level limits are not appropriate for the object types, the self defined level limits are to be set in the model calibration; redo the calibration again. Possibly you’ll iterate this process and also you may need to increase the fitting accuracy threshold in order to get a BRDF model appropriate to your image.

7. Now, you’re read to go for the full image, using the model you just created. Check the outputs, possibly do a mosaicking for the analysis.

8. If everything is correct, the whole campaign may be proceed using this calibrated model.
Chapter 5

Description of Modules

For most ATCOR modules a convenient graphical user interface is available, but batch jobs can also be submitted. If the atcor4 binary `atcor4.sav` is opened by the IDL virtual machine or when "atcor4" is typed on the IDL command line a menu with pull-down buttons pops up see Figure 5.1, with a thematic grouping of modules. A detailed discussion of the interactive panel-driven modules is given hereafter whereas a description of the batch commands can be found in chapter 6.

![Figure 5.1: Top level menu of the airborne ATCOR.](image)

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5.1 Menu: File

The menu ‘File’ offers some collaborative tools for handling of the data and ENVI files. Below, a short description of the individual functions is given:

**Figure 5.2: The File Menu**

5.1.1 Display ENVI File

Use this function for displaying a band sequential ENVI-formatted file in a simple way.

An initial dialog allows to select the band(s) to display - either a true color, CIR color, or a single band mode may be selected.

After band selection, the following options are available within the file viewer from the menu:

- **File: Show ENVI Header** Displays the ENVI header of the current image in a new editable window. This allows to make changes to the ENVI header. Note that the file needs to be loaded from scratch if changes have been made.

- **File: Band Selection** Allows to select a new combination of spectral bands and updates the display.

- **File: Display ENVI Image** Displays an additional ENVI image in a new window.

- **File: Display TIFF Image** Loads a multi-band TIFF image in a new window.

---

1The ENVI format is a raw binary file, accompanied by an ASCII header *.hdr; in ATCOR it should be stored in band sequential order. Note: this ENVI file display routine may be used separately using the GLIMPS software, available free of charge from ReSe Applications.
CHAPTER 5. DESCRIPTION OF MODULES

Figure 5.3: Band selection dialog for ENVI file display

**File:** Plot Spectral Library  Displays the spectra of an ENVI spectral library.

**File:** Show Text File  Displays an ASCII file.

**File:** Close  Closes the window.

**Edit:** Equalize Image  Performs a histogram equalization on the three bands.

**Edit:** Scale Image  Applies standard linear scaling on the imagery on 5 levels.

**Edit:** Scale Zoom  Applies standard linear scaling on the imagery on 5 levels based on the statistics of the Zoom Window.

**Edit:** No Scaling  Reverts to unscaled display of the image.

**Edit:** Scale to Range  Scales a single-band image linearly to a range entered as lower and upper limit (only applicable in single band displays).

**Edit:** Load Color Table  Loads the standard color table (applicable for single band display only).

**Calculate:** Band Index  Allows to calculate a spectral band index from the currently displayed image such as band ratio or NDVI-type of indices.

**Calculate:** Spectral Classification  Allows to calculate a spectral quantification/classification using the spectral angle mapper or spectral fitting technique.

**Calculate:** Statistics  Calculate the statistics per spectral band from the current image and stores the result as either an SLB or TEXT file.

**Profile:** Horizontal  Opens a window for a horizontal profile through the image (of the first band only). The profile is updated for the cursor location in the zoom window whenever the zoom window is clicked.

**Profile:** Vertical  Opens a window for a vertical profile through the image (of the first band only).
CHAPTER 5. DESCRIPTION OF MODULES

Profile:Spectrum Opens a window for a spectrum of the image (for images with 4 and more bands only); use the middle mouse button to show the spectrum directly.

Export Allows to export the currently displayed image of either the zoom or the full image window to one of the given image data formats. The displayed image may be exported as a scaled 8bit/24bit image to the available standard image formats.

Note: when middle-clicking in the zoom window, the current image value and location is displayed, or a small plot of the spectrum at this pixel location is created (same as the function Profile:Spectrum of above). The menu in the such loaded window allows to save the spectrum to an ASCII table, to adapt the graph’s properties and font size, configure the display and to output the graph to an appropriate graphics format.

The position field may be used to jump to a given coordinate by entering the coordinates manually and using the 'Enter' key on the keyboard. The button Get Position is used to get the currently stored coordinates from clipboard, whereas the button Push Position lets you push the currently showns coordinates to clipboard.

Figure 5.4: Display of ENVI imagery
5.1.2 Show Textfile

Use this function if you need to edit a plain text file which comes together with the data to be processed. The file is opened in a simple editor and may be changed and saved. The function comes handy if an ENVI header needs to be checked or updated (e.g.). Selecting the 'Save' or the 'Save As' function in the submenu will allow to overwrite the file or to create a new one.

![Figure 5.5: Simple text editor to edit plain text ASCII files](image)

5.1.3 Select Input Image

This function allows to select the basis input image (i.e., the band sequential uncorrected image data in ENVI format). It is useful to define the starting point including the default paths of the images for further processing.

5.1.4 Resize Input Image

This tool allows to resize some georeferenced ATCOR input data to new spatial dimensions (without resampling).

The following inputs are necessary:

**Inputs Files:**

- Input Image Name: name of input data file to be resized
- Input DEM Name: name of input elevation data file (this line may be left empty if no DEM is available)

**Dimensions:**

- Upper Left corner: coordinates of upper left corner pixel (ENVI convention: lower left edge of upper left corner pixel is pixel 1/1) - the button "Get UL Position" may be used to get the upper left position of the image subset from a coordinate stored with the 'Push Position' button in an ENVI file display window
• Use Reference File: the coordinates of a reference file may be used
• Number of Pixels: size of output file in pixels in x and y direction (ncols, nrows) starting at upper left corner.
• Output name Basis: name of output main file; auxiliary files are stored according to ATCOR conventions, i.e. *_ele.bsq etc.

Output: Based on the output name (Basis) the auxiliary files are stored according to ATCOR conventions, i.e. *_ele.bsq etc. Possible outputs are:
• at sensor radiance image (ATCOR main input image)
• scan angle file ( *_sca.bsq if available)
• illumination file (*_ilu.bsq if available)
• elevation file (if set)
• side layers of topography: *_slp, *_asp, *_sky.

Figure 5.6: Resize ATCOR input imagery

5.1.5 Rename File Group
Use this function to rename a full group of ATCOR files. ATCOR processing relies on a significant number of files which can be renamed all at once, including the entries in the *.inn file. The such renamed data set is ready for further ATCOR processing.
CHAPTER 5. DESCRIPTION OF MODULES

5.1.6 Import

Some standard formats are supported for importing data layers to an ENVI band sequential format, for efficient use with ATCOR-4:

**Geo-TIFF** Multi-Band geotiff in band ascending order.

**RGBN-Geo-TIFF** Geotiff in 4-band configuration, storage order R-G-B-N (e.g. for photogrammetric data)

**NRGB-Geo-TIFF** Geotiff in 4-band configuration, storage order N-R-G-B

**JPEG2000-Geo** Import of JPEG2000 file (extension .jp2) with embedded georeferencing information in geotiff or gml format.

**ENVI BIP Image** Imports and transforms an ENVI image in BIP (band interleaved by pixel) format to the ATCOR default BSQ (band sequential) format.

**ENVI BIL Image** Imports and transforms an ENVI image in BIL (band interleaved by line) format to the ATCOR default BSQ (band sequential) format.

**Erdas Imagine** Imports an uncompressed image in ERDAS Imagine format to ENVI format (specifically suited for DEM data import).

**AVIRIS** Imports the orthorectified AVIRIS imagery and creates an appropriate DEM and scan angle file for atmospheric correction.

**PARGE Status** Imports the orthorectified PARGE imagery and reads all required information from the status file (*.gcs and ENVI image file header). A correct *.inn file for further processing is created.

![Figure 5.7: Import AVIRIS imagery from JPL standard format.](image)
Figure 5.8: Import PARGE status and meta data information.
5.1.7 Export

Transformation and export of standard ENVI format outputs to:

**ENVI BIP Image** Transforms and Exports an ENVI BSQ (band sequential) image to BIP format.

**ENVI BIL Image** Transforms and Exports an ENVI BSQ (band sequential) image to BIL format.

**Multiband TIFF** creates an universal multiband TIFF image from the input ENVI file.

Use the function 'Display ENVI file' (see section 5.1.1) to export JPG, PNG or TIFF imagery in RGB or grayscale for illustration purposes.

5.1.8 Plot Sensor Response

In the panel *Plot Sensor Response*, one may select the respective sensor response from within the available response functions in the ATCOR installation (or elsewhere). When selecting a response file, the related bands are loaded automatically and the total number of bands is displayed. The band range for display can be adjusted manually afterwards.

Normalization of the curves can be such that the area below the curves is constant (same weight of the functions) or the maximum is at 1 for all curves. The displayed graph may be adjusted in appearance and size and finally being exported to a standard graphics file for further use.

![Figure 5.9: Plotting the explicit sensor response functions](image-url)
5.1.9 Plot Calibration File

When selecting this function, the dialog defaults to the atcor installation for the selection of a *.cal file to be displayed. Both, gain and offset are then plotted in the same graph to get an overview of their relative values.

![Plotting a calibration file](image)

Figure 5.10: Plotting a calibration file

5.1.10 Show System File

This is the same function as 'Show Textfile' but defaults always to the ATCOR installation in order to allow to select an ATCOR system file from within the installation, such as the cal-files, the solar

![Displaying a calibration file](image)

Figure 5.11: Displaying a calibration file (same file as in Fig. 5.10)
reference files, the sensor definition files. The function then allows to adjust and save the respective text contents of the selected file.
5.1.11 Edit Preferences

The default settings of ATCOR may be edited through this panel. The updated preferences are then written to the ASCII file as displayed at the top of the panel. The preferences persist for the user who started the ATCOR application the next time the system is started and also for batch processing.

NOTE: Preferences are set when one of the ATCOR modules has been opened. So, one should select one of the modules from within the menu ‘ATCOR’ before editing the preferences. For a description of the preferences, please refer to Section 9.4.

![Figure 5.12: Panel to edit the ATCOR preferences.](image-url)
5.2 Menu: Sensor

The menu ‘Sensor’ is used to create a new sensor from calibration information, if the sensor is not supported as standard sensor by ATCOR.

![New Sensor Menu](image)

**Figure 5.13: The ‘New Sensor’ Menu**

Fig.5.14 shows the three required steps to include a new (hyperspectral) sensor to ATCOR. The example uses a sensor with 96 spectral bands, denoted as ‘sensor_x96’. Using the function ‘Define Sensor Parameters’, the following steps are done in the background, which also could be done manually: a sub-directory of ‘.../atcor/sensor/’ is created (named ‘x96’) and the three files as displayed in Fig. 5.14 are placed in this sub-directory. After execution of these steps, the new sensor will be automatically detected when ATCOR is started. Details about the sensor definition files are explained in chapter 4.6. Template files of several sensors are included in the distribution. After sensor definition, the module RESLUT (see section 5.2.5) is to be run.

![Sensor Definition Files Diagram](image)

**Figure 5.14: Sensor definition files: the three files on the left have to be provided/created by the user.**

5.2.1 Define Sensor Parameters

This panel is the first step if a new sensor is to be defined. The panel as displayed in Fig. 5.15 allows the below options.
CHAPTER 5. DESCRIPTION OF MODULES

Figure 5.15: Definition of a new sensor

**Sensor Selection**  Select any of the already defined sensors from within ATCOR - a sensor is selected by its sensor definition file (sensor.dat) within a sensor directory. Use the function 'New Sensor' in case your sensor has not yet been defined.

**Inputs**

**Sensor Type**  This is to be selected first, smile sensor and thermal sensors require additional inputs (as displayed in the panel).

**Sensor Total FOV [deg]**  : edge to edge FOV in across track direction in degrees.

**Number of Across Track Pixels:**  : Nominal number of pixels in the unrectified data.

**First, last Reflective Band:**  Band numbers (starting at one) - none: 0

**First, last Mid IR Band:**  Band numbers (starting at one) - none: 0

**First, last Thermal IR Band:**  Band numbers (starting at one) - none: 0
CHAPTER 5. DESCRIPTION OF MODULES

Applied scaling factor: Enter the constant scaling factor which had been applied to the data to fit it to an integer data type from \( \text{mW/(cm}^2\text{sr} \cdot \mu\text{m)} \). Typical values are 100, 500 or 1000. The *.cal file will be created with constant values for all bands according to this value. If the scaling factor is not constant, the *.cal file is to be edited manually. 

NOTE: a scale factor of zero or below inhibits to write a cal-file.

Calibration Pressure: absolute pressure in hPa of instrument during spectral calibration in laboratory.

Instrument Pressure: pressure of instrument during operations
- 'Absolute': pressurized instrument with constant pressure as of this value
- 'Relative': pressure difference to in-flight ambient pressure (in hPa; value of 0 hPa is ambient pressure)

Thermal Sensor: Temperature Band Number: Spectral band used for temperature retrieval algorithm (Optional -according to sensor type)

Smile Sensor: Smile Sensor Response Type, this selection may be required for convolution when sensor definition is not given explicitly: 'Butterworth Order 1 (slow drop-off)', 'Butterworth Order 2 (close to Gauss)', 'Butterworth Order 3 (between Gauss/Rect)', 'Butterworth Order 4 (close to Rectangular)', 'Gaussian', 'Rectangular', 'Triangular', 'Decreasing Binning (from Rectangular to Triangular)', or ' - Arbitrary - (as of *.rsp files) '. The last option is the recommended default for smile correction.

Button: Import Smile File: Use this button to import a smile file *.dat as created by the ATCOR smile detection routine into the sensor definition to the required file smile_poly_ord4.dat.

Actions

New Sensor: A new sensor is created within the ATCOR installation, which results in a new directory in the 'sensor' directory of the installation.

Delete: Allows to delete any sensor directory and all of its contents.

Update Sensor: The sensor parameters of the selected sensor definition are updated according to the settings.

Rename: The current sensor is renamed (both directory and sensor*.dat file)

Outputs: A new sensor*_*.dat file, and possibly 'sensor' directory is created.
ATTENTION: this routine requires write-access to the sensor directory of the ATCOR installation.

5.2.2 Generate Spectral Filter Functions

ATCOR requires the spectral response of all bands of a sensor being present as a spectral filter file (response file) *.rsp . For spectroscopic instruments, the band characteristics are often only available by band center and width at FWHM (Full width half maximum). This function creates the response curves from the latter information (compare Fig. 5.16).
CHAPTER 5. DESCRIPTION OF MODULES

Inputs

**Wavelength File** An ENVI header file (standard .hdr) or a wavelength reference file.

Format of ASCII File: 3 columns, no header; column 1: band number; column 2: center wavelength; column 3: band width. Unit: [nm] or [µm] (same for columns 2 and 3).

Note: if the FWHM is not contained in the ENVI header or in the ASCII file, a bandwidth resolution is assumed which corresponds to 1.2 times the spectral sampling interval.

**Type of Filter Function** The type defines the basic shape of each of the created response curves.

Options are:
- Butterworth Order 1 (slow drop-off)
- Butterworth Order 2 (close to Gauss)
- Butterworth Order 3 (between Gauss/Rect)
- Butterworth Order 4 (close to Rectangular)
- Gaussian
- Rectangular
- Triangular
- Decreasing Binning (from Rectangular to Triangular)

**Spectral Binning Factor** This factor allows binning of channels, e.g., a factor 4 will combine four spectral response functions to produce the equivalent new channel filter function. This is a convenient feature for programmable instruments (e.g., HySpex or Specim AISA). For using this function, a wavelength reference file containing the unbinned spectral response is required as an input.

Outputs

A numbered series of band_*.rsp files are written to the path indicated by the output directory. The files contain wavelength reference and the relative response in two ASCII formatted columns.

5.2.3 Apply Spectral Shift to Sensor

This is a tool to shift the spectral response of a predefined sensor or to change the FWHM values of the sensor based spectral position and FWHM detection results from the ATCOR smile detection routine.

**Inputs**

- input sensor definition file (sensor_*.dat)
- smile file (from smile detection routine) - or FWHM file (smile_poly_ord4.dat or smile_poly_ord4_fwhm.dat)
- New Sensor Name: directory name of new sensor.

**Outputs**

A new sensor definition is created which is shifted against the original sensor by the values given in the smile definition file. The shift is calculated as across track average of the band-wise polynomial. All components of old sensor other than the response files are copied to the new sensor directory. One may use this function for a constant offset by creation of a text file with the number of lines = number of bands and the second column containing the spectral offset to be applied.
5.2.4 BBCALC : Blackbody Function

This routine calculates the blackbody function as described in section 10.1.5, weighted by the spectral response curve of the thermal band used for the temperature retrieval (compare Fig. 5.18).

Inputs
**Spectral response file** Select the *.rsp file of the spectral band in the thermal IR to be used for temperature retrieval.

**Exponential Fit - Limits** The lower and the higher limit of the temperatures for which a fitting function should be created.

**Unit of radiance output** Select the unit either per micron or without normalization.

**Outputs** A file *.hs.bbbfit is created containing the fitting parameters.

5.2.5 **RESLUT : Resample Atm. LUTS from Database**

The monochromatic database of atmospheric LUTs has to be resampled for the specific channel filter functions of each sensor. Details are given in chapters 4.6 to 9.2. Figure 5.19 repeats the panels of the LUT-resampling program RESLUT. The resampling has to be done separately for the reflective and thermal region. Only the required range of flight altitudes and aerosol types should be selected.

*Figure 5.18: Black body function calculation panel*
Figure 5.19: Panels of RESLUT for resampling the atmospheric LUTs.
5.3 Menu: Topographic

The "Topographic" menu contains programs for the calculation of slope/aspect images from a digital elevation model, the skyview factor, and topographic shadow. Furthermore, it supports the smoothing of DEMs and its related layers.

![Topographic modules](Image)

*Figure 5.20: Topographic modules.*
5.3.1 DEM Import

Import Global DEM

Import Global (Lat/Lon) DEM from SRTM data to UTM WGS-84 coordinates. The approx. 200m (0.5 arc min) resolution global DEM data is available for download through the help menu.

Inputs

Longitude range: range in longitude direction; pixel edges - in decimal degrees only!

Latitude range: range in latitude direction; pixel edges [deg]

Output Resolution: output pixel size (default: 200m)

Output DEM: name of DEM (*.ele.bsq) to be written.

Restrictions: The DEM named DEM_SRTM30_WORLD.DEM has to be available in the demo_data directory of the ATCOR installation.

Output: converted DEM as ENVI file in UTM coordinates.

---

![Import Global DEM](image)

*Figure 5.21: Import DEM from global elevation data (SRTM).*

Import Geo TIFF

This function import a Geo Tiff file to an ENVI formatted elevation data file.

Import ARC GRID

This Procedure reads a standard ARC GRID digital elevation model.
CHAPTER 5. DESCRIPTION OF MODULES

Inputs:
**Filename:** Name of DEM to be read (usually extension *.asc)

**Default output value for found missing data:** Value to be written to the pixels as defined by the NODATA keyword in the GRID file - if output value is less than zero, the mean of the read data is put to the missing pixels

Actions:
**View Parameters** Shows the header information of the GRID file.

**Read DEM** Reads the DEM based on the given definition and stores it to a single channel ENVI formatted file (if required).

Output: converted DEM as ENVI file in arbitrary coordinates.

![Read ARC GRID Digital Elevation Model](image)

**Figure 5.22:** Import DEM from ARC GRID ASCII.

5.3.2 DEM Preparation

The purpose of this program is to provide a tool to resize a DEM to an image cube's dimensions. This menu function as depicted in Fig. 5.23 allows to calculate both layers in one step.

![Prepare and Resize a DEM](image)

**Figure 5.23:** DEM Preparation
CHAPTER 5. DESCRIPTION OF MODULES

Inputs:

**Input DEM file** Input DEM (typically larger than image).

**Input Image file** This is the target image (georeferenced). The DEM is automatically resized to its dimensions.

**Window Diameter for DEM Processor** Size of the kernel in number of pixels; the slope and aspect and sky view side outputs are calculated using this kernel size and the output DEM is smoothed by this kernel size.

**Options** The writing of side outputs (i.e. _slp and _asp and _sky may be triggered by deselecting either of the first two options. The option "Fill in Missing Background Values" is used if the DEM is incomplete, the missing parts are filled with an average value from the DEM.

Outputs: An ATCOR elevation file _ele.bsq is written of the same size and location as the reference Image. Optionally, files of slope and aspect and sky view are created, same size as DEM. Attention: Both the DEM and the image should be represented in the same coordinate system and cover the same area. If the DEM is too small, the remaining parts of the image are filled with a zero value.

Note: this routine is only meant for rectified imagery. If working in raw geometry data format, the PARGE routine for inversion of DEM related layers to raw geometry is required.

5.3.3 Slope/Aspect

Slope and aspect are to be calculated from the DEM before ATCOR is run for rugged terrain. This menu function as depicted in Fig. 5.24 allows to calculate both layers in one step.

![Figure 5.24: Slope/Aspect Calculation panel](image-url)
CHAPTER 5. DESCRIPTION OF MODULES

Inputs

**Input DEM file** The standard DEM file used for atmospheric correction. This DEM should be in meters.

**Output file names** The names for the output files are entered automatically and can’t be changed as ATCOR asks these files to be named exactly according to the conventions.

**Kernel Size Box** Size of the kernel in number of pixels; the slope and aspect is calculated as gradient of the pixels at the edges of this box - the default value is 3 pixels (i.e. direct neighbors of center pixel).

**DEM resolution** This is the pixel size, a default of 30m is assumed. This needs to be entered manually.

**DEM height unit** The unit of the values in the DEM, usually a DEM is stored in meters, but sometimes an integer DEM is stored as dm or cm data in order to preserve disk space.

Outputs The two files of slope and aspect are created, same size as DEM, integer data.

5.3.4 Skyview Factor

The sky view factor of a DEM is calculated with a ray tracing program and ranges from $v_{sky} = 0$ to 1, with 1 indicating a full hemispherical view. Data in the sky view file are scaled from 0 to 100% and coded as byte. The sky view factor determines the fraction of the hemispherical diffuse sky flux and $1-v_{sky}(x,y)$ determines the fraction of radiation reflected from surrounding mountains onto the considered pixel, see chapter 10.1.1. This program is also available in the batch mode, see chapter 6.3.

Input parameters besides the DEM file are:

- DEM horizontal resolution in meters: the x and y resolution must be the same.
- DEM height unit: supported units are m, dm, and cm.
- Angular resolution (degrees) in azimuth and elevation.
- The undersampling factor of the DEM in pixels. For large DEM’s the skyview processing may be very time consuming unless an undersampling is chosen here.

Figure 5.25 shows the GUI panel, and figure 5.26 presents a skyview image derived from a DEM image. An angular azimuth / elevation resolution of 10 degrees / 5 degrees is recommended. For large images it causes a high execution time which can be reduced by selecting an undersampling factor of 3 pixels. A high angular resolution is more important than a low undersampling factor.

5.3.5 Cast Shadow Mask

The calculation of the cast shadow map is done by ATCOR after reading the DEM files. If the shadow map is computed “on-the-fly” it is kept in memory, and it is not stored as a separate file. If the user wants to inspect the DEM shadow map the program ”shadow” has to be started before running ATCOR. The program accepts float values of the solar zenith and azimuth angles. The output file name of the DEM cast shadow map includes the zenith and azimuth angles rounded to integer values.
CHAPTER 5. DESCRIPTION OF MODULES

The DEM cast shadow map is a binary file where shadow pixels are coded with 0 and sunlit pixels with 1. It includes self shadowing and cast shadow effects. Self shadowing consists of pixels oriented away from the sun with slopes steeper than the solar elevation angle. The cast shadow calculation is based on a ray tracing algorithm and includes shadow regions caused by higher surrounding mountains. Figure 5.27 shows the GUI panel.

5.3.6 Image Based Shadows

This routine detects the fractional shadows in the image using a multi-index based approach. A floating point illumination file is saved to be used as input for atmospheric correction. Figure 5.28 shows the GUI panel. NOTE: this function is found in the menu 'Filter' for ATCOR versions without support for terrain correction.

**Input Files:**

- **infile**: file to be analysed, requires at least 4 bands (Blue, Green, Red, NIR)
- **calfile**: calibration file for current input file (usually found in the sensor or cal directory of the ATCOR installation).
- **e0_solar**: solar irradiance file for current input, this file is also situated in the sensor or cal directory
- **outfile**: output file of the processing

---

**Figure 5.25: Panel of SKYVIEW.**

**Figure 5.27** shows the GUI panel.
Figure 5.26: Example of a DEM (left) with the corresponding sky view image (right).

Figure 5.27: Panel of Cast Shadow Mask Calculation (SHADOW).

Options:

- Include Terrain Illumination - calculates the illumination based on slope and aspect files and the solar illumination angles as defined below
- Calculate Skyview Estimate - calculates an estimate of the local skyview fraction on the basis of a linear model.
- Apply Shade Pixel Filter - filters single pixels within full cast shadow areas
- Write all Side Layers - creates a file containing all indices calculated during the processing in 5-7 layers. This function may be useful to find the appropriate range limits for a sensor system or imaging situation
- Limits of Index: - The lower and upper boundary can be set in order to derive a continuous fractional shadow from index. Larger values will result in large fractions of shadows being detected. The range can not exceed 0 to 2 in any case (default values are 0.5 to 1)

Parameters used for terrain illumination and skyview estimate:
• Slope File Name: Name of input Slope file to be used for illumination calculation (the corresponding _asp.bsq file needs also to be present)

• Solar Zenith/Azimuth: solar angles (zenith/azimuth) for this data set and optionally the flight altitude above ground (airborne ATCOR version only)

A combined index is created based on NIR brightness, and two red/blue and green/blue indices which results in a scaled number between 0 for full cast shadow and 1 for full illumination which is proportional to the cast shadow.

5.3.7 DEM Smoothing

Smooth a DEM (or any other single-band image) in order to remove artifacts in the atmospherically corrected imagery. All related DEM-layers are automatically smoothed as well (e.g. slope / aspect / skyview). Alternatively, this task could be done with any image processing software.

Inputs:

Input DEM/File Name Usually a DEM _* ele.bsq is selected here; but any other single band ENVI image or the _* _ilu.bsq file is also accepted. The routine searches automatically for related files (i.e. _*sky, _*slp, and/or _*asp) and smoothes them with the same parameters.

Diameter of DEM Filter: Size of filter box in pixels (diameter)

Output Name: Name of Elevation file output (auxiliary layer names will be derived from that)
Outputs: ENVI file(s) smoothed or filtered by the given factor and method.

Actions:

Smooth: uses the standard smoothing (i.e., lowpass-filter) in the spatial domain.

Median: uses a median filter for data correction, e.g., to remove noise or outliers from the DEM.

ATTENTION: The *.ilu file is not smoothed automatically by this routine. If the ilu has already been calculated before, it should be either removed or be smoothed separately.

\[ f = \begin{cases} 1 - (1 - \cos \Theta_S / \cos \beta)w & \text{if } \lambda < 1.1 \mu m \\ \cos \Theta_S / \cos \beta & \text{if } \lambda \geq 1.1 \mu m \end{cases} \] (5.1)

\[ f = \cos \Theta_S / \cos \beta \] (5.2)

\( \Theta_S \) is the solar zenith angle of the scene. For \( \lambda \geq 1.1 \mu m \) the diffuse flux is neglected, i.e., \( w = 1 \).

The method was compared with the standard Minnaert correction (eq. 5.4) and was superior in most cases. Figure 5.30 shows the GUI panel.

\[ f = \left( \frac{\cos \Theta_S}{\cos \beta} \right)^{\log(\cos \Theta_S / \cos \beta)} \] (5.4)
**Figure 5.30:** Topographic correction only, no atmospheric correction.
5.4 Menu: ATCOR

The menu ‘ATCOR’ contains the main processing modules of ATCOR, i.e., the panels for ATCOR4 for flat and rugged terrain.

![Figure 5.31: The ‘Atm. Correction’ Menu](image)

5.4.1 Haze Removal

The atmospheric haze in imagery may be removed in a statistical procedure before going into atmospheric correction routines. Figure 5.32 shows the parameter settings which can be chosen for this procedure.
Figure 5.32: ATCOR haze removal module.
5.4.2 The ATCOR main panel

Figure 5.33 (top) shows the input parameters required for ATCOR. The lower part of the panel contains buttons for selecting SPECTRA, determining the aerosol type, employing in-flight radiometric CALIBRATION, and starting the image processing. The processing options are shown in the separate panel as described in section 5.4.10. The trivial panels, e.g., band selection, spatial subimage etc. will not be shown here. The panels should be filled or clicked in the top-down direction. The message widget at the bottom will display hints, warnings, or errors.

<table>
<thead>
<tr>
<th>Input Image File</th>
<th>Output Image File</th>
</tr>
</thead>
<tbody>
<tr>
<td>[Image]</td>
<td>[Image]</td>
</tr>
</tbody>
</table>

**Figure 5.33: ATCOR panel.**

Options that are not allowed for a specific sensor will appear insensitive. If the haze removal option is selected in combination with "Variable Visibility" the visibility index (proportional to total optical thickness) map is coded with the values 0-182. The value visindex=0 corresponds to visibility 190 km, each integer step of 1 corresponds to an AOT increase of 0.006. The array serves as a fast method of addressing the radiative transfer quantities (transmittance, path radiance etc) in case of a spatially varying visibility, i.e., in combination with the DDV algorithm. IF the "Haze or Sunglint Removal" button is selected the next panel will ask for haze removal over land (option 1), haze or sunglint removal over water (option 2), or haze removal over land and water (option 3).

In case of the rugged terrain version of ATCOR the panel for the DEM files has to be specified in addition (Figure 5.34). It pops up after the input file has been specified. A quick quality check is performed on the DEM files. The solar illumination file is calculated and if its standard deviation is large the panel of Figure 5.35 pops up with a warning. In this case the DEM elevation file (and the derived files of DEM slope, aspect etc) probably have a lot of large steps. The DEM resolution is often not appropriate for high spatial resolution imagery, and integer coded DEM’s might have
to be resampled and stored as float data. Appropriate action for resampling or low pass filtering is recommended in these cases, see the tips in chapter 9.6. Figure 5.36 shows an example in terms of the DEM illumination. The top image is obtained after low pass filtering the original elevation file, the central image is the illumination based on the original DEM, and the bottom shows a 100 pixel transsect of the original elevation data revealing the steps. The original DEM had a resolution of 30 m, was coded as 16 bit integer, and initially resampled to the 6 m pixel size of the image with integer arithmetic. After reprocessing the elevation file the other DEM derived files should also be reprocessed.

![Figure 5.34: Panel for DEM files.](image)

The pixel size of the DEM files must be the same as the image pixel size specified on the main panel, see figure 5.33. The physical units of pixel size (m) and adjacency range (km) are also used to calculate the equivalent number of pixels needed to cover the adjacency range.

### 5.4.3 ATCOR4f: flat terrain

The menus for ATCOR-4 share the same functionalities as described above in both flat and rugged terrain options. The ATCOR4f variant is recommended to speed up processing time and for fast
checks as hyperspectral image processing may be very time consuming in rugged terrain.

5.4.4 ATCOR4r: rugged terrain

This routine is to be taken if highest accuracy is required in terrain for imaging spectroscopy instruments. The functionality is analogous as described for the other panels.
5.4.5 SPECTRA module

The SPECTRA module (see figure 5.37) serves to extract spectra of different targets of the scene as a function of the visibility. These spectra can be compared to field spectra or library spectra to estimate the visibility. Scene-derived spectra also may indicate calibration errors in certain channels. In that case, a copy of the sensor calibration file can be edited to match the retrieved reflectance with the field / library spectrum or the inflight calibration module may be employed (see chapter 5.4.8).

In most cases it is useful to check some scene-derived target spectra, e.g. water, vegetation, or soils, before starting the processing of the image cube at the risk of "blind" processing and obtaining wrong results.

To obtain a target spectrum of the scene click at any position in the image. In figure 5.37 the solid white-line spectrum at the top shows a meadow signature, the green line represents a reference spectrum taken from the spec_lib directory, already resampled for the HyMap sensor. A reference spectrum can be loaded when clicking the corresponding button at the top right of the panel. An exact match of scene spectra and library spectra cannot be expected when they are measured at different times and locations, but the trends (spectral shapes) should be similar. The bottom
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graphics shows an asphalt spectrum taken from the aircraft runway in the lower part of the image.

The parameter Visibility can be set in the corresponding widget near the center of the panel and its influence on the retrieved spectrum will be shown when placing the target box in the scene. For hyperspectral sensors, the water vapor column corresponding to the last spectrum is given in the "message" box (upper left of panel). In addition, there is the choice of selecting the global neighborhood (average over whole image) or the local neighborhood (adjacency range as specified on the main panel, see figure 5.33) for the correction of the adjacency effect. The "global" option is faster, because it has to be calculated only once per channel. The "local" option is more accurate and slower because it is always updated, but depending on the scene content, the difference between both options may be small.

The button "Save last spectrum" (upper right corner of figure 5.37) can be used to save the selected surface reflectance spectrum. A sequence of target DN spectra can also be generated here that is required as input to the spectral calibration module (see chapters 2.2, 5.8.4). Up to 9 targets can be defined to be used in the spectral calibration. They have to be labeled consecutively, e.g. "target1", "target2", etc. These output file names (without file name extension) have to be specified after pressing the button "Save last spectrum". For each target name three files will be generated, e.g., "target1.dat" (surface reflectance spectrum), "target1.txt" (a description file), and "target1_dn1.dat" (the DN spectrum). The sequence of "target*_dn*.dat" files is used in the spectral calibration module.

5.4.6 Aerosol Type

The aerosol type is a parameter which is fixed for atmospheric correction. This routine searches automatically for the best suited aerosol type for the currently selected image. This type can then be used when selecting the atmospheric file.

5.4.7 Visibility Estimate

ATCOR uses the Dark Dense Vegetation (DDV) approach to calculate the best visibility for an image. This button allows to calculate the visibility estimate for the whole image without going into the image processing. Note, that for the 'Variable Visibility' option, the visibility map will be calculated anyway from the image during the image processing.

5.4.8 Inflight radiometric calibration module

This chapter presents the GUI for the inflight calibration, which may also be invoked from one of the four possible ATCOR main panels. The purpose is the calculation of the radiometric calibration coefficients for spectral bands in the solar region based on measured ground reflectance spectra. The user should be familiar with chapter 2.4 before using this function.

Note: a ground reflectance spectrum from a field spectrometer has to be resampled with the channel filter curves of the selected sensor. The field spectrometer file format should be converted into a simple ASCII file containing the wavelength (nm or \(\mu m\)) in the first column and reflectance in the second column. The resampling can then be done with the sequence "Filter", "Resample a Spectrum" from the ATCOR main panel. The result is an ASCII file with 2 columns: the first contains the channel center wavelength (the nm and \(\mu m\) unit is allowed), the second contains the resampled reflectance value (either in the 0 - 1 or 0 - 100% range).
If "target1" is the name of the target, ATCOR provides three ASCII files with information on target / background properties and the derived calibration file. These are the output of the "c1" option of ATCOR’s calibration module,

- File "target1.adj" contains the original target $DN_1$, the adjacency corrected $DN_1^*$, and the ground reflectance data for each band.

- File "target1.rdn" (radiance versus digital number) contains the band center wavelength, target radiance $L_1$ and corresponding digital number. This file can be used as input to a regression program "cal_regress" that allows the calculation of the calibration coefficients with a least squares fit in case of multiple targets (more than two).

- File "target1.cal" contains three columns: band center wavelength, offset or bias set to zero ($c_0 = 0$), and $c_1$ according to equation (2.22).

Remark: a bright target should be used here, because for a dark target any error in the ground reflectance data will have a large impact on the accuracy of $c_1$.

![Figure 5.38: Radiometric calibration: target specification panel.](image)

At the top line of the menu of figure 5.38 the mode of calibration is specified: one or two targets can be employed by selecting the button ‘c1’ or ‘c0 & c1’, respectively. If a calibration is intended for $n > 2$ targets, each target has to be specified separately in the ‘c1’ mode, which creates a file ‘target_i.rdn’ (i=1, 2, ..., n) with the name ‘target_i’ specified by the user. These files contain the radiance and corresponding digital number spectrum as a 2-column ASCII table, and they are employed in the batch module cal_regress (see chapter 5) to calculate a least squares regression for the calibration coefficients. Next the target box size and the corresponding ground reflectance
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file have to be specified. The button for the file name of target 2 is insensitive, because the single target mode was selected here. Then the file name for the calibration results should be specified. The default name is "test.cal". However, it is recommended to include the name of the ground target here. Now the target(s) can be clicked in the zoom window that pops up automatically. Target 1 has to be clicked with mouse button 1 (mb1, left), target 2 with mouse button 2 (mb2, center). The zoom window is moved in the main window by pressing mb1 (for target 1) and mb2 (for target 2). Alternatively, the target coordinates (x / y = column / line) can be specified.

In addition to the file "xxx.cal" the files "xxx.rdn" (radiance versus digital number), and "xxx.adj" (original and adjacency corrected DN’s) are automatically created.

![Figure 5.39: Radiometric CALIBRATION module.](image)

The appearance of the inflight calibration module is similar to the SPECTRA module. In the left part the image is loaded. A zoom window can be created and two contrast stretching options (Gaussian and histogram equalization) are available. In the right part two windows are provided to display the c1 spectrum and the box-averaged target DN spectrum. The ymin / ymax widgets allow the user to scale the graphical display. The parameters visibility and adjacency range can be varied and their influence on the calibration display can be studied.
5.4.9 Shadow removal panels

The interactive session of the de-shadowing method enables the setting of three parameters that influence the results, compare Figures 5.40, 5.41:

1. a threshold $\Phi_T$ for the unscaled shadow function $\Phi = \Phi_{\text{Uni}}$ to define the core size of the shadow regions, see chapter 10.5.6 for details.

2. the maximum range $\Phi_{\text{max}}$ for re-scaling the unscaled shadow function $\Phi = \Phi_{\text{Uni}}$ into the $(0,1)$ interval of the scaled shadow function.

3. the last parameter sets the minimum value of the scaled shadow function $\Phi^* = \Phi_{\text{S}}$, typically in the range $\Phi_{\text{S}} = 0.02 - 0.10$, i.e., the darkest shadow pixels of the scene are treated as being illuminated with a fraction $\Phi_{\text{S}}$ of the direct solar irradiance.

$\Phi_{\text{min}}$ $\Phi_{\text{max}}$ $\Phi_{\text{uni}}$

The first two parameters are most important. The last parameter is not very critical, and a default value in the range $\Phi_{\text{S}} = 0.04 - 0.08$ covers most cases of interest.

The linear type of re-scaling of $\Phi_{\text{uni}}$ is recommended. The "exponential" option exaggerates the de-shadowing results, but it can be used to enhance the trends for a quick visual inspection.

When pressing the button "Run Interactive De-Shadowing", a reduced size image of the original scene, the shadow mask, and the de-shadowed image will pop up with the histogram of $\Phi_{\text{uni}}$ containing the threshold for core areas (point 1 of Fig. 5.41), the range of re-scaling of $\Phi_{\text{uni}}$ (point 2), and the current value for $\Phi_{\text{S}}$ (point 3). Now these three parameters can be modified, and the results will again be shown as the corresponding quicklook image and histogram. When good results have been obtained the parameters can be saved and the final image processing can take place.

Figure 5.42 presents an example with two iterations. Results for iteration 1 contain too many shadow pixels (black areas in the central section of the image), therefore the threshold $= -0.15$ was decreased to $-0.38$ (parameter 1 of Fig. 5.41). After de-shadowing, most areas in the shadow mask were overcorrected in iteration 1, therefore the maximum range $= 0.40$ was decreased to 0.19 (parameter 2 of Fig. 5.41, see diagonal line from lower left to upper right in histogram of Fig. 5.42). The shadow mask for iteration 2 is appropriate, and no overcorrection effects can be observed.
Figure 5.41: Panel to define the parameters for interactive de-shadowing.

Note: when leaving the panel of Fig. 5.41 it is possible to edit the cloud shadow map before continuing the scene processing, using any available image processing software. Then the edited map is employed for the processing. This provides some flexibility because it is difficult to calculate a satisfactory shadow map in all cases.
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Figure 5.42: Quicklook of de-shadowing results.
Top left: histogram of PhiU (threshold=-0.15, range=0.40), iteration 1. Top right: histogram of PhiU (threshold= -0.38, range=0.19), iteration 2. Center: results for iteration 1, left-to-right: original, shadow mask, de-shadowed image. Bottom: results for iteration 2.
5.4.10 Panels for Image Processing

When pressing the button "IMAGE PROCESSING" in one of the main panel (figure 5.33) some additional panels will pop up. First, the processing options are to be selected (see figure 5.43).

Options that are not allowed for a specific sensor will appear insensitive. If the haze removal option is selected in combination with "Variable Visibility" the visibility index (proportional to total optical thickness) map is coded with the values 0-182. The value visindex=0 corresponds to visibility 190 km, each integer step of 1 corresponds to an AOT increase of 0.006. The array serves as a fast method of addressing the radiative transfer quantities (transmittance, path radiance etc) in case of a spatially varying visibility, i.e., in combination with the DDV algorithm. IF the "Haze or Sunglint Removal" button is selected the next panel will ask for haze removal over land (option 1), haze or sunglint removal over water (option 2), or haze removal over land and water (option 3).

In case of thermal bands, an emissivity selection panel will appear:

The first two options are available for instruments with only a single thermal band, the NEM and ISAC options require multiple thermal bands and are not shown if there are less than 5 thermal bands. The surface classes (water, vegetation, soil etc) are calculated on-the-fly employing the surface reflectance spectra from the reflective bands.

Figure 5.45 shows the panel with the two options for haze over land processing as explained in chapter 10.5.3.

The panel of figure 5.46 pops up when the spatially varying visibility option was selected and if the sensor has a 2.2 µm (or 1.6 µm) band (or at least a red and NIR band) required for the automatic masking of the dark reference pixels, compare chapter 10.4.2.
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Figure 5.44: Emissivity selection panel.

Figure 5.45: Options for haze processing.

Figure 5.46: Reflectance ratio panel for dark reference pixels.
The panel of figure 5.47 pops up for rugged terrain and contains the input parameters to incidence BRDF compensation due to terrain variations as discussed in chapter 2.6.

![Incidence BRDF compensation panel.](image)

Figures 5.48 and 5.49 are associated with the value added products as described in chapter 7. This value added file contains up to 10 channels if the sensor has thermal bands. In case of a flat terrain, the air temperature has to be specified. For a rugged terrain, air temperature at some base elevation and the air temperature gradient as well as water vapor have to be defined.

If the value added option is selected another panel pops up (Figure 5.50). It contains parameters for the leaf area index (LAI model) and FPAR model as described in chapter 7.

Finally, a job status window indicates the processing progress.

**Note:**
The job status window of ATCOR shows the percentage of processed image data and the estimated remaining time. The time estimate is based on the processing time for the current band. The time per band increases for channels in atmospheric water vapor regions, it decreases in regions where interpolation is applied, e.g., around 1400 nm. However, the time also depends on other factors such as the overall CPU load in case of multi-user machines, or the traffic on net-worked machines. Accordingly, the estimate for the remaining time is not always continuously decreasing, but may increase sometimes.
**Figure 5.48:** Value added panel for a flat terrain.

**Figure 5.49:** Value added panel for a rugged terrain.
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Figure 5.50: LAI / FPAR panel

Figure 5.51: Job status window.
5.4.11 Start ATCOR Process (Tiled / from *.inn)

This is a way to start a tiled process of ATCOR from within the ATCOR GUI (instead of the standard batch-based process ‘atcor_tile’).

The process requires that an *.inn file has been created before by going through the atcor GUI or by editing a respective ASCII file manually or by using the routine write_atcor_inn_file.pro provided in the directory docu of the ATCOR installation.

The below parameters are to be entered for processing:

- Input file name: name of image data cube to be processed. The file must be accompanied by a valid *.inn file for processing.
- Name of output cube to be created.
- Number of tiles in X and Y dimensions - the total number of tiles to process is then X x Y tiles.
- ATCOR method: selection between flat processing and rugged processing - for the latter the DEM has to be prepared and ready.

After starting the process using the ‘run’ button, the messages are printed to the prompt (or the console) if available. Error and status messages may also be found in the *.log file during and after processing.

![Figure 5.52: ATCOR Tiled Processing](image)
5.5 Menu: BRDF

The "BRDF" menu provides access to the simple nadir normalization method and the more advanced BREFCOR BRDF correction.

![Image](image.png)

*Figure 5.53: BRDF top Menu.*

5.5.1 BREFCOR Correction

This module calculates an observer BRDF correction using a model based approach. (see chapters 2.6, 10.6.3). Figure 5.54 shows the corresponding GUI panel.

The BREFCOR is implemented for the airborne version of ATCOR for sensors with a large FOV. It works best if a number of scenes can be taken for model calibration, but it may also lead to good results for single scenes as long as the statistics are uniform within scene.

The Ross Thick-Li-Sparse reciprocal BRDF model kernels are used for correction of the imagery. The various viewing angles of the individual images provided are used as reference points to calibrate the kernel weighting factors and the isotropic component of the BRDF models. This model requires a sufficient number of images for good calibration as a variety of incidence and viewing angles are required.

The following inputs are necessary:

**Inputs Files:** A list of atmospherically corrected input file is to be compiled (*_atm.bsq). The ATCOR *.inn file is used for the meta data information. It should be named according to the ATCOR convention and situated at the same place as the *_atm file. Additionally, the viewing angles are read from the scan angle file, which should be named *_sca.bsq. Files are added using the 'Add File(s)' Button on top, while entries may be removed by the 'Remove Entry' Button. Further files considered are:

- .ilu file is used to exclude shadow pixels (if floating point ilu file is available according to ATCOR conventions)
- .hch file is used to exclude hazy/cloudy pixels from calculation.

**Model Options:**

- Roujean Geometric Kernel: the Roujean Kernel is used for the geometric part of the model (instead of the Li Sparse model).
• Maignan Hot Spot Geometry: use the improved hot spot geometry for the volumetric kernel as proposed by Maignan (RSE, 2003).

• Spectral Smoothing factor (number of bands): Smooth the weighting functions in the spectral dimension - this option is useful for hyperspectral instruments only to avoid spectral artifacts. A spectral smoothing in a range of 20-30nm is recommended.

• Leave water ‘as is’: The first level (water) is not touched by the BRDF correction, i.e. the kernel weights are set to zero for both geometric and volumetric kernel; this may be useful because water can not be easily modeled by the Ross-Li formulation.

• Write ANIF outputs: By default, the corrected image is written. Use this option to get the side outputs; i.e. the files '_anif' (anisotropy map) and the '_bci' which is the BRDF correction index used to discriminate the BRDF levels.

Calibration Granularity: For calibration of the model, the image is stratified in a number of BRDF levels. The number of pre-defined levels can be selected; calibration on a bigger number of levels leads to better accuracy of the correction, while it requires good image statistics. A smaller number of levels leads to higher stability of the results.
The stratification thresholds for the calibration may be tuned manually by the 'Self Defined' option - use the define button to adjust the levels. Please check the BCI image to find suited calibration limits.

**Reflectance scale factor:** This is a constant factor which is to be applied to the input image (which has to be directional reflectance HDRF) to convert the image DNs to absolute reflectance values (between 0 and 1), i.e. if the reflectance in [\%] is scaled by 100 to store the data, the factor is 10'000.

**Fitting accuracy threshold [\%]:** Limit to fitting accuracy to make an across track fitting function accepted. Increasing the limit will lead to more correction functions. For large FOV instruments, the threshold may be increased to up to 30\% in certain cases.

**Output Definition:** The directory for all outputs, and the file name appendix can be defined here. If the directory is not given, the outputs are stored together with the input imagery using the given extension. The Ext field is the extension which will be added to all input files to store the output of the calculation. The BRDF model file name can be defined or selected here. The latter applies if the option 'Use precalculated Model' is selected above. It will be stored in the directory defined above.

**Band Numbers:** By default all bands are corrected. For hyperspectral imagery, it may be useful to check the performance on an RGB sample first.

The button 'Plot Weights' allows to show the found kernel factors in relation to the BCI.

**Actions:**
- Calibrate Model: the code calibration file is created without applying a correction to the images.
- Do Image Correction: the Model based BRDF correction is performed. If the selected model file is available, it is used for the correction.

### 5.5.2 Nadir normalization (Wide FOV Imagery)

This module (NADIR,REFL) performs an empirical BRDF correction by normalizing the across-track radiance gradients to the nadir brightness value (see chapters 2.6, 10.6.1). Figure 5.55 shows the corresponding GUI panel.

The statistical nadir normalization works best for wide FOV airborne data and requires a minimum field-of-view of 20°.

### 5.5.3 BRDF Model Analysis

This tool is to analyze the model parameters which have been derived by best-fit approach from the imagery. An example is shown in Fig. 5.56. The top section shows the meta parameters of the currently selected model file (i.e., the output from the model calibration procedure).
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5.5.4 BRDF Model Plot

This tool is to plot a BRDF model as it has been used for BREFCOR correction. An example is shown in Fig. 5.58.

Sliders and Inputs:

Select: This lets you select a model file (IDL save file) which has been created by the BREFCOR model calibration procedure.

Input File Slider: If calibration has been done on multiple files at once, the files may be selected here. Setting this slider to '0' displays the averaged results from all files.

Spectral Band/BCI Levels Slider: Lets you select the respective band number and BCI level number.
**Figure 5.56: BRDF model analysis panel.**

**BRF Range:** Defines the scaling of the graphics in absolute BRF values or in ANIF values (if selected in the model Options below). The button 'AutoRange' will reset to the min/max of the available data.

**Max Zenith:** Maximum sensor zenith angle to be displayed (default: 45 degrees off nadir)

**rho_iso/f_geo/f_vol:** Factors of 3-parameter BRF-model to be plotted (these factors may be edited for test purposes; on return, the plot is updated)

**Solar Zenith:** Solar zenith angle for the plot in degrees.

**Options:** Allows to select variations of the BRDF model and to plot Anisotropy instead of BRF values.

The currently displayed graphics can be exported to an EPS file using the 'Export Graphics' button.
Figure 5.57: BRDF model fitting analysis panel.
Figure 5.58: BRDF model plot.
5.5.5 Mosaicking

Mosaic a number of georectified scenes into one (in an efficient way...). Figure 5.59 shows the parameter settings which can be chosen for this procedure.

Inputs

Input Files (first file on top): list of files to be mosaicked. The files are stacked in the order of appearance (i.e. the first file in the list is on top of the mosaicked output)

Buttons:

Add File(s): adds one or more new files
Remove Entry: removes the selected file(s)

emphMove: moves the file one position up (or rotates if already at top)

Range: range in x and y direction to edge of pixels of the mosaicked product should be entered. The coordinates refer to the pixel borders (edges) on either side of the image.

Button From Files: reads the maximum extend of all selected files as of the list.

Pixel Size: Size of output pixels in meters

Note: the mosaicking uses a bilinear interpolation; no aggregation is done.

Cut Options: treatment of image borders in overlap area

Edge Overlay: The mosaicking is such that the first file is in the list is strictly on top Center Cropped: While mosaicking, the routine tries to find the middle of the overlap area between the new image and all the images mosaicked so far as a cut line.

number of bands to process Button Select Bands: lets you select the bands to mosaic - the number of currently selected bands is displayed beneath the button

Output File Name: name of output file to be written.

Actions

Preview: the first of the selected bands is mosaicked at a resolution reduced by a factor of 2 and the result is displayed.

Run Process: The mosaicking is performed.

Restrictions

This routines requires georeferenced data with:
- same coordinate system
- no rotation in ENVI header
- background coded with 0
- all files should have the same number of bands for mosaicing

The input resolutions of the imagery may vary.
**Figure 5.59:** Mosaicking Tool.
5.6 Menu: Filter

The "Filter" menu provides spectral filtering of single spectra (reflectance, emissivity, radiance) provided as ASCII files, spectral filtering of image cubes, and spectral polishing.

5.6.1 Resample a Spectrum

This program serves for the general purpose of resampling. It requires an ASCII file with two columns as input. The first column is wavelength (nm or µm unit), the second is reflectance, or emissivity, or something else, e.g. spectral radiance. The reflectance range can be 0 - 1 (the intrinsic reflectance unit) or the percent range (0 - 100). Figure 5.61 shows the graphical user interface. The input spectrum is high spectral resolution data. After specifying the first band of the sensor the resampling is performed for all spectral bands, and the result is written to an output file, again with two columns (wavelength and resampled spectrum).

5.6.2 Low pass filter a Spectrum

This program serves to prepare target reference spectra for IFCALI inflight calibration. Unwanted spectral noise is removed by lowpass filter (i.e., spectral smoothing). It requires an ASCII file with
two columns as input. The first column is wavelength (nm or µm unit), the second is reflectance, or emissivity, or something else, e.g. spectral radiance. The reflectance range can be 0 - 1 (the intrinsic reflectance unit) or the percent range (0 - 100). Figure 5.62 shows the graphical user interface. The input spectrum is high spectral resolution data. The result has the same format and dimensions as the input file.

Figure 5.62: Low pass filtering of a (reflectance) spectrum.

5.6.3 Spectral Polishing: Statistical Filter

Remove spectral artifacts in high spectral resolution imaging spectroscopy data.

Inputs:

Input file name A hyperspectral image cube, usually the output of atmospheric correction *_atm.bsq

Sensor Spectral Response Defines the first band of the sensor response *rsp. This field may be left empty - in that case, the wavelength information from the ENVI header is used if the wavelengths tag is properly given. If no wavelength reference is there, a spectrally equidistant succession of the spectral bands is assumed.

Note: the Savitzky-Golay filter is not wavelength-aware and uses always the assumption of equidistant, constantly increasing bands.

Number of polishing bands on each side Adjacent bands to be used for calculation on each side of the target band; e.g. factor 3 uses 7 bands for polishing (3 on each side plus central band).

Smoothing Factor smoothing applied stand alone or in combination with the derivative filter:

0: no smoothing
1: slight smoothing (filter: 1/4/1)
2: moderate smoothing (filter: 1/2/1)
3: standard smoothing (filter: 1/1/1)
4 and more: standard smoothing with moving average

Polishing Filter Type Four options are available for statistical spectral polishing: Derivative Filter: all spectral bands of the given window size are taken into account to calculate derivative used to reconstruct the value of the center band.

Neighbour Derivative: all spectral bands except for the center itself are taken into account to
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calculate derivative used to reconstruct the value of the center band.

Lowpass Filter: Only the smoothing is performed, no derivatives are calculated.

Savitzky-Golay: Filter to perform a numerical polynomial fit of 4th degree through the selected total window size.

Output:
A cube containing the spectrally filtered copy of the original image data cube is generated (compare Paper Earsel, SIG-IS Workshop, Edinburgh, 2011).

![Figure 5.63: Statistical spectral polishing.](image)

5.6.4 Spectral Polishing: Radiometric Variation

A module that was originally developed for the airborne version of ATCOR is the spectral polishing. The algorithm is only intended for hyperspectral imagery.

![Figure 5.64: Radiometric spectral polishing.](image)

Input to the spectral polishing program is the reflectance cube calculated with ATCOR. It employs the vegetation index \( 0 < NDVI < 0.33, NDVI = (\rho_{NIR} - \rho_{RED})/(\rho_{NIR} + \rho_{RED}) \) to mask soil pixels. A soil spectrum is a slowly varying function of wavelength, therefore a spectral smoothing will only remove spikes without disturbing the spectral shape. Then the average reflectance spectrum over all soil pixels is calculated and smoothed with a 5-channel filter except for
the atmospheric water vapor regions where a linear interpolation is performed. The ratio of the filtered to the original soil spectrum is the spectral polishing function applied to all image pixels. If "xxx_atm.bsq" is the atmospherically corrected input image, then "xxx_atm_polish.bsq" is the polished output reflectance cube, and the spectral polishing function is stored in "xxx_atm_polish.dat", an ASCII file with two columns containing the center wavelength of each channel and the polishing factor. Figure 5.64 shows the GUI panel.

5.6.5 Flat Field Polishing

This routine is to remove spectral artifacts from atmospherically corrected imaging spectroscopy data. Average residual gains and offsets are calculated by comparison of assumed flat field data values to the measured values for each spectral band to find a gain correction factor. The flat field is searched automatically within an image as the spectrally least variable part of the image.

![ATCOR Flat Field Polishing](image)

**Figure 5.65:** Flat field radiometric polishing.

**Inputs:**
- Input file name (usually output of atmospheric correction _atm.bsq), can be any kind of image.
- Type of Correction Function:
  - Gain and Offset: calculate average residual gain and offset for each pixel and apply them as correction function
  - Gain only: constrain the residual gain to an offset of 0; this is the typical situation.

**Output:**
- A cube containing the spectrally filtered copy of the original image data cube is created.

5.6.6 Pushbroom Polishing / Destriping

This routine treats each detector pixel of a pushbroom imaging spectrometer separately and derives gain and optional offset values in comparison to its direct neighbors. The routine may be used for both, spectral polishing of residual gain/offset errors and for destriping of pushbroom imagery.

**Inputs:**
- **Input file name** A hyperspectral image cube, usually the output of atmospheric correction _atm.bsq
- **Interpolation Distance** Number of pixels from center pixel, i.e., a factor of 2 uses 2 pixels on each side for calculation of residual gains. The distance should be in a range of the width of visible striping artefacts.
Polishing Filter Type  Three options are available for pushbroom spectral polishing: Spectral: Polishing is done in spectral dimension only; one adjacent band on each spectral side is taken for residual calculation.
Spatial: Only the spatial filter is applied; use this option for destriping of imagery.
2D Filter: Do both dimensions (spatial/spectral) simultaneously the filter size only applies to the spatial dimension, however..

Type of Correction Function  For each detector pixel, correction parameters are generated:
Gain and Offset: Calculate average residual gain and offset for each pixel and apply them as correction function.
Gain Only: Constrain the residual gain to an offset of 0; this is the typical situation.

Output:
A cube containing the spectrally filtered copy of the original image data cube is created. As a side output, a gain file is written (name: _gain.bsq'), containing the following three layers: 1: offset of linear recalibration offset function
2: gain of linear recalibration offset function
3: gain of linear recalibration offset function if function is constrained to zero offset such that the corrected spectral band is: \( L_{polish} = L_{ori} - (L_{offs} + \text{gain} \ast L_{ori,smooth}) \)

5.6.7 Spectral Smile Interpolation
For sensors affected by "spectral smile" the surface reflectance cube is calculated accounting for the smile shift function in the column (across-track) direction of the detector array. The smile shift is specified as a 4th-order polynomial function, i.e. the file "smile_poly_ord4.dat" in the corresponding sensor folder, see chapter 4.7. Due to the smile shift the wavelength values of a spectral channel vary slightly in across-track direction. The "smile interpolation" function allows the specification of a common center wavelength for each channel. Then for each channel all pixel reflectances are interpolated to this new reference wavelength. Since the smile shift between adjacent bands does not vary significantly, a linear interpolation can be applied. If \( \lambda_j(i) \) denotes the center wavelength of band i and column j, and \( \rho_j(i) \) the surface reflectance of a column j pixel, then the new interpolated
reflectance is

\[
\rho_{j}^{(\text{new})}(i, \lambda_{\text{ref}}(i)) = \rho_{j}(i) + \frac{(\lambda_{\text{ref}}(i) - \lambda_{j}(i))}{\lambda_{j}(i+1) - \lambda_{j}(i-1)} (\rho_{j}(i+1) - \rho_{j}(i-1))
\]  \quad (5.5)

where \(\lambda_{\text{ref}}(i)\) is the user-defined reference center wavelength for band \(i\). There are three options for the reference wavelength grid:

1. use wavelength corresponding to the center of the detector array,
2. use average wavelength over all detector columns (per band),
3. use nominal wavelength specified in the (ENVI) header of the reflectance cube.

This tool is available in the interactive mode (main menu, then "Filter", then "Spectral Smile Interpolation (Image Cube)" and in the batch mode ("smile_interp3_batch", see chapter 6.3).

This routine is used after smile-dependent atmospheric correction. It applies a linear interpolation on the reflectance data in order to bring the spectral bands to a common reference in across track direction. The inputs are as follows (see Fig. 5.67):

**Inputs:**

**Input Image** A hyperspectral image cube, usually the output of atmospheric correction in smile mode *atm.bsq*

**Smile polynomial file** The file 'smile_poly_ord4.dat' as of the sensor definition used for the smile-aware atmospheric correction.
CHAPTER 5. DESCRIPTION OF MODULES

Options
Three options for the spectral reference wavelength grid to be used for interpolation may be selected:

center of detector array: The spectral position of the center pixel in across track direction of the detector is taken as the reference wavelength for each spectral band.

average over all detector columns: For each spectral band, the average of all (smiled) center wavelengths is calculated and used as the new reference wavelength.

nominal position (ENVI header) the nominal position as provided in the ENVI header file is taken as the reference.

Output:
A cube containing the spectrally interpolated image data is generated and the ENVI header is updated (for options 1 and 2).

5.6.8 Cast Shadow Border Removal

This routine is used after cast shadow correction in case the borders of the shadows show dark or bright artifacts. It corrects the imagery by adjusting the brightness to the direct neighbor pixels. Using the illumination file, all pixels at the cast shadow borders are first identified and a buffer along the borders is calculated using the border width. Secondly, a brightness correction factor is calculated for each pixel in the border lines using the relative brightness in comparison to its neighbours. The pixels are then corrected using this relative difference in a multiplicative way.

The inputs are as follows (see Fig. 5.67):

Inputs:

Input File A hyperspectral image cube, usually the output of atmospheric correction *_atm.bsq

Illumination/Cast Shadow File illumination file containing a shadow mask which had been applied to the image during atmospheric correction.

Options
Two options for alternate processing are available (try those in case the artifacts are not well removed):

adjust spectral mean: An average additional correction factor is applied to the border pixels in order to correct for spectral variation of the brightness difference.

Interpolate Borders: The border pixels are interpolated from their neighbours, overwriting the original values.

Shadow Border Width: the width of the border to be corrected/interpolated.

Output:
A cube containing the filtered image data is generated and the ENVI header is copied to the new file.
Figure 5.68: Shadow border removal tool
CHAPTER 5. DESCRIPTION OF MODULES

5.7 Menu: Simulation

The "Simulation" menu provides programs for the simulation of at-sensor radiance scenes based on surface reflectance (or emissivity and temperature) images.

5.7.1 TOA/At-Sensor Radiance Cube

This routine calculates an At-Sensor Radiance Cube from an reflectance image cube. All parameters used for the processing are generated from the *.inn file of the input cube. If the function is called, the cube is opened and the *.inn file is read, which results in an at-sensor cube *toarad.bsq. Note that this routine does not consider adjacency effects and is a simple forward propagation based on the given parameters and the given standard model.

No specific panel is displayed. The routine asks for the input reflectance image. All other information is taken from the *.inn file. Please make sure that the reflectance image spectral definition corresponds exactly to the chosen atmospheric library and sensor definition as of the *.inn file.

5.7.2 TOA/At-Sensor Thermal Radiance

This routine calculates an At-Sensor Thermal Radiance Cube from an emissivity/temperature cube (where temperature is in the last band). All parameters used for the processing are generated from the *.inn file of the input cube. If the function is called, the cube is opened and the *.inn file is read, which results in an at-sensor cube *toarad.bsq

5.7.3 At-Sensor Apparent Reflectance

This routine calculates an at-sensor apparent reflectance from a calibrated at-sensor radiance image cube. This routines calculates for each image band the following output:

\[ \rho_{app} = (DN \times c_1 + c_0) \times \pi \times d^2 / (E_0 \times \cos(\theta_0)) \]  (5.6)

where:
- \( DN \): stored data values
- \( c_1 \): gain for conversion of DN to at sensor radiance
- \( c_0 \): offset for conversion to at sensor radiance
- \( d \): relative sun-earth distance (average: \( d = 1 \))
- \( E_0 \): solar irradiance, top of atmosphere (NOT at aircraft altitude)
- \( \theta_0 \): solar zenith angle
CHAPTER 5. DESCRIPTION OF MODULES

Inputs:

- input file name
- calibration file name (*.cal)
- solar radiation file (e0_solar_*.spc)
- output file name
- scale factor (see below)
- date of the year, given exactly as 'day/month', eg. 26/7 for July, 26th, used for sun-earth distance calculation
- solar zenith angle (use 'Tools : Solar Zenith and Azimuth' for its calculation)

Output: A cube containing the scaled apparent reflectance in [%] is stored. The data format is driven by the scaling factor as follows:

- scale < 10 : byte
- scale >= 10 : integer
- scale >= 500: unsigned integer
- scale <= 1.: floating point (effective value, unit: [%]) (wavelength reference and FWHM are taken from the file e0_solar_*.spc)

5.7.4 Top of Atmosphere Radiance Calculation

This routine allows to calculate at sensor signals from spectral measurements of bottom of atmosphere reflectance under consideration of irradiance measurements.
Inputs:

- BOA Reflectance File: ENVI spectral library containing the bottom of atmosphere re-
  flectance. All spectra in the file are used for TOA Radiance calculation.
- BOA irradiance File: ENVI spectral library containing the irradiance on the ground in W/(m2
  sr nm)
- Option: Use BOA irradiance data values directly: for TOA radiance calculation, one may
  use the BOA irradiance instead of modeled irradiance.

Scene Specific Parameters:
The parameter may be read from an ATCOR *.inn file as far as they are available.

- Sensor: current sensor definition (directory name)
- Atmosphere: current atmospheric file (for ATCOR-4: be aware to select the correct flight
  altitude)
- Visibility: current visibility.
- Date, Solar Zenith, Azimuth, and Ground Altitude, should be entered. The values are read
  from the *.inn file if available.
- Sensor View Zenith: this value is to be entered manually; zero degrees corresponds to nadir
  view.

Actions: Upon availability of irradiance spectra, the atmospheric parameters may be inverted
from the irradiance spectra (first two actions):

Find Best Atmospheric Model searches for the applicable atmospheric file (i.e. aerosol model
and water vapor content)

Estimate Visibility searches for the best visibility fitting to the irradiance spectrum.

Calculate TOA Radiance Performs the calculation and creates the output library.

Plot TOA Radiance Starts the internal spectral library viewer for plotting the outputs.

Outputs: Output File: An ENVI spectral library file is written which contains all spectra as of
the input file transformed to at-sensor radiance. all relevant information is written to the ENVI
file header.

Note: Use the tool 'Simulation:Spectral Validation' for validation of the such created spectral
libraries in comparison to the ENVI at-sensor radiance imagery.

5.7.5 ATCOR Vicarious Validation Tool

This routine allows to validate imagery in comparison to simulated at-sensor radiance spectra.
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Figure 5.71: TOA radiance calculation from ground reference reflectance spectral library

Inputs:

- **TOA Radiance SLB**: ENVI spectral library containing the tof of atmosphere radiance (as created by the tool ‘Simulation:TOA Radiance’)
  - using the button ‘Plot TOA Radiance’ allows to plot the currently selected file

- **Validation Image**: Calibrated at sensor radiance image (ATCOR input image) to be used for cross validation
  - using the button ‘Show Image’ starts an image display of the currently selected image file.

- **Sample Position**: Coordinates of sample spectrum in image (may be either pixel coordinates or geographic coordinates)
  - using the ‘Get’ button transfers the position coordinates from the ATCOR file viewer (after ‘Push’ has been pressed there).

- **Sample Diameter**: diameter of square sample in pixels around the sample position (uneven numbers). Statistics for the validation are taken from this sample. NOTE: the sample size should be at least 3x3 clean pixels; this means that the total size of the target should be at least 7x7 pixels (compare [62]).

- **Adjacency Range**: The adjacency range is given in image pixels. The default is the adjacency range in km divided by the standard pixel size.

- **Parameters**: sensor, atmosphere, visibility, solar zenith, and ground altitude are read from slb ENVI header at first. If selecting an image file; the values are read from the *.inn file if available.

- **CAL file for TOA Validation**: This file can be selected, specifically for analysis using the ‘Validation Plot’ tool.
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Actions:

Validate Calculates both the validation spectra including adjacency correction and writes an ascii report of the calculations.

Validation Plot Creates a plot of the sample statistics and the corresponding reference spectrum, scaled by the parameters from the currently selected *.cal file.

Write CAL File Lets you select a number of validation output files and will create a new *.cal file by linear regression of image data to simulated data for all selected files.

Outputs: Output File: an idl save dump is written upon pressing the 'Validate' button; in parallel a validation report with all spectra is written named [out].eval.txt. The calibration reports may be used for derivation of CAL file by linear regression.

![Image of sensor vicarious validation interface](image)

Figure 5.72: TOA radiance validation against calibrated image spectra

5.7.6 Resample Image Cube

This routine allows to simulate a multispectral image from imaging spectroscopy data. A detailed description of this routine is given in chapter 8.
Figure 5.73: Plot sample of validation against calibrated image spectra
5.8 Menu: Tools

The "Tools" menu contains a collection of useful routines such as the calculation of the solar zenith and azimuth angles, spectral classification, nadir normalization for wide field-of-view imagery, adding of a synthetic blue channel for multispectral sensors with a blue band (e.g. SPOT) which is done for the atmospherically corrected surface reflectance image, spectral calibration, conversion of the monochromatic atmospheric database from one to another solar irradiance spectrum, and BIL to BSQ conversion.

**Figure 5.74:** The tools menu.

5.8.1 Solar Zenith and Azimuth

The routine SOLAR_GEOMETRY is used to calculate the zenith and azimuth angle for the image location and acquisition time. All explanations concerning the definition of angles are included in the panel (Fig. 5.75).

**Figure 5.75:** Calculation of sun angles.
5.8.2 Classification of Surface Reflectance Signatures

The spectral classification (SPECL) is a hierarchical classifier based on surface reflectance data employing the Landsat Thematic Mapper (TM) wavelengths. It is not a land use classification, but a classification into certain spectral categories, e.g. dark and bright bare soil, and different vegetation classes, see figure 5.76. The following set of rules is used where b1, b2, b3, b4, b5 and b7 indicate the surface reflectance in the TM bands 0.48, 0.56, 0.66, 0.84, 1.65, 2.2 \( \mu m \), respectively, or the nearest corresponding channel:

- snow: \( b4/b3 \leq 1.3 \) and \( b3 \geq 0.2 \) and \( b5 \leq 0.12 \)
- cloud: \( b4 \geq 0.25 \) and \( 0.85 \leq b1/b4 \leq 1.15 \) and \( b4/b5 \geq 0.9 \) and \( b5 \geq 0.2 \)
- bright bare soil: \( b4 \geq 0.15 \) and \( 1.3 \leq b4/b3 \leq 3.0 \)
- dark bare soil: \( b4 \geq 0.15 \) and \( 1.3 \leq b4/b3 \leq 3.0 \) and \( b2 \leq 0.10 \)
- average vegetation: \( b4/b3 \geq 3.0 \) and \( (b2/b3 \geq 0.8 \) or \( b3 \leq 0.15) \) and \( 0.28 \leq b4 \leq 0.45 \)
- bright vegetation: \( b4/b3 \geq 3.0 \) and \( (b2/b3 \geq 0.8 \) or \( b3 \leq 0.15) \) and \( b4 \geq 0.45 \)
- dark vegetation: \( b4/b3 \geq 3.0 \) and \( (b2/b3 \geq 0.8 \) or \( b3 \leq 0.15) \) and \( b3 \leq 0.08 \) and \( b4 \leq 0.28 \)
- yellow vegetation: \( b4/b3 \geq 2.0 \) and \( b2 \geq b3 \) and \( b3 \geq 0.08 \) and \( b4/b5 \geq 1.5 \)
- mix veg/soil: \( 2.0 \leq b4/b3 \leq 3.0 \) and \( 0.05 \leq b3 \leq 0.15 \) and \( b4 \geq 0.15 \)
- asphalt/dark sand: \( b4/b3 \leq 1.6 \) and \( 0.05 \leq b3 \leq 0.20 \) and \( 0.05 \leq b4 \leq 0.20 \) and \( 0.05 \leq b5 \leq 0.25 \) and \( b5/b4 \geq 0.7 \)
- sand/bare soil/cloud: \( b4/b3 \leq 2.0 \) and \( b4 \geq 0.15 \) and \( b5 \geq 0.15 \)
- bright sand/bare soil/cloud: \( b4/b3 \leq 2.0 \) and \( b4 \geq 0.15 \) and \( (b4 \geq 0.25 \) or \( b5 \geq 0.30) \)
- dry vegetation/soil: \((1.7 \leq b4/b3 \leq 2.0 \) and \( b4 \geq 0.25) \) or \((1.4 \leq b4/b3 \leq 2.0 \) and \( b7/b5 \leq 0.83) \)
- sparse vegetation/soil: \((1.4 \leq b4/b3 \leq 1.7 \) and \( b4 \geq 0.25) \) or \((1.4 \leq b4/b3 \leq 2.0 \) and \( b7/b5 \leq 0.83 \) and \( b5/b4 \geq 1.2) \)
- turbid water: \( b4 \leq 0.11 \) and \( b5 \leq 0.05 \)
- clear water: \( b4 \leq 0.02 \) and \( b5 \leq 0.02 \)
- clear water over sand: \( b3 \geq 0.02 \) and \( b3 \geq b4 + 0.005 \) and \( b5 \leq 0.02 \)

Figures 5.77, 5.78 show the panel of the SPECL program and a sample output. When the button "emissivity classes and emissivity values" is clicked the emissivity classes of table 10.1 are assigned the emissivity values as defined in file "emissivity.dat" in the atcor4 directory. So this file can be edited to modify the emissivity values for the 15 classes.
Figure 5.77: SPECL: spectral classification of reflectance cube.

5.8.3 Spectral Smile Detection

This routine uses sharp atmospheric absorption features and Fraunhofer lines for inflight smile (i.e., spectral across-track non-uniformity) detection. The calculation is done by correlation analysis of a number of spectral bands in the vicinity of selected absorption features. The outputs may be used for smile-aware atmospheric correction.

Initially, the smile characterization for each spectrometer channel is derived from laboratory measurements. From such data, the wavelength shift with respect to the center pixel of the detector array can be parametrized using a 4th order polynomial fit. However, in case of instrument changes during the mission, a spectral re-calibration might be necessary from the image data or from onboard calibration facilities using well-defined absorption features. Onboard spectral calibration devices such as interference or rare earth filters would be well suited for this purpose. However, such devices are often not available in sensor systems. Therefore, atmospheric gas absorption features or solar Fraunhofer lines have to be taken as a reference from the imagery itself. The processing steps are:

1. A calibrated image is averaged in along track direction, leading to a signature image of the size of the detector array.
2. The surface reflectance is calculated (atmospheric correction) and smoothed.
3. The spectral bands within the spectral matching range are selected.
4. Spectral shifts with intervals between 0.01-0.05 nm are calculated and applied to the selected spectral band response functions.

5. An appropriate pre-calculated fine-spectral resolution atmospheric LUT is selected which serves for the calculation of at-sensor radiance values for the series of spectrally shifted response functions using the surface reflectance spectrum from step (2).

6. The derived spectral signatures are then correlated to the observed column-averaged signal in the image, such that the best fitting spectral shift $\Delta \lambda_j = \Delta_j$ can be found for each image column $j$, i.e., the $\Delta_j$ with the highest Pearson’s correlation coefficient is selected. This is equivalent to minimizing the merit function

$$
\chi^2(\Delta_j) = \sum_{\Delta_j = \lambda_k - 5nm}^{\lambda_k + 5nm} [L_I(j,k) - L_R(\lambda_k + \Delta_j,k)]^2,
$$

(5.7)

where $L_I(j,k)$ is the average at-sensor radiance of the image for column $j$ and channel $k$, and $L_R(\lambda_k + \Delta_j,k)$ is the corresponding reference radiance for a wavelength shift $\Delta_j$ within a 5 nm interval around $\lambda_k$. By using the continuum removed scaled radiances, the stability of the routine is enhanced.

7. A 4th order polynomial is fitted through the calculated spectral points and the respective polynomial parameters of eq. 4.1 are stored.

8. The polynomial parameters are interpolated and optionally extrapolated to all other bands within the same detector or spectrometer unit. Optionally, the polynomial coefficients can be set to zero in atmospheric absorption regions to expedite the processing.
The same approach is used for FWHM detection, with the difference that not the position of the spectral bands is varied, but the FWHM of the spectral bands is scaled in a systematic way. Once the coefficients have been determined, they are converted into the required file format and are placed in the respective sensor folder for a subsequent fully automatic radiometric and atmospheric processing. Fig. 5.79 shows the panel of the smile detection module.

**Inputs:**

**Input file**: calibrated ENVI image file. The file should be statistically as uniform as possible in order to get valid averages.

**Atmospheric Database File**: non-resampled atmospheric database file most appropriate for the selected image; take care of the flight altitude and aerosol model.

**Sensor spectral response**: First band of sensor spectral response file(s) *.rsp. The corresponding pressure definition file (pressure.dat) is selected automatically from the same directory as the *.rsp files, if available.

**Output smile coefficients**: file name of ASCII file which is written and contains the 4th order polynomial coefficients. This file may be copied to the sensor definition as ’smile_poly_ord4.dat’ for use as smile definition with a sensor.

**Detection resolution**: finest spectral resolution used for correlation analysis - results will be resolved by this resolution.

**Search range**: maximum total spectral range used for smile detection (ie., 20 nm is +- 10 nm search distance). For FWHM detection, it is the maximum factor to be applied. A factor of 2, e.g. means to search between FWHM scaling from 0.5 to 2.

**Band range**: bands which shall be used for smile detection and for which the coefficients are written to the output file.

**Split band**: First band of second detector for sensor having more than one detector (starting numbering at 1). NOTE: for imagers with more than two detector it is recommended to derive the smile separately for each detector to avoid artefacts in the transition range.

**Visibility**: horizontal visibility as of modtran conventions [km]

**Solar Zenith**: angle measured from zenith to sun

**Mean Ground Elevation**: Ground altitude in km.

**Flight Altitude**: in km a.s.l.

**Feature wavelength**: 17 selectable features used for smile detection. The per-band smile is interpolated from these feature wavelengths.

**Interpolation type**: interpolation used to expand the feature wavelength results to all spectral bands

**Extrapolation type**: specifies how the bands outside of the selected features are treated. Repeat: repeats the last value toward the borders. ’to zero’: gradually decrease to zero at border of detector.
Actions:

**Detect Smile** The module will perform the smile detection calculation.

**Detect FWHM** The FWHM variations are calculated instead of smile variations by the same technique.

**Plot Smile** Starts a plotting window to check the smile and lets you select suitable features for calibration.

**Save Report** Saves an informational report about the smile detection.

Outputs:

An ASCII-file which may be used as smile (or FWHM) description file in the respective sensor directory. Note that this file should be named "smile_poly_ord4.dat" or "smile_poly_ord4_fwhm.dat" in order to be automatically recognized by ATCOR.

As a side output, an IDL save dump (*.sav) is written in parallel which contains all used parameters and the effectively calculated smile results in array "smileresult(4,ncols,17)". Here the first dimension (4) contains the center wavelength (nm), smile shift (nm), atmospheric transmittance, and correlation coefficient. The second dimension (ncols) refers to the number of image columns, and the last dimension (17) contains results for the 17 feature wavelengths. For example, the center wavelengths for all across-track pixels and the 760 nm feature are stored in smileresult[0,*,5], because this feature is the third one and IDL arrays start with index 0.

![Figure 5.79: Spectral smile detection](image-url)
5.8.4 Spectral Calibration (Atm. Absorption Features)

The program (SPECTRAL_CAL) is only intended for hyperspectral sensors and employs atmospheric absorption features to detect and remove possible wavelength calibration errors (see chapter 2.2). For this purpose, a certain number of target spectra have to be selected in the SPECTRA module.

NOTE: Alternatively to this routine, the spectral calibration can be done by first using the spectral smile detection routine (see section 5.8.3) and then applying the shift to the sensor as described in section 5.2.3.

Input to the spectral calibration module are the DN spectra of selected fields saved as ASCII files in the SPECTRA module by pressing the button "Save last spectrum". The files should be numbered consecutively, starting with a name such as "location_target1", without extension. The next target has to be named "location_target2" etc. For each target field three files will generated, for example:

- "location_target1.dat" : contains surface reflectance spectrum,
- "location_target1.txt" : contains target coordinates, and processing parameters (visibility, water vapor column, etc.),
- "location_target1_dn1.dat" : contains the DN spectrum.

For a given location (scene) up to 9 targets can be extracted and used for the spectral calibration. The geometry, scene visibility and average water vapor content of the targets enter as parameters to the spectral calibration, see Fig. 5.80. The water vapor content has to be averaged from the values found in the "location_target*.txt" files. The first target DN file has to be selected by the user, the remaining target files are automatically found provided the nomenclature with a consecutive numbering is applied. The result of the spectral calibration are files with the spectral shifts per spectrometer and the new center wavelengths of all channels. The spectral bandwidth of channels is not modified.

The DN spectra will be employed in an optimization procedure that minimizes the spikes of the derived surface reflectance spectra in the atmospheric absorption regions. The first target DN file has to be entered at the top left button of the GUI panel (Figure 5.80). The other target files are automatically searched and employed if the nomenclature of chapter 2.2 is employed. Further input data are the sensor definition, the range of bands per spectrometer, solar geometry and atmospheric parameters. Output is a file with the spectral channel center wavelength shifts per spectrometer, and a new wavelength file containing the updated wavelengths for each channel.

The results of the spectral shift are summarized in a file "spectral_calibration_results*.txt" where the wavelength shift is listed for each spectrometer and each target. The final shift is taken as the average of all target wavelength shifts. In addition, a new wavelength file "sensor_new.wvl" is created containing the channel center wavelengths and the FWHMs (bandwidth as full width at half maximum). A copy of the original radiometric calibration file (e.g., "xxx.cal") is provided for convenience (e.g., "xxx_new.cal") which contains the original radiometric calibration coefficients and the updated channel center wavelengths.

In case of originally non-Gaussian filter curves, the output FWHM values of "sensor_new.wvl" represent the equivalent Gaussian FWHM values, even though the spectral re-calibration is based on the original (non-Gaussian) filter curves. The corresponding sensor with the new spectral calibration has to be added to the list of existing sensors (see chapter 4.6) to process imagery.
with the updated spectral calibration. In case of non-Gaussian filter curves, the original channel response files ("band*.rsp") should be copied to the new sensor directory applying the appropriate wavelength shifts. For sensors with Gaussian filter curves the 'gauss_rsp’ module (see chapter 5) can be applied to the "sensor_new.wvl" file to generate the corresponding "band*.rsp" files. Note that a change of the spectral calibration usually requires a radiometric re-calibration.

### 5.8.5 Calibration Coefficients with Regression

This routine employs the "*.rdn" files obtained during the single-target calibration (the "c1 option" of ATCOR’s calibration module to create a calibration file by linear regression.

![CAL_REGRESS.png](attachment://CAL_REGRESS.png)

*Figure 5.81: CAL_REGRESS.: radiometric calibration with more than one target*

So for n > 2 the single-target calibration is to be performed n times and the resulting "*.rdn" files (radiance versus digital number) are offered as input to the "cal_regress" program.

![SPECTRAL_CAL.png](attachment://SPECTRAL_CAL.png)

*Figure 5.80: SPECTRAL_CAL.: spectral calibration*
Inputs:

**Number of calibration targets:** A maximum of 9 calibration targets may be selected. The files *.rdn should having been calculated beforehand and the need to be calculated consecutively, e.g., calib1.rdn, calib2.rdn, ...

**First *.rdn file:** Name of the first *.rdn file of the series to be used for regression

**Output Name** Name of the calibration output to be created.

**Output:** The output of this program is an ASCII file "name.cal" with three columns (center wavelength, \(c_0\), \(c_1\)) where "name" is specified by the user.

**Note:** If several calibration targets are employed, care should be taken to select targets without spectral intersections, since calibration values at intersection bands are not reliable. If intersections of spectra cannot be avoided, a larger number of spectra should be used, if possible, to increase the reliability of the calibration.
5.8.6 Convert High Res. Database (New Solar Irradiance)

The standard solar irradiance curve used with ATCOR is the Fontenla database (2011). However, there’s some uncertainty about the solar irradiance and people may want to use ATCOR with a different solar reference function. This module (CONVERT_DB3) converts the complete high resolution atmospheric database from the current definition to a new irradiance function. Normally, the standard database is converted (this function does not apply to the thermal IR), but also the specific CHRIS database may be selected. In the panel (see Fig. 5.82), the two databases may be selected on the basis of the directory (f1) and the new reference function e0_solar*.dat.

![Figure 5.82: Convert monochromatic database to new solar reference function](image)

5.8.7 Convert .atm for another Irradiance Spectrum

The conversion as described in module 5.8.6 can be applied to a sensor-specific atmospheric library of a self-defined sensor using this function. In the panel as of Fig. 5.83, the sensor has first to be entered and the new solar function e0_solar*.dat is to be selected before the conversion may be applied.
Figure 5.83: Convert atmlib to new solar reference function
5.8.8 Thermal Spectral Calibration (Atm. Features)

This program is intended for hyperspectral thermal sensors and employs atmospheric absorption features to detect and remove possible wavelength calibration errors (see chapter 3). It is the counterpart to the spectral calibration in the solar reflective region. At least 20 channels in the 8.5 - 13.5 µm region are required.

Input to the thermal spectral calibration module is the original (non-geocoded) scene and its '*.inn' file from which the necessary parameters (sensor name, atmospheric LUTs etc) are taken. Then 10 spectra are evaluated from 10 image lines located at the image center (nadir) and the corresponding wavelength shifts are calculated for the specified surface temperature range. In addition, an averaging pixel box size can be specified to reduce noise.

Figure 5.84 presents the interactive GUI. Output is a file with the new shifted channel center wavelengths. This feature is also available for bath processing (‘sp_calth’), see chapter 6.3.

Another module provides the capability of radiometric calibration for thermal band imagery (‘thermalcal’) if the scene contains water bodies. In this case the theoretical spectral water emissivity is used and the user has to specify the file name of the input scene, the water target pixel coordinates (xpos, ypos) and the box size. Output is the new calibration gain (c1), assuming that the offset (c0) is zero for all bands, see chapter 6.3.

A related subject is the estimation of the water vapor column from purely hyperspectral imagery. Here the water vapor is estimated by automatically selecting 10 pixel spectra from the scene covering a temperature interval (Tmin, Tmax), but avoiding extreme temperatures. Then the difference between measured and simulated at-sensor radiance spectra for these 10 pixels is calculated and...
the water vapor LUT which minimizes the difference is recommended for the processing, see chapter 6.3.

5.8.9 Create Scan Angles

Creates a scan angle file from input geometry. Two procedures are supported:

(a) (Preferred) If a MAP file or an ENVI GLT file is available for the imagery, the scan angle file may be created from the geometric information contained in that file. This results in a pixel accurate scan angle files.

(b) If no MAP/GLT is available, the file is to be created from scratch on the basis of the flightpath direction. The given parameters are used to to interpolate a guess of the scan angle file under assumption of stable and straight flight conditions between (and beyond) a given starting point and ending point.

**Accepted Inputs:**

- Create SCA file...: select either option (a) or (b) from above
- Name of Reference File: File which defines the spatial dimensions of the output scan angle file to be created. - this is the GLT/MAP file if the respective option is selected, but may also be a DEM of output resolution if flight path parameters are used.

OPTION (a):

- Flight Heading: direction of the flight path with respect to north, east is 90 degrees.
- Flight Altitude: average flight altitude in meters.

OPTION (b):

- Starting Point:
  one initial point in pixel coordinates and meters altitude of nadir line the pixel coordinates are the row and column numbers as of the reference file selected in this window.

- Ending Point:
  one final point in pixel coordinates and meters altitude of nadir line.

**NOTE:** the two points may be anywhere on the flight path, they are just used to find the nadir line within the imagery. The y coordinates are entered in bottom-up system, i.e. the lower left pixel of the image is pixel number 0/0. Use the function `File:Display ENVI file 5.1.1` to find the correct coordinates.

(END OPTION b)

- Sensor total Field of View:
  total across track FOV of the sensor in degrees (default: as defined in sensor model)
- Scan Angle Output:
  output file name (to be created):

The routine creates a standard 3 layers scan angle file (ENVI format) as it is used by ATCOR for radiometric processing, containing:

- pixel zenith (angle in degrees * 100)
- pixel azimuth (angle in degrees from north towards east * 10)
- sensor altitude (meters asl.)

Figure 5.85: Scan angle creation panel; option (a): top, option (b): bottom.
5.8.10 MTF, PSF, and effective GIFOV

A useful representation of the spatial resolution of a sensor is the transfer function, describable in terms of a modulation transfer function (MTF) and a phase transfer function. The transfer function describes how the system modifies the amplitude (MTF) and shifts the phase of the input spatial frequencies. Usually, only the MTF is given as a figure-of-merit or its counterpart, the point-spread-function (PSF). The MTF is the 2-D Fourier transform of the PSF [87].

Fig. 5.86 presents the GUI for the MTF / PSF evaluation. The image is loaded in the left part. The user should click with the left mouse button in the area where the MTF evaluation is intended. This area will be shown in top middle part of the GUI. Then the MTF can be evaluated for a line or an edge, yielding the line-spread function (LSF) or edge-spread function (ESF), respectively. The user is responsible to select an appropriate feature by clicking the start and end points of the line with the left and center mouse buttons, respectively.

After clicking ”Calculate MTF”, the MTF is plotted below for up to four fixed channels as well as the effective GIFOV (Ground Instantaneous Field-of-View) defined as one half of the reciprocal of the spatial frequency at which the MTF is 0.5. The effective GIFOV is also called effective instantaneous field-of-view EIFOV and it is specified in pixels. Low values of the EIFOV represent a good capability at resolving high spatial frequencies. The slider ”Line averaging” can be adjusted between 3 and 9 to evaluate up to 9 parallel lines (centered on the user-defined line) to average the phase effect.

Results (MTF, LSF, DN_profile, Effective GIFOV) for all bands can be saved as ENVI spectral library files (”*.slb”). The ”Status” widget indicates the mode of calculation (LSF or ESF).

5.8.11 FODIS Processing

The reader is referred to chapter 4.13 for the description and presentation of the graphical user interface for FODIS (Fiber Optic Downwelling Irradiance Sensor) data processing.
Figure 5.86: MTF and effective GIFOV.
5.9 Menu: Help

Finally, the "Help" menu allows browsing of the ATCOR user manual, provides a link to the web resources, contains the software update center, and displays license information.

5.9.1 Help Options

The options of the help menu are listed below:

**Browse Manual** Opens this manual in the default PDF display application of your machine.

**Web Resources** Opens the html document `atcor3_webresources.htm` in the systems default applications for viewing HTML-documents.

**About** Provides basic information about the copyright and also displays the build-number of the software (please provide the build-number for debugging purposes in case the ATCOR support is contacted).

**Check for Updates** Connects to the ReSe Web server to check if a new build/release of the software is available. The software update is downloaded automatically if available and may be installed thereafter.

**Install Components** This is your software update center - atmospheric database files may be downloaded and updated directly through this update tool upon availability.

**Your License** Provides information about the licensed features in your license key and also displays which license key is currently active.

*Figure 5.87: The help menu.*
Chapter 6

Batch Processing Reference

For most ATCOR modules a convenient graphical user interface is available, but batch jobs can also be submitted. A detailed discussion of the interactive panel-driven modules is given in chapter 5. Running ATCOR in batch mode can be done in two ways: either from the operating system console directly or from within IDL. The latter provides a higher flexibility whereas the first is well suited for integration of ATCOR in a processing environment.

6.1 Starting ATCOR from console

ATCOR can be started directly from a console or from a different processing environment using the standard call to IDL run time environment. This call will show a splash screen when starting IDL unless an IDL RT or a full IDL license is installed on the processing computer. RT licenses can be acquired from exelisvis.com at a significantly reduced price than full IDL licenses. This makes this method useful for operational processing with ATCOR.

The call in a windows environment is:

```
[idlpath]\idlrt.exe -rt="[atcorpath]atcor_4\bin\atcor4.sav"
-args input R|F|E [output] [logfile] [elefile] [factor]
```

where [idlpath] is the path to the idl installation, typically something like ..\idl84\bin\bin.x86.64\.

On a unix/macOSX system, the call syntax is as follows:

```
[idlpath]/idl -rt=".../bin/atcor4.sav"
-args input R/F/E [output] [logfile] [elefile] [factor]
```

The [idlpath] in this case is something like ../exelis/idl84/bin/.

Arguments:

input: Input file to be processed or input reference file, respectively

R/F/E: Flag for processing option:// R: rugged terrain processing (atcor4r_batch)// F: flat terrain processing (atcor4f_batch)// E: elevation data preprocessing (at_prepele)

output: Name of output file to be created.
logfile: name of log file to be used or created.

elefile: input elevation file for at_prepele

factor: factor for elevation data processing.

NOTE: files names other than the input file may be set to 'dum' in order to process default file names. After execution, the idl session quits and throws an error status of '1' if an error occurred during processing.

ATTENTION: These routines are overwriting existing outputs.

6.2 Using the batch mode from within IDL

ATCOR can process scenes in the batch mode. For large scenes the tiling option is also available which splits a big scene into a number of smaller sub-scenes, processes the sub-scenes, and finally merges them into one file. A prerequisite for the tiling is that enough IDL memory is available to keep one image channel and the sub-scene channel in memory.

The batch mode can be accessed after the processing parameters have been specified in the interactive graphical user interface (GUI) panel, i.e., after the SPECTRA module has been accessed or after one of the image processing options has been selected. Thus, the GUI panel creates an '*.inn' file containing all input parameters.

The batch mode can be started after quitting the interactive session, using the same IDL window. It can also be started in a new IDL session after typing 'atcor4' on the IDL command line. Then continue with:

```
>atcor4f_batch, input='/data1/examples/example_image.bsq' (case of flat terrain) or
>atcor4r_batch, input='/data1/examples/example_image.bsq' (case of rugged terrain)
```

At this stage, all required input parameters are already available in the '*.inn' file, in this specific case 'example_image.inn'.

The submitted job is a 'quasi batch' job, the corresponding IDL window is used for error and status messages and it may not be closed during the run time of the job. A log file is created during processing, e.g. 'example_image_atm.log', which contains information about the job status. It contains three message levels (I=Info, W=Warning, E=Error) followed by a two digit number (between 0 and 99) and a space, e.g. 'W19 ', followed by the appropriate information. These three message levels can easily be parsed by a separate user program if desired. Other information in the log file is marked with the hashmark symbol in the first column.

In the tiling mode the user has to specify the number of tiles in x=column direction (ntx) and in y=line direction (nty), e.g.,

```
>atcor4f_tile, input='/data1/examples/example_image.bsq', ntx=3, nty=2
```

In this case the image is split into 3*2=6 tiles, each tile is processed separately, finally all tiles are merged into one output file, and the sub-scenes are deleted. The maximum size of each tile depends on the available memory for a specific machine because ATCOR performs most calculations in memory loading one or two complete bands of the scene. A typical tile size for modern machines is 3000*3000 pixels = 9 Mpixels to 5000*5000 pixels = 25 Mpixels. The user has to try
and find out the appropriate tile size. As an example, with a 9 Mpixel tile size and a 30 Mpixel scene the image has to be split into 4 sub-scenes. Assuming that the number of image columns and lines is approximately the same, one would choose the keywords 'ntx=2, nty=2' in this example. Of course, processing of much smaller tiles is also possible, e.g., 'ntx=20, nty=10', but this is not recommended because of potential image border effects, i.e., larger visibility differences for the small tiles might lead to seams at the tile borders.

An optional keyword output can be used to define the output directory and name of the output reflectance file. If the keyword specifies only the output path, which is recommended, then all output files are written to the specified output directory, and the reflectance output file name is the name of the input file with "_atm.bsq" appended.

The optional keyword 'vis' can be used to overwrite the visibility value in the "*.inn" file. For a constant visibility per scene (npref=0 in the "*.inn" file), the input 'vis' value is the start value that will be iterated as described in chapter 10.4.1. In case of a variable scene visibility (npref=1) the 'vis' parameter is ignored if the scene contains enough dark reference pixels. If not, the program switches to the constant visibility mode and 'vis' is used as a start value.

An IDL routine called 'write_atcort4.inn_file' is available to users who want to generate the "*.inn" file without the ATCOR GUI.

Note:
On the IDL command line, the command 'atcort4' has to be typed first to load the 'atcort4.sav' file. Then the 'atcort4f_tile' or 'atcort4r_tile' commands will execute the tile processing. A simple trick can be used to start the 'atcort4_tile' programs directly on the IDL command line without having to type 'atcort4' first: just copy the 'atcort4.sav' file to 'atcort4f_tile.sav' and 'atcort4r_tile.sav'. The same can be done for 'atcort4f_batch.sav' and 'atcort4r_batch.sav'. For the Linux/Unix operation systems a symbolic link is sufficient, e.g., ln -s atcort4.sav atcort4f_batch.sav.

For Linux/Unix users with a full IDL license a batch job can be started directly from the shell, e.g.: idl -e "atcort4f_batch, input='/export/data/data7/atcort4/hymap07/flight71.bsq'"

Most of the modules are available in both modes (interactive and batch). If the 'atcort4.sav' file is copied to 'atcort4f_batch.sav' and 'atcort4r_batch.sav', a batch job can be started immediately from the IDL command line, otherwise 'atcort4' has to be typed first. We begin with a description of the batch modules and keyword-driven modules.

6.3 Batch modules, keyword-driven modules
A number of modules can also immediately be started as batch jobs. You have to type "atcort4" on the IDL command line, then the ATCOR GUI selection panel pops up. Disregard this panel and continue on the IDL command line with the name of the batch job module where all the input parameters have to be specified via key words. Current batch programs are:

- slopasp_batch, input=filename, [ pixelsize=10.0, kernel=3, dem_unit=0 ]

The 'filename' should have the last four characters as "_ele" and the extension "_.bsq". Two output files (slope and aspect) are generated from the elevation file, e.g.,
"example_DEM25m_slp.bsq" and "example_DEM25m_asp.bsq". The values are coded in degrees. The keyword pixelsize is not required if this information is included in the map info of the ENVI header. The keywords kernel and dem_unit can be omitted if the default values kernel=3, and dem_unit=0 are used. The unit of pixelsize is 'meter'. For the elevation height unit three options exist: dem_unit=0 : height unit is 'meters', =1 for 'dm', =2 for 'cm'.

Note: Before running ATCOR with a DEM, please check the results of the slope image. We often encounter severe horizontal and vertical striping in the slope image in case of low quality DEMs or if coarse DEMs of 25 m have to be resampled to say 5 m. Additional appropriate filtering is required in these cases! A simple way might be to try a larger kernel size, e.g., kernel=5 or kernel=7. A simple quality check on the derived DEM solar illumination file is also performed at the start of ATCOR, see the discussion below.

- **skyview_batch**, input=filename, pixelsize=10.0, [dem_unit=0, unders=unders, azi_inc=azi_inc, ele_inc=ele_inc]
  'filename' is the full file name including the path, 'filename' should have the last four characters as "_ele" and the extension ".bsq" to indicate a digital elevation file and to enable an automatic processing, e.g., "example_DEM25m_ele.bsq". 'pixelsize' is specified in meters, 'dem_unit' is the integer code for the DEM height unit (0 represents [m], 1 means [dm], 2 means [cm]). The option dem_unit=0 is default and can be omitted. The keyword "unders" specifies the undersampling factor (in pixels) to reduce the execution time for very large files. The default angular azimuth resolution is azi_inc=10 degrees and the default elevation increment is ele_inc=30 degrees. However, the recommended resolution is 10 degrees for azimuth and 5 degrees for elevation. In case of large files, an undersampling factor > 1 can be specified to reduce the execution time. Usually an undersampling factor of 3 is sufficient. A high angular resolution is more important than a low undersampling factor. The output file replaces the ending "_ele" with the ending "_sky", e.g., "example_DEM25m_sky.bsq".

- **shadow_batch**, input=filename, pixelsize=10.0, solze=30.5, solaz=160.8, dem_unit=0,
  The keywords have the same meaning as for skyview_batch, 'solze' is the solar zenith angle [degr.], and 'solaz' is the solar azimuth angle [degr.]. In particular, 'filename' should have the last four characters as "_ele" and the "_bsq" extension. The output file replaces the ending "_ele" with the ending "_shd" e.g., "example_DEM25m_zen31 azi161_shd.bsq". The rounded zenith and azimuth angles will be included in the "*shd" file name.
  Note: The shadow and skyview calculations can be omitted in gently undulated terrain. Example: for maximum slopes of 25°and a solar zenith angle of 40°no DEM shadow is possible. Also, the local trigonometric sky view factor, employed if the "_sky.bsq" file is missing, is sufficiently accurate, compare figure 10.5.

- **atcor4f_batch**, input=filename [, output=file, vis=vis] or
  atcor4f_tile, input=filename, ntx=3, nty=2 [, output=file, vis=vis]
  The "f" in atcor4f_batch means the code for flat terrain, i.e. no DEM is employed. The 'filename' must be fully qualified, i.e. it includes the path, e.g., "/data2/project1/image1.bsq". The file should have the band sequential (BSQ) file structure. A corresponding "*.inn" file, e.g., "/data2/project1/image1.ini" must be available that contains all processing parameters. This file will be generated during the interactive session. It may be also be created by the user, e.g. employing the program "write_atcor4_inn_file.pro" that is available on request. The default output file name without the output keyword specification is the input name with an "_atm.bsq" appended, e.g., "/data2/project1/image1_atm.bsq".
  The keyword output can be used to specify the full output name, or only the output path,
the latter option is recommended. In that case, all output files are written to the specified output directory, and the reflectance output file name is the name of the input file with "atm.bsq" appended. Example: output="/data4/project1/" then the output reflectance file will be "/data4/project1/image1_atm.bsq".

The corresponding tile program atcor4f_tile in this example is called to split the image into 3 sub-images in x direction, and 2 in y direction, compare chapter 6.2.

The optional keyword 'vis' can be used to overwrite the visibility value in the ".inn" file. For a constant visibility per scene (npref=0 in the ".inn" file), the input 'vis' value is the start value that will be iterated as described in chapter 10.4.1. In case of a variable scene visibility (npref=1) the 'vis' parameter is ignored if the scene contains enough dark reference pixels. If not, the program switches to the constant visibility mode and 'vis' is used as a start value. A negative vis value means the value abs(vis) is used for processing even if it causes a large percentage of negative reflectance pixels.

- atcor4r_batch, input=filename [, output=file, vis=vis] or atcor4r_tile, input=filename, ntx=3, nty=2 [, output=file, vis=vis]
The "r" in atcor4r_batch means the code for rugged terrain, i.e. a DEM is employed as well as other DEM related files, e.g., slope, aspect, skyview. Otherwise, the same explanations hold as for the "flat" terrain ATCOR.
The corresponding tile program atcor4r_tile in this example is called to split the image into 3 sub-images in x direction, and 2 in y direction, compare chapter 6.2. The keywords output and vis are described in atcor4f_batch above.

- Note: optional keywords for atcor4f_batch, atcor4r_batch, atcor4f_tile, atcor4r_tile.
There are four keywords concerning spectral interpolation to overwrite the interpolation settings in file "preference_parameters.dat":
  - i725=-1: no interpolation for 725/820 nm channels, i725=1: interpolation.
  - i760=-1: no interpolation for 760 nm channels, i760=1: interpolation.
  - i940=-1: no interpolation for 940 nm channels, i940=1: nonlinear interpolation, i940=2: linear.
  - i1400=-1: no interpolation for 1400/1900 nm channels, i1400=1: nonlinear interpolation, i1400=2: linear.

- toarad, input=filename, [ pixelsize=pixelsize, sz=solar_zenith, atmfile=atmfile, elev=elevation, vis=visibility, adjrange=adjrange, scalef=scalef ]
The keywords in brackets are optional, the meaning of all keywords is described in chapter 8. Information on all missing keywords is taken from the corresponding ".ini" file. If the keyword elev is missing and the corresponding "*.inn" file contains the DEM files (elevation, slope, aspect) then the simulation is performed for a rugged terrain, otherwise for a flat terrain, compare chapter 8.

- toarad2, input=filename, [ pixelsize=pixelsize, atmfile=atmfile, elev=elevation, vis=visibility ]
Similar to program toarad, but for thermal channels, compare chapter 8.
• `cal_regress, ntargets=4, outfile='regression4'`
  This program uses the "*.rdn" files to calculate a regression for the c0, c1 radiometric calibration, see chapters 2.4, 5.4.8. The above example is for the case of n=4 targets, and the output file will be "regression4.cal" in the directory of the "*.rdn" files which are prompted with a dialog pickfile panel. A graphical user interface for this program is available in the "Tools" pull-down menu of ATCOR labeled "Calibration Coefficients with Regression".

• `sp_calth, input=filename, trange=[280,320], box=3`
  Spectral calibration in the thermal region, the temperature range (Kelvin) and box (pixels) parameters are optional. Ten emissivity spectra in nadir direction are evaluated, evenly spaced between the image lines. Spectral shifts smaller than FWHM/30 usually can be neglected and do not required an update of the calculation of the sensor-specific atmospheric LUTs. This module can also be started from the main menu of ATCOR: 'Tools', 'Thermal Spectral Calibration (Atm.Features)'.

• `thermalcal, input=filename, xpos=xpos, ypos=ypos, box=box`
  Radiometric calibration for thermal band imagery if the scene contains water bodies. xpos, ypos are the user-specified pixel coordinates, box is the window size for averaging over box*box pixels centered around xpos, ypos. If box is not specified, then box=1 is taken. This module calculates the radiometric gain (c1), assuming that the offset (c0) is zero. For this purpose, the theoretical spectral emissivity of water is used (file 'eps_water_7-14um.dat' in the ATCOR4 folder).

• `estimate_wv, input=filename`
  This module estimates the water vapor column from purely hyperspectral imagery. It automatically selects 10 pixel spectra from the scene covering a temperature interval (Tmin, Tmax), but avoiding extreme temperatures. Then the difference between measured and simulated at-sensor radiance spectra for these 10 pixels is calculated and the water vapor LUT which minimizes the difference is recommended for processing.

• `spec_l_batch, input=filename, sensor='xx' or spec_l_tile, input=filename, sensor='xx', ntx=ntx, nty=nty`
  The spectral classification based on template reflectance spectra is also available in the batch mode and with the tiling option. The 'xx' is a keyword for the sensor type, e.g., 'xx'='hymap04'. The complete list of sensor keywords is shown when typing `spec_l_batch` on the IDL command line without the sensor specification. The ntx, nty keywords have the meaning explained for the ATCOR tile programs above.

• `smile_interp4_batch, input=filename, fpoly=fpname, option=number [, silent=silent]`
  Purpose: The atmospheric correction accounts for the column-dependent smile shift as specified in the "smile_poly_ord4.dat" of the corresponding sensor folder, but the image columns of each band belong to slightly different wavelengths.
  This function interpolates the pixel reflectance values for each band to a specified reference wavelength. Three options exist for the reference wavelength grid:

  1. use wavelength corresponding to the center of the detector array,
  2. use average wavelength over all detector columns (per band),
  3. use nominal wavelength specified in the (ENVI) header of the reflectance cube.
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The new reference center wavelengths are included in the header of the output file. If the input filename is "/path1/image_atm.bsq" the output name is "/path1/image_atm_smcorr.bsq" indicating the "smile corrected" common wavelength grid.

Function parameters are: "filename" is the full name of the surface reflectance file, "fpname" is the full name of "smile_poly_ord4.dat", i.e., including the path, 'number' is the above option number (1 - 3), and if the keyword 'silent' is set the progress about the band processing is not issued to the command line.

This module is also available in the interactive mode (see main menu, 'Filter', "Spectral Smile Interpolation (Image Cube)", chapter 4).

• dehaze, input=filename [/ water=iw, ipm=ipm, ihm=ihm, keep=keep, nodem=nodem, nocirrus=nocirrus, fraction_haze=fh]

Here the filename is the name of the image (level 1 data, original DN, stored as ENVI bsq, band-sequential) including the path. The sensor name is read from the scene '.inn' file, e.g. sensor='ads40'. If keyword water=0 (or not keyword set) then only land pixels are dehazed, with water=1 also water pixels will be dehazed. Keyword "ipm" specifies the interpolation method:

ipm = 1 is default and uses bilinear interpolation for the bright areas (very fast)

ipm = 2 uses triangulation (very slow)

ihm = 0, 1, 2 specifies the dehazing option, ihm=1: standard dehazing, ihm=2: stronger dehazing, ihm=0: both options are executed, the results contain the identifier 'dh1' and 'dh2', respectively. Afterward, the better result is renamed with 'dh' and the inferior result is deleted if keep=0 or if this keyword is not specified.

If 'scene.bsq' is the name of the input image, then the output file name is "scene_dh_bilin.bsq", and "scene_dh_trian.bsq" for ipm=1, 2, respectively. The 'dh' indicates the dehazing and the interpolation method is also included. Additionally, a file '* haze_map.bsq' is created containing the classes haze/land, haze/water, clear, water, and geocoded background. If the keyword water=1 is specified, then the corresponding names are 'dhw', 'dh1w' etc. Typing the name of the module without parameters will yield the list and description of the parameters. This program can also be invoked from the ATCOR main panel.

If a DEM file is in the folder of the input scene with the name ending '.ele.bsq' then it will be taken into account in the dehazing algorithm. Then the cirrus band information is not used for high mountains above 2000 m, because the water vapor column is low and the cirrus band signal might be due to ground reflected radiation. However, keyword nodem=1 may be set to ignore a DEM file.

Keyword nocirrus=1 may be set to ignore the cirrus channel info, e.g. in case this channel is corrupt.

If the keyword fraction_haze is not specified, then fh=0.01 (1 percent of scene), i.e. the program will not perform the dehazing, if the fraction of haze pixels is less than fw.

• reslut_batch, sensor='xxx', [aero=aero, h1s=h1s, h2s=h2s, ith=ith] / 

Here 'xxx' is the sensor name corresponding to the '../atcor4/sensor/xxx/' folder. The keyword aero can have the values 'rura', 'urba', mari or 'dese'. If not specified aero='rura' is the default. The keywords h1s and h2s specify the lower and upper flight levels in (km),
e.g. h1s=3 and h2s=5. For the processing of thermal band LUTs the keyword ith=1 has to be set.

• at_derpolish, infile, outfile, nbin, respfile \*.rsp, smooth .., /lowpass, /adj

Derivative polishing routine.

PARAMETERS:
infile: file of reflectances to be filtered
outfile: name of output file to be created
nbin: number of adjacent bands to use for filtering (nbin=1)

KEYWORDS:
respfile: response file used for wavelength reference (default: ENVI header values)
lowpass: perform lowpass filtering only
smooth: smooth the outputs by a lowpass filter of size ‘smooth’ after derivative filtering
adj: use only adjacent bands (excluding current) fro derivatives

• at_pushpoli, infile, outfile, gainfile, spatial .., /nospectral, /zero_offset

Spectral polishing and post-calibration/destriping on the basis of pixel wise deviations from local average.

PARAMETERS:
infile: file of reflectances to be filtered
outfile: name of output file to be created
gainfile: optional output file containing gains and offsets (will be stored)

KEYWORDS:
spatial: use the spatial dimensions with a filter size of the keyword for filtering
nospectral: don’t apply spectral interpolation (by default, spectral interpolation with 3 bands is used to get systematic offsets)
zero_offset: apply linear function through zero (no offset)

• at_prepele, infile, demfile, ofile, [kernelsize .., /fillzero, log=.., /slopasp, /skyview]

Program to resize and prepare a DEM and all its layers for ATCOR processing.

PARAMETERS:
infile: ENVI formatted BSQ file (to be processed with ATCOR)
demfile: ENVI DEM covering the area of ifile (same coordinate system, can be different resolution/extent)
ofile: name of primary elevation file

KEYWORDS:
kernelsize: size of kernel to calculate slope/aspect (default: 5)
fillzero: fill in zero values at edges of image (shrink image)
slopasp: set to calculate slope/aspect files
skyview: set to calculate skyview factor file
• \texttt{at\_rhoapp, infile, calfile, e0solar [outfile, scale= .., zen=.., date=..]}

Apparent reflectance calculation.

PARAMETERS:
calfile: ATCOR calibration file to be used for conversion of the cube
e0solar: File containing the solar irradiance for the sensor (atcor file)
outfile: name of output

KEYWORDS:
scale: scale for processing (same convention as for ATCOR, scale=1.0 is floating point output)
zen: solar zenith angle (default: 0 degrees), in degrees
date: date as two-element array [day,month]

• \texttt{at\_shadowdetect, infile, calfile, e0solar [outfile, solangles= .., slopefile= .., /skyview, min= .., /all\_layers , /pixfilter , range= ..,}

Image based shadow detection/classification.

PARAMETERS:
infile: file to be analysed
calfile: calibration file for current input file
e0solar: solar irradiance file for current input
outfile: output file of the processing

KEYWORDS:
castshadow: if keyword is set, an existing cast shadow file is provided in order to find the shadows through this keyword.
slopefile: use slope and aspect and include them in the illumination.
solangles: solar angles [zenith/azimuth]; only required for slopefile option
skyview: calculate additional skyview layer (based on illumination) requires keyword solangles and slopefile being set
range: range of index (default 0.5-1.0)
min: minimum value of output (default:0)
pixfilter: set to filter single pixels from dark areas (to avoid artifacts in full cast shadows)
all\_layers: write all layers instead of the illumination file only.

• \texttt{at\_shadowfilter, infile, ilufile, outfile[, smfact ¯.., /interp,/meanadjust, min..]}

Filter dark/bright borders of shadows after cast shadow correction.

PARAMETERS:
reflfile: file to be filtered (reflectance file)
ilufile: illumination file containing a shadow mask which had been applied to the image.
outfile: output file of the processing

KEYWORDS:
smfact: width of shadows to be corrected
interp: shadow borders are replaced by interpolations (instead of brightness adjustment)
meanadjust: The mean brightness across all spectral bands is adjusted (not only band wise)
min: minimum threshold for border pixel detection (default: 0.02)
• at_scalefwhm, sensorin, sensout_dir, fwhmdat, meanshift=..

• at_shiftresp, sensorin, sensout_dir, smiledat, meanshift=..

Apply FWHM or smile detection results to a sensor.

PARAMETERS:
sensorin: input sensor definition file sensor*.dat
sensout_dir: output sensor name; to be created, path to directory in sensor directory of atcor installation
fwhmdat/smiledat: files smile_poly_ord4.dat to be applied to sensor

• at_fwhmdetect, incube, dbfile, respfile, resol, outfile, featureflags=../, vis=.., zen=.., ele=.., alti=.., chlist=.., results=.., /spline, zeroborder=[0/1/2], range=.., splitband=..,/overwrite

• at_smiledetect, incube, dbfile, respfile, resol, outfile, featureflags=../, vis=.., zen=.., ele=.., alti=.., chlist=.., results=.., /spline, zeroborder=[0/1/2], range=.., splitband=..,/overwrite

Smile detection routine.

PARAMETERS:
incube: input data cube
dbfile: raw database file to be used for convolution (no height interpolation)
respfile: response file (e.g., band001.rsp)
resol: internal resolution for the calculation
outfile: name of output file for smile coefficients

KEYWORDS:
featureflags: bytarr(n_feat), 15 feature regions, featureflags[i] = 1 if feature is set, else 0;
vis: visibility [km]
zen: solar zenith angle [deg]; zenith at 0 deg.
ele: average ground elevation [km]
alti: flight altitude elevation [km] for pressure compensation
chlist: list of bands which are used for smile detection and for interpolation of the results numbering starting at 0.
results: write idl save dump of all results in a file named *.sav together with the regular output
spline = 1, spline channel interpolation, =0 linear channel interpolation of smile coefficients
zeroborder : 2 set smile coefficients to 0 at spectral borders
(first, last channel) 1 repeat smile coefficients outside of interpolated values
range : search range, default =20 nm
splitband: splitchannel index between two detectors (starting at 0) - first band of second detector. overwrite: silently overwrites the older output

• at_smoothdem, infile, dist, outfile, /median

DEM smoothing routine.

PARAMETERS:
infile: input data cube (single band ENVI image)
dist: size of smoothing filter
outfile: name of output file
KEYWORDS:
median: use median filter instead of default low pass filter.

- bil_2_bsq, infile[, outfile]
- bip_2_bsq, infile[, outfile]

Conversion of ENVI band interleaved by line (BIL) or by pixel (BIP) to ATCOR standard band sequential storage order (BSQ).

PARAMETERS:
infile: input data cube in BIL or BIP format (ENVI image)
outfile: name of output file to be created (optional; default *.img.bsq).

- multires, sensor=sensor, psens=psens, pspec=pspec
  Resampling of multiple spectra stored as separate ASCII files (*.dat') in the path pspec. The spectral channel response functions (*.rsp') are stored in the path psens, and the keyword sensor specifies the sensor name. The ASCII files (*.dat') contain 2 columns: wavelength (µm or nm), spectral value. Empty lines (in between or at the end) are not allowed. The spectrum can have any unit (radiance, reflectance, emissivity, ...). Example:
multires, sensor='xxx', psens='data7/sensor/xxx/', pspec='data7/spectra/'
The resampled spectra are written to a new folder 'data7/spectra/xxx/'.

- multires_slb, sensor=sensor, psens=psens, slbfile=slbfile
  Same as multires, but all input spectra are stored in a single ENVI spectral library file (.slb) and the keyword slbfile specifies this file (with path). Example:
multires_slb, sensor='xxx', slbfile='data7/spectra/field1.slb'
The resampled spectra are written to file 'data7/spectra/field1_xxx.slb'.

- spect_xx, input=input, xc=xc, yc=yc, box=box, spnam=spnam
  Extraction of a reflectance spectrum from a Level-1 image (DN image), file name=input. The spectrum is taken from the center coordinates xc, yc, and averaged over a square box of box*box pixels. The name of the spectrum is specified with the keyword spnam. The corresponding *.inn' file must exist and all parameters for the calculation of the surface reflectance are taken from this file. If box is an even number it is replaced with the next higher odd number to uniquely define the box center. If box is not specified then box=1 is taken.

NOTE: The above spect_xx is a symbolic placeholder. There are four implementations (flat and rugged terrain, satellite and airborne ATCOR)

- spect_a2 : satellite version, flat terrain
- spect_a3 : satellite version, rugged terrain
- spect_4f : airborne version, flat terrain
- spect_4r : airborne version, rugged terrain
So an example of invoking this feature is:

spect_4r, input='/data1/mission2/scene15.bsq', xc=500, yc=600, box=3, spname='target15_n1'

Two ASCII output files will be created in the folder '/data1/mission2/' :
- 'target15_n1.dat' : the surface reflectance spectrum, 2 columns: wavelength, reflectance (%)
- 'target15_n1.rdn' : 4 columns: wavelength, at-sensor radiance, DN, standard dev(DN).
Radiance is in ($mWcm^{-2}sr^{-1}\mu m^{-1}$). The standard deviation indicates the spatial uniformity within the box.

- **angles=sun(month=..,day=.., hour=..,minute=..,lat=..,lon=..)**
  Calculate solar position from parameters; the keyword are self-explanatory. The output 'angles' is an array containing [zen,azi,doy], Zenith angle and azimuth angle in degrees and day of the year as a number.

- **at_cresca, reffile, startpoint, endpoint, fov, [rawdims, scafile = ..]**
  Create scan angle file from parameters.

  **PARAMETERS:**
  - reffile: Reference file for output dimensions
  - startpoint: first point of flightpath (x,y,z)
  - endpoint: last point of flightpath (x,y,z)
  - fov: total FOV of image in degrees
  - rawdims: n_pixels/n_lines of raw image as two element array

  **KEYWORDS:**
  - scafile: name of output scan angle file (default [input]_sca.bsq)

- **gc_glttosca, gltfile, fov, heading, alt, scafile = ..**
  Create scan angle file from GLT.

  **PARAMETERS:**
  - gltfile: GLT reference file in BIL format (or MAP file in BSQ format)
  - fov: total FOV of image in degrees
  - heading: average heading to direction north of flight (north: 0; east: 90)
  - alt: altitude of aircraft for first and last line (two element vector; if only one value is provided it is assumed to be constant)

  **KEYWORDS:**
  - scafile: name of output scan angle file (default [input]_sca.bsq)

- **br_calmodel, filelist, outfile, rfact = .., inpath = .. /roujean, /maignan, /interp, /constraint, /anifurite, smooth = .., scalist = .., chlist = .., accuracy = .., config = .., logfile = ..**
  Do BRDF model calibration based on a list of files.

  **PARAMETERS:**
  - filelist: list of files to process; convention: the following files should be present for each file in
file list:
infile = [*].xxx.bsq (usually *_atm.bsq)
scaang = [*].sca.bsq (path may be changed using inpath keyword)
in = [*].inn (path may be changed using inpath keyword)
outfile: name of output file to be created (default brdf_model.sav)

KEYWORDS:
roujean: calculate Roujean/Ross-Li Model Kernels for geometric kernel
maignan: Use improved hotspot geometry for volumetric kernel
chlist: list of bands to process
rfact: scaling factor to reflectance [-]
accuracy: fitting accuracy threshold [%] anifwrite: write BCI side output (*_bci.bsq)
interp: perform interpolation on incomplete BCI levels (not recommended)
smooth: spectral smoothing of BRDF model (factor to be given as number of bands)
config: standard configurations or array of BCI limits (only required if no brdf model is given)
inpath: path of side inputs (SCA/INN) if different from file.
scalist: list of scan angle files to be used; same order as filelist
logfile: write log to that file

• br_brefcor.infile,scafile,outfile,[brdfsav=.., /anifwrite,/constrain,/roujean,/maignan,/anif]
  Perform BREFCOR brdf correction.

PARAMETERS:
infile: input file
scafile: scan angle file *_sca.bsq
outfile: name of output file

KEYWORDS:
brdfsav: use save set of brdf parameters (default: calculate from this image only)
constrain: constrain correction to vegetation only such that no correction is done for BCI LE -1.2
anifwrite: write anif and BCI side output (*_anif.bsq and *_bci.bsq)
roujean: calculate Roujean/Ross-Li Model Kernels for geometric kernel
maignan: Use improved hotspot geometry for volumetric kernel
chlist: list of bands to process
rfact: scaling factor to reflectance [-]
config: standard configurations or array of BCI limits (only required if no brdf model is given)
logfile: write log to that file

• gc_glttosca, gltfile, fov, heading, alt, scafile = ..
  Create scan angle file from GLT.

PARAMETERS:
gltfile: GLT reference file in BIL format (or MAP file in BSQ format)
fov: total FOV of image in degrees
heading: average heading to direction north of flight (north: 0; east: 90)
alt: altitude of aircraft for first and last line (two element vector; if only one value is provided it is assumed to be constant)
KEYWORDS:
scafile: name of output scan angle file (default [input]_{sca.bsq})
Chapter 7

Value Added Products

As a “by-product” of atmospheric correction a number of useful quantities can readily be calculated. The first group of value added products include vegetation indices (based on surface reflectance instead of at-sensor radiance), simple parametrizations of the leaf area index, and wavelength-integrated reflectance (albedo). The second group comprises quantities relevant for surface energy balance investigations which are a useful supplement for studies in landscape ecology and related fields, e.g., as input for regional modeling of evapotranspiration. These include global radiation on the ground, absorbed solar radiation, net radiation, and heat fluxes. Emphasis is put on simple models based on the reflectance / temperature cube derived during the atmospheric correction. No additional data (with the exception of air temperature) is taken into account. All value added products are written to a file with up to 11 bands. The file structure is band sequential. If the input file name is "example.bsq", the output reflectance file name is "example_atm.bsq", and the value added file name is "example_atm_flx.bsq", the ‘flx’ indicating the most important part of the calculation, i.e., the radiation and heat fluxes.

7.1 LAI, FPAR, Albedo

Many vegetation indices have been introduced in the literature. Only two are presented here, because these are often used for simple parametrizations of the leaf area index (LAI), the fraction of absorbed photosynthetically active radiation (FPAR), and surface energy fluxes (Baret and Guyot 1991, Choudary 1994). The normalized difference vegetation index (NDVI) is defined as

\[ NDVI = \frac{\rho_{850} - \rho_{650}}{\rho_{850} + \rho_{650}} \]  (7.1)

where \( \rho_{650} \) and \( \rho_{850} \) are surface reflectance values in the red (650 nm) and NIR (850 nm) region, respectively. The soil-adjusted vegetation index (SAVI) is defined as (Huete 1988, Baret and Guyot 1991, with \( L=0.5 \) ) :

\[ SAVI = \frac{(\rho_{850} - \rho_{650}) + 1.5}{(\rho_{850} + \rho_{650} + 0.5)} \]  (7.2)

The leaf area index (LAI) can often be approximated with an empirical three-parameter relationship employing a vegetation index (VI=SAVI or VI=NDVI)

\[ VI = a_0 - a_1 \exp(-a_2 \text{LAI}) \]  (7.3)
Solving for LAI we obtain

\begin{equation}
LAI = -\frac{1}{a_2} \ln\left(\frac{a_0 - VI}{a_1}\right)
\end{equation}

(7.4)

Sample sets of parameters are \(a_0=0.82, a_1=0.78, a_2=0.6\) (cotton with varied soil types), \(a_0=0.68, a_1=0.50, a_2=0.55\) (corn), and \(a_0=0.72, a_1=0.61, a_2=0.65\) (soybean) with VI=SAVI (Choudury et al. 1994).

**Note:** Since it is difficult to take into account the parameters for different fields and different seasons it is suggested to use a fixed set of these three parameters for multitemporal studies. Then, the absolute values of LAI may not be correct, but the seasonal trend can be captured.

Plants absorb solar radiation mainly in the 0.4 - 0.7 \(\mu\)m region, also called PAR region (photosynthetically active radiation, ASRAR 1989). The absorbed photosynthetically active radiation is called APAR, and the fraction of absorbed photosynthetically active radiation is abbreviated as FPAR. These terms are associated with the green phytomass and crop productivity. A three-parameter model can be employed to approximate APAR and FPAR (Asrar et al. 1984, Asrar 1989, Wiegand et al, 1990, 1991).

\begin{equation}
FPAR = C[1 - A \exp(-B LAI)]
\end{equation}

(7.5)

Typical values are \(C=1, A=1, B=0.4\). Again, since it is difficult to account for the crop- and seasonal dependence of these parameters, a constant set may be used for multitemporal datasets to get the typical FPAR course as a function of time.

The wavelength-integrated surface reflectance (in a strict sense the hemispherical - directional reflectance) weighted with the global flux on the ground \(E_g\) is used as a substitute for the surface albedo (bi-hemispherical reflectance). It is calculated as:

\[
a = \frac{\int_{0.3 \mu m}^{2.5 \mu m} \rho(\lambda)E_g(\lambda)d\lambda}{\int_{0.3 \mu m}^{2.5 \mu m} E_g(\lambda)d\lambda}
\]

(7.6)

Since most airborne sensors cover only part of the 0.3 - 2.5 \(\mu\)m region the following assumptions are being made for extrapolation. Extrapolation for the 0.30-0.40 \(\mu\)m region:

- \(\rho_{0.3-0.4\mu m} = 0.8\ \rho_{0.45-0.50\mu m}\), if blue a band (0.45-0.50 \(\mu\)m) exists.
- \(\rho_{0.3-0.4\mu m} = 0.8\ \rho_{0.52-0.58\mu m}\), green band, no blue band available.

Extrapolation for the 0.40-0.45 \(\mu\)m region:

- \(\rho_{0.4-0.45\mu m} = 0.9\ \rho_{0.45-0.50\mu m}\), if a blue band (0.45-0.50 \(\mu\)m) exists.
- \(\rho_{0.4-0.52\mu m} = 0.9\ \rho_{0.52-0.58\mu m}\), green band, no blue band available.

The reflectance reduction factors in the blue part of the spectrum account for the decrease of surface reflection for most land covers (soils, vegetation). The extrapolation to longer wavelengths is computed as:

- If a 1.6 \(\mu\)m band exists
  - \(\rho_{2.0-2.5\mu m} = 0.5\ \rho_{1.6\mu m}\), if \(\rho_{850}/\rho_{650} > 3\) (vegetation)
If no bands at 1.6 μm and 2.2 μm are available the contribution for these regions is estimated as:

- \( \rho_{1.5-1.8\mu m} = 0.50 \rho_{0.85\mu m} \), if \( \rho_{850}/\rho_{650} > 3 \) (vegetation)
- \( \rho_{2.0-2.5\mu m} = 0.25 \rho_{0.85\mu m} \), if \( \rho_{850}/\rho_{650} > 3 \)
- \( \rho_{1.5-1.8\mu m} = \rho_{0.85\mu m} \), (else)
- \( \rho_{2.0-2.5\mu m} = \rho_{0.85\mu m} \), (else)

At least three bands in the green, red, and near-infrared are required to derive the albedo product. Wavelength gap regions are supplemented with interpolation. The contribution of the 2.5 - 3.0 μm spectral region can be neglected, since the atmosphere is almost completely opaque and absorbs all solar radiation. The output "flx" file contains the channels SAVI, LAI, FPAR, and albedo coded as 16 bit integer with the following scale factors:

- SAVI: range 0-1000, scale factor 1000, e.g., scaled SAVI=500 corresponds to SAVI=0.5.
- LAI: range 0-10,000, scale factor 1000, e.g., scaled LAI=5000 corresponds to LAI=5.0.
- FPAR: range 0-1000, scale factor 1000, e.g., scaled FPAR=500 corresponds to FPAR=0.5.
- Albedo: range 0-1000, scale factor 10, e.g., scaled albedo=500 corresponds to albedo=50%.

The next section presents a simplified treatment of the radiation and heat fluxes in the energy balance.

### 7.2 Surface energy balance

Surface energy balance is an essential part of climatology. The energy balance equation applicable to most land surfaces can be written as (Asrar 1989):

\[
R_n = G + H + LE
\]  

(7.7)

where, \( R_n \) is the net radiant energy absorbed by the surface. The net energy is dissipated by conduction into the ground (G), convection to the atmosphere (H) and available as latent heat of evaporation (LE). The amount of energy employed in photosynthesis in case of vegetated surfaces is usually small compared to the other terms. Therefore, it is neglected here.

The terms on the right hand side of equation (7.7) are called heat fluxes. The soil or ground heat flux (G) typically ranges from 10% to 50% of net radiation. Convection to the atmosphere is called sensible heat flux (H). It may warm or cool the surface depending on whether the air is warmer or cooler than the surface. The energy available to evaporate water from the surface (LE) is usually obtained as the residual to balance the net radiation with the dissipation terms. Net radiation is expressed as the sum of three radiation components:

\[
R_n = R_{solar} + R_{atm} - R_{surface}
\]  

(7.8)

where \( R_{solar} \) is the absorbed shortwave solar radiation (0.3 - 3 μm, or 0.3 - 2.5 μm), \( R_{atm} \) is the longwave radiation (3 - 14 μm) emitted from the atmosphere toward the surface, and \( R_{surface} \) is the longwave radiation emitted from the surface into the atmosphere. Downwelling radiation is
counted with a positive sign, the upwelling thermal surface radiation has a negative sign. The absorbed solar radiation can be calculated as:

\[
R_{\text{solar}} = \int_{0.3\mu m}^{2.5\mu m} \{1 - \rho(\lambda)\} E_g(\lambda) \, d\lambda
\]  

(7.9)

where \(\rho(\lambda)\) is the ground reflectance, \(1 - \rho(\lambda)\) is the absorbed fraction of radiation, and \(E_g(\lambda)\) is the global radiation (direct and diffuse solar flux) on the ground. The numerical calculation of equation (7.9) is based on the same assumptions regarding the extrapolation of bands and interpolation of gap regions as discussed in chapter 7.1 dealing with the surface albedo. If the airborne imagery contains no thermal band(s) from which a map of ground temperature can be derived, then \(R_{\text{solar}}\) is the only surface energy component that can be evaluated. In case of flat terrain with spatially varying visibility conditions or rugged terrain imagery, a map of the global radiation is included as an additional value added channel.

\[
E_g = \int_{0.3\mu m}^{2.5\mu m} E_g(\lambda) \, d\lambda
\]  

(7.10)

For flat terrain imagery with constant atmospheric conditions the global radiation is a scalar quantity and its value can be found in the "log" file accompanying each output reflectance file. For rugged terrain imagery, the global radiation accounts for the slope/aspect orientation of a DEM surface element.

With thermal bands a ground temperature or at least a ground brightness temperature image can be derived. Then the emitted surface radiation is calculated as

\[
R_{\text{surface}} = \epsilon_s \sigma T_s^4
\]  

(7.11)

where \(\epsilon_s\) is the surface emissivity, \(\sigma = 5.669 \times 10^{-8} \text{ Wm}^{-2}\text{K}^{-4}\) is the Stefan-Boltzmann constant, and \(T_s\) is the kinetic surface temperature. For sensors with a single thermal band such as Landsat TM an assumption has to be make about the surface emissivity to obtain the surface temperature. Usually, \(\epsilon_s\) is selected in the range 0.95 - 1, and the corresponding temperature is a brightness temperature. A choice of \(\epsilon_s = 0.97\) or \(\epsilon_s = 0.98\) is often used for spectral bands in the 10 - 12 \(\mu\)m region. It introduces an acceptable small temperature error of about 1-2°C for surfaces in the emissivity range 0.95 - 1. Examples are vegetated or partially vegetated fields (\(\epsilon = 0.96 - 0.99\)), agricultural soil (\(\epsilon = 0.95 - 0.97\)), water (\(\epsilon = 0.98\)), and asphalt / concrete (\(\epsilon = 0.95 - 0.96\)). Emissivities of various surfaces are documented in the literature (Buettner and Kern 1965, Wolfe and Zissis 1985, Sutherland 1986, Salisbury and D’Aria 1992).

The atmospheric longwave radiation \(R_{\text{atm}}\) emitted from the atmosphere toward the ground can be written as

\[
R_{\text{atm}} = \epsilon_a \sigma T_a^4
\]  

(7.12)

where \(\epsilon_a\) is the air emissivity, and \(T_a\) is the air temperature at screen height (2 m above ground), sometimes 50 m above ground are recommended. For cloud-free conditions, Brutsaert’s (1975) equation can be used to predict the effective air emissivity:

\[
\epsilon_a = 1.24 \left\{ \frac{p_{wc}}{T_a} \right\}^{1/7}
\]  

(7.13)
Here, $p_{wv}$ is the water vapor partial pressure (millibars = hPa), and $T_a$ is the air temperature (K). Figure 7.1 shows $p_{wv}$ as a function of air temperature for relative humidities of 20 - 100%. The partial pressure is computed as:

$$p_{wv} = RH \frac{e_s}{100} \tag{7.14}$$

where RH is the relative humidity in per cent, and $e_s$ is the water vapor partial pressure in saturated air (Murray 1967):

$$e_s(T_a) = e_{s0} \exp\left\{a\frac{(T_a - 273.16)}{T_a - b}\right\} \tag{7.15}$$

The constants are $a = 17.26939$, $b = 35.86$, and $e_{s0} = e_s(273.16K) = 6.1078$ hPa. An alternative to equation (7.13) is the following approximation (Idso and Jackson 1969) which does not explicitly include the water vapor and holds for average humidity conditions, compare Figure 7.2.

$$\epsilon_a = 1 - 0.261 \exp\left\{-7.77 \times 10^{-4} (273 - T_a)^2\right\} \tag{7.16}$$

![Figure 7.1: Water vapor partial pressure as a function of air temperature and humidity. Relative humidities are 20% to 100% with a 10% increment, bottom to top curves, respectively (eq. 7.14).](image)

The calculation of the heat fluxes G, H, and LE on the right hand side of equation 7.7 requires different models for vegetated and man-made surfaces. For vegetated or partially vegetated surfaces, we employ a simple parametrization with the SAVI and scaled NDVI indices (Choudury 1994, Carlson et al. 1995):

$$G = 0.4 R_n \frac{(SAVI_m - SAVI)}{SAVI_m} \tag{7.17}$$

where $SAVI_m = 0.814$ represents full vegetation cover. The sensible heat flux is computed as:

$$H = B (T_s - T_a)^n \tag{7.18}$$

$$B = 286 \left(0.0109 + 0.051 \text{ NDVI}^*\right) \tag{7.19}$$
Figure 7.2: Air emissivity after Brutsaert (eq. 7.13) and Idso/Jackson (eq. 7.16).

\[ n = 1.067 - 0.372 \, NDVI^* \]  
\[ NDVI^* = \frac{\rho_{850} - \rho_{650}}{0.75 \times (\rho_{850} - \rho_{650})} \]

Equation (7.18) corresponds to equation (1a) of Carlson et al. (1995), because G is neglected there, and so \( R_n - G \) represents the energy left for evapotranspiration. The factor 286 in equation (7.19) converts the unit \((\text{cm/day})\) into \((\text{Wm}^{-2})\). \( NDVI^* \) is the scaled NDVI ranging between 0 and 1, and truncated at 1 if necessary. Equation (7.21) corresponds to equation (3) of Carlson et al. (1995) with \( NDVI_0 = 0 \) (bare soil) and \( NDVI_s = 0.75 \) (full vegetation cover). The latent heat flux \( LE \) is computed as the residual:

\[ LE = R_n - G - H \]  

A different heat flux model is employed for urban areas with man-made surfaces (asphalt, concrete, roofs, etc.). These are defined here with the reflectance criteria

\[ \rho_{650} \geq 0.10 \quad \text{and} \quad \rho_{850} \geq 0.10 \quad \text{and} \quad \rho_{850} > \rho_{850} \times 0.7 \]

representing low vegetation indices with \( NDVI < 0.176 \). This simple spectral definition is not unambiguous, it might also apply to soils. For urban areas the latent heat is usually very small, and the fluxes G and H dominate. Therefore, the terms G, LE, and H are approximated by the following three equations (Parlow 1998):

\[ G = 0.4 \, R_n \]  
\[ LE = 0.15 \, (R_n - G) \]  
\[ H = R_n - G - LE \]

For low vegetation indices \((SAVI < 0.1)\) the ground heat flux \( G \) from equation (7.17), i.e., the vegetation model, agrees well with \( G \) from equation (7.24), i.e., the urban model. However, major
differences exist for the LE and H terms, see table 7.1. Parameters for this table are: \( E_g = 800 \), \( R_n = 600 \), \( R_{atm} + R_{surface} = -100 \text{ Wm}^{-2} \), \( T_s = 30^\circ \text{C} \), and \( T_a = 20^\circ \text{C} \). The (veg) and (urb) indicate the heat fluxes derived from the vegetation and urban model, respectively. For the urban surfaces (asphalt, concrete) the G(veg), H(veg) and LE(veg) values are given in brackets for comparison, but the corresponding "urban" heat fluxes are valid because the "urban" criterion (equations 7.23, \( \rho_{650} \geq 0.10 \), \( \rho_{850} \geq 0.10 \), and \( \rho_{650} > \rho_{850} \ast 0.7 \)) applies. The last row repeats the concrete case for \( R_{solar} = 800 \ast (1 - 0.36) = 512 \), \( R_n = R_{solar} + R_{atm} + R_{surface} = 512 - 100 = 412 \text{ Wm}^{-2} \), a realistic reduced \( R_n \) value (compared to the asphalt where \( E_g = 800, R_{solar} = 800 \ast (1 - 0.12) = 700, R_n = 700 - 100 = 600 \text{ Wm}^{-2} \)).

<table>
<thead>
<tr>
<th>surface</th>
<th>( \rho_{650} )</th>
<th>( \rho_{850} )</th>
<th>NDVI</th>
<th>G(veg)</th>
<th>H(veg)</th>
<th>LE(veg)</th>
<th>G(urb)</th>
<th>H(urb)</th>
<th>LE(urb)</th>
</tr>
</thead>
<tbody>
<tr>
<td>full veget.</td>
<td>0.05</td>
<td>0.40</td>
<td>0.78</td>
<td>77</td>
<td>87</td>
<td>435</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>partial veget.</td>
<td>0.10</td>
<td>0.20</td>
<td>0.33</td>
<td>185</td>
<td>76</td>
<td>338</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>dark asphalt</td>
<td>0.11</td>
<td>0.13</td>
<td>0.09</td>
<td>(228)</td>
<td>(50)</td>
<td>(322)</td>
<td>240</td>
<td>306</td>
<td>54</td>
</tr>
<tr>
<td>bright concrete</td>
<td>0.35</td>
<td>0.40</td>
<td>0.07</td>
<td>(222)</td>
<td>(48)</td>
<td>(330)</td>
<td>240</td>
<td>306</td>
<td>54</td>
</tr>
<tr>
<td>bright concrete (*)</td>
<td>0.35</td>
<td>0.40</td>
<td>0.07</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>164</td>
<td>210</td>
<td>37</td>
</tr>
</tbody>
</table>

Table 7.1: Heat fluxes for the vegetation and urban model. All fluxes in [Wm\(^{-2}\)].

All radiation and heat fluxes are calculated in units of \( \text{Wm}^{-2} \). They represent instantaneous flux values. For applications, where daily (24 h) LE values are required the following equation can be used for unit conversion:

\[
LE_{\text{[cm/day]}} = \frac{1}{286} \times LE_{\text{[Wm}^{-2}\text{]}}
\]  

(7.27)

The latent heat flux LE is frequently called evapotranspiration (ET). Although LE and ET are used interchangeably the unit (cm/day) or (mm/day) is mostly employed for ET. For water surfaces the distribution of net radiation into G, LE, and H is difficult to determine, because it depends on several other parameters. Therefore, G and H are set to zero here, and so LE equals \( R_n \).

Spatial maps (files) of air temperature and air emissivity can also be included in the processing. Usually, isolated point-like measurements of air temperature are available from meteorological stations. These have to be interpolated to generate a spatial map coregistered to the image prior to applying the ATCOR model. Data in the file containing the air temperature must have the Celsius unit, data of the air emissivity file must range between 0 and 1. Future improvements to the ATCOR model will include an air temperature map derived from the image ("triangle" or "trapezoidal" method employing the thermal band surface temperature and NDVI, Carlson et al. 1995; Moran et al. 1994).

In case of mountainous terrain, the air temperature \( T_a(z_0) \) and water vapor partial pressure \( p_{wv}(z_0) \) at a reference elevation \( z_0 \) have to be specified. The height dependence of air temperature is then obtained with linear extrapolation employing a user-specified adiabatic temperature gradient \( \partial T/\partial z \):

\[
T_a(z) = T_a(z_0) + \frac{\partial T}{\partial z}(z_0 - z)
\]

(7.28)

where \( \partial T/\partial z \) is typically in the range 0.65 - 0.9 (Celsius / 100 m). The water vapor partial pressure is extrapolated exponentially according to

\[
p_{wv}(z) = p_{wv}(z_0) \ast 10^{-\left(\frac{z - z_0}{z_s}\right)}
\]

(7.29)

where \( z_s \) is the water vapor scale height (default 6.3 km). The list of all output channels of the value added "*.flx.bsq" file is:
1. Soil adjusted vegetation index (SAVI), scaled with factor 1000,
2. Leaf area index (LAI), scaled with 1000,
3. Fraction of photosynthetically active radiation FPAR, scaled with 1000,
4. Surface albedo (integrated reflectance from 0.3-2.5 µm), per cent * 10,
5. Absorbed solar radiation flux $R_{solar} \ [Wm^{-2}]$,
6. Global radiation $E_g \ [Wm^{-2}]$, omitted for constant visibility in flat terrain because it is a scalar which is put into the "*log" file. The next channels are only available in case of at least one thermal band:
7. Thermal air-surface flux difference $R_{therm} = R_{atm} - R_{surface} \ [Wm^{-2}]$,
8. Ground heat flux $G \ [Wm^{-2}]$,
9. Sensible heat flux $H \ [Wm^{-2}]$,
10. Latent heat $LE \ [Wm^{-2}]$,
11. Net radiation $R_n \ [Wm^{-2}]$. 
Chapter 8

Sensor simulation of hyper/multispectral imagery

After atmospheric correction the surface reflectance and temperature/emissivity cubes can be used to simulate new products which might be of interest:

- at-sensor radiance cubes in the solar region (0.4 - 2.5 µm) for different solar geometries and atmospheric conditions,
- at-sensor radiance cubes in the thermal region (8 - 14 µm) for different atmospheric conditions, e.g. for satellite sensor studies,
- resampling of the surface reflectance cube to an existing or new multispectral sensor,
- resampling of the surface emissivity cube to an existing or new multispectral sensor.

This is a convenient way to simulate realistic data for a spaceborne version of an airborne instrument, to obtain radiance data at different flight levels, or to compare hyperspectral (hs) data with broad-band multispectral (ms) data. As a restriction, the TOA (top-of-atmosphere) or at-sensor radiance calculated with the "TOARAD" program assumes a nadir view. The "HS2MS" (hyperspectral-to-multispectral) program requires the hs and ms center wavelengths and the ms channel filter curves for resampling. In addition, noise of the ms sensor can be included as Gaussian noise with a specified amplitude, either as noise-equivalent radiance (NER) or as noise-equivalent reflectance (NEΔρ). The hs contribution to a certain ms band is weighted with the value of the ms response curve at the corresponding hs wavelength, compare Fig. 8.1.

After summing all contributions the result is normalized with the sum of the hs filter values

\[ L_{ms}(i) = \frac{\sum_{k=1}^{n_i} L_{hs}^{(k)} R_{ms}^{(i)}(\lambda_{hs}^{(k)})}{\sum_{k=1}^{n_i} R_{ms}(\lambda_{hs}^{(k)})} \]  

(8.1)

where \( L \) denotes at-sensor (or TOA) radiance, \( R_{ms}^{(i)} \) the ms response function of channel i, and \( n_i \) is the number of hs channels covered by the i-th ms filter function. A similar equation is used for the resampling of surface reflectance or emissivity. The weight factors \( w_k \) for each hs channel are calculated with eq. 8.2, and they are documented in the corresponding "log" file created by program "HS2MS".
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Figure 8.1: Weight factors of hyperspectral bands.
The solid curve shows the response function $R_{\text{ms}}^{(i)}$ of a ms channel, and the dashed lines indicate the hs center wavelengths $\lambda_{\text{hs}}^{(j)}$.

$$w_k(i) = \frac{R_{\text{ms}}^{(i)}(\lambda_{\text{hs}}^{(k)})}{\sum_{j=1}^{n} R_{\text{ms}}^{(i)}(\lambda_{\text{hs}}^{(j)})}$$  \hspace{1cm} (8.2)

Fig. 8.2 describes the sequence of processing for the sensor simulation in the solar region. After atmospheric correction with ATCOR the ”image_atm.bsq” contains the surface reflectance cube. Program ”TOARAD” then calculates the at-sensor radiance for a different flight altitude, solar geometry, or atmospheric parameters. All parameters not specified as keywords (see list of keywords below) are taken from the ”image.inn” file created by ATCOR.

The program ”HS2MS” can be started to resample the radiance cube to a ms image by specifying the ms sensor (i.e., channel filter functions) and the ms noise equivalent radiance (NER). NER = 0 is allowed, so the resampled image product will only include the noise of the hs scene, which will be reduced due to the inherent integration over several hs bands. A channel-constant NER or a file with channel-dependent NER values may also be employed.

Figure 8.3 shows the GUI panel of program ”HS2MS”. Although the input image will usually be a hyperspectral scene with $n > 50$ channels and the output a multispectral scene with $m << n$ channels, this program can also be employed for the case of a multispectral input image with $n < 10$ channels and a panchromatic sensor with $m=1$ band. The program supports the four cases of resampling mentioned above, i.e., solar or thermal at-sensor radiance, surface reflectance, or emissivity.

Fig. 8.4 describes the sequence of processing for the sensor simulation in the thermal region. The file ”image.bsq” consists of thermal band data and possibly also reflective band data. After atmospheric correction, the file ”image.atm.bsq” starts with the surface reflectance channels (if existing), continues with the surface radiance for the thermal channels, and contains the surface temperature as the last channel. The corresponding emissivity data is stored as a separate file (”image_atm.emiss.bsq”, 16 bit integer) which is scaled with a factor of 1,000. Program ”TOARAD2” corresponds to ”TOARAD”, but calculates thermal at-sensor radiance imagery. Again, program ”HS2MS” converts the hyperspectral imagery into multispectral scenes. However, the noise is now
specified in terms of $NE\Delta T$ (noise equivalent delta temperature) which is specified in [K=Kelvin]. For the emissivity resampling, the noise equivalent emissivity $NE\Delta \epsilon$ is internally computed from the $NE\Delta T$:

$$NE\Delta \epsilon = NE\Delta T \frac{\partial L_{bb}}{\partial T}$$

(8.3)

Since the temperature-radiance relationship for thermal bands is calculated with an exponential fit equation (see eq. 10.36)

$$L_{bb} = \exp\left(\frac{1}{T} - a\right)/b$$

(8.4)

we can express $NE\Delta \epsilon$ (using $T=300$ K) as

$$NE\Delta \epsilon = \frac{NE\Delta T}{b T^2} = \frac{NE\Delta T}{b 300^2}$$

(8.5)

where $b$ is a channel-dependent coefficient.

Keywords for batch programs "toarad, toarad2"
On the IDL command line, program names can be written in lower case letters, so as an example "toarad" instead of "TOARAD" is used synonymously in this context. If "toarad" or "toarad2" is submitted as a batch job, the following keywords can be specified:

- **toarad**, **input=filename**, [ **pixelsize=pixelsize**, **sz=solar_zenith**, **atmfile=atmfile**, **elev=elevation**, **vis=visibility**, **adjrange=adjrange**, **scalef=scalef** ]

  The input file name must include the path, and the keywords in brackets indicate optional parameters. A detailed description is given below.

- **toarad2**, **input=filename**, [ **temfile=temfile**, **elev=elevation**, **vis=visibility** ]

Similar to "toarad", temfile is the atmospheric LUT file name for the thermal region. The solar zenith angle is not required in the thermal region. In addition, the pixel size and adjacency range are missing, because the adjacency effect can be neglected. The output thermal radiance is in the unit $\text{mW m}^{-2} \text{ sr}^{-1} \text{ µm}^{-1}$. For ambient surface temperatures the radiance range is from about 5,000 to 15,000 in this unit, and data is always stored as float, i.e., the scale factor is scalef=1. Besides the input "image_atm.bsq" the file "image_atm_emiss.bsq" is automatically loaded as well and all processing parameters will be taken from the "image.inn" generated by the ATCOR run. If the keyword temfile is not specified, the file "h99000_wv10.tem" is used. The default visibility is 23 km.
If a keyword is set, it will overwrite the corresponding parameter from the "inn" file, compare chapters 6.3 and 9.5. To perform a TOA/at-sensor radiance simulation for a given airborne scene, the user has to resample files from the monochromatic atmospheric database:

- for the chosen new airborne altitude to be simulated (if not already available),
- for the altitude 99,000 m that serves as "flight" altitude for space sensors, see chapter 9.1.

After running ATCOR4 for a certain scene and sensor, a surface reflectance cube is obtained which is input to the TOA/at-sensor simulation that can be performed for a flat or a mountainous terrain. A detailed description of the "toarad" keywords follows:

- input='/data1/image_atm.bsq', the '_atm.bsq' indicates a surface reflectance file which is the output of an ATCOR run. The input file to ATCOR was '/data1/image.bsq', and toarad extracts some information from the corresponding file '/data1/image.inn', for example the sensor name. The output file name is '/data1/image_toarad.bsq'.

- atmfile='h99000_wv29_rura', this is an example of an atmospheric look-up-table file with a rural aerosol and a water vapor column of 2.9 gcm$^{-2}$, see chapter 9.1. If the keyword atmfile is not specified, then 'h99000_wv10_rura' will be taken.

- elev=500, an example of a ground elevation at 500 m above sea level. If elev is not specified, then elev=0 is assumed. However, if the keyword elev is not specified and the 'input'
FILE contains file names for the DEM elevation, slope, and aspect, then the DEM files are taken, and the TOA calculation is performed for a rugged terrain. If the keyword `elev` is specified the simulation is always performed for a flat terrain regardless of any possible DEM file names in the ‘*.ini’ file.

- `sz=35.5`, an example of a solar zenith angle of 35.5° (not used for ‘toarad2’).
- `vis=25`, an example of a visibility of 25 km.
- `pixelsz=4.5`, an example of a pixel size of 4.5 m.
- `adjrange=500`, an example of an adjacency range of 500 m.
- `scalef=10,000`, scale factor for the TOA radiance. The default is `scalef=1.0` which provides the output file as float data of TOA radiance in units of \[mW \text{ cm}^{-2} \text{ sr}^{-1} \mu m^{-1}\]. If `scalef > 1`, e.g. `scalef=10,000`, the output TOA radiance is stored as 16 bit unsigned integer multiplied with the scale factor. The advantage is a smaller output file (compared to the 32 bit float), the drawback is that radiances will be truncated at 65,000 which might happen for bright surfaces (e.g., snow, vegetation in the NIR) with `scalef=10,000`, see Figure 8.5. Therefore, the easiest way to avoid scale problems is to use the default `scalef=1.0` and have a float radiance output cube.

![Figure 8.5: TOA radiances for three albedos and a solar zenith angle of 30°.
MODTRAN calculation for a mid-latitude summer atmosphere, rural aerosol, visibility 15 km, ground elevation 500 m above sea level.](image)

For convenience, a ‘*.log’ and an ‘*.ini’ file are created for the documentation of the processing parameters, e.g., ‘/data1/image_toarad.ini’. In addition, the corresponding sensor calibration file
will be created. Example: sensor='casi96', scalef=1000, then file 'casi96_scalef1000.cal' will be created on the '../atcor4/sensor/casi96/' directory with the radiometric calibration coefficient \( c_1 = 0.001 \) for each band.
Chapter 9

Implementation Reference and Sensor Specifics

This chapter discusses miscellaneous topics associated with the current implementation of ATCOR. First the user is acquainted with the structure and handling of the atmospheric database. Second, the supported input/output file types are given. The next item discusses the preference parameters, e.g. the definition of thresholds employed for the masking of cloud and water areas, and options for interpolating certain spectral regions. Then the parameters of the "*.inn" file are described which is employed for the interactive and batch processing. Last but not least a section on problems and tips is included.

9.1 Monochromatic atmospheric database

This chapter presents the technical details of the atmospheric database.

To be capable of handling typical hyperspectral sensors with arbitrary spectral bands in the solar and thermal spectral regions a large database of atmospheric LUTs was compiled with the MODTRAN® R⃝5 radiative transfer code in 2010. The database is called "monochromatic" because of its high spectral resolution, compare figure 9.1. The size is currently about 6.2 GB. After resampling with the spectral response functions of any sensor a typical size of the sensor-specific database is 10-50 MB. Chapter 9.2 contains a description of the resampling program RESLUT.

In the solar spectral region (0.34 - 2.56 µm) MODTRAN® was run with different wavenumber spacings to achieve a wavelength grid spacing of approximately 0.4 nm (except for the 1400 nm and 1800 nm regions). This required the use of MODTRAN®’s "p1_2008" database (i.e., 0.1 cm⁻¹) in the 2.1 - 2.5 µm region. In addition, different RT algorithms were used: in atmospheric window regions the scaled DISORT algorithm with 8 streams (SD-8) was employed, in absorption regions the more accurate SD-8 with the correlated k algorithm was selected [31]. Since the wavenumber grid is not equidistant in wavelength, the LUTs were resampled with an equidistant 0.4 nm grid of Gaussian filter functions of FWHM=0.4 nm to speed up subsequent calculations. So the new LUT database should be sufficient for instruments with bandwidths > 2 nm covering the solar spectral region from 340 to 2540 nm.

The thermal high-resolution database employs a spectral sampling distance of SSD=0.4 cm⁻¹ for the wavelength region 7 - 10 µm, i.e. corresponding to a wavelength SSD=2 - 4 nm, and SSD=0.3 cm⁻¹ for the wavelength region 10 - 14.9 µm, i.e. corresponding to a wavelength SSD=3 - 5.5 nm. A triangular weight function is used with a spectral bandwidth of twice the SSD. The Isaacs’s
2-stream method is employed including the correlated k algorithm. The Isaacs’s algorithm is much faster than DISORT, and yields the same results for our cases in the thermal region. All files ("*.bt7") are calculated for view or scan angles from 0° (nadir) to 40° off-nadir with a 5° increment to enable an accurate interpolation.

![Diagram](image)

**Figure 9.1: Monochromatic atmospheric database.**

The database comprises the flight altitudes 0.1, 1, 2, 3, 4, 5, 10, and 20 km above sea level, and the aerosol types rural, urban, maritime, and desert. In the altitude regime 1 - 5 km the increment is 1 km. A larger increment is adequate for the high altitudes of 10 and 20 km because of the lower optical depth increment. For flight altitudes in between ATCOR will interpolate if requested by the user. In addition, the database contains LUTs for the 99 km altitude which can be used for satellite sensors.

Imagery of the commercially available standard satellite sensors such as Landsat TM, SPOT, or IRS-LISS are already treated in the satellite ATCOR environment.

For each flight altitude and aerosol type files with five water vapor columns are available (W=0.4, 1.0, 2.0, 2.9, and 4.0 cm or g cm$^{-2}$, sea level-to-space values). These represent dry to humid atmospheric conditions [63, 8]. They are needed for the water vapor retrieval to create interpolated / extrapolated values for the range W=0.3 - 4.5 cm. In spectral regions where water vapor absorbs, the accuracy of the surface reflectance retrieval depends on the number of water vapor grid points and the interpolation method (full range of W or sub-interval pertaining to a pixel [70]). The CO$_2$ mixing ratio of the atmosphere is set at 400 ppmv, the ozone column is fixed at 330 DU (Dobson Units, equivalent to the former 0.33 atm-cm) for a ground at sea level.

The file names for the solar region include the altitude, the aerosol type, and the water vapor content. They have the extension ".atm". Example: "h03000_we04_rura.atm" represents a file with flight altitude 3000 m (above sea level), water vapor column 0.4 cm, and the rural aerosol. Each ".atm" file contains the look-up tables for the visibility range 5 - 120 km, solar zenith angles 0 - 70°, and ground elevations 0 - 2500 m (increment 500 m). If the flight altitude is lower than 2500 m the maximum ground elevation is set to 100 m below flight altitude.

In the solar spectral region the database was compiled for a nadir view. However, the atmospheric transmittance will be calculated in ATCOR depending on the actual scan angle for each image pixel employing the nadir value. For the path radiance, the nadir value is not sufficient to predict the off-nadir values. Therefore, the scan-angle dependence of the path radiance was approximated by 2nd order polynomials obtained by a least squares fit. The polynomial coefficients depend on the aerosol type, view and illumination geometry, and wavelength. In general, they are different for the left and right part of a scanline.

The extension of the files for the thermal spectral region is ".tem". Since the aerosol type is of negligible influence in the thermal region, these files do not include an aerosol type identifier in
their name. They were compiled for the rural aerosol. So, the file name corresponding to the 3000 m altitude and the 0.4 cm water vapor column is called "h03000_wv04_tem". The thermal LUTs were compiled for the scan angle range 0 - 40° with an increment of 5° to keep the interpolation error of the radiance and transmittance values smaller than one percent. Transmittance and path radiance values for scan angles above 40° will be extrapolated.

9.1.1 Database update with solar irradiance

In the solar region any high spectral resolution database of LUTs is based on the specification of an extraterrestrial spectral solar irradiance, because the values of path radiance, direct and diffuse solar fluxes depend on solar irradiance. Other quantities (direct and diffuse atmospheric transmittances and spherical albedo) are independent of the solar spectrum. ATCOR’s standard atmospheric database is calculated for a certain irradiance $E_1(\lambda)$, and the corresponding file 'e0_solar_xxx.dat' is included in the directory 'atm_database'.

Beginning with the ATCOR-2011 release there is an option to switch from one extraterrestrial solar irradiance source $E_1(\lambda)$ to another one $E_2(\lambda)$. The delivered high spectral resolution database of atmospheric LUTs is based on the Fontenla-2011 solar irradiance spectrum (Fontenla et al. 2009, 2011 [20, 21]). It represents the solar irradiance for a quiet or “low activity” sun and is recommended as the standard spectrum. The original 0.1 cm$^{-1}$ resolution spectrum is convolved with Gaussian filter functions (FWHM=0.4 nm) and mapped on an equidistant 0.4 nm grid. The file name of this spectrum $E_1(\lambda)$ is 'e0_solar_fonten2011_04nm.dat'. If $R_1$ denotes the set of quantities (path radiance, direct, diffuse solar flux) based on $E_1(\lambda)$, then the new set $R_2$ with the irradiance spectrum $E_2(\lambda)$ is calculated as:

$$R_2(\lambda) = R_1(\lambda) \frac{E_2(\lambda)}{E_1(\lambda)}$$  \hspace{1cm} (9.1)

Figure 9.2 presents a schematic sketch of this conversion. The folder 'sun_irradiance' contains a number of solar irradiance files that can be selected. The folder of the atmospheric database $DB_1$ pertaining to $E_1(\lambda)$ includes the corresponding irradiance file (e.g. 'e0_solar_fonten2011_04nm.dat') and the calculated new database $DB_2$ includes the $E_2(\lambda)$ file (e.g., 'e0_solar_kurucz2005_04nm.dat'). The standard or "active" database is named 'atm_database', while the new database includes 10 characters from the $E_2$ file name, e.g. 'atm_database_kurucz2005'.

![Figure 9.2: Solar irradiance database.](image)
The ATCOR "tools" panel contains the program to convert from one to another spectral irradiance database (see Figure 9.3). It enables an update of the "monochromatic" atmospheric database without the need to repeat the time-consuming MODTRAN® 5 computations (involving the correlated k algorithm in some spectral regions). Figure 9.3 shows an example for a reduced height set of files, the complete conversion for all height levels requires about 3 minutes. The user can also provide additional solar irradiance files to the 'sun_ irradiance' folder provided the spectral range, increment, and irradiance unit agree with the template spectra.

Attention: ATCOR will always work with files in the "active" folder 'atm_database', therefore the old 'atm_database' has to be renamed (or deleted) and the folder with the new database has to be renamed as 'atm_database' before applying the sensor-specific resampling program (RESLUT). Since each 'atm_database' folder contains its corresponding solar irradiance spectrum a unique identification is always possible. Previously generated channel-resampled "*.atm" files do not have a reference to their solar irradiance file, but they are based on the 'e0_solar_kurucz1997_06nm.dat' irradiance. Beginning with the 2011 release the directory of the "*.atm" files contains an ASCII file named 'irradiance_source.txt' identifying the underlying solar irradiance file.

![Figure 9.3: User interface to convert database from one to another solar irradiance.](image)

### 9.2 Sensor-specific atmospheric database

This database is created by resampling the files of the monochromatic database with the sensor’s spectral response functions employing program RESLUT, see figure 9.4. Usually, only a subset of the total number of files of the monochromatic database is needed for a new sensor. The sensor-specific database will contain the altitudes and aerosol types that were specified during the resampling with program RESLUT. There are 5 water vapor files per altitude and aerosol type.
that have to be resampled. The folder with the ".atm" files also contains a file ".irrad_source.txt" identifying the underlying solar irradiance spectrum.

Attention: Do not resample all flight altitudes and aerosol types contained in the monochromatic database, because the program might run an hour, but restrict the resampling to the cases you need. If you want to resample a larger number of files let the program run over night.

Example: A sensor operates exclusively in the altitude region from 1 - 3 km in areas where rural and maritime aerosols are likely to occur. In this case, it makes no sense to resample the atmospheric files for the altitude regions above 3 km. Also, a restriction to the files with rural and maritime aerosols will reduce the computation time of the RESLUT program considerably.

In case of sensors with solar and thermal bands the option "Thermal Region" on the first GUI panel of RESLUT has to be selected as well. The second GUI panel will look slightly different, because the aerosol type selection is missing. In the thermal region, the aerosol type can usually be neglected. Therefore, all files in the monochromatic database were calculated with the rural aerosol, but the aerosol identifier is not included in the file name, because only the rural aerosol is available.

Note: Any altitude-interpolated files that were created by ATCOR will be deleted as soon as the user exits the program. This is done intentionally to avoid "garbage collection". Any interpolated file is generated again within a few seconds if necessary. Interpolated ".atm" files for the solar region have the extension ".atmi", for the thermal region the interpolated ".tem" files have the extension ".temi".

**Figure 9.4:** GUI panels of program RESLUT.

### 9.2.1 Resample sensor-specific atmospheric LUTs with another solar irradiance

It is also possible to resample existing sensor-specific LUTs (".atm" files) with another solar irradiance spectrum. Input is a sensor from the "atcor/sensor/" folder (example name 'xxx') with the corresponding spectral response files (".rsp") and a high-resolution solar irradiance file from the "atcor/sun_irradiance/" directory (example: 'e0_solar_kurucz2005_04nm.dat').
Output is a new sensor subdirectory (example: "sensor/xxx_kurucz2005/") where the first 10 characters of the 'e0_solar_kurucz2005_04nm' (starting after the 'e0_solar_') are appended to the input sensor name. The contents of the input 'atcor/sensor/xxx/' are copied to the output directory, 'e0_solar_xxx.spc' is deleted (in the output directory) and replaced by the new 'e0_solar_xxx_kurucz2005.spc'. A comparison of 'e0_solar_xxx.spc' with 'e0_solar_xxx_kurucz2005.spc' shows the influence of the change of the irradiance spectrum. In addition, a new 'atm_lib/xxx_kurucz2005/' is created where all the LUTs '.atm' from the input 'atm_lib/xxx/' are replaced with the resampled selected irradiance spectrum. This new folder also contains a file "irrad_source.txt" identifying the selected irradiance source.

9.3 Supported I/O file types

Below, a list is compiled of all input and output files to the atcor main process.

9.3.1 Main Input

The input image to ATCOR-4 must have the band sequential (BSQ) ENVI format. Several data types exist for the encoding. The following data types of an input image are supported:

- byte, or unsigned 8 bit integer, ENVI data type = 1;
- signed 16 bit integer, ENVI data type = 2;
- unsigned 16 bit integer, ENVI data type = 12;
- signed 32 bit long integer, ENVI data type = 3;
- float (32 bit), ENVI data type = 4.

9.3.2 Side inputs

Before running the main processor, certain files have to be provided (the DEM derived files are not required for a flat terrain):

**Parameter file** extension: *.inn

This file is written automatically while using the graphical user interface. It may also be written externally for use in batch processing. See Section 9.4 for a detailed description.

**Scan angle file** extension: *[image]*_sca.bsq

Data layers defined in map geometry or in raw geometry with scan zenith, scan azimuth angle, and the absolute distance from aircraft to the pixel for each geocoded pixel. The first band of this image is used as standard input for ATCOR-4 radiometric processing whereas the second band is only required for a potential future BRDF correction. Only band 1 is mandatory for use with ATCOR-4.

Format: 3 bands 16-bit integer ENVI image.

1: Sensor zenith angle (degree * 100), 0 deg. is downward looking (nadir),
- unknown/invalid/background values are greater than 9000 (assigned value: 9100),
- pixels to the right* with respect to the flight direction have negative values
this is sometimes confusing in raw data formats as it depends on the scan direction of the instrument...
2: Absolute azimuth angle to direction north (degree * 10, unknowns are lower zero)
3: Height of airplane above ground for each pixel [meters], unknowns are 0
4: Optional: pixel distance: distance in meters from aircraft to each pixel.
   Optional: Relative azimuth angle to flight direction (would be in channel 2 while shifting
   channels 2/3 to 3/4)

Elevation file (DEM) extension: [dem]_ele.bsq
   Elevation file containing the digital terrain model used for processing.
   Format: single band ENVI file in either integer, unsigned integer, or floating point format.
   Default unit: meters.

Slope file extension: [dem]_slp.bsq
   Slope angle for each image pixel.
   Format: single band ENVI file byte or 16-bit integer or float data type. Unit: degrees.

Aspect file extension: [dem]_asp.bsq
   Aspect angle for each image pixel with respect to north.
   Format: single band ENVI file 16-bit integer or float data type. Unit: degrees.

Skyview factor file extension: [dem]_sky.bsq
   Skyview Factor
   Format: single band ENVI file byte data type. Unit: % (percentage 0-100)

Cast shadow mask extension: [dem+angles]_shd.bsq
   Shadow flag (0: cast shadow, 2: no shadow, 1: border region) for each image pixel.
   Format: single band ENVI file byte data type.

Illumination file extension: [image]_ilu.bsq
   External Illumination file, Format: ENVI floating point data file - external illumination files
   are only read if they are in floating point format and if they are in the same size as the
   imagery. Contents: a value of zero is a complete cast shadow whereas a value of 1 is in
   full illumination, corresponding to the definition of the \( \cos(\beta) \) of the incidence angle, where
   \( \beta = 0 \) deg is a 90 degree incidence direction.

Calibration file extension: [sensor].cal
   Calibration file, containing wavelength, c0 and c1 for each spectral band for conversion of
   image data to calibrated radiance \( L = c0 + c1 \ast DN \), units \( mW/(cm^2 sr \mu m) \)
   Format: 3 column ASCII, one header row

Response Function extension: [bandxxx].rsp
   Spectral response function for one spectral band with wavelength reference and relative re-
   sponse
   Format: 2 column ASCII, no header

Sensor Description extension: sensor_[type].dat
   Sensor description file as created by the Function ‘File:New Sensor’ Format: ASCII.

Solar Reference extension: e0_solar_[type].dat
   Solar reference function for the sensor as created when compiling the atmospheric LUT for a
   sensor. Format: ASCII, three columns: center wavelength, bandwidth, extraterrestrial solar
   irradiance.
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9.3.3 Main output

The default output image data type is byte if the input is byte data. Then a scale factor \( s=4 \) is employed, i.e., the per cent reflectance value of each pixel is multiplied with \( s=4 \) and rounded to byte. As an example, a surface reflectance value of 20.2\% will be coded as 81. However, the user can modify the scale factor (on ATCOR’s main panel). A value of \( s=10 \) to \( s=100 \) causes the output file to be coded as signed 16 bit integer, i.e., with two bytes per pixel. The specification \( s=1.0 \) produces a float output image, i.e., with 4 bytes per pixel.

**Attention:** The float output could be used for testing on small images. For large files and an input data type of 2 bytes per pixel the output float image would require twice the disk space of the input image.

The default output data type is signed 16 bit integer for all integer (and float) input data employing the scale factor \( s=100 \). The scale factor is always included in the output ENVI header file.

**Note:** Any positive value of the scale factor \( s \) is accompanied with a truncation of surface reflectance values at 0 in the output cube. So a negative reflectance, e.g., caused by a wrong choice of visibility or inaccurate radiometric calibration, will be reset to zero in the output image. In the "SPECTRA" module no truncation is applied. If a user wants the output reflectance cube without zero-truncation, the scale factor \( s \) should be specified with a negative value, e.g., \( s=-1 \) will provide a float output surface reflectance retaining negative reflectance values, \( s=-100 \) will provide a 16 bit integer output file. The byte scale (\(-10 < s < -1\) and output data range 0-255) cannot be used to represent negative values. The negative scale factor should only be used for test purposes, since the results do not make a physical sense, and some further processing options or modules are excluded in this case, e.g. the value added calculation of surface energy balance components, the automatic spectral classification (SPECL), BRDF corrections, or top-of-atmosphere radiance (TOARAD).

Summary of output data types:

- byte (default surface reflectance scale factor = 4.0)
- 16 bit signed integer (scale factor \( \geq 10.0 \), typically 10 or 100)
- float (scale factor = 1.0)

9.3.4 Side outputs

A number of side outputs is written by default after successful execution whereas some outputs are optional (marked with an '*').

**Log file** Name: [outputname].log  
Standard log file containing all necessary information about the data processing.  
Format: ASCII

**Error log file** Name: atcor_error.log  
An error log file is written whenever an uncaught error occurred. Please send this file to the software supplier in case of malfunction.  
Format: ASCII

**Aerosol optical thickness** Name: [outputname]_atm_aot.bsq  
Aerosol optical thickness map (scale facor 1,000).  
Format: 1 channel, binary, 16-bit signed integer
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**DDV classification** Name: [outputname]_atm_ddv.bsq
Classification of dark dense vegetation, cast shadows, and water pixels map used for aerosol retrieval.
Format: 1-band ENVI byte file containing the three classes

**Visibility index** Name: [outputname]_atm_visindex.bsq
Index of Visibility used for atmospheric correction (if variable aerosol distribution has been selected as processing option).
Format: 1-band ENVI byte file containing the indices

**Water vapor** Name: [outputname]_wv.bsq
Columnar amount of water vapor calculated from image
Format: ENVI single band file, 16-bit signed integer; unit: cm (i.e., g/cm²)*1,000, i.e. scale factor 1,000.

**Illumination** Name: [outputname]_ilu.bsq
Illumination file created during processing (in case no external floating *_ilu has been provided).
Format: Scaled illumination in byte format from 0 to 100%, single band.

**Haze/cloud/water** Name: [outputname]_out_hcw.bsq
Haze, cloud, and water mask.
Format: ENVI byte image; classes.

**Fractional shadow** Name: [outputname]_fshd.bsq
Fractional cloud/building shadow (optional)
Format: ENVI Integer image; Scaled shadow between 0 and 1000.

**Diffuse irradiance** Name: [outputname]_edif.bsq
Diffuse irradiance component, 1 band, 16-bit signed integer, unit (Wm⁻²nm⁻¹).

**Direct irradiance** Name: [outputname]_edir.bsq
Direct irradiance component, 1 band, 16-bit signed integer, unit (Wm⁻²nm⁻¹).

**Global irradiance** Name: [outputname]_eglo.bsq
Global irradiance (sum of edir and edif).
1 band, 16-bit signed integer, unit (Wm⁻²nm⁻¹).

**Value Added Vegetation** Name: [outputname]_atm_flx.bsq
Multi Layer file containing side outputs for vegetation, i.e. flux, fapar, savi, etc..
Format: ENVI signed 16-bit integer with scale factors as specified in the header.

### 9.4 Preference parameters for ATCOR

The preference parameters are located in a user specific HOME directory "~/.idl/rese/atcor4/", so multiple users of the same license can retain their personal preference settings. The settings of the last input image is saved separately together with GUI-related parameter settings in a save file called "atcor4_settings.sav". In addition, the file "preference_parameters.dat" contains a number of default parameters that can be adjusted to scene properties. This file contains the parameters with a short description; a graphical view of the preferences can be found from the menu ¿File:Edit Preferences¿.

More details are given here.
A choice to set the water vapor values for water pixels:
1 = average water vapor value of land pixels is assigned to water pixels,
Option available for iwv_model = 1 and iwv_model = 2, see section 9.5.
2 = line average of water vapor of land pixels is assigned to water pixels.
Option only available with iwv_model = 1, see the job control parameter section 9.5.

A cloud reflectance threshold $T_c$ (%) in the blue-green region to define a cloud mask.
Pixels belong to the cloud mask if:
(a1) $\rho^\ast(\text{blue}) > T_c$ or (a2) $\rho^\ast(\text{green}) > T_c$ (asterisk: apparent reflectance)
Typical values for $T_c$ range from 15 - 35% .
If the cloud reflectance threshold is too high, clouds will be included in the haze mask.
This will reduce the performance of the haze removal algorithm.

A surface reflectance threshold $\rho_{w1}$ (%) for water in the NIR band.
Pixels belong to the water mask if $\rho(\text{NIR}) < \rho_{w1}$ (only NIR band available).

A surface reflectance threshold $\rho_{w2}$ for water in the 1600 nm region (if band exists).
Pixels belong to the water mask if $\rho(\text{NIR}) < \rho_{w1}$ and $\rho_{1600} < \rho_{w2}$.
The defaults $\rho_{w1} = 5\%$ and $\rho_{w2} = 3\%$ allow some margin for turbid water.

interpolate bands in 760 nm oxygen region (0=no, 1=yes)
interpolate bands in 725 / 825 nm water region (0=no, 1=yes)
interpolate bands in 940 /1130 nm water region (0=no, 1=nonlinear, 2=linear)
smooth water vapor map (box 50m*50m) (0=no, 1=yes)
interpolate bands in 1400/1900 nm nm water region (0=no, 1=yes)
cut-off limit for max. surface reflectance (default 150%)
".out_hcw.bsq" file (haze/cloud/water/land) (0=no, 1=yes, 2=hcw + quality file)
water vapor threshold to switch off the cirrus algorithm, unit [cm]
define saturation with factor b: $DN(\text{saturated}) \geq b \times DN(\text{max})$, b=0.9 to 1.
include non-linear influence of vegetation in water vapor calculation (yes/no)
Only for water vapor retrieval with regression (iwv_model=2)
start/stop wavelengths for interpolation in the 940 nm region
start/stop wavelengths for interpolation in the 1130 nm region
start/stop wavelengths for interpolation in the 1400 nm region
start/stop wavelengths for interpolation in the 1900 nm region
haze/sun glint over water: (apparent) NIR reflectance $T_1(\text{clear})$, $T_2(\text{haze})$
reduce over/under-correction in cast shadow (0=no, 1=yes)
keep ".atmi" files in sensor-specific "atm_lib" (0=no, 1=yes)
adapt path radiance for blue bands (visib > 0, no DDV pixels) (0=no, 1=yes)

Note on the non-linear influence of vegetation in water vapor calculations:
This option applies to the APDA-regression algorithm and only if the 940 nm region is selected
for the water vapor retrieval. The retrieval is based on a linear interpolation across the absorption
region, and errors can occur due to the non-linear behavior of the reflectance of vegetated surfaces
in this region. A simple empirical correction to the water vapor map $W$ is applied using the NDVI
calculated with the apparent reflectances in the red/NIR channels:

$$W(\text{new}) = W(\text{old}) - 0.1 \times (NDVI/0.7) \quad (cm)$$

(9.2)
The correction is only performed for pixels with $NDVI \geq 0.25$ and values $NDVI > 0.7$ are reset to 0.7.

**Note on cut-off limit for surface reflectance**

The default limit is set to a high value (150% reflectance) to avoid a truncation in spectra of high reflectance surfaces, e.g. snow and/or surfaces with specular reflectance. The previous default cut-off (before 2012) was set to 90%.

**Note on factor b:**

Factor b is a relative saturation factor applied to the maximum radiometric encoding, e.g. for 8 bit data and b=1 all pixels with $DN=255$ will be marked as saturated in the "image.hcw.bsq" file (color coding: red). Setting b=0.9 implies pixels with $DN > 230$ will be considered as saturated or in the non-linear radiometric range close to saturation. This factor is only used for 8 and 16 bit (signed or unsigned) data, not for float or 32 bit data.

**Note on the cloud mask:**

The criterion (a1) or (a2) is also coupled with the conditions: $\rho^*(NIR)/\rho^*(red) < 2$ and $\rho^*(NIR)/\rho^*(SWIR1) > 1$ and $NDSI < 0.7,$ where NDSI is the normalized difference snow index. A quality or probability mask for clouds is generated with the above three conditions and different apparent reflectance thresholds $T_c$ in the blue/green spectral region. For $T_c = 15\%$ we define a low probability cloud, $T_c = 25\%$ a medium probability, and $T_c = 35\%$ a high probability cloud. The result is put into a cloud map file named "image_quality.bsq" if the input scene is named "image.bsq". The cloud mask in the "image_out.hcw.bsq" file is based on the user-defined threshold $T_c$ (in the preference parameter file) and the above three conditions.

The band interpolation options are only intended for hyperspectral imagery. Linear interpolation is employed in the 760, 725, and 825 nm regions. Non-linear interpolation as a function of the vegetation index is applied in the 940 and 1130 nm parts of the spectrum to account for the leaf water content in plants. Interpolation in the strong atmospheric water vapor absorption regions around 1400 nm and 1900 nm is recommended because of the low signal and large influence of sensor noise. However, interpolation can be disabled if required, i.e., for test purposes. If enabled, non-linear interpolation is performed in the 1400 / 1900 nm regions by fitting the surface reflectance curves with a hull of a template vegetation or soil spectrum. All interpolated channels are marked with an '*' in the ENVI header of the output reflectance cube.

Haze or sunglint removal over water: the default apparent reflectance thresholds in the NIR channel for clear water and haze are $T_1(\text{clear}) = 0.04$ (or 4%) and $T_2(\text{haze}) = 0.12$ (or 12%), respectively. Pixels with values less than $T_1(\text{clear})$ are defined as clear water, pixels with values between the thresholds $T_1(\text{clear})$ and $T_2(\text{haze})$ are assigned as haze (or sunglint). A lower value (i.e., $T_1(\text{clear}) < 0.04$) can be be specified, but might cause a wrong classification of bright coastal water (sandy bottoms and bleached coral waters). If the threshold $T_1(\text{clear})$ is too high, haze pixels might erroneously be put into the clear water category.

Cast shadow areas (mountainous terrain): these may contain over- and/or undercorrected pixels during the standard empirical BRDF correction. A reduction of these pixels is tried with the following steps:

- bright pixels in the NIR with $DN > DN(mean) - \sigma$, $\sigma=$standard deviation, are removed from the list of shadow pixels.
• a 7 x 7 pixel convolution filter is being applied to the shadow mask.

• a transition region shadow/sunlit is introduced.

• already dark pixels ($DN < DN_{\text{mean}} - \sigma$ in the NIR) are not reduced in brightness during BRDF correction. The threshold is evaluated for a NIR channel, but the non-reduction of brightness (reflectance) is applied to all channels.

Note on keeping the ‘*.atmi’ files

In the interactive mode, the temporary ‘*.atmi’ files in the sensor-specific sub-directory of the ‘atm_lib’ are deleted to avoid an accumulation of these temporary files. However, in the batch mode it might be necessary to keep these files if subsequent runs with the same flight altitude exist: these need the files and they might have been deleted by the previous run. Then, the user should delete the ‘*.atmi’ files after processing the imagery of the flight campaign. The GUI ‘Edit Preferences’ should be used to change the default setting.

Note on ”adapt path radiance for blue bands”

This flag is set to 1 (enabled) as a default. It is only used if the option of constant scene visibility is selected, and if the visibility in the ”*.inn” file is positive. Then the program might change the input visibility if negative reflectance pixels (in the red and NIR bands) occur, while a negative visibility in the ”*.inn” file will not be changed even in case of negative surface reflectance values. Therefore, if this flag is set and the constant (positive) visibility is selected, the following steps are executed:

• Iteratively increase the visibility in the shortest wavelength blue band (up to vis=120 km), if negative surface reflectance pixels occur.

• Re-scale the path radiance using the shortest blue band and the red band (0.66 $\mu$m) as tie bands, using exponential interpolation.

• If no negative surface reflectance pixels occur (in the shortest blue band, for the selected input visibility) then search for dark pixels with ($NDVI > 0.4$, $\rho(NIR) > 0.1$).

• If the percentage of dark pixels is less than 4% of the scene, then terminate, else continue:

• Calculate the surface reflectance $\rho(\text{blue})$ and $\rho(\text{red})$ for these dark pixels.

• If $\rho(\text{blue,average}) > \rho(\text{red,average})$ then increase the visibility iteratively (until vis=120 km).

• Re-scale the path radiance with the final visibility using the shortest blue band and the red band (0.66 $\mu$m) as tie bands, using exponential interpolation.

9.5 Job control parameters of the ”inn” file

If the file name of the input image is ”example_image.bsq” then a file ”example_image.inn” is created during the interactive ATCOR session. When all image processing parameters have been defined by the user this ”inn” file is written to the directory of the corresponding image. When the image is re-loaded during a later session, or when a batch job is submitted, all input parameters are read from this file. It is suggested not to edit this file, because it might create inconsistent input data. The file might contain empty lines when input is not required for a specific case. The IDL routine
for writing this file is available on request for users who want to run ATCOR batch jobs without employing the interactive GUI panels. The contents of an "inn" file are:

**line 1:**
20/08/1989 Date (dd/mm/year) i.e., 20/08/1989 occupies the first 10 columns.

**line 2:**
100.0 ; scale factor reflectance (0-100% range)

The default reflectance scale factor in the output file is $s=100$ for 16 bit data and $s=4$ for 8 bit data, i.e. a reflectance value of say 21.75% is coded as 2175 and 87 for $s=100$ and $s=4$, respectively. For 8 bit input data and $s=4$ reflectance values above 63.75% are truncated at 255. If higher values are expected, a scale factor of $s=100$ should be selected. For $s=1$ the output file will be coded as 32 bit (float) data requiring twice the disk storage as for $s=100$. With $s > 0$, negative reflectance values are truncated to 0. With $s=-100$ or $s=-1$, the negative values are not truncated. This option may sometimes be useful to identify areas in the scene where overcorrection exists. This is usually caused by a too low visibility or in shadow areas.

**line 3:**
5.0 ; pixel size [m]

**line 4:**
casi04, casi04, ; name of "/atcor4/sensor/casi04" sub-directory, twice, separated by a comma and space

**line 5:**
1.0 ; gain setting
Any positive value is accepted, this gain setting $g$ is used to replace the $c_1$ in the corresponding ".cal" file with $c_1/g$, where $g$ is the same for all channels.

**line 6:**
calfile ; calibration file name

**line 7:**
scapix ; scan angle file "*.sca", empty line, if image is not geocoded

**line 8:**
0.9500, 0 ; iemiss, dem_unit (surface emissivity, DEM height unit)
iemiss = surface emissivity option or value, disregarded if no thermal band exists.
iemiss = 0 invokes $\epsilon = 0.98$ to be consistent with the definition of earlier ATCOR versions. Since iemiss=1 is reserved for the cover-dependent emissivity setting below, $\epsilon=1.0$ has to be approximated as iemiss=0.999 or iemiss=0.9999.
In case of multiple thermal bands this $\epsilon$ holds for the thermal band $itemp$ employed for the surface temperature evaluation, see chapter 9.2.
iemiss = 1 : fixed values of surface emissivity = 0.98 (water), 0.97 (vegetation), 0.96 (soil).
iemiss = 2 : surface emissivity map calculated with SPECL, compare chapters 10.1.5, 2.5.
iemiss = 3 : NEM or ANEM method, requires multiple thermal bands, see chapter 10.1.5.
iemiss = 4 : ISAC method, requires multiple thermal bands, see chapter 10.1.5.
iemiss = 5 : both NEM and ISAC, but ISAC is currently only supported for flat terrain imagery.
dem_unit : 0 = [m], 1 = [dm], 2 = [cm] DEM height unit
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line 9:
fele ; DEM elevation file name (empty line for a flat terrain calculation)

line 10:
fslp ; DEM slope file name (empty line for a flat terrain calculation)

line 11:
fasp ; DEM aspect file name (empty line for a flat terrain calculation)

line 12:
fsky ; DEM skyview file name (empty line for a flat terrain calculation)

line 13:
fshd ; DEM cast shadow file name (empty line for a flat terrain calculation, rugged terrain: empty if calculated ”on-the-fly”)

line 14:
atmfile ; atmospheric LUT file name (reflective region)

• If the automatic aerosol type retrieval is intended for batch jobs, the usual aerosol identifier in the file name (e.g. ‘rura’) has to be replaced with ‘auto’. Example: file name (without path) is ‘h03000_wv10_rura.atm’, replace it with ‘h03000_wv10_auto.atm’. The program then uses all aerosol types for the 3000 m altitude in the aerosol type estimate, and selects the one with the closest match, compare chapter 10.4.2. Of course, the user has to provide the desired aerosol types using the RESLUT program. If only one aerosol type is available for the specified altitude, the program will use this atmosphere and quit the automatic aerosol mode. The automatic aerosol type retrieval requires the parameter npref=1 (variable visibility, see line 20 below). If npref=0, it is reset to npref=1.

In the interactive mode the user can just press the ‘Aerosol Type’ button on ATCOR’s main panel to execute the aerosol type retrieval, irrespective of the ‘*.atm’ name in the ‘.inn’ file.

line 15:
temfile ; atmospheric LUT file name (thermal region, empty if no thermal band)

line 16:
1.0 ; adjacency range [km]

line 17:
35.0 ; visibility [km]

line 18:
0.7 ; mean ground elevation ([km asl] not used in case of rugged terrain, where elevation file applies

line 19:
33.0, 178.0 ; solar zenith, solar azimuth angle [degr.]

line 20:
4.2, 230.0 ; flight altitude [km asl], heading [deg. north]

line 21:
0, 0, 1, 0, 0, 0, 1 ; npref, iwaterwv, ihaze, iwat_shd, ksolflux, ishadow, icl_shadow
seven parameters controlling the processing options:
npref = 0 : constant visibility
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npref = 1 : variable visibility, based on dark reference areas in the scene
npref = -1 : variable visibility for each sub-image during batch job with tiling
iwaterwv = 0 : no water vapor correction or no water vapor bands available
iwaterwv = 1 : water vapor correction using bands in the 940 nm region
iwaterwv = 2 : water vapor correction using bands in the 1130 nm region
iwaterwv = 3 : 940 and 1130 nm bands are employed

- Haze removal is enabled by setting the parameter $ihaze>0$, no haze removal is specified with $ihaze=0$. Separate parameter values define haze removal over land, haze/sun glint removal over water, and the combination. Some criteria exist to check whether the haze/land removal is likely to yield good results. The haze/land algorithm is switched off if those criteria are not passed ($ihaze=1$). However, as these criteria might fail in certain cases, there is the option of setting $ihaze=-1$ which enforces the haze removal disregarding the termination criteria.

$ihaze = 0$ : no haze correction
$ihaze = 1$ : haze/land correction, might be switched off if quality check criteria are not passed
$ihaze = 2$ : haze over water removal (requires clear water pixels)
$ihaze = 3$ : haze/land and haze/water removal
$ihaze = 4$ : sun glint removal over water
$ihaze = -1$ : haze/land correction is executed disregarding quality checks
$ihaze = -2$ : is treated as $ihaze=2$
$ihaze = -3$ : haze/land removal is forced, haze/water removal needs clear water pixels

Haze removal is performed for the visible bands, sun glint removal for all bands.

$iwat\_shd = 0$ : water pixels are excluded from de-shadowing (land), default
$iwat\_shd = 1$ : water pixels are included in de-shadowing (land)
The option $iwat\_shd = 1$ might be useful if the internal water classification based on purely spectral criteria fails, and dark land pixels are classified as water (which is excluded from de-shadowing). So this flag is only used if the de-shadowing option is set and if no external water map is supplied. Example: scene is "image1.bsq", and a file "image1\_water\_map.bsq" or "image1\_hcw.bsq" exist in the same folder, then the flag $iwat\_shd$ is ignored because an external water map always has the first priority.

$ksolflux = 0$ : file with value added channels not calculated
$ksolflux = 1$ : value added channels are calculated (*.flx file)

$ishadow = 0$ and $fshd="$ (empty string): no DEM cast shadow map is used
$ishadow = 0$ and $fshd="valid file name": pre-calculated DEM cast shadow file is used
$ishadow = 1$ : DEM cast shadow mask is calculated "on-the-fly"
The pre-calculated map avoids repeated on-the-fly calculations.

$icl\_shadow = 0$ : no cloud/building shadow correction
$icl\_shadow > 0$ : cloud/building shadow correction is performed
   = 1 : de-shadowed output as DN image corresponding to input DN scene
   = 2 : de-shadowed output as surface reflectance image
   = 3 : de-shadowed output as surface reflectance and DN image

If a float '*ilu.bsq’ file exists in the scene folder, then the de-shadowing is performed with this file, i.e. the matched filter algorithm is not applied. Otherwise this '*ilu.bsq’ has to be renamed (temporarily) if the matched filter method shall be executed.
line 22:
0, 0.5, 0.5 : itriang, ratio_red_swir, ratio_blu_red
itriang = 0 : average vis. index of reference areas is employed for non-reference pixels
itriang = 1 : triangular interpolation of visibility index of reference areas
ratio_red_swir : ratio of surface reflectance of red to 2.2 µm band for the reference pixels
If no 2.2 µm band exists, but a 1.6 µm band, the ratio holds for the red-to-1.6 µm band.
If only VNIR bands exist (400 - 1000 nm) the ratio holds for the red-to-NIR band.
ratio_blu_red : ratio of surface reflectance of blue band to red band for the reference pixels.

line 23:
0, 65.0, 0.25 : ibrdf, beta_thr, thr_g, parameters for BRDF correction in rugged terrain
For a flat terrain, these parameters are not used.

- ibrdf = 0 : no empirical BRDF correction (or flat terrain)
- ibrdf = 1 : correction with cosine of local solar zenith angle (eq. 10.118 with b=1)
- ibrdf = 2 : correction with sqrt(cos) of local solar zenith angle (eq. 10.118 with b=1/2)
- ibrdf = 11 : correction with cosine of local solar zenith angle (eq. 10.118 with b=1), for soil/sand. Vegetation: eq. 10.118 with exponent b=3/4 and b=1/3 for λ < 720 nm and λ > 720 nm, respectively, i.e., option (a) in the BRDF panel, see Figure 5.47 ("weak" correction),
- ibrdf = 12 : correction with cosine of local solar zenith angle (eq. 10.118 with b=1), for soil/sand. Vegetation: eq. 10.118 with exponent b=3/4 and b=1 for λ < 720 nm and λ > 720 nm, respectively, i.e., option (b) in the BRDF panel, see Figure 5.47 ("strong" correction),
- ibrdf = 21 : correction with sqrt(cos) of local solar zenith angle (eq. 10.118 with b=1/2), for soil/sand. Vegetation: eq. 10.118 with exponent b=3/4 and b=1/3 for λ < 720 nm and λ > 720 nm, respectively, i.e., option (a) in the BRDF panel, see Figure 5.47 ("weak" correction). This is the recommended standard yielding good results in most cases.
- ibrdf = 22 : correction with sqrt(cos) of local solar zenith angle (eq. 10.118 with b=1/2), for soil/sand. Vegetation: eq. 10.118 with exponent b=3/4 and b=1 for λ < 720 nm and λ > 720 nm, respectively, i.e., option (b) in the BRDF panel, see Figure 5.47 ("strong" correction).
- beta_thr : threshold local solar illumination angle β_T where BRDF correction starts. If beta_thr=0 (and ibrdf > 0) then the angle β_T is calculated in ATCOR depending on the solar zenith angle and its value can be found in the corresponding "*_atm.log" file.
- thr_g : g, lower boundary of BRDF correction factor, see chapter 10.6.2, eq. (10.118, 10.119).

line 24:
1, 0.820, 0.780, 0.600 : lai_model, a0_vi, a1_vi, a2_vi
Parameters for the LAI model to be used if ksolflux > 0, see chapter 7

line 25:
0.900, 0.950, 0.380 : c_fpar, a_fpar, b_fpar
parameters for the fpar model to be used if ksolflux > 0, see chapter 7
26. 20.0, 0.83 ; air temperature (C), air emissivity, see chapter 7
Parameters for the net flux calculation (used for flat terrain, ignored for rugged terrain)

27. 20.0, 0.50, 0.65, 15.0, 6.3 ; t_air, z0_ref, tgradient, p_wv, zh_pwv, chapter 7
\(t_{\text{air}}\) : air temperature (Celsius) at elevation \(z_{0 \text{ref}}\)
\(z_{0 \text{ref}}\) : reference elevation for \(t_{\text{air}}\) [km asl]
\(t_{\text{gradient}}\) : air temperature gradients (Celsius per 100 m height)
\(p_{\text{wv}}\) : ([mb or hPa], default, water vapor partial pressure at \(z_{0 \text{ref}}\))
\(zh_{\text{pwv}}\) : ([km], scale height of water vapor, exponential decrease falls to 1/e value)
Parameters for the net flux calculation (rugged terrain, \(k_{\text{soilflux}} > 0\))
These are dummy values (not used) if \(k_{\text{soilflux}}=0\), or for a flat terrain.

28. 2, 2 ; ihot_mask, ihot_dynr , parameters for haze correction
\(\text{ihot\_mask} = 1\) : small area haze mask
\(\text{ihot\_mask} = 2\) : large area haze mask
\(\text{ihot\_dynr} = 1\) : thin to medium haze levels are corrected
\(\text{ihot\_dynr} = 2\) : thin to thick haze levels are corrected

29. 2, -0.500, 0.12, 0.08, 1 ; iclshad_mask, thr_shad, phi_unscl_max, phi_scl_min, istretch_type
Parameters for correction of cloud/building shadow effects, if icl_shadow > 0
Default values are put in this line even if icl_shadow=0
\(\text{iclshad\_mask} = 1,2,3\) : small, medium, large cloud shadow mask
\(\text{thr\_shad} = -0.500\) : threshold for core shadow areas, \(-999\) means threshold is calculated from image histogram
\(\text{phi\_unscl\_max}\) : max of unscaled shadow function \(\Phi_{\text{max}}\), see chapters 2.5, 10.5.6..
\(\text{phi\_scl\_min}\) : min of scaled shadow function \(\Phi_{\text{min}}^*\), see chapters 2.5, 10.5.6..
\(\text{istretch\_type}\) : 1=linear stretching, 2=exponential stretching of \(\Phi\) into \(\Phi^*\).

30. \text{ch940}(1:6) vector with 6 channel numbers for the 940 nm water vapor retrieval
\text{ch940}(1) : left window channel (850 - 890 nm)
\text{ch940}(2) : right window channel (850 - 890 nm)
\text{ch940}(3) : left absorption channel (920 - 970 nm)
\text{ch940}(4) : right absorption channel (920 - 970 nm)
\text{ch940}(5) : left window channel (1000 - 1040 nm)
\text{ch940}(6) : right window channel (1000 - 1040 nm)
The left and right channel numbers for each window or absorption region may be the same. Put in a zero channel number if not applicable. If the 820 nm water vapor region is selected the vector \text{ch940} holds the corresponding window and absorption channels for this region.

31. \text{ch1130}(1:6) vector with 6 channel numbers for the 1130 nm water vapor retrieval
\text{ch1130}(1) : left window channel (1050 - 1090 nm)
\text{ch1130}(2) : right window channel (1050 - 1090 nm)
\text{ch1130}(3) : left absorption channel (1110 - 1155 nm)
\text{ch1130}(4) : right absorption channel (1110 - 1155 nm)
\text{ch1130}(5) : left window channel (1200 - 1250 nm)
The left and right channel numbers for each window or absorption region may be the same. Put in a zero channel number if not applicable.

**line 32:**
chth\_w1, chth\_a1, chth\_a2, chth\_w2  
  bands for thermal water vapor retrieval (10 - 12 \( \mu m \))
chth\_w1 : left window channel (SWCVR method, see chapter 10.1.5)
chth\_w2 : right window channel
chth\_a1 : left absorption channel
chth\_a2 : right absorption channel

**line 33:**
e\_water, e\_veget, e\_soil, e\_sand  
  surface emissivities (adjusted NEM, channel with Tmax)

**line 34:**
0  iwv\_model  
  (water vapor retrieval: 1=no band regression, 2=band regression)
  The choice iwv\_model = 0 indicates the water vapor retrieval is disabled. Option 1 means the water vapor retrieval is performed for the selected bands, and in case of several measurement bands the one with the smallest standard deviation is selected (per 940 and 1130 nm region). Finally, if both regions are active, the average of the water vapor maps of both regions is taken. Option 2 employs a linear regression of bands which yields better results for the water vapor map if the data is noisy or not accurately calibrated. If iwv\_model = 2 and channels in the 940 nm and 1130 nm regions are specified, then only the 940 nm region is evaluated with a linear regression of bands. If the regression is intended for the 1130 nm region, then the 940 nm channels (all channels in line 30) have to be specified as 0.

**line 35:**
0  icirrus  
  flag for cirrus removal (0=disabled, 1=enabled, -1=forced))
  The value icirrus = -1 enforces cirrus detection and removal, i.e., termination criteria are ignored.

**line 36:**
0  irrad0  
  flag for solar flux on ground (0=disabled, 1=enabled)
  For irrad0=2 the surface reflected (leaving) radiance is calculated additionally.

For a flat terrain, ASCII spectra of the direct, diffuse, and global flux on the ground are provided in the folder of the input scene, see chapter 10.1.4. In case of a flat terrain the global (i.e. direct plus diffuse) flux image is calculated. For a rugged terrain the images of the direct and diffuse fluxes are calculated. Note: as the flux files have to use a float encoding (32bits/pixel) the file size is twice or four times the size of the input scene for a 16bit/pixel and 8bit/pixel input scene, respectively.

Notice concerning visibility iterations:
ATCOR will automatically iterate the initial visibility (parameter visib set in the ‘.inn’ file) if the number of negative reflectance pixels is larger than 1% of the scene for the red band (around 650 nm, vegetation is checked here) or the NIR band (around 850 nm, water is checked here). The specified visibility is always kept if the visibility is set to a negative value, i.e., visib=-20 means the program performs the calculation with visib=20 km and does not iterate even if a large number of negative reflectance pixels occurs.

If the parameter npref is set to 1 the program computes the visibility map based on dark reference pixels in the scene and npref=1 overwrites the initial value of the visib parameter. With npref=1 the program still iterates the average visibility of the visibility map by checking for water pixels in the NIR band unless the specified visib is negative. A constant scene visibility is employed for npref=0. In case of scene tiling and npref=0 or npref=1 the iterated visibility obtained for sub-scene 1 is also applied to all other sub-scenes to avoid brightness steps for the merged sub-scenes.
caused by potentially different visibilities.

Attention: If scene tiling has to be performed and the aerosol map is requested for each sub-image then specify npref=-1, but this could cause different average visibilities in the sub-scenes and potentially brightness steps at the sub-scene borders.

9.6 Problems and Hints

- **Distinction of haze and cloud, when can the haze removal algorithm be applied?**
  Ground surface information under haze areas can still be recognized in the 600-900 nm region, but the brightness contrast is low. The haze removal is only applied for channels in the 400-800 nm region. However, for cloud areas no ground information can be observed. If in doubt whether a certain area should be assessed as haze or cloud covered, take a look at the scene in the NIR (around 850 nm) channel: if surface features can be seen in this area, the haze algorithm might be applied. If not, the area is cloud covered and a haze removal run will not be successful.

- **The cloud mask is not appropriate.**
  The cloud mask might contain too few pixels: then decrease the cloud reflectance threshold in the ../atcor/preferences/preference_parameters.dat file. The default threshold is 25% reflectance in the blue-green spectrum. The opposite case can also happen: if the cloud mask comprises too many pixels the threshold has to be raised.

- **The haze mask is not appropriate.**
  This problem may be connected with the setting of the cloud threshold. If the cloud threshold is too high, cloud pixels are counted as haze, and the results of the haze removal are bad because the haze algorithm is applied to clouds.

- **The water mask is not appropriate.**
  This might lead to problems for the haze removal as well as the cloud/building shadow removal, i.e., water is erroneously counted as land and included in the land mask. Check the two water reflectance thresholds (NIR, SWIR region) in the preference_parameters.dat and modify them appropriately. Enable the output of a haze/cloud/water map, compare Fig. 4.13, to check the water mask. Read chapter 4.8. If necessary, modify the NIR/SWIR1 reflectance thresholds for water, an increase/decrease of the thresholds will increase/decrease the number of water pixels.

- **Rugged terrain: the slope and illumination maps show strong horizontal and vertical stripes.**
  Strong artifacts in the DEM files will immediately be visible in the atmospherically / topographically corrected surface reflectance image. This problem frequently occurs for resampled DEMs, e.g. the original DEM resolution is 30 m, which is resampled to a 5 m pixel size. Artifacts will be enhanced, especially if the stepsize of the original DEM (height resolution) is coded as integer. Float data would have smoother transitions. A simple way to get better results is the use of a larger kernel size for the slope/aspect calculation, e.g., kernel=5 or kernel=7 instead of the default kernel=3 (pixels), but this approach causes a reduction of the high frequency spatial information.

  Attention: in addition to using float data before resampling, it is recommended to calculate the slope/aspect maps on the original (coarse spatial resolution) data, followed by the high-resolution resampling step for all DEM files (elevation, slope, aspect). Do not employ...
the sequence of resampling the elevation, followed by a slope/aspect calculation of the high-resolution elevation map, because this approach enhances artifacts!

Steps to reduce slope/aspect striping:

1. FLOAT(ELEVATION file) (if it is stored as integer)
2. Calculate SLOPE (and also ASPECT) with a low pass filter of 5x5 pixels
3. Resize SLOPE (ASPECT) file: factor 4 larger
4. Apply a lowpass filter (7x7 pixels)
5. Resize with factor 0.25 using nearest neighbor to obtain original size.
Chapter 10

Theoretical Background

Standard books on optical remote sensing contain an extensive presentation on sensors, spectral signatures, and atmospheric effects where the interested reader is referred to (Slater 1980 [90], Asrar 1989 [4], Schowengerdt 2007 [87]). This chapter contains a description of the concepts and equations employed for the atmospheric correction.

We start with the basic equations in the solar and thermal spectral region for clear sky conditions (standard case), then move on to non-standard conditions comprising bidirectional reflectance (BRDF) effects, hazy scenes, and a treatment of shadow areas caused by clouds or buildings. Standard atmospheric conditions include the option of a constant visibility (aerosol optical thickness) and water vapor content per scene, as well as the retrieval of a visibility and water vapor map if the required spectral bands are available for the specific sensor. Water vapor correction on a pixel-by-pixel basis is usually necessary for hyperspectral imagery.

The section on the non-standard conditions contains a short discussion on empirical correction methods for bidirectional effects. It continues with the description of a statistical haze removal method. The third section presents a technique to compensate shadow effects, i.e. cloud or building shadow areas are masked and de-shadowed. Then, an overview is presented of all major processing steps involved in the atmospheric correction.

After atmospheric correction, the surface reflectance cube can be used for classification. A simple automatic method is included here based on template reflectance spectra of different surface covers. Finally, the accuracy of the atmospheric correction is discussed.

Before going into details, a brief overview of the main processing steps during atmospheric correction is described in the next two flow charts. Figure 10.1 contains a compact summary of the main processing elements: after reading the sensor-specific LUTs a masking and preclassification is conducted to obtain land, water, haze, cloud, and shadow areas. Then an optional haze or cirrus removal is conducted followed by an optional shadow removal. The next step calculates the visibility or aerosol optical thickness map using the dense dark vegetation (DDV) method. This is followed by an update of the aerosol model (path radiance behavior in the blue to red bands) if a blue band exists and the update option (ratio,blu,red > 0) is enabled. Otherwise the selected aerosol model is not changed. After calculation of the water vapor map the iterative surface reflectance retrieval is conducted accounting for adjacency and spherical albedo effects. After atmospheric correction, a spectral polishing (for hyperspectral instruments) and BRDF correction might be performed.

The visibility/ AOT(550 nm) retrieval flow chart describes the case with a SWIR band at 2.2 $\mu m$. It starts with a low reflectance threshold T1=0.05 and performs the masking in this SWIR band to obtain the darkest pixels, excluding water. If the number of reference pixels is less than 1% of
the scene pixels then the threshold $T_1$ is increased until threshold $T_2 = 0.12$ is reached. If not enough reference pixels are found then a constant VIS=23 km is used otherwise the visibility for each reference pixel is determined as the intersection of modeled and measured radiance in the red channel.

Then the NIR channel is checked concerning negative reflectance pixels (mainly water, shadow). If the percentage of negative reflectance pixels is higher than 1% of the scene pixels, then the visibility is iteratively increased up to 80 km. Finally, the visibility index and AOT(550nm) are calculated and the nn-reference pixels are assigned the average visibility or optionally a spatial triangular interpolation can be performed. If the aerosol type (rural, urban, maritime, desert) is not fixed by the user, the flow chart is executed for these four types, and the type closest to the scene-estimated type is used, see chapter 10.4.2 for more details.

**Figure 10.1:** Main processing steps during atmospheric correction.
10.1 Basics on radiative transfer

This chapter presents the basic concepts and the terminology. The full set of equations is documented here as implemented in ATCOR. We start with the radiative transfer equation in the solar spectral region (0.4 - 2.5 \( \mu m \)) for a flat terrain under clear sky conditions. First, the equation for an infinite plane of uniform reflectance is presented. Then the case of a small uniform surface embedded in a large homogeneous background of different reflectance is discussed. We continue with the rugged terrain, and finally discuss the equations for the thermal spectral region (8-14 \( \mu m \)).

### 10.1.1 Solar spectral region

For a cloud-free sky and a uniform ground of reflectance \( \rho \), the radiance signal received at the sensor consists of scattered solar radiation and ground reflected radiation. The scattered radiation component is also called path radiance. It depends on the solar and viewing geometry as sketched in Fig. 10.3. In case of a flat terrain, the at–sensor radiance \( L \) can be written as (Asrar 1989, chapter 9):

\[
L = L_p(\Theta_v, \Theta_s, \phi) + \tau_v(\Theta_v) \frac{\rho}{\pi} \frac{E_o(0)}{1 - \rho_v s}
\]  
(10.1)
CHAPTER 10. THEORETICAL BACKGROUND

Figure 10.3: Radiation components, illumination and viewing geometry.

$L$  at-sensor radiance for surface reflectance $\rho$ ;
$L_p$  path radiance ;
$\tau_v$  total ground-to-sensor atmospheric transmittance, sum of direct $\tau_{\text{dir}}$ and diffuse $\tau_{\text{dif}}$ transmittance;
$E_g$  global flux on a horizontal surface, sum of direct ($E_{\text{dir}}$) and diffuse ($E_{\text{dif}}$) flux, $E_g(0)$ is calculated for a ground surface with $\rho = 0$ ;
$\rho_r$  large scale reference background reflectance determining the effective global flux ($\rho_r=0.15$ is used for ATCOR) ;
s  spherical albedo of the atmosphere, accounts for atmospheric backscattering to the ground.

The geometry is described by the angles $\Theta_v$ (view zenith), and $\Theta_s$, $\phi$ (solar zenith and relative azimuth angles), compare figure 10.3. Since $\rho$ and $\rho_r$ are not known for image data and can vary within a scene, equation (10.1) has to be solved for $\rho$ iteratively, compare equations (10.9 - 10.15).

In a strict sense, the reflectance $\rho$ used here should be called hemispherical-directional reflectance factor (HDFR), because most surfaces show an anisotropic reflectance behavior characterized by the bidirectional reflectance distribution function (BRDF, Nicodemus 1970, Slater, 1985). The ground is illuminated hemispherically by the direct and diffuse solar flux and the reflected radiation is recorded from a certain direction, i.e., hemispherical input radiation, directional reflected radiation. Since the reflected radiation is always measured in a small cone, the term hemispherical-conical reflectance factor HCRF is also used, but for small instantaneous field-of-view sensors, "directional" is a sufficiently accurate geometrical description. However, for simplicity we will use the abbreviation reflectance in this manual.

In spectral regions dominated by scattering effects, the terms of equation (10.1) are calculated with the scaled DISORT option (discrete ordinate radiative transfer [55]), in regions with strong atmospheric absorption the more accurate correlated k algorithm is used in combination with
DISORT [7]. The results are stored in look-up tables (LUT). Since MODTRAN calculates the path radiance including the diffuse reflected ground radiation in the form
\[ L_{\text{path}}(\rho) = L_{\text{path}}(0) + \frac{\tau_{\text{dif}} E_g(0)}{1 - \rho_s} \rho / \pi = L_{\text{path}}(0) + \tau_{\text{dif}} E_g(\rho) \rho / \pi \] (10.2)
two MODTRAN runs with surface reflectance \( \rho = 0 \) and \( \rho_r = 0.15 \) are required to calculate the diffuse ground-to-sensor transmittance \( \tau_{\text{dif}} \) and spherical albedo \( s \) from equation (10.2)
\[ \tau_{\text{dif}} = \frac{[L_{\text{path}}(\rho_r) - L_{\text{path}}(0)] \pi}{\rho_r E_g(\rho_r)} \] (10.3)
\[ E_g(\rho_r) = \frac{E_g(\rho = 0)}{1 - \rho_r s} \] (10.4)
\[ s = \frac{[1 - E_g(0)]/\rho_r}{E_g(\rho_r)} \] (10.5)
For image data, the pixel reflectance \( \rho \) may differ from the background reflectance \( \bar{\rho} \). In this case the signal at the sensor consists of three components as sketched in Fig. 10.4:
- component 1: scattered radiance, path radiance,
- component 2: radiation reflected from pixel under consideration,
- component 3: radiation reflected from the neighborhood and scattered into the viewing direction (“adjacency” effect).

\[ L = c_0 + c_1 \text{DN} \]

Only component 2 contains information on the surface properties of the pixel, the other components have to be removed during the atmospheric correction. As detailed in [69] the adjacency radiation \( L_3 \) consists of two components (atmospheric backscattering and volume scattering) which are combined into one component in Fig. 10.4.
The radiometric calibration assigns to each digital number (DN) the corresponding at-sensor radiance \( L \)
\[
L(k) = c_0(k) + c_1(k) DN(k)
\]
where \( k \) indicates the channel number, and \( c_0 \), \( c_1 \) are the calibration coefficients (offset and slope). For sensors with adjustable gain settings the equation is
\[
L(k) = c_0(k) + c_1(k) DN(k)/g(k)
\]
where \( g(k) \) is the gain setting in channel \( k \). The atmospheric correction has to be performed iteratively, since the surface reflectance and large-scale (0.5-1 km neighborhood) background reflectance are not known. So three steps are employed in the ground reflectance calculation:

Step 1: The influence of the neighborhood (adjacency effect) is neglected and the surface reflectance is obtained from
\[
\rho^{(1)} = \frac{\pi [d^2 (c_0 + c_1 DN) - L_p]}{\tau \varepsilon_g (\rho_r = 0.15)}
\]
where the spectral band index is omitted for clarity. The factor \( d^2 \) takes into account the sun-to-earth distance (\( d \) is in astronomical units) since the LUTs with path radiance and global flux are calculated for \( d=1 \) in ATCOR.

Step 2: The second step calculates the average reflectance in a large neighborhood of each pixel (range \( R=0.5-1 \) km)
\[
\bar{\rho} = \frac{1}{N^2} \sum_{i,j=1}^{N} \rho_{i,j}^{(1)}
\]
where \( N \) corresponds to the number of pixels for the selected range \( R \) of the adjacency effect [62], [69]. The exact choice of \( R \) is not critical since the adjacency influence is a second-order effect. Instead of the range-independent weighting in eq. (10.9), a range-dependent function can be selected with an exponential decrease of the weighting coefficients [63]. The range-dependent case requires more execution time, of course. Except for special geometries, the difference between both approaches is small, because the average reflectance in a large neighborhood usually does not vary much and the influence is a second-order effect.

\[
\rho^{(2)}(x, y) = \rho^{(1)}(x, y) + q(\rho^{(1)} - \bar{\rho}(x, y))
\]
The function \( q \) indicates the strength of the adjacency effect. It is the ratio of the diffuse to direct ground-to-sensor transmittance. The range-dependent version of eq. (10.10) is:
\[
\rho^{(2)}(x, y) = \rho^{(1)}(x, y) + q(\rho^{(1)}(x, y) - \int_0^R \rho^{(1)}(r) A(r) \exp(-r/r_s) dr)
\]

Here, \( R \) is the range where the intensity of the adjacency effect has dropped to the 10% level (i.e. \( r=R=2.3x r_s \), where \( r_s \) is a scale range, typically \( r_s=0.2-0.4 \) km, \( R=0.5-1 \) km). \( \rho(r) \) is the reflectance at range \( r \) from the \((x,y)\) position and \( A(r) \) is the area of a circular zone from \( r \) to \( r+dr \). Now we approximate the circular regions by square regions to obtain the discrete version of eq. (10.11) with exponentially decreasing weighting coefficients \( w_i \):
\[
\rho^{(2)}(x, y) = \rho^{(1)}(x, y) + q(\rho^{(1)}(x, y) - \sum_{i=1}^{n_R} \bar{\rho}_i w_i)
\]
\[
w_i = \frac{1}{\sum_{i=1}^{n} W_i} \quad \text{and} \quad W_i = \int_{r_{i-1}}^{r_i} A(r) \exp(-r) dr \approx \int_{r_{i-1}}^{r_i} (2r)^2 \exp(-r) dr \quad (10.13)
\]

ATCOR supports up to \( n_R = 5 \) regions. Since the sequence of moving digital low pass filters works with square filters of size \( 2r_i \times 2r_i \), the area \( A(r) \) is approximated as the corresponding square region \( A(r) = (2r)^2 \).

Step 3: it includes the spherical albedo effect on the global flux that was initially calculated with the reference background reflectance \( \rho_r = 0.15 \) and is finally adapted to the scene-dependent value \( \bar{\rho} \) by correcting with the difference \( \bar{\rho} - \rho_r \):

\[
\rho^{(3)}(x, y) = \rho^{(2)}(x, y)[1 - (\bar{\rho}(x, y) - \rho_r)s]
\quad (10.14)
\]

Radiation components in rugged terrain: Figure 10.5 shows a sketch of the radiation components in a rugged terrain [63]. Compared to the flat terrain one additional component is taken into account in the ATCOR model. It is an approximation of the terrain reflected radiation. It is obtained by weighting the reflected radiation in a 0.5 km surrounding of a pixel with the terrain view factor. The terrain view factor is \( V_{\text{terrain}}(x, y) = 1 - V_{\text{sky}}(x, y) \), and the sky view factor \( V_{\text{sky}}(x, y) \) is calculated from the DEM as explained below. The sky view factor is normalized to 1 for a flat terrain.

The reflectance is calculated iteratively. The first step neglects the adjacency effect and starts with a fixed terrain reflectance of \( \rho^{(0)}_{\text{terrain}} = 0.1 \) [66]:

\[
\rho^{(i)}(x, y) = \frac{\pi [d^2(c_0 + c_1 \text{DN}(x, y)) - L_p(z, \Theta_v, \Phi)]}{\tau_s(z, \Theta_v) [b(x, y) E_s \tau_s(z) \cos \beta(x, y) + E_d^*(x, y, z) + E_i^{(i-1)}(z, \rho_r) \rho^{(i-1)}_{\text{terrain}} V_{\text{terrain}}(x, y)]} \quad (10.15)
\]

The terms are defined as:

- \( x, y \): horizontal coordinates, corresponding to the georeferenced pixel positions;
- \( z \): vertical coordinate, containing the elevation information from the DEM;
- \( \text{DN}(x,y) \): digital number of georeferenced pixel;
- \( L_p(z, \theta_v, \phi) \): path radiance, dependent on elevation and viewing geometry;
- \( \tau_v(z, \Theta_v) \): ground-to-sensor view angle transmittance, direct plus diffuse components;
- \( \tau_s(z) \): Sun-to-ground beam (direct) transmittance;
- \( \beta(x, y) \): angle between the solar ray and the surface normal (illumination angle);
- \( b(x,y) \): binary factor: \( b=1 \) if pixel receives direct solar beam, otherwise \( b=0 \);
- \( E_s \): extraterrestrial solar irradiance (earth-sun distance \( d=1 \) astronomical unit);
- \( E_d^*(x, y, z) \): diffuse solar flux on an inclined plane (see equation 10.18);
- \( E_g(z) \): global flux (direct plus diffuse solar flux on a horizontal surf. at elevation \( z \));
- \( E_i^{(i)}(z) \): radiation incident upon adjacent slopes;
- \( \rho_{\text{terrain}}^{(0)} \): initial value of average terrain reflectance;
- \( \rho_{\text{terrain}}^{(i)}(x, y) \): locally varying average terrain reflectance, calculated iteratively \( i=1,2,3 \);
- \( V_{\text{terrain}}(x, y) \): terrain view factor (range 0-1).

The solar and DEM geometry is shown in figure 10.6 as well as the three solar radiation components taken into account for rugged terrain: direct and circumsolar irradiance, and diffuse hemispherical sky flux. It can be shown that these three components are equivalent to the direct and diffuse solar flux components in flat terrain. In case of a shadow pixel the direct and circumsolar components...
Figure 10.5: Radiation components in rugged terrain, sky view factor.


are set to zero, i.e. the binary factor \( b = 0 \).

The next step iterates eq. 10.15 averaging the reflected terrain radiation over a square box of 1.0 \( \times \) 1.0 km. If equation (10.15) is used with \( E_t = E_g \) then three iterations are usually sufficient to be independent of the start value of the terrain reflectance \([63]\). However, for highly reflective surfaces, e.g. snow, and high terrain view factors, more than three iterations are necessary, and a faster convergence of \( \bar{\rho}_\text{terrain}^{(i)} \) can be achieved with a geometric series for the terrain reflected radiation \( E_t \) as proposed in \([89]\):

\[
E_t^{(i)} = E_g \cdot \frac{\bar{\rho}_\text{terrain}^{(i-1)} V_{\text{terrain}}}{1 - \bar{\rho}_\text{terrain}^{(i-1)} V_{\text{terrain}}} \tag{10.16}
\]

The next steps include the adjacency correction (eq. 10.9, 10.10) and the spherical albedo effect (eq. 10.14).

If \( \Theta_s, \Theta_n, \phi_s, \phi_n \) denote solar zenith angle, terrain slope, solar azimuth and topographic azimuth, respectively, the illumination angle \( \beta \) can be obtained from the DEM slope and aspect angles and the solar geometry:

\[
\cos \beta(x, y) = \cos \Theta_s \cos \Theta_n(x, y) + \sin \Theta_s \sin \Theta_n(x, y) \cos \{\phi_s - \phi_n(x, y)\} \tag{10.17}
\]
The illumination image \( \cos \beta(x, y) \) is calculated within ATCOR and stored as a separate map. The diffuse solar flux on an inclined plane is calculated with Hay’s model (Hay and McKay 1985, also see Richter 1998 for the enhancement with the binary factor b):

\[
E^*_d(x, y, z) = E_d(z)[b \tau_s(z) \cos \beta(x, y) / \cos \Theta_s + \{1 - b \tau_s(z)\}V_{sky}(x, y)]
\]  
(10.18)

The sky view factor can be computed from local information as \( V_{sky}(x, y) = \cos^2(\Theta_n(x, y)/2) \) based on the local DEM slope angle \( \Theta_n \). ATCOR uses the horizon algorithm that provides a more accurate value of the sky view factor by considering the terrain neighborhood of each pixel (Dozier et al. 1981). \( V_{sky} \) and \( V_{terrain} \) are related by:

\[
V_{sky}(x, y) = 1 - V_{terrain}(x, y)
\]  
(10.19)

\textit{Figure 10.6:} Solar illumination geometry and radiation components.
10.1.2 Illumination based shadow detection and correction

For high resolution imagery, the correction of cast shadows and illumination on the basis of a surface model does not lead to useful results as the surface representation with respect to the radiometry is never accurate enough. Severe over- and under-correction artifacts are observed due to these inaccuracies in the resulting images.

The detection and correction of cast shadows has been widely studied, specifically for space borne high resolution instruments [3, 88]. A new method for cast shadow detection has been implemented for the ATCOR case. It produces a continuous shadow field and relies on the fact that all areas in cast shadows are illuminated by diffuse irradiance only. The diffuse illumination is caused by scattering and thus exhibits very specific spectral characteristics if compared to the direct irradiance. Specifically, the signal in the blue spectral band is significantly higher in cast shadow areas than in directly illuminated areas. For the shadow quantification, the brightness in the NIR spectral band is first calculated using the solar illumination. Secondly, two blue indices have been defined as the band ratios green/blue and red/blue, respectively. These three measures are combined such that a value equivalent to the illumination between 0 and 1 is created (0 being a cast shadow area). The shadow fraction parameter is then defined as:

\[
 p_{\text{shad}} = \max\left[5 \cdot \left(\frac{L_{\text{red}}}{L_{\text{blue}}} - 0.35\right), 8 \cdot \left(\frac{L_{\text{green}}}{L_{\text{blue}}} - 0.42\right), 6 \cdot \rho_{\text{app,nir}}\right] 
\]

(10.20)

where \(L_{\text{blue}}, L_{\text{green}}, \) and \(L_{\text{red}}\) are the at-sensor radiance values in the true color bands and \(\rho_{\text{app,nir}}\) is the apparent at-sensor reflectance in the near infrared band. The scaling factors (here: 5,8, and 6) are chosen such that all three parameters are within the same range (i.e., a value range between 0 and 2 and optimized for full cast shadow at 0.5). The parameter \(p_{\text{shad}}\) is then scaled to a shadow fraction number \(f_{\text{shad}}\) between 0 (full cast shadow) and 1 (no cast shadow) using empirically found limits. These limits may be variable between sensors. In a second step, this map is combined with the standard geometrically derived illumination field (see Fig.10.7). The resulting illumination map serves as an input to the ATCOR method.

**Skyview Factor Estimate**

The skyview factor \(V_{\text{sky}}\) describes the relative amount of the unobstructed sky hemisphere. This factor is highly variable on small scales, e.g., in vicinity of forest borders. The skyview factor is approximated from the cast shadow fraction such that all areas in complete cast shadows get a reduced skyview factor as:

\[
 V_{\text{sky}} = \left(1 - \frac{\theta_s}{180^\circ} + f_{\text{shad}} \frac{\theta_s}{180^\circ}\right) \cdot 100\% < V_{\text{sky,geom}}, \quad (10.21)
\]
where $\theta_s$ is the solar zenith angle and the geometrical skyview factor $V_{\text{sky,geom}}$ had been calculated based solely on the digital terrain model.

**Application in ATCOR Workflow**

In ATCOR-4, the local illumination angle $\varphi$ is first calculated on the basis of the terrain model using an efficient vector algebra based method. The topographically corrected reflectance $\rho_{\text{topo}}$ is then retrieved from the atmospherically corrected ground leaving excitance ($M_g$) using the direct irradiance $I_{\text{dir}}$, the diffuse illumination field $I_{\text{dif}}$, and the terrain illumination $I_{\text{ter}}$, as:

$$\rho_{\text{topo}} = \frac{M_g}{I_{\text{dir}} \cos(\varphi) + 0.1 I_{\text{dif}} \cos(\varphi) + 0.9 I_{\text{dif}} V_{\text{sky}} + I_{\text{ter}}},$$

(10.22)

where the factors 0.1 and 0.9 account for the relative amount of circumsolar irradiance, estimated to be 10% of the total diffuse irradiance. The local illumination factor $\cos(\varphi)$ is now enhanced by the cast shadow fraction, such that shaded areas are not affected by direct and circumsolar irradiance.

![Figure 10.8: Effect of combined topographic / cast shadow correction: left: original RGB image; right: corrected image (data source: Leica ADS, central Switzerland 2008, courtesy of swisstopo).](image)

After cast shadow correction, the border pixels of the shadows are often under/overcorrected, which is visible as black or bright borders in the cast shadow areas. A filter has to be applied to remove this artifact (compare Fig. 10.9). A simple approach to this problem is an interpolation of all border pixels. However, a considerable data loss may be the result of such a process. Thus a more sophisticated approach is required. As the border pixels are usually isolated, a filter approach has been used which compares the pixel brightness $\rho_{i,j}$ to the brightness of its direct neighbors $\rho_{\text{prox}}$ in a 3x3 or 5x5 box, respectively. The brightness of the shadow border pixel is then adjusted by the relative brightness difference of the whole spectrum such that:

$$\rho_{\text{filt},i,j} = \rho_{i,j} \frac{\rho_{\text{prox}}}{\rho_{i,j}}.$$

(10.23)

This method proved to successfully remove shadow borders for high resolution imagery and an urban environment. However, for terrain shadows on a larger scale, the border pixels are not such clearly isolated and often cannot be corrected using this method, or only after increasing the size of the border pixel filter significantly.
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Figure 10.9: Effect of cast shadow correction (middle) and shadow border removal (right) for building shadows.

The updated processing leads to improved terrain correction as displayed in Fig. 10.8. Advantages of this method are:

- terrain and forest cast shadows are corrected,
- water and cast shadows are discerned (in most cases),
- operational usability has been proven on large ADS data sets, and
- a consistent physically based method applicable to photogrammetry and imaging spectroscopy is implemented.

10.1.3 Integrated Radiometric Correction (IRC)

The IRC method was published by Kobayashi and Sanga-Ngoie [43, 44] to provide a combined atmospheric and topographic correction. The algorithm is briefly outlined here, more details can be found in the original papers. Although it was originally proposed for satellite imagery, it can also be applied to airborne scenes.

The first step is the orthorectification of the scene using a digital elevation model (DEM). Then the slope and aspect maps are calculated. The next step is the calculation of the sky view factor, see chapter 10.1.1. The original paper uses the simple equation based solely on the slope angle, but with ATCOR a more accurate calculation based on a ray tracing can also be used in case of a steep terrain. Then the following quantities are computed (keeping the original notation of Kobayashi in most cases):

\[ h_0 = \frac{\left( \pi + 2\theta_s \right)}{2\pi} \] (10.24)

Here \( \theta_s \) is the solar zenith angle in radian. IF \( s \) denotes the slope map (in radian) then the simple version of the skyview is obtained with

\[ h = 1 - s/\pi \] (10.25)

The cosine of the local solar zenith (illumination angle \( \beta \)) is given in eq. 10.17. Then the surface radiance for each channel \( L_s \) is calculated by subtracting the path radiance \( L_p \) from the at-sensor radiance \( L \):

\[ L_s(x, y) = L(x, y) - L_p(x, y, z) \] (10.26)
In the ATCOR version of the IRC algorithm the path radiance varies spatially, particularly due to the DEM height variation, while a constant value (per channel) is used in the original IRC paper. Then a regression analysis (per channel) of $L_s$ versus $\cos \beta$ is applied to calculate the slope $m$ and intercept $b$. After defining $C = m/b$ the topographic correction map $A$ is calculated:

$$ A(x, y) = \frac{\cos \theta_s + C/h_0}{\cos \beta(x, y) + C \bar{h}(x, y)/h_0} $$

(10.27)

Finally, the surface reflectance $\rho$ is computed according to:

$$ \rho(x, y) = \frac{\pi L_s(x, y, z) A(x, y)}{T(x, y, z) \{ E_{dir}(x, y, z) \cos \theta_s + E_{dif}(x, y, z) \}} $$

(10.28)

where $T$ is the total ground-to-sensor transmittance, and $E_{dir}$, $E_{dif}$ are the direct irradiance and diffuse solar flux on the ground, respectively.

So the ATCOR version of IRC contains some improvements with respect to the original method: the path radiance varies spatially, mainly caused by terrain height variations, possibly also due to visibility variations, and the sky view factor can be provided from a ray tracing analysis instead of the local slope angle.

Note: the IRC method usually performs well. However, due to the statistical evaluation of the regression analysis unphysically large ($\rho > 1$ reflectance unit) or small ($\rho < 0$) surface reflectance values might happen for some pixels, usually in areas with topographic shadow or low local sun elevations.

### 10.1.4 Spectral solar flux, reflected surface radiance

The spectral solar fluxes on the ground can be calculated by setting the parameter $irrad0 = 1$ in the .inn file or using the graphical user interface. The fluxes depend on solar geometry, terrain elevation, topography, and atmospheric conditions. All fluxes and the surface reflected radiance of this section are evaluated for the current earth-sun distance. For a flat terrain, ATCOR provides spectra of the direct, diffuse, and global flux for the selected visibility / water vapor. In case of variable visibility / water vapor the spectra are calculated for the average scene visibility / water vapor. The direct flux is just the beam irradiance on the ground times the cosine of the local solar zenith angle. The diffuse flux spectrum $E_{dif}$ is evaluated for a surface reflectance of $\rho = 0$, and the global flux for $\rho = 0.15$, i.e., $E_g = (E_{dir} + E_{dif}(0))/(1 - s \cdot 0.15)$, where $s$ is the spherical albedo.

The spectral band index is omitted for brevity. For a flat terrain these fluxes are provided in the directory of the input file (e.g. 'scene.bsq'):

- the direct spectral flux on the ground: 'scene_edir.dat'
- the diffuse spectral flux on the ground: 'scene_edif.dat' for surface reflectance $\rho = 0$.
- the global spectral flux on the ground: 'scene_eglo.dat' for a typical average surface reflectance $\rho = 0.15$.

These spectra will already give a realistic description for a flat terrain, but they lack the dependence on the spectral reflectance variations in the scene. Therefore, an image of the global flux is also provided that accounts for the spatial reflectance and visibility / water vapor patterns (VIS), named 'scene_eglobal.bsq':

$$ E_g(x, y) = \frac{E_{dir}(VIS(x, y)) + E_{dif}(\rho = 0, VIS(x, y))}{1 - s(x, y) \bar{\rho}(x, y)} $$

(10.29)
Here, $\bar{\rho}$ indicates a spatial averaging with a filter size corresponding to the specified adjacency range. The unit of the global flux is $mWcm^{-2}\mu m^{-1}$ and it is stored as float data (32 bits/pixel). Therefore, its file size will be twice or four times the size of the input scene if the scene is encoded as 16bit/pixel and 8bits/pixel, respectively.

For a rugged terrain, images of the direct and diffuse fluxes will be calculated using the available DEM information on height ($z$), slope and aspect (i.e. local solar illumination angle $\beta$), and atmospheric conditions (visibility / water vapor VIS). The direct flux on the ground is:

$$E_{dir}(x, y) = b(x, y) E_0 T_{sun}(VIS(x, y, z)) \cos\beta(x, y)$$  

(10.30)

where $E_0$, $T_{sun}$ are extraterrestrial solar irradiance and sun-to-ground transmittance, respectively, and $b$ is the topographic shadow mask ($0=$shadow, $1=$sunlit pixel).

The diffuse flux in mountainous terrain accounts for the adjacency effect and multiple reflection effects from the surrounding topography. Using the terrain view factor $V_t$ from the last section and the effective terrain reflectance $\rho_t = V_t(x, y) \bar{\rho}(x, y)$ and $\bar{\rho} = \tilde{V}_t(x, y) \tilde{\rho}(x, y)$ the diffuse flux is approximated as:

$$E_{dif}(x, y) = \frac{E_{dif,flat}(x, y)}{\pi} + \frac{E_{dir,flat}(x, y, z) + E_{dif,flat}(x, y, z)}{1 - \rho_t(x, y)}$$  

(10.31)

The first line describes the anisotropic and isotropic components of the diffuse flux, the second line accounts for multiple terrain reflection effects.

Related quantities to the global spectral solar flux on the ground are the wavelength-integrated global flux and the absorbed solar flux ($Wm^{-2}$). These play a role in the surface energy balance and they are available as part of the value added channels, see chapter 7.2, equations 7.9, 7.10.

Surface reflected radiance

The ground reflected (or ground leaving) radiance per band can be obtained in addition to the spectral solar fluxes by setting the parameter $irrad0=2$. It is calculated corresponding to the surface reflectance cube $\rho(x, y)$, named ‘scene_surfrad.bsq’. For a flat terrain it is:

$$L(\text{surf}, x, y) = E(\text{global}) \rho(x, y) / \pi$$  

(10.32)

In case of a mountainous terrain the direct and diffuse reflected radiation maps from the equations 10.30 and 10.31 are used:

$$L(\text{surf}, \text{dir}, x, y) = (E_{dir} + E_{dif})\rho(x, y) / \pi$$  

(10.33)

Again, the same output file name is used (’scene_surfrad.bsq’).

10.1.5 Thermal spectral region

Similar to the solar region, there are three radiation components: thermal path radiance ($L_1$), i.e., photons emitted by the atmospheric layers, emitted surface radiance ($L_2$), and reflected radiance ($L_3$). The short form of the radiance equation in the thermal region can be written as [39] :

$$L = L_p + \tau \varepsilon L_{BB}(T) + \tau (1 - \varepsilon) F/\pi$$  

(10.34)

where
**Figure 10.10:** Radiation components in the thermal region.

$L_1 = L_P$, $L_2 = \tau \epsilon L_{BB}(T)$, $L_3 = \tau (1 - \epsilon) F/\pi$.

$L$ at-sensor radiance,
$L_1 = L_P$ thermal path radiance,
$\tau$ ground-to-sensor atmospheric transmittance,
$\epsilon$ surface emissivity,
$T$ surface temperature,
$L_{BB}$ blackbody radiance at temperature $T$, weighted with the channel’s filter curve,
$F$ thermal downwelling flux on the ground.

The second term on the right-hand side of equation (10.34) is emitted surface radiance reaching the sensor, the third term is the atmospheric radiance reflected at the surface and attenuated by the surface-to-sensor path. The spectral band index, elevation, and angular dependence is omitted for brevity.

The $L_{BB}(T)$ term is Planck’s blackbody radiance $B(\lambda, T)$ weighted with the spectral channel response function $R(\lambda)$:

$$L_{BB}(T) = \frac{\int_{\lambda_1}^{\lambda_2} B(\lambda, T) R(\lambda) d\lambda}{\int_{\lambda_1}^{\lambda_2} R(\lambda) d\lambda} \quad (10.35)$$

For a discrete temperature interval $T = (T_1, T_2)$ and increment (e.g. $T_1 = 200$ K, $T_2 = 350$ K, increment 1 K) equation (10.35) is solved numerically. Temperature and radiance are then approximated by an exponential fit function function with channel dependent coefficients $a_1, a_2$:

$$L_{BB} = \frac{a_1}{\exp(a_2/T - 1)} \quad (10.36)$$

$$T = \frac{a_2}{\ln(a_1/L_{BB} + 1)} \quad (10.37)$$
For convenience, an offset \( a_0 \) is introduced with default \( a_0 = 0 \). The offset term can be used to adjust a temperature bias in a scene. Example: if scene temperatures are too low by 3K they can be raised by setting \( a_0 = 3 \).

\[
T = a_0 + \frac{a_2}{\ln(a_1/L_{BB} + 1)} \quad (10.38)
\]

**Remark:**

The computer implementation of the channel-resampled radiance equations is coded to minimize spectral resampling effects [64], [65].

**Temperature / emissivity separation**

For a sensor with \( n \) thermal channels there are \( n \) equations of (10.34) with \( n+1 \) unknowns, namely the \( n \) surface emissivities plus a surface temperature. So, the system of equations (10.34) is always underdetermined. Several possibilities exist to address this problem (Gillespie et al. 1986, 1998 [27, 28]). Five options are offered by the airborne version of ATCOR:

- a constant emissivity (default \( \epsilon = 0.98 \) independent of surface cover type, 10-12 \( \mu \)m region, for sensors with a single thermal channel. For sensors with multiple thermal bands the parameter \( \text{itemp}_{-}\text{band} \) described in chapter 4.6) defines the channel employed for the surface temperature calculation.

- fixed emissivity values assigned for 3 classes for the selected surface temperature band (parameter \( \text{itemp}_{-}\text{band} \): \( \epsilon(\text{soil}) = 0.96 \), \( \epsilon(\text{vegetation}) = 0.97 \), else \( \epsilon = 0.98 \) (water and undefined class). The assignment to the vegetation / soil class is performed on-the-fly in memory employing the vegetation index (red and NIR bands required), and the 3-class emissivity map is also available (file ‘image_atm_emis3..bsq’, compare chapter 4.5).

- emissivity map based on a pre-classification performed with coregistered reflective bands [59]: to each class the user can assign an \( \epsilon \) value as shown in Table 10.1. The classification algorithm itself is described in chapter 5.8.2. The file name nomenclature is described in chapter 4.5. The channel for the emissivity map is defined by the user (parameter \( \text{itemp}_{-}\text{band} \) described in chapter 4.6).

- for multispectral thermal bands the normalized emissivity method (NEM) or adjusted NEM are also implemented. In the NEM [27], the surface temperature is calculated for all channels with a constant user-defined emissivity, and for each pixel the channel with the highest temperature is finally selected. In the adjusted NEM (ANEM) [14] the assigned emissivity is surface cover dependent. Here, we define four surface cover classes (water, vegetation, soil/dry vegetation, sand/asphalt) based on the following criteria:

  - vegetation: \( \rho_{\text{nir}}/\rho_{\text{red}} > 2 \) and \( \rho_{\text{nir}} > 0.20 \).
  
  - soil/dry vegetation: \( \rho_{\text{nir}}/\rho_{\text{red}} \geq 1.4 \) and \( \rho_{\text{nir}}/\rho_{\text{red}} < 2.0 \) and \( \rho_{\text{red}} > 0.09 \).
  
  - sand/asphalt: \( \rho_{\text{nir}}/\rho_{\text{red}} < 1.4 \) and \( \rho_{\text{red}} > 0.09 \).
  
  - water: \( \rho_{\text{nir}} < 0.05 \) and \( \rho_{1.6\mu\text{m}} < 0.03 \).
To each class the user can assign an emissivity valid for the channel with the highest temperature. There is only one common emissivity class in case of night data or data from purely thermal channels.

The ANEM method provides accurate channel emissivities and surface temperatures if the classes are assigned correctly and the emissivity value assigned to the channel with the maximum temperature is close to the actual channel emissivity. Maximum surface emissivities usually lie in the 10.5 - 13 μm region. After calculating the surface temperature, the emissivities for all channels are computed.

- the ASTER TES algorithm is available for sensors with at least 5 channels in the thermal interval \((\lambda_x, \lambda_y)\) with \(\lambda_x - \text{FWHM}_x/2 \geq 8.08 \mu \text{m}\) and \(\lambda_y + \text{FWHM}_y/2 \leq 13.0 \mu \text{m}\) to exclude channels in strong absorption regions. The TES algorithm implemented here consists of 3 major parts (Gillespie et al. 1998 [28]):
  - the NEM algorithm described above.
  - the ratio module:
    It calculates relative emissivities \(\beta_i\) (channel i) by ratioing the NEM emissivity values \(\varepsilon_i\) to their average:
    \[
    \beta_i = \frac{n \sum_{i=1}^{n} \varepsilon_i}{\sum_{i=1}^{n} \varepsilon_i} \quad (i = 1, n)
    \] (10.39)
    Here \(n\) is the number of channels in the allowed thermal interval.
  - maximum minimum distance (MMD module):
    The maximum and minimum values of the relative emissivities (\(\beta\) spectrum) are calculated to find the spectral contrast:
    \[
    \text{MMD} = \max(\beta) - \min(\beta)
    \] (10.40)
    Then an empirical relationship is used to predict the actual emissivities \(\varepsilon_i\) from the MMD employing a regression with laboratory or field emissivity spectra:
    \[
    \varepsilon_{\text{min}} = a + b \times \text{MMD}^c
    \] (10.41)
    For small values \(\text{MMD} < 0.03\) (e.g. water, vegetation) the value \(\varepsilon_{\text{min}}\) is set to 0.983.
    The 3 parameters \((a, b, c)\) of the regression can be specified by the user, the default values are taken from [28], i.e., \(a=0.994, b=-0.687, c=0.737\) . The final step calculates the actual emissivities using the \(\beta\) spectrum and \(\varepsilon_{\text{min}}\):
    \[
    \varepsilon_i = \beta_i \times \frac{\varepsilon_{\text{min}}}{\min(\beta)} \quad (i = 1, n)
    \] (10.42)

- for thermal band imagery with at least 5 channels the ISAC (In-Scene Atmospheric Compensation) method is available. A detailed description is given by Young et al. [100]. The method does not require ancillary meteorological data or atmospheric modeling. It neglects the downwelling thermal flux and employs the equation
  \[
  L = L_p + \tau \varepsilon L_{BB}(T) = L_p + \tau L_{\text{surface}}
  \] (10.43)
  This approximation is justified for pixels with a high emissivity close to 1, i.e. "blackbody" pixels. First, the highest brightness temperature \(T_{\text{sensor}}\) for each pixel in each channel is computed based on the at-sensor radiance \(L\) converted into brightness temperature. In the
current implementation, only channels in the spectral region 8 - 13 \( \mu m \) are employed for the maximum brightness temperature search, because the spectral regions \( \lambda < 8 \mu m \) and \( \lambda > 13 \mu m \) are strongly affected by atmospheric water vapor absorption. Next, a reference channel is defined where most pixels with maximum brightness temperature occur. Only those blackbody pixels are retained which have the maximum brightness temperature in this reference channel ("most hits" method). For these selected blackbody pixels the scatterplot of measured at-sensor radiance \( L \) versus blackbody radiance corresponding to \( L_{BB}(T_{max}^{sensor}) \) is computed for each channel. This means the surface radiance of eq. 10.43 is approximated as \( L_{surface} = L_{BB}(T_{max}^{sensor}) \). The final step is a least squares regression of the scatterplot data \( L \) versus \( L_{surface} \), yielding the intercept (path radiance \( L_p \)) and slope (transmittance \( \tau \)) of eq. 10.43. Care has to be taken to apply the regression only to the points near the top edge of all cluster points, but allow some margin, so the fitting line is allowed to sink by an amount of the sensor noise equivalent spectral radiance (NESR). The quality of the regression is significantly increased by allowing only those pixels in the scatterplot that had their maximum temperatures in the reference channel.

Two comments: first, because of the involved assumptions, the obtained intercept is not the physical path radiance, and the slope not the physical atmospheric transmittance. Both quantities may be negative in some channels, therefore they are referred to as unscaled path radiance \( L_p^{(u)} \) and unscaled transmittance \( \tau^{(u)} \). They might be rescaled to proper atmospheric path radiance and transmittance spectra, e.g. using a radiative transfer code. Second: the ISAC method requires an adequate spread in surface temperatures in the scene, and surface temperatures higher than the atmospheric radiation temperature. So, results for night-time imagery will likely be degraded.

The compensated unscaled surface radiance spectrum is calculated as

\[
L_{surface}^{(u)}(\lambda) = \frac{L(\lambda) - L_p^{(u)}(\lambda)}{\tau^{(u)}(\lambda)}
\]  

(10.44)

and the unscaled ISAC surface emissivity can be obtained with

\[
epsilon_{isac}(\lambda) = \frac{L_{surface}^{(u)}(\lambda)}{L_{BB}(\lambda, T_{ref})}
\]  

(10.45)

where \( T_{ref} \) is the brightness temperature image in the reference channel. The compensated surface radiance spectrum \( L_{surface}^{(u)}(\lambda) \) can be converted into the equivalent compensated brightness temperature spectrum where most of the atmospheric absorption features are removed. Both the compensated surface radiance and compensated brightness temperature are spectrally consistent with the data and represent the best estimate for the spectral shape. The emissivity spectrum \( \epsilon_{isac}(\lambda) \) may exceed the value 1 in certain channels if the maximum brightness temperature of a pixel does not occur in the selected reference channel. However, a common reference channel is needed in this method to obtain a consistent pixel-independent spectrum of unscaled path radiance \( L_p^{(u)} \) and transmittance \( \tau^{(u)} \).

As soon as the emissivity is set in a certain thermal band, this band can be used for the surface temperature retrieval. In case of multispectral thermal bands the remaining channels can be used to retrieve the emissivity spectrum. The advantage of the classification approach is a flexible assignment of emissivity classes and values. This approach also works for sensors with only a single
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<table>
<thead>
<tr>
<th>label</th>
<th>description</th>
<th>emissivity</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>not classified</td>
<td>0.980</td>
</tr>
<tr>
<td>1</td>
<td>clear water</td>
<td>0.991</td>
</tr>
<tr>
<td>2</td>
<td>dark vegetation</td>
<td>0.980</td>
</tr>
<tr>
<td>3</td>
<td>average veget.</td>
<td>0.975</td>
</tr>
<tr>
<td>4</td>
<td>bright veget.</td>
<td>0.980</td>
</tr>
<tr>
<td>5</td>
<td>yellow veget.</td>
<td>0.980</td>
</tr>
<tr>
<td>6</td>
<td>mixed veg./soil</td>
<td>0.975</td>
</tr>
<tr>
<td>7</td>
<td>dark bare soil</td>
<td>0.970</td>
</tr>
<tr>
<td>8</td>
<td>bare soil</td>
<td>0.970</td>
</tr>
<tr>
<td>9</td>
<td>asphalt</td>
<td>0.955</td>
</tr>
<tr>
<td>10</td>
<td>sand / soil</td>
<td>0.970</td>
</tr>
<tr>
<td>11</td>
<td>bright sand/soil</td>
<td>0.970</td>
</tr>
<tr>
<td>12</td>
<td>dry veg./soil</td>
<td>0.975</td>
</tr>
<tr>
<td>13</td>
<td>sparse veg./soil</td>
<td>0.975</td>
</tr>
<tr>
<td>14</td>
<td>snow</td>
<td>0.980</td>
</tr>
<tr>
<td>15</td>
<td>cloud</td>
<td>0.980</td>
</tr>
<tr>
<td>16</td>
<td>turbid water</td>
<td>0.984</td>
</tr>
</tbody>
</table>

Table 10.1: Example of emissivity values for a 11 µm channel.

thermal band such as Daedalus 1268 (10 reflective, 1 thermal band). The drawback is that it cannot be employed for night-time data or sensors with only thermal bands. In this case, a fixed emissivity has to be specified for one channel, e.g., $\epsilon = 0.97$. In the ATCOR model the following radiance unit $mW \ cm^{-2} \ sr^{-1} \ \mu m^{-1}$ is employed, equivalent to $\mu W \ cm^{-2} \ sr^{-1} \ nm^{-1}$.

Split-window covariance-variance ratio SWCVR

The method derives the water vapor map from thermal band imagery [42, 47, 38]. The water vapor content $W$ can be retrieved as a function of the ratio $R_{j,i}$ of transmittances $\tau_i$, $\tau_j$ in two thermal bands $i$ and $j$:

$$W = a + b \ R_{j,i}$$ (10.46)

with

$$R_{j,i} = \frac{\epsilon_j \ \tau_j}{\epsilon_i \ \tau_i} = \frac{\sum_{k=1}^{N} (T_{i,k} - \bar{T}_i) \ (T_{j,k} - \bar{T}_j)}{\sum_{k=1}^{N} (T_{i,k} - \bar{T}_i)^2}$$ (10.47)

where $N$ is the number of pixels in a moving window box centered at pixel $k$, $\bar{T}_i$ is the average brightness temperature in this box, and $\epsilon$ is the land surface emissivity. Equation 10.47 is the ratio of covariance to variance accounting for the abbreviation SWCVR. The two selected channels should be in the 10.5 - 12.5 µm region, where the emissivity of most land surfaces changes only slightly, yielding an emissivity ratio $\epsilon_j/\epsilon_i$ close to 1 yielding $R_{j,i} = \tau_j/\tau_i$. Then the parameters "a" and "b" in eq. 10.46 can be calculated from a regression of channel transmittances versus water vapor content using the pre-calculated LUTs. The moving averaging window box is selected at 100 m × 100 m. The method requires moderate to high temperature contrasts in the moving window, otherwise results are not reliable. Therefore, it is preferable to retrieve the water vapor map from channels in the solar reflective region if possible.
10.2 Masks for haze, cloud, water, snow

A useful first step before executing an atmospheric correction is the calculation of a pixel map for haze, cloud, water, snow, etc. Such a pre-classification has a long history in atmospheric correction methods [22, 40, 59, 60, 61, 41, 49]. It is also employed as part of NASA’s automatic processing chain for MODIS [1] using the classes land, water, snow/ice, cloud, shadow, thin cirrus, sun glint, etc. A similar approach is taken here.

The calculation is done "on-the-fly" and if the scene is named "scene.bsq", then the corresponding map is named "scene_out_hcw.bsq". There is also the possibility to provide this information from an external source: if a file "scene_hcw.bsq" exists in the same folder as the "scene.bsq" then this information is taken and the internal ATCOR calculations for this map are skipped. In this case, the coding of the surface types has to agree with the ATCOR class label definition, of course, see Table 10.2. This file is written if the corresponding flag is set to 1, see chapter 9.4 and figure 5.12 in chapter 4.

<table>
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<tr>
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<th>class</th>
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</thead>
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<td>geocoded background</td>
</tr>
<tr>
<td>1</td>
<td>shadow</td>
</tr>
<tr>
<td>2</td>
<td>thin cirrus over water</td>
</tr>
<tr>
<td>3</td>
<td>medium cirrus over water</td>
</tr>
<tr>
<td>4</td>
<td>thick cirrus over water</td>
</tr>
<tr>
<td>5</td>
<td>land</td>
</tr>
<tr>
<td>6</td>
<td>saturated (blue/green band)</td>
</tr>
<tr>
<td>7</td>
<td>snow / ice</td>
</tr>
<tr>
<td>8</td>
<td>thin cirrus over land</td>
</tr>
<tr>
<td>9</td>
<td>medium cirrus over land</td>
</tr>
<tr>
<td>10</td>
<td>thick cirrus over land</td>
</tr>
<tr>
<td>11</td>
<td>thin haze over land</td>
</tr>
<tr>
<td>12</td>
<td>medium haze over land</td>
</tr>
<tr>
<td>13</td>
<td>thin haze over water</td>
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<td>14</td>
<td>medium haze over water</td>
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<td>15</td>
<td>cloud over land</td>
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<tr>
<td>16</td>
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<td>18</td>
<td>cirrus cloud</td>
</tr>
<tr>
<td>19</td>
<td>cirrus cloud thick</td>
</tr>
</tbody>
</table>

*Table 10.2: Class labels in the hcw file.*

Depending on the available spectral channels, it may not be possible to assign certain classes. Table 10.2 contains one class for cloud (over land, meaning water cloud), whereas the low optical thickness cloud is put into the thin and medium thickness haze class. Thin and medium haze can often be corrected successfully. Of course, there is no clear distinction between thick haze and cloud. We take a pragmatic view, and if the haze removal is successful in areas with thick haze, then these pixels can be included in the haze mask. Since this is not clear at the beginning, it might be necessary to run the program twice, with and without haze removal. A check of the results will reveal whether the haze removal was successful. ATCOR contains a number of criteria to assess
the probability of a successful haze removal, and will switch off the haze option if the chances are not good. This automatic haze termination works in most cases, but a success cannot always be guaranteed. The two cirrus cloud classes are merged with the (normal) cloud class for the surface reflectance retrieval during the treatment of the adjacency effect: here the reflectance of the cloud pixels is replaced with the average reflectance of the non-cloud pixels to avoid an overestimation of the adjacency effect. The classes are currently defined with the following criteria:

Water class:
If the surface elevation of a pixel is lower than 1.2 km above sea level and the flight altitude higher than 10 km, then a water pixel has to fulfill the criteria:

\[
\rho^*(\text{blue}) \leq 0.20 \quad \text{and} \quad \rho^*(\text{blue}) > \rho^*(\text{green}) - 0.03 \\
\rho^*(\text{NIR}) < \rho^*(\text{green}) \quad \text{and} \quad \rho^*(1.6\mu m) < T_{\text{water,SWIR1}}
\]

where \( T_{\text{water,SWIR1}} \) is the water threshold reflectance in the SWIR1 band (around 1.6 \( \mu m \)) as defined in the preference parameter file, see chapter 9.4. Basically, the gradient of the apparent water reflectance has to be negative.

If the pixel elevation is higher than 1.2 km or the flight altitude lower than 10 km, the criterion of a negative gradient for the apparent reflectance does not properly work (as the path radiance in the visible, especially in the blue, becomes small), and the following criterion based on surface reflectance instead of apparent reflectance is used:

\[
\rho(\text{NIR}) \leq |T_{\text{water,NIR}}| \quad \text{and} \quad \rho(\text{SWIR1}) \leq |T_{\text{water,SWIR1}}|
\]

where \( T_{\text{water,NIR}} \) is the water reflectance threshold for the NIR band (around 850 nm). Equation 10.49 is also applied if any threshold \( T_{\text{water,NIR}} \) or \( T_{\text{water,SWIR1}} \) is set to a negative value. In this case the elevation (pixel below 1.2 km) and flight altitude (higher than 10 km) criteria are overruled.

Saturated pixels:
These pixels fulfill the criterion

\[
DN(\text{blue}) \geq T_{\text{saturation}}
\]

where \( DN(\text{blue}) \) is the digital number in a blue band (around 470 nm) and the threshold \( T_{\text{saturation}} \) is defined in the preference parameter file. If a blue band does not exist, a green band (around 550 m) is used as a substitute. If a green band also does not exist, a red band (around 650 nm) is used. \( T_{\text{saturation}} = b \cdot \text{encoding} \), default \( b=1.0 \), e.g. \( 1.0*255=255 \) for 8 bit sensors with \( \text{encoding}=255 \). A lower value, e.g. \( b=0.9 \), might be used because nonlinear effects might already occur at lower radiance levels. This would provide some safety margin and indicate situations near saturation. Setting \( b=0.9 \) for an 8-bit sensor means that pixels exceeding \( DN = 0.9*255=230 \) will be marked as (nearly) saturated.

For a 32 bit encoding (integer or float) no saturation threshold is defined. As saturation usually occurs in the blue-to-red part of the spectrum, channels in this region are checked and assigned to the class ‘saturated’, false color coded red in the "*.out_hcw.bsq" file. However, the "*.atm.log" file contains the percentage of saturated pixels for each channel.

Cloud over land:
Pixels must satisfy the conditions:
\[
\rho^*(\text{blue}) > T_c \quad \text{and} \quad \rho^*(\text{red}) > 0.15 \quad \text{and} \quad \rho^*(\text{NIR}) / \rho^*(\text{red}) < 2 \\
\quad \text{and} \quad \rho^*(\text{NIR}) > 0.8 \rho^*(\text{red}) \quad \text{and} \quad \rho^*(\text{NIR}) / \rho^*(\text{SWIR}_1) > 1 \\
\quad \text{and} \quad \text{NDSI} < 0.7 \quad \text{or} \quad \text{DN(\text{blue})} > T_{\text{saturation}}
\]
(10.51)

where \(\rho^*(\text{blue})\) is the apparent reflectance in a blue band, \(T_c\) is the cloud threshold as defined in the preference parameter file, and \(\text{DN(\text{blue})}\) is the corresponding digital number. If no blue band is available, a green band (around 550 nm) is taken as a substitute. If no green band exists, a red band (around 650 nm) is taken. NDSI is the normalized difference snow index:
\[
\text{NDSI} = \frac{\rho^*(\text{green}) - \rho^*(\text{SWIR}_1)}{\rho^*(\text{green}) + \rho^*(\text{SWIR}_1)}
\]
(10.52)

Note that saturated pixels in visible bands are automatically counted as cloud although they might be something else (e.g., snow, or a specular reflection from a surface). If a thermal band exists, the following cloud criterion must also be fulfilled:
\[
(1 - \rho^*(\text{SWIR}_1)) \ T_{\text{bb}} < 225 \ (\text{Kelvin}) \quad \text{and exclude} \quad T_{\text{bb}} > 300 \ (\text{Kelvin})
\]
(10.53)

where \(T_{\text{bb}}\) is the at-sensor blackbody temperature in the selected thermal band.

Cloud over water:

The following criteria have to be fulfilled:
\[
0.20 < \rho^*(\text{blue}) < 0.40 \quad , \quad \rho^*(\text{green}) < \rho^*(\text{blue}) \\
\rho^*(\text{NIR}) < \rho^*(\text{green}) \quad , \quad \rho^*(\text{SWIR}_1) < 0.15 \quad , \quad \text{NDSI} < 0.2
\]
(10.54)

For optically thick clouds it is not possible to distinguish clouds over water from clouds over land if only spectral criteria are used.

Cloud shadow:

Pixels must satisfy the conditions:
\[
0.04 < \rho^*(\text{NIR}) < 0.12 \quad \text{and} \quad \rho^*(\text{SWIR}_1) < 0.20
\]
(10.55)

and they should not belong to the water class. This may also include building shadow pixels.

Snow / ice:

Pixels must satisfy the conditions:
\[
\rho^*(\text{blue}) > 0.22 \quad \text{and} \quad \text{NDSI} > 0.6 \quad \text{and} \quad \text{DN(\text{blue})} < T_{\text{saturation}}
\]
(10.56)

The condition \(\text{DN(\text{blue})} < T_{\text{saturation}}\) means that saturated pixels in the blue spectral band are not included in the snow mask, instead they are put into the cloud class. If no blue band exists, a green band (around 550 nm) is taken. However, if the blue or green band is saturated and \(\text{NDSI} > 0.7\) then this pixel is assigned to the snow class. If a green band and a SWIR2 band (around 2.2 \(\mu\)m) exist the following relationships are used:
\[
\text{DN(\text{blue})} < T_{\text{saturation}} \quad \text{and} \quad (\rho^*(\text{blue}) > 0.22, \quad \text{NDSI} > 0.6) \\
\quad \text{or} \quad (\rho^*(\text{green}) > 0.22, \quad \text{NDSI} > 0.25, \quad \rho^*(\text{SWIR}_2)/\rho^*(\text{green}) < 0.5)
\]
(10.57)
Again, if the blue or green band is saturated and \( NDSI > 0.7 \) then the snow class is assigned.

**Cirrus over land and water:**

The apparent cirrus reflectance is calculated in the cirrus band (1.38 \( \mu m \)). Cirrus classes are defined according to the apparent reflectance. Thin cirrus over land is calculated with

\[
1.0\% < \rho^*(\text{cirrus}) < 1.5\%
\]

employing the percent reflectance unit. Medium thickness cirrus is calculated as

\[
1.5\% \geq \rho^*(\text{cirrus}) < 2.5\%
\]

and the thick cirrus class consists of pixels with

\[
2.5\% \geq \rho^*(\text{cirrus}) < 4.0\%
\]

The same definition is used for cirrus over water if the land/water distinction is still possible based on the selected spectral criteria.

Still higher apparent reflectance values are defined as `cirrus cloud’ if

\[
4.0\% \geq \rho^*(\text{cirrus}) < 5.0\%
\]

and `thick cirrus cloud’ if

\[
\rho^*(\text{cirrus}) \geq 5.0\%
\]

and no distinction concerning land/water is made for the last two classes.

Cirrus detection is is switched off in the following cases:

- no water vapor map available: if DEM height \( < 2000 \) m
- water vapor map \( W \) available: if \( W < 1 \) cm or \( W < Twv \) where \( Twv \) is the water vapor threshold specified in the file `preference\_parameters.dat`

**Haze over land:** see chapter 10.5.3

The the mean of the tasseled cap transformation (TC) is calculated. Clear pixels are those with \( TC < \text{mean}(TC) \) and \( \rho^*(\text{blue}) < T_c \) (cloud over land threshold) and \( \rho^*(\text{NIR}) > T_{\text{water}}(\text{NIR}) \) (water reflectance threshold, defined in ”preference\_parameters.dat”). Next, the mean and standard deviation \( \sigma \) of the HOT transformation are calculated. Pixels are assigned to the compact haze mask if \( HOT > \text{mean}(HOT) \), and to the large haze mask if \( HOT > \text{mean}(HOT) - 0.5 \sigma(HOT) \). Then the HOT histogram of all haze pixels is calculated. Pixels with values less than 40% of the cumulative histogram are assigned to thin-medium haze, pixels with higher values to medium-thick haze. This distinction is arbitrary and has no effect on the subsequent processing.

**Haze over water:**

Pixels must belong to the water mask, and the NIR apparent reflectance \( \rho^*(\text{NIR}) \) must be greater than the NIR clear water threshold \( T_{\text{clear\_water\_NIR}} \) defined in the preference parameter file (chapter 9.4). Thin haze over water is defined as:

\[
T_{\text{clear\_water\_NIR}} \geq \rho^*(\text{NIR}) \geq 0.06
\]
Medium haze over water is defined as:

\[ 0.06 < \rho^*(\text{NIR}) \leq T_2 \]  

(10.64)

where \( T_2 \) (default=0.12) is another editable parameter in the preference file. The method of haze removal over water is described in chapter 10.5.4. The same technique is also employed to remove sun glint.

### 10.3 Quality layers

The previous section defined a coarse pixel classification which is useful for an atmospheric correction. In addition, it supports an assessment of the quality of the processing. For example, a large error in the radiometric calibration could cause a scene classification with all pixels labeled as water. In this case, a user can immediately identify the problem. Of course, a more detailed assessment is possible with an analysis of the reflectance spectra. Nevertheless, the classification map (land, water, haze, cloud, etc) is a useful product and the quality of the atmospheric correction may depend on the correct class assignment, at least for some classes.

The previous (haze, cloud, water, land) pixel classifier is a binary decision: a pixel belongs to a certain class or not. In reality the decision is typically not unique and a class assignment has only a certain probability. As the absolute probability of a class assignment is very difficult to assess, we define three probability levels: low, medium, and high coded 30, 60, 90, respectively. These numbers might be interpreted as a percent probability, but the numbers are relative and arbitrary. Currently there are three quality layers, cloud, water, and snow, which are solely calculated with spectral criteria. The quality file is written if the corresponding flag is set to 2, see chapter 9.4 and figure 5.12 in chapter 4.

#### Cloud probability:

- **low cloud probability** (coded 30):

\[
\rho^*(\text{blue}) > 0.15 \quad \text{and} \quad \rho^*(\text{red}) > 0.15 \quad \text{and} \quad \rho^*(\text{NIR}) / \rho^*(\text{red}) < 2 \\
\text{and} \quad \rho^*(\text{NIR}) > 0.8 \quad \rho^*(\text{red}) \quad \text{and} \quad \rho^*(\text{NIR}) / \rho^*(\text{SWIR1}) > 1 \\
\text{and} \quad \text{NDSI} < 0.7 \quad \text{or} \quad \text{DN}(\text{blue}) > T_{saturation}
\]

(10.65)

where \( \rho^*(\text{blue}) \) is the apparent reflectance in a blue band, and \( \text{DN}(\text{blue}) \) is the corresponding digital number. If no blue band is available, a green band (around 550 nm) is taken as a substitute. If no green band exists, a red band (around 650 nm) is taken. Note that saturated pixels in visible bands are counted as cloud although they might be something else (e.g., snow, or a specular reflection from a surface). Only saturated pixels with a very high \( \text{NDSI} > 0.7 \) are assigned to the snow class.

- **medium cloud probability** (coded 60): same as for low probability, but with

\[
\rho^*(\text{blue}) > 0.25 \quad \text{and} \quad \rho^*(\text{red}) > 0.18
\]

(10.66)

This is similar to the standard cloud assignment in the "*_out_hcw.bsq" file where \( \rho^*(\text{blue}) > T_c, \) \( T_c = 0.25 \) (or 25%), and \( \rho^*(\text{red}) > 0.15. \) The 25% reflectance threshold in the blue (or green) band is the default value in the preference parameter file.
• high cloud probability (coded 90): same as for medium probability, but with

\[ \rho^* (\text{blue}) > 0.35 \quad \text{and} \quad \rho^*(\text{red}) > 0.25. \]  

(10.67)

If a thermal band exists, the relationships 10.53 must also be fulfilled.

**Water probability:**

The criteria for the water class are described in the previous section. The following water probability rules are employed:

• low water probability (coded 30): water pixels fulfilling the above criteria (eq.’s 10.48, 10.49). This is the water assignment in the “*.out.hcw.bsq” file.

• medium water probability (coded 60): same as for low probability, but with a more stringent NIR reflectance threshold. If no SWIR1 band (around 1.6 \( \mu m \)) exists, the criterion is:

\[ \rho^*(\text{NIR}) \leq 0.04 \]  

(10.68)

Note: the default threshold \( T_{\text{water,NIR}} \) is 0.05 (or 5% in the reflectance percent unit), defined in the preference parameter file, yielding more low probability water pixels than medium probability pixels (eq. 10.68). If a SWIR1 band exists the apparent NIR reflectance threshold is relaxed (first line of eq. 10.69) because of the additional SWIR1 surface reflectance threshold.

\[ \rho^*(\text{NIR}) \leq 0.05 \quad \text{and} \quad \rho(\text{SWIR1}) \leq 0.03 \quad \text{or} \]
\[ \rho^*(\text{NIR}) \leq 0.03 \quad \text{or} \quad \rho(\text{SWIR1}) \leq 0.03 \]  

(10.69)

The relationships on the second line of eq. 10.69 assign the water probability based on the smaller reflectance value in the NIR or SWIR1.

• high water probability (coded 90): same as for medium probability, but with lower NIR, SWIR1 thresholds.

\[ \rho^*(\text{NIR}) \leq 0.03 \quad (\text{no SWIR1 band}) \]
\[ \rho^*(\text{NIR}) \leq 0.03 \quad \text{or} \quad \rho(\text{SWIR1}) \leq 0.02 \]  

(10.70)

Note: the default threshold \( T_{\text{water,SWIR1}} \) is 0.03 (or 3% in the reflectance percent unit), defined in the preference parameter file.

**Snow / ice probability:**

The criteria for the snow-ice class are described in the previous section. As mentioned before, if pixels are saturated in the blue-green spectral bands they are counted as cloud unless the \( NDSI > 0.7 \). The following probability rules are employed for snow:
• low snow/ice probability (coded 30):

\[ \rho^*(\text{blue}) > 0.22 \quad \text{and} \quad NDSI > 0.4 \quad \text{and} \quad DN(\text{blue}) < T_{saturation} \quad (10.71) \]

If no blue band exists, a green band is used as a substitute. If a green band and a SWIR2 band exist, the rules are:

\[ DN(\text{blue}) < T_{saturation} \quad \text{and} \quad ( (\rho^*(\text{blue}) > 0.22 , \quad NDSI > 0.4) \]
\[ \quad \text{or} \quad (\rho^*(\text{green}) > 0.22 , \quad NDSI > 0.25 , \quad \rho^*(\text{SWIR2})/\rho^*(\text{green}) < 0.5 ) ) \quad (10.72) \]

• medium snow/ice probability (coded 60): same as for low probability, but with a more stringent NDSI threshold of 0.6. This is the snow assignment in the "hcw.bsq" file.

\[ \rho^*(\text{blue}) > 0.22 \quad \text{and} \quad NDSI > 0.6 \quad \text{and} \quad DN(\text{blue}) < T_{saturation} \quad (10.73) \]

If no blue band exists, a green band is used as a substitute. If a green band and a SWIR2 band exist, the rules are:

\[ DN(\text{blue}) < T_{saturation} \quad \text{and} \quad ( (\rho^*(\text{blue}) > 0.22 \quad \text{and} \quad NDSI > 0.6) \]
\[ \quad \text{or} \quad (\rho^*(\text{green}) > 0.22 \quad \text{and} \quad \rho^*(\text{SWIR2})/\rho^*(\text{green}) < 0.3 ) ) \quad (10.74) \]

This is very similar to the snow assignment in the "hcw.bsq" file, except for the threshold for \( \rho^*(\text{SWIR2})/\rho^*(\text{green}) \).

• high snow/ice probability (coded 90): same as for medium probability, but with a more stringent NDSI threshold of 0.7.

\[ \rho^*(\text{blue}) > 0.22 \quad \text{and} \quad NDSI > 0.7 \quad \text{and} \quad DN(\text{blue}) < T_{saturation} \quad (10.75) \]

If no blue band exists, a green band is used as a substitute. Again, if a green band and a SWIR2 band exist, the rules are:

\[ DN(\text{blue}) < T_{saturation} \quad \text{and} \quad ( (\rho^*(\text{blue}) > 0.22 \quad \text{and} \quad NDSI > 0.7) \]
\[ \quad \text{or} \quad (\rho^*(\text{green}) > 0.22 \quad \text{and} \quad \rho^*(\text{SWIR2})/\rho^*(\text{green}) < 0.2 ) ) \quad (10.76) \]

10.4 Standard atmospheric conditions

Standard conditions comprise scenes taken under a clear sky atmosphere. This means the visibility (aerosol optical thickness) can be assumed as constant over a scene, or it might vary within a certain range (excluding haze) and a visibility map can be calculated. It also includes situations with constant or spatially varying water vapor column contents.
10.4.1 Constant visibility (aerosol) and atmospheric water vapor

This is the easiest case for atmospheric correction. Still, it can often be applied if homogeneous atmospheric conditions exist. These might be encountered for small area scenes, i.e., high spatial resolution imagery. If the sensor has no channels in atmospheric water vapor regions, results of atmospheric correction are not sensitive with respect to the selected water vapor content, and a climatological value (e.g., midlatitude summer, US standard, or tropical water vapor profile) is usually sufficient. For hyperspectral instruments, the processing has to include the image-derived pixel-by-pixel water vapor map.

The program performs a check whether the specified visibility leads to negative reflectance pixels for dark surfaces in the red band (660 nm, vegetation) and NIR band (850 nm, water). If this is the case, the visibility is iteratively increased (up to VIS=120 km) to reduce the percentage of negative reflectance pixels below 1% of the scene pixels. During an interactive ATCOR session the user is notified, and can continue with the recommended visibility update or with the initial visibility (if the input visibility in the .inn file is positive). For a negative visibility (in the '*.inn' file) no visibility iterations are performed. A corresponding notice on the final visibility is given in the '*atm.log' output file. The set of visibility grid point is given in Table 10.3.

The upper visibility threshold of 80 km is a trade-off: although higher visibilities are possible they are not very likely, and even if a situation with a higher visibility (say VIS=120 km) is encountered, results of a calculation with vis = 80 km do not differ much from the results with VIS = 120 km. So the iteration capability is most important for low visibility start values.

<table>
<thead>
<tr>
<th>visibility [km]</th>
<th>vis. increment [km]</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>3</td>
</tr>
<tr>
<td>8</td>
<td>3</td>
</tr>
<tr>
<td>11</td>
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<td>100</td>
<td>20</td>
</tr>
<tr>
<td>120</td>
<td>max VIS=120 km</td>
</tr>
</tbody>
</table>

*Table 10.3: Visibility iterations on negative reflectance pixels (red, NIR bands).*

10.4.2 Aerosol retrieval and visibility map

If a sensor has the appropriate spectral bands the aerosol type and visibility or optical thickness of the atmosphere can be derived, provided the scene contains reference areas of known reflectance
behavior (Kaufman and Sendra 1988, Kaufman et al. 1997). The minimum requirements are spectral bands in the red and near IR. If the scene contains dense dark vegetation (coniferous type) the reflectance values in the red band can be obtained from a correlation with the SWIR band reflectance as detailed below. The visibility of each reference pixel can then be calculated in the red band as the intersection of the modeled at-sensor radiance curve with the measured radiance, see figure 10.11.

The measured radiance for a reference pixel of digital number DN is $L = c_0 + c_1 DN$, which is a constant value indicated by the dashed line in figure 10.11. The curve indicates the modeled radiance. It employs the reflectance of the reference surface (e.g., $\rho_{ref} = 0.02$) and uses values of path radiance, atmospheric transmittance, and global flux for the current solar and viewing geometry stored in precalculated LUTs.

\begin{equation}
L = L_p + \tau \rho_{ref} \frac{E_g}{\pi}
\end{equation}

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{figure10_11.png}
\caption{Schematic sketch of visibility determination with reference pixel.}
\end{figure}

Automatic masking of reference areas (1.6 or 2.2 $\mu$m band required, or at least red/NIR bands)

If the sensor has a SWIR band (at 1.6 or 2.2 $\mu$m), then the scene can be searched for dark pixels in this band and a correlation of the SWIR reflectance with the reflectance in the red and blue band can be employed to estimate the visibility automatically (Kaufman et al. 1997). For this purpose, we use a modified version of the original idea for the following algorithm. If a SWIR band exists the SWIR reflectance is calculated assuming a visibility of 23 km (instead of the original version of top of atmosphere reflectance). Then, water pixels are excluded by employing only those pixels with SWIR reflectance values above 1% and an NDVI > 0.1. For the 2.2 $\mu$m band the upper threshold of the reflectance of the dark pixels is selected as 5%. If the number of reference pixels is less then 1% of the image pixels, then the upper threshold is increased to 10% or finally 12%. If a 1.6 $\mu$m band exists, but no 2.2 $\mu$m band, the corresponding upper thresholds are selected as 10% and 15%, or finally 18%, respectively. The reflectance ratios for the red and blue band are then calculated as:

\begin{equation}
\rho_{red} = 0.5 \rho_{2.2} \quad \text{and} \quad \rho_{blue} = 0.5 \rho_{red} + 0.005
\end{equation}

(10.77)
\[ \rho_{\text{red}} = 0.25 \rho_{1.6} \quad \text{and} \quad \rho_{\text{blue}} = 0.5 \rho_{\text{red}} + 0.005 \] (10.78)

The offset 0.005 for the blue band yields a better correlation with ground measurements than a zero offset [46]. This situation is sketched in figure 10.12. The correlation factor of 0.5 between the 2.2 \( \mu m \) and the red region is not a universal constant, but may typically vary between 0.4 and 0.6. The correlation actually also works for dark soils. So the dark pixels may also include soil areas. For narrow band hyperspectral sensors a band close to 2.13 \( \mu m \) is used instead of a 2.20 \( \mu m \) band.

![Figure 10.12: Correlation of reflectance in different spectral regions.](image)

The red band is then used to calculate the visibility (compare figure 10.11) as the intersection of the measured radiance with the simulated visibility-dependent at-sensor radiance curve. Since the same visibility is employed for the blue spectral band this provides an opportunity to adjust the spectral behavior of the path radiance (which is essentially the aerosol path radiance, since the Rayleigh path radiance is known) in the blue spectral region.

\[ L_{p,\text{blue}}^{\text{update}} = L_{\text{blue}} - \tau_{\text{blue}} \rho_{\text{blue}} E_{g,\text{blue}} / \pi \] (10.79)

The question of an automatic aerosol type calculation is addressed next.

### Aerosol type estimation

After calculation of the scene path radiance in the blue and red region (as total minus reflected radiance, using the average values obtained for the dark reference pixels) the ratio of \( L_p(\text{blue, scene}) \) to \( L_p(\text{red, scene}) \) can be compared to the corresponding ratio for the MODTRAN standard aerosols (rural, urban, maritime, desert):

\[ d_p = \frac{L_p(\text{blue, scene})/L_p(\text{red, scene})}{L_p(\text{blue, MODTRAN})/L_p(\text{red, MODTRAN})} \] (10.80)

The aerosol type for which the double ratio \( (d_p) \) is closest to 1 is the best approximation for the scene. It approximates the corresponding MODTRAN aerosol type. However, some fine tuning is subsequently performed to be able to modify the wavelength behavior of the path radiance compared to the standard aerosol types. If \( L_p(\text{blue, scene}) \) deviates more than 5% from \( L_p(\text{blue, MODTRAN}) \) then \( L_p(\text{blue, scene}) \) is used as the valid path radiance. In addition, the
path radiance for any other bands in the blue to red region is linearly re-scaled with the factor \( \frac{L_p(\text{blue, scene})}{L_p(\text{blue, MODTRAN})} \), see Figure 10.13. Here, the path radiance in the red band is used as a fixed tie point. For wavelengths greater than 700 nm a possible typical 10% difference in path radiance between the selected aerosol type (after fine tuning) and the actual aerosol is usually not important, because path radiance contributes only a small fraction to the total radiance.

If the sensor has no blue spectral band, but a green band, than the green band is substituted and for the dense dark vegetation the surface reflectance relationship is used:

\[
\rho(\text{green}) = 1.3 \, \rho(\text{red})
\]  
(10.81)

Now eq. 10.79 is again employed for the green band instead of the blue band to calculate the path radiance, the best match to a MODTRAN aerosol type, and possibly a fine tuning of the path radiance.

\[
\begin{align*}
\text{Figure 10.13: Rescaling of the path radiance with the blue and red band.} \\
\text{After subtraction of the reflected radiance from the total radiance in the blue band the remaining} \\
\text{signal is the updated path radiance in the blue band. The path radiance of bands in the blue to} \\
\text{red region is then rescaled with interpolation.}
\end{align*}
\]

**Aerosol retrieval for VNIR sensors**

If no SWIR bands exist, but at least a red band (around 660 nm) and a NIR band (around 850 nm) a different approach has to be taken, see reference [68] for details. It starts with the assumption of average clear atmospheric conditions (visibility VIS=23 km) to calculate the surface reflectance in the red and NIR bands which is appropriate for situations of clear atmospheres (VIS = 15 - 40 km). The second step derives a mask of dark vegetation pixels using the ratio vegetation index \( \text{rvi} \) of the red and near-infrared surface reflectance, \( \text{rvi} = \rho_{\text{nir}}/\rho_{\text{red}} \), and multiple reflectance thresholds:

- The mask pixels have to fulfill: \( \text{rvi} \geq 3 \) and \( \rho_{\text{nir}} \geq 0.10 \) and \( \rho_{\text{nir}} \leq 0.25 \) and \( \rho_{\text{red}} \leq 0.04 \).

Water pixels are automatically excluded from this mask because of the \( \rho_{\text{nir}} \geq 0.10 \) condition, and soil pixels are excluded with the combination of all four conditions. If the percentage of reference pixels is smaller than 2% of the scene the search is iterated with VIS=120 km (covering the very clear conditions of visibility = 40 - 60 km). Again, if the percentage is smaller than 2%, the search is iterated with VIS=10 km to cover higher aerosol loadings (VIS = 8 - 15 km). Each visibility iteration is supplemented with an iteration of the threshold \( \rho_{\text{red}} \) which is decreased in steps of 0.005.
down to $\rho_{\text{red}} = 0.025$ to include only the darkest vegetation pixels (see [68] for details). Currently, the algorithm terminates if less than 2% reference pixels are found after these two iterations. In this case, the user has to employ the constant visibility option specifying the value of the visibility for the scene. During batch mode operation the program takes the specified visibility (from the ".inn" file). Then a check for negative reflectance pixels is performed with dark pixels in the red band (660 nm, vegetation) and the NIR band (850 nm, water), and the visibility is iteratively increased (up to VIS=60 km) to reduce the percentage of negative reflectance pixels below 1% of the scene pixels. A corresponding notice is given in the "*atm.log" output file.

The third step calculates the surface reflectance in the red band as a fraction $\alpha$ of the NIR band reflectance:

$$\rho_{\text{red}} = \alpha \cdot \rho_{\text{nir}} = 0.1 \cdot \rho_{\text{nir}}$$ (10.82)

Similar to the empirical SWIR relationships the coefficient $\alpha = 0.1$ is an average empirical value yielding results in close agreement with the SWIR method in many cases. However, deviations from the nominal value $\alpha = 0.1$ can vary about 30% depending on biome. Before the final step of atmospheric correction takes place the visibility of non-reference pixels in the scene can be set to the average value of the reference pixels or a spatial interpolation can be applied.

The visibility calculated for each reference pixel (range 5 - 190 km in ATCOR) is converted into an integer, called visibility index $vi$, with range 0-182. The visibility index is closely related to the total optical thickness $\delta$ at 550 nm, the equidistant optical thickness spacing is 0.006 for a ground at sea level, and smaller for increasing elevations.

$$\delta = 0.185 + 0.006 \cdot vi$$ (10.83)

It is easy to calculate the aerosol optical thickness (AOT) from a known total optical thickness by subtracting the Rayleigh optical thickness and a very small trace gas optical thickness, compare Fig. 2.1 in chapter 2.

With the MODTRAN code the AOT (at 550 nm) can be calculated from a given visibility VIS (km) as

$$AOT = \exp(a(z) + b(z) \ln(VIS))$$ (10.84)

where $z$ is the surface elevation, and $a(z)$, $b(z)$ are coefficients obtained from a linear regression of $\ln(AOT)$ versus $\ln(VIS)$. 
10.4.3 Water vapor retrieval

A water vapor retrieval can be included after the aerosol retrieval because the aerosol retrieval does not use water vapor sensitive spectral bands, but the water vapor algorithm (employing bands around 940 or 1130 nm) depends on aerosol properties. The water vapor retrieval over land is performed with the APDA (atmospheric precorrected differential absorption) algorithm [82]. In its simplest form, the technique uses three channels, one in the atmospheric water vapor absorption region around 940 or 1130 nm (the "measurement" channel), the others in the neighboring window regions ("reference" channels). The depth of the absorption feature is a measure of the water vapor column content, see figure 10.15.

In case of three bands the standard method calculates the water vapor dependent APDA ratio as:

\[
R_{\text{APDA}}(\rho, u) = \frac{L_2(\rho, u) - L_{2,p}(u)}{w_1(L_1(\rho_1) - L_{1,p}) + w_3(L_3(\rho_3) - L_{3,p})} \tag{10.85}
\]

where the index 1 and 3 indicates window channels (e.g. in the 850-890 nm region and 1010-1050 nm region), respectively. Index 2 indicates a channel in the absorption region (e.g., 910-950 nm). \(L\) and \(L_p\) are the total at-sensor radiance and path radiance, respectively. The symbol \(u\) indicates the water vapor column. The weight factors are determined from

\[
w_1 = (\lambda_3 - \lambda_2)/(\lambda_3 - \lambda_1) \quad \text{and} \quad w_3 = (\lambda_2 - \lambda_1)/(\lambda_3 - \lambda_1) \tag{10.86}
\]

The problem is the estimation of the surface reflectance \(\rho_2\) in the absorption band (eq. 10.85). The technique tries to estimate the reflectance \(\rho_2\) with a linear interpolation of the surface reflectance values in the window channels (ch. 1, 3) that are not or only slightly influenced by the water vapor content: Therefore, the reflectance \(\rho_2\) is calculated as

\[
\rho_2 = w_1\rho_1 + w_3\rho_3 \tag{10.87}
\]

Then equation (10.85) can be written as

\[
R_{\text{APDA}}(u) = \frac{\rho_2 \tau_2(u) E_{g2}(u)}{\rho_2 \tau_2(u = 0) E_{g2}(u = 0)} = \frac{\tau_2(u) E_{g2}(u)}{\tau_2(u = 0) E_{g2}(u = 0)} \tag{10.88}
\]
where $E_{g2}(u)$ is the global flux on the ground for the measurement channel (index 2). ATCOR employs 4 to 5 water vapor columns ($u=0.4, 1.0, 2.0, 2.9, 4.0$ cm, sea-level-to space geometry) to calculate an exponential fit function

$$R_{APDA}(u) = \exp(-\alpha + \beta \sqrt{u}) \quad (10.89)$$

which can be solved for the water vapor column $u$, see Fig. 10.16, where the diamonds in the figure mark the calculated water vapor grid points ($u= 0.4, 1.0, 2.0, 2.9$ cm):

$$u = \left( \frac{\alpha + \ln R_{APDA}}{\beta} \right)^2 \quad (10.90)$$

Equations (10.85, 10.87 to 10.90) are iterated, starting with $u=1.0$ cm, calculating $R_{APDA}$, updating $u$, $L_{i,p}(u)$, $\rho_1$, $\rho_3$ and repeating the cycle. A minimum of two channels (one reference, one measurement channel) is required. The advanced APDA method can take into account multiple absorption channels in the 910-960 nm and 1110-1150 nm regions. Two water vapor retrieval algorithms are available in ATCOR (compare chapter 9.5, parameter iwv_model = 1, 2):

1. The water vapor maps with the smallest standard deviation in the 940 nm and 1130 nm region are selected. Finally, if both regions are available, the average of these two water vapor maps is taken (parameter iwv_model=1 in the ”.inn” file). The scan angle dependence of the path radiance is taken into account.

2. A linear regression ratio (LIRR) is applied to multiple bands (parameter iwv_model=2). This water vapor map might be more accurate, because the regression reduces sensor noise and may partially compensate calibration problems in lucky cases. Although the water vapor map might be less noisy, the retrieved surface reflectance spectrum will always retain any channel calibration problems. The scan angle dependence of the path radiance is not accounted for.

Remarks:
1. The APDA algorithm is relatively fast. Its disadvantage is that it is not stable numerically for very low reflectance targets (water, shadow regions). The transmittance slope ratio method [76] might work better in these cases, so it is an interesting alternative water vapor algorithm. However, since the required processing time is much higher than for the APDA method, it is currently not implemented in the ATCOR environment. In addition, the method requires data with a very accurate spectral and radiometric calibration, otherwise its potential advantage will be lost.

2. Five water vapor grid points at 0.4, 1.0, 2.0, 2.9, and 4.0 cm are sufficient to cover the 0.5 - 5.0 cm range with an accuracy of about 5-10 % [70].

### 10.5 Non-standard conditions

The non-standard situations refer to scenes with a substantial amount of haze and shadow areas. The non-standard atmospheric conditions treat the haze removal and de-shadowing employing spectral and statistical algorithms. Although bidirectional surface reflectance effects are independent of the atmospheric conditions, the subject is included here, because the isotropic reflector is used for the standard conditions. We present some methods of BRDF correction in flat and rugged terrain.

#### 10.5.1 Haze removal

In many cases of satellite imagery the scene contains haze and cloud areas. The optical thickness of cloud areas is so high that the ground surfaces cannot be seen, whereas in hazy regions some information from the ground is still recognizable. In ATCOR the scene is partitioned into clear, hazy, and cloud regions. Here we will treat the low altitude boundary layer (0 - 3 km) haze as opposed to high altitude cirrus. Thin boundary layer haze can be detected with broad-band multispectral instruments, while a detection of thin cirrus requires specific narrow bands around 1.38 μm or 1.88 μm, compare chapter 10.5.5. As a first approximation, haze is an additive component to the radiance signal at the sensor. It can be estimated and removed as described below. Cloud
areas have to be masked to exclude them from haze areas and to enable a successful haze removal. The treatment of cloud shadow regions is discussed in chapter 10.5.6.

Two de-hazing algorithms are available: method 1 is based on the haze thickness map [51], method 2 employs the haze optimized transform [101]. Method 1 can be submitted from a GUI (graphical user interface) or as a batch job and performs the de-hazing on the original digital number (DN) data. This is an optional pre-processing step to ATCOR. An atmospheric correction can be performed as an independent next step to obtain surface reflectance data. Method 2 performs a de-hazing embedded in ATCOR, so it is always combined with the atmospheric correction. It is available in the GUI and batch modes.

10.5.2 Haze removal method 1

This method automatically detects and removes haze in medium and high spatial resolution multispectral images. The dark-object subtraction algorithm is further developed to calculate a haze thickness map. The haze thickness is computed for each spectral band excluding very bright object, because these can be misinterpreted as hazy areas. The haze thickness for these bright object pixels is approximated by a spatial interpolation. The first step is the search for dark pixels using a small local nonoverlapping window box (w=3 x 3 pixels) for the calculation of the haze thickness map (HTM). For this purpose a blue spectral channel is employed, because it is most sensitive to haze. If no blue band exists, the green band is taken. The next step calculates an additional HTM map with a moderately large window size (e.g. w= 21 x 21 pixels). It is used to label haze and haze-free regions by thresholding this HTM map. The third step calculates the correlation of the band-specific HTM(\lambda) maps, and re-scales these maps in the interval (1,0) for \lambda = (\lambda_{blue}, 2.2\mu m). More details can be found in the reference paper [51].

10.5.3 Haze removal method 2

The method 2 haze removal algorithm runs fully automatic. It is a combination of the improved methods [61], [101] and consists of five major steps:

1. Masking of clear and hazy areas with the tasseled cap haze transformation [17].

\[ TC = x_1 \ast BLUE + x_2 \ast RED \]  \hspace{1cm} (10.91)

where BLUE, RED, \( x_1 \), and \( x_2 \) are the blue band, red band, and weighting coefficients, respectively. The clear area pixels are taken as those pixels where TC is less than the mean value of TC.

2. Calculation of the regression between the blue and red band for clear areas ("clear line" slope angle \( \alpha \)), see figure 10.17. If no blue band exists, but a green spectral band, then the green band is used as a substitute.

3. Haze areas are orthogonal to the "clear line", i.e., a haze optimized transform (HOT) can be defined as (Zhang et al. 2002):

\[ HOT = BLUE \ast sin\alpha - RED \ast cos\alpha \]  \hspace{1cm} (10.92)

4. Calculation of the histogram of HOT for the haze areas.
5. For bands below 800 nm the histograms are calculated for each HOT level \( j \). The haze signal \( \Delta \) to be subtracted is computed as the DN corresponding to HOT(level \( j \)) minus the DN corresponding to the 2% lower histogram threshold of the HOT(haze areas). The de-hazed new digital number is (see figure 10.17):

\[
DN(new) = DN - \Delta
\]  

(10.93)

\*Figure 10.17: Haze removal method. Left: regression between red and blue band for clear areas. Right: calculation of Delta=\( \Delta \) as a function of the HOT haze level (example Landsat TM band 1).

So the haze removal is performed before the surface reflectance calculation. Two options are available: the use of a large area haze mask (eq. 10.94), which is superior in most cases, or a compact smaller area haze mask (eq. 10.95).

\[
HOT > \text{mean}(HOT) - 0.5 \times \text{stdev}(HOT)
\]  

(10.94)

\[
HOT > \text{mean}(HOT)
\]  

(10.95)

In addition, the user can select between haze removal of ”thin / medium haze” or ”thin to moderately thick haze”, the last option is superior in most cases.

The algorithm only works for land pixels, so the near infrared band (NIR) is used to exclude water pixels. The current implementation provides a mask for haze-over-land (see the ‘*out_hcw.bsq’ file). The haze-over-water mask is treated in the next section.

Figure 10.18 shows an example of a subset of an Ikonos scene of Dresden where the haze removal algorithm was applied. More images with the results of the haze removal method are shown on ATCOR’s web page, http://www.rese.ch or http://www.op.dlr.de/atcor.

10.5.4 Haze or sun glint removal over water

The haze removal over water uses a near infrared (NIR) band to estimate the spatial distribution of haze. The principal method is described in [78]. We use a modified version of this approach without an interactive definition of haze polygons. First, the water pixels are masked either using spectral criteria or taking an external water map. If the scene is named "scene1.bsq" the external map must be named "scene1_water_map.bsq", a 1-channel 8-bit/pixel or 16-bit/pixel file where water is coded
with an arbitrary positive number. The external map is automatically taken if it is placed in the same folder as the scene. The second step is the definition of clear water pixels using the apparent reflectance in the NIR band. Pixels are labeled as clear if

$$\rho^*(\text{NIR}) < T_1$$  \textit{clear pixels}  \tag{10.96}$$

The default value is $T_1 = 0.04$ (i.e. 4%). The value is one of the editable preference parameters (see chapter 9.4). Thin haze over water is defined as:

$$T_1 \leq \rho^*(\text{NIR}) \leq 0.06$$  \textit{thin haze}  \tag{10.97}$$

Medium haze over water is defined as:

$$0.06 < \rho^*(\text{NIR}) \leq T_2$$  \textit{medium haze}  \tag{10.98}$$

The default value is $T_2 = 0.12$ (i.e. 12%). This value is also one of the editable preference parameters. The third step is a linear regression between haze pixels in the NIR band and each other (reflective) band. The regression is iterated with only those pixels deviating less than half a standard deviation from the average. If $\alpha_j$ and $\beta_j$ denote offset and slope of the regression line, respectively, the de-hazed pixel for each channel $j$ can be calculated as

$$DN(\text{corrected},j) = DN(\text{original},j) - (\alpha_j + \beta_j DN_{\text{NIR}} - DN(\text{clear},j))$$  \tag{10.99}$$

where $DN(\text{clear},j)$ is the average of all clear water pixels in channel $j$. The same technique is also employed to remove sun glint. The main problem is the specification of the clear water threshold. If the threshold is too low, clear water pixels are included in the haze mask, if it is set too high haze or sun glint pixels will be included in the clear pixel class. There is no unique solution, because sandy bottoms over shallow water can have a similar spectral reflectance behavior as haze, so the clear water threshold is scene-dependent. In addition, the upper threshold defining haze (or sun glint) might be scene-dependent. However, the default values usually provide good results and a solid basis for a possible iteration of these two parameters.
10.5.5 Cirrus removal

On the first glance, images contaminated by cirrus appear similar to hazy scenes discussed in the previous section. However, haze usually occurs in the lower troposphere (0-3 km) while cirrus clouds exist in the upper troposphere and lower stratosphere (8 - 16 km). The effect of boundary layer haze can be observed in the visible region, but seldom in longer wavelength channels > 850 nm. However, cirrus also affects the NIR and SWIR spectral regions. Thin cirrus clouds are difficult to detect with broad-band multispectral satellite sensors in the atmospheric window regions, especially over land, because land scenes are spatially inhomogeneous and this type of cloud is partially transparent. On the other hand, water vapor dominates in the lower troposphere and usually 90% or more of the atmospheric water vapor column is located in the 0 - 5 km altitude layer. Therefore, if a narrow spectral band is selected in a spectral region of very strong water vapor absorption, e.g., around 1.38 $\mu$m or 1.88 $\mu$m, the ground reflected signal will be totally absorbed, but the scattered cirrus signal will be received at a satellite sensor or a sensor in a high-altitude aircraft (e.g., 20 km AVIRIS scenes).

So a narrow channel at 1.38 $\mu$m is able to detect cirrus clouds, and if a correlation of the cirrus signal at this wavelength and other wavelengths in the VNIR and SWIR region can be found, then the cirrus contribution can be removed from the radiance signal to obtain a cirrus-corrected scene. The basic ideas of cirrus correction were presented in several papers ([23], [24], [26], [75]). The algorithm differs for water and land pixels. For water, a scatterplot of the 1.38 $\mu$m versus the 1.24 $\mu$m channel is used, for land the band-correlation is determined from a scatterplot of the 1.38 $\mu$m versus a red channel (around 0.66 $\mu$m). To obtain a high sensitivity, only vegetation pixels are taken because they have a low reflectance in the red spectral region, so the cirrus contribution is easily traced. The scatterplot is computed in terms of the apparent (TOA or at-sensor) reflectance of $\rho_{1.38}$ versus $\rho_{\text{red}}$, where the apparent reflectance is defined as:

$$\rho^* = \frac{\pi L}{E_s \cos \theta_s}$$ (10.100)

where L is the recorded radiance signal, $E_s$ the extraterrestrial solar irradiance for the selected band, and $\theta_s$ is the solar zenith angle. Following [23] the method can be described by the following set of equations:

$$\rho^*(\lambda) = \rho_c(\lambda) + \frac{T_c(\lambda)}{1 - s_c(\lambda)} \rho(\lambda)$$ (10.101)

Here, $\rho_c$ is the reflectance of the cirrus cloud, $T_c$ the two-way transmittance (direct plus diffuse) through the cloud, $\rho$ the reflectance of the "virtual" surface (land or water surface including all effects of molecular and aerosol scattering below the cirrus), and $s_c$ is the cloud base reflectance of upward radiation. Eq. 10.101 can be simplified, because of $s_c \rho << 1$, yielding

$$\rho^*(\lambda) = \rho_c(\lambda) + T_c(\lambda) \rho(\lambda)$$ (10.102)

With the assumption that the cirrus reflectance $\rho_c(\lambda)$ is linearly related to the cirrus reflectance at 1.38 $\mu$m we obtain

$$\rho^*(\lambda) = \frac{\rho_c(1.38\mu m)}{\gamma}$$ (10.103)

where $\gamma$ is an empirical parameter derived from the scene scatterplot of $\rho_{1.38}$ versus $\rho_{\text{red}}$ (land) or $\rho_{1.24}$ (water). It depends on the scene content, cirrus cloud height, and solar and viewing angles. Fig. 10.19 shows an example of such a scatterplot. The red line is the left-side boundary of
data points that are not influenced by ground surface reflection, i.e. cirrus-contaminated pixels are clustered around this line, and its slope represents the correlation coefficient $\gamma$ (the blue line represents the first of several iterations). Papers on the cirrus algorithm often restrict eq. 10.103 to the wavelength interval $0.4 < \lambda < 1$ $\mu$m, but we will extend this relationship into the SWIR region. Substituting eq. 10.103 into eq. 10.102 yields

$$T_c(\lambda) \rho(\lambda) = \rho^*(\lambda) - \rho_c(1.38\mu m) / \gamma$$  \hspace{1cm} (10.104)

![Figure 10.19: Scatterplot of apparent reflectance of cirrus (1.38 $\mu$m) band versus red band.](image)

Neglecting the cirrus transmittance $T_c$ (i.e., setting $T_c = 1$), we obtain the "cirrus path radiance corrected" apparent reflectance image (index ‘cc’):

$$\rho_{cc}^*(\lambda) = \rho^*(\lambda) - \rho_c(1.38\mu m) / \gamma$$  \hspace{1cm} (10.105)

As the cirrus is almost on top of the atmosphere we have $\rho_c(1.38\mu m) = \rho_c^*(1.38\mu m)$ and the apparent cirrus reflectance can be calculated with eq. 10.100. Cirrus removal is conducted as the first step during atmospheric correction, followed by the aerosol and water vapor retrievals. If the average water vapor column $W$ of a scene is less than some threshold (default $W=0.6$ cm) then the cirrus removal algorithm is switched off, to avoid a misinterpretation of bright surfaces as cirrus in the 1.38 $\mu$m channel. Normally, atmospheric water vapor completely absorbs surface features in the 1.38 $\mu$m channel, but the channel might become partly transparent to surface features for very low water vapor values. This water vapor threshold can be set by the user, see chapter 9.4.

The file ‘xxx.out_hcw.bsq’ (haze, cloud, water) corresponding to a scene ‘xxx.bsq’ contains three relative levels of cirrus optical thickness (thin, medium, and high). The corresponding thresholds are arbitrarily set depending on the statistics (mean, standard deviation) of the apparent reflectance $\rho_c^*(1.38\mu m)$ map. The file ‘xxx.out_hcw.bsq’ is intended as a visual aid or quicklook, therefore, the cirrus level maps of different scenes cannot be compared quantitatively. As an example, a certain scene setting could be:

- thin cirrus thickness, color coded as light yellow, with $0.010 < \rho^*(\text{cirrus}) \leq 0.015$,
- medium thickness, color coded as darker yellow, with $0.015 < \rho^*(\text{cirrus}) \leq 0.025$,
- high thickness, color coded as bright yellow, with $\rho^*(\text{cirrus}) > 0.025$ (reflectance units).
In addition to the 1.38 μm cirrus channel, another channel (index w1) around 1.24 μm (or as a substitute a NIR channel from the 800 to 900 nm region) is employed with a ratio criterion to define cirrus pixels:

\[ \frac{\rho^*(\text{cirrus})}{\rho^*(w1)} > T(\text{cir}) \]  

Reference [25] proposes a threshold of \( T(\text{cir}) = 0.3 \) to distinguish tropospheric aerosols due to dust storms from cirrus clouds. However, in the absence of dust storms, this threshold is too high and predicted no cirrus in a number of test scenes containing a lot of cirrus clouds. Therefore, we use much lower values of \( T(\text{cir}) \) ranging from 0.01 (for water vapor columns \( W > 1 \) cm) to \( T(\text{cir}) = 0.15 \) (for \( W < 0.5 \) cm). So with these thresholds, tropospheric aerosols might be misclassified as cirrus in situations with dust storms, but this is a necessary trade-off. In any case, those cloud areas are excluded from the map of pixels employed for the aerosol retrieval, which is the main purpose.

The cirrus and (boundary layer) haze removal options are exclusive, i.e., only one of them can be selected per run.

### 10.5.6 De-shadowing with matched filter

Remotely sensed optical imagery of the Earth’s surface is often contaminated with cloud and cloud shadow areas. Surface information under cloud covered regions cannot be retrieved with optical sensors, because the signal contains no radiation component being reflected from the ground. In shadow areas, however, the ground-reflected solar radiance is always a small non-zero signal, because the total radiation signal at the sensor contains a direct (beam) and a diffuse (reflected skylight) component. Even if the direct solar beam is completely blocked in shadow regions, the reflected diffuse flux will remain, see Figure (10.20). Therefore, an estimate of the fraction of direct solar irradiance for a fully or partially shadowed pixel can be the basis of a compensation process called de-shadowing or shadow removal. The method can be applied to shadow areas cast by clouds or buildings.

This section describes a de-shadowing method based on the matched filter approach, which is complementary to the scene-based method described in section 10.1.2.

The proposed de-shadowing technique works for multispectral and hyperspectral imagery over land acquired by satellite / airborne sensors. The method requires a channel in the visible and at least one spectral band in the near-infrared (0.8-1 μm) region, but performs much better if bands in the short-wave infrared region (around 1.6 and 2.2 μm) are available as well. The algorithm consists of these major components: (i) the calculation of the covariance matrix and zero-reflectance matched filter vector, (ii) the derivation of the unscaled and scaled shadow function, (iii) a histogram thresholding of the unscaled shadow function to define the core shadow areas, (iv) a region growing to include the surroundings of the core shadow areas for a smooth shadow/clear transition, and (v) the de-shadowing of the pixels in the final shadow mask. Details are published in [67].

The method starts with a calculation of the surface reflectance image cube \( \rho_i = \rho(\lambda_i) \), where three spectral bands around \( \lambda_i = 0.85, 1.6, \) and 2.2 μm are selected. These bands from the near and shortwave infrared region are very sensitive to cloud shadow effects, because the direct part of the downwelling solar radiation flux at the ground level is typically 80% or more of the total downwelling flux. Channels in the blue-to-red region (0.4-0.7 μm) are not used for the detection of shadow regions because they receive a much larger diffuse radiation component, making them less sensitive to partial shadow effects. Instead, visible channels serve to define a potential cloud mask.
CHAPTER 10. THEORETICAL BACKGROUND

The surface reflectance is first computed with the assumption of full solar illumination, i.e., the global flux on the ground consists of the direct ($E_{\text{dir}}$) and diffuse ($E_{\text{dif}}$) component. If $\text{DN}_i$ denotes the digital number of a pixel, $L_P$ the path radiance, and $\tau$ the atmospheric transmittance (ground-to-sensor) the surface reflectance can be obtained as:

$$\rho_i(x,y) = \frac{\pi d^2 \{c_0(i) + c_1(i)\text{DN}_i(x,y)\}}{\tau_i \{E_{\text{dir},i} + E_{\text{dif},i}\}}$$

(10.107)

Here, $d$ is the Earth-Sun distance at the image acquisition time in astronomical units, $c_0$ and $c_1$ are the radiometric calibration coefficients (offset and slope) to convert the digital number into the corresponding at-sensor radiance $L$, i.e., $L = c_0 + c_1 \text{DN}_i$, and $i$ is the channel index.

The proposed de-shadowing algorithm consists of a sequence of eight processing steps as sketched in Fig. 10.21. It starts with the atmospheric correction. The next step is the masking of water bodies and cloud areas with simple spectral criteria as detailed below. Water pixels have to be excluded as far as possible to avoid their assignment as shadow pixels. Step 3 calculates the covariance matrix $C(\rho)$ where $\rho$ is the surface reflectance vector comprising only the non-water and non-cloud pixels. For each pixel, this vector holds the reflectance values in the 3 selected channels (around 0.85, 1.6, 2.2 $\mu$m). The matched filter is a vector tuned to a certain target reflectance spectrum $\rho_t$ to be detected [2]:

$$V_{mf} = \frac{C^{-1}(\rho_t - \bar{\rho})}{(\rho_t - \bar{\rho})^T C^{-1} (\rho_t - \bar{\rho})}$$

(10.108)

Here, $\bar{\rho}$ is the scene-average spectrum, without the water/cloud pixels. Selecting $\rho_t = 0$ for a shadow target yields a special simplified form of the matched filter, where the 'sh' index symbolizes shadow:

$$V_{\text{sh}} = -\frac{C^{-1} \bar{\rho}}{\bar{\rho}^T C^{-1} \bar{\rho}}$$

(10.109)

The shadow matched filter vector is then applied to the non-water/non-cloud part of the scene and yields the still un-normalized values $\phi$ that are a relative measure of the fractional direct illumination, also called unscaled shadow function here:

$$\Phi(x,y) = V_{\text{sh}}^T (\rho(x,y) - \bar{\rho})$$

(10.110)
The matched filter calculates a minimum RMS shadow target abundance for the entire (non-water/non-cloud) scene. Therefore, the values of $\Phi$ are positive and negative numbers. The arbitrary, image-depending range of $\Phi$ has to be rescaled to the physical range from 0 to 1, where 0 indicates no direct illumination (full shadow), and 1 means full direct illumination. The histogram of $\Phi$ is used to rescale the image data. Fig. 10.22 shows a schematic sketch of such a histogram with a smaller peak (at $\Phi_2$) representing the shadow pixels and the main peak (at $\Phi_{\text{max}}$) representing the majority of the fully illuminated areas. The statistical assumption is used that full direct solar illumination is already obtained for pixels with $\Phi(x,y) = \Phi_{\text{max}}$. Then the values $\Phi$ are linearly mapped from the unscaled ($\Phi_{\text{min}}, \Phi_{\text{max}}$) interval onto the physically scaled (0,1) interval, where the scaled shadow function is named $\Phi^*$:

$$\Phi^* = \frac{\Phi - \Phi_{\text{min}}}{\Phi_{\text{max}} - \Phi_{\text{min}}} \quad if \quad \Phi \leq \Phi_{\text{max}}$$

$$\Phi = 1 \quad if \quad \Phi > \Phi_{\text{max}}$$

The smallest value of the scaled shadow function is $\Phi^*_{\text{min}} = 0$, which means no direct illumination. However, to avoid overcorrection and to cope with scenes containing merely partial shadow areas, it is advisable to set $\Phi^*_{\text{min}}$ at a small positive value. This value of $\Phi^*_{\text{min}}$, i.e., the minimum fractional direct illumination (deepest shadow in a scene, typically ranging between 0.05 and 0.10) is scene-dependent, see the detailed discussion below.

In principle, the de-shadowing could now be performed with the physically scaled function $\Phi^*$, which represents the fraction of the direct illumination for each pixel in the $\rho$ vector, i.e., the complete scene without cloud and water pixels. However, since the matched filter is not a perfect shadow transformation, it is much better to restrict its application to the potential, most-likely shadow areas. This is an important processing step to reduce the number of mis-classifications or false-alarms. If omitted it will cause strange 'shadow' pixels scattered all over the image. An example can be found in the central part of Fig. 10.23 where the standard shadow map contains a lot of artifact shadow areas.
Therefore, the proposed method tries to find the core shadow areas in a scene, and subsequently expands the core regions to obtain the final mask that includes a smooth shadow/clear transition. The physically scaled shadow function $\Phi^*$ is then applied only to the pixels in the final mask.

The histogram of the unscaled shadow function $\Phi$ can be employed to separate regions of low values of $\Phi$ from the moderate-to-high values, compare Fig. 10.22. A threshold $\Phi_T$ can be set in the vicinity of the local histogram minimum ($\Phi_1$) and the core shadow mask is defined by those pixels with $\Phi(x,y) < \Phi_T$. The details of the choice of $\Phi_T$ are discussed below. As always with thresholding, some arbitrariness is involved in the final selection.

Once the core shadow mask has been defined, it is expanded to include the surrounding shadow/clear transition zone of 100 m width. De-shadowing with the scaled shadow function $\Phi^*$ is then exclusively applied to the pixels in this final mask. This means the direct solar flux ($E_{\text{dir}}$ term in eq. 10.107) has to be multiplied with $\Phi^*(x,y)$:

$$\rho_i(x,y) = \frac{\pi(d^2\{c_0(i) + c_1(i)DN_i(x,y)\} - L_{p,i})}{\tau_i\{E_{\text{dir},i}\Phi^*(x,y) + E_{\text{dif},i}\}}$$

(10.113)

In equations (10.107, 10.113) the aerosol optical thickness or visibility required for the atmospheric terms (path radiance, transmittance, direct and diffuse flux) can be derived from the image provided the necessary bands in the visible and shortwave infrared region exist and the scene contains dark reference areas [41]. Otherwise, the user has to specify an estimated visibility. The second important atmospheric parameter is the water vapour column. For instruments with bands in the atmospheric water vapour regions this information can be derived from the image data [82], otherwise an estimate has to be provided by the user. In summary, three channels (around 0.85, 1.6, and 2.2 $\mu$m) are used to define a matched filter vector with three elements per pixel. For each image pixel the surface reflectance in these three channels and the scene-average reflectance of these channels are calculated to obtain the unscaled shadow function, and finally the scaled shadow function. The same shadow function is employed to de-shadow the imagery not only in the initial three channels but for all channels of the sensor (eq. 10.113).
Figure 10.23: Cloud shadow maps of a HyMap scene.

Left: surface reflectance image of HyMap at Chinchon, Spain, 12 July 2003. Colour coding: RGB=878, 646, 462 nm channels. Center: standard shadow map showing a lot of artifact shadow areas (grey patches) which do not appear with the core shadow approach (right part). Right: improved cloud shadow map derived from core shadow regions.

One of the most important parameters is the available number of spectral channels during the covariance matrix and matched filter part of the algorithm. The minimum requirement is a band in the near-infrared region (0.8 - 1.0 µm). The performance usually increases significantly if two additional bands (at 1.6 µm and at 2.2 µm) are available, i.e., a Landsat TM type of multispectral sensor. Even for hyperspectral imagery these three bands (around 0.85, 1.6, 2.2 µm) are sufficient for the matched filter calculation. The usage of a hundred bands would not be helpful, but only cause numerical problems during the inversion of the covariance matrix (eq. 10.109).

Spectral channels from the visible region are merely employed for the masking of cloud regions, not for the matched filter part, because water, vegetation, dark soils, and shadowed pixels all range within a few percent reflectance. In addition, the visible region is not very sensitive to partial shadow effects, because of its larger fraction of diffuse radiation component as compared to wavelengths longer than 0.8 µm.

The distinction of water bodies from cloud shadow areas may be difficult or impossible if it is based merely on spectral reflectance shape and amplitude information. Water bodies should be excluded as far as possible to improve the performance of the de-shadowing algorithm. Currently, water and cloud pixels are masked with the spectral criteria defined in the file 'preference_parameters.dat', see chapter 9.4, e.g.,

\[
\rho(0.85\mu m) \leq 5\% \quad \text{and} \quad \rho(1.6\mu m) \leq 3\% \quad (\text{water}) \\
\rho(0.48\mu m) \geq 25\% \quad \text{and} \quad \rho(1.6\mu m) \geq 25\% \quad (\text{cloud})
\] (10.114) (10.115)
If no channel in the blue region is available, a channel in the green (0.5-0.6 \( \mu m \)) or red part of the spectrum (0.6-0.68 \( \mu m \)) could be used as a substitute. Both criteria do not uniquely define the corresponding class. The water criteria allow some margin for turbid water in the NIR region. The more restrictive criterion \( \rho(0.85 \mu m) < 3\% \) would perform better for clear water bodies. However, it would fail for moderately turbid or muddy waters. Other common water classification criteria such as average reflectance over all bands \( \bar{\rho} \leq 3\% \) or \( \rho(0.4 - 0.6 \mu m) < 6\% \) may also fail. So one has to compromise and tolerate a certain amount of misclassification for a fully automatic algorithm. The scaled shadow map \( \Phi^*(x, y) \) is written to an output file.

The histogram of the unscaled shadow function \( \Phi \) (Fig. 10.22) typically has a main peak at \( \Phi_{max} \), a smaller secondary peak (at \( \Phi_2 \)) due to shadow pixels, and a local minimum (at \( \Phi_1 \)). The secondary peak can be determined by level-slicing the normalized histogram. We arbitrarily define a threshold \( \Phi_T \) as the intersection of this slice line at the level of \( h(\Phi_2) \) with the normalized histogram \( h(\Phi) \) for \( \Phi_1 < \Phi < \Phi_{max} \). The approach with a main peak and a smaller secondary peak is restricted to cases where the percentage of shadow pixels in the scene is less than about 25\%. This applies to the fully automatic processing mode. If the secondary peak at \( \Phi_2 \) is not clearly defined numerically, i.e., no local minimum found at \( \Phi_1 \), or histogram difference \( h(\Phi_2) - h(\Phi_1) < 0.03 \), then \( \Phi_T \) is defined as the intersection of the slice level 0.10 with \( h(\Phi) \) for \( \Phi < \Phi_{max} \). More flexibility exists in the interactive mode, see chapter 2.5, figure 5.41.

Masking of the core shadow areas with \( \Phi < \Phi_T \) (Fig. 10.22) is critical like any thresholding process: a large threshold could potentially include non-shadow areas, a low threshold could miss shadow areas. The current automatic algorithm has the three user-selectable options of a small, medium, or large core shadow mask corresponding to thresholds set at \( \Phi_T - 0.1, \Phi_T, \) and \( \Phi_T + 0.1 \), respectively. The default value for the fully automatic algorithm is the medium-size mask. In addition, an interactive mode for adjusting the threshold is also available.

A second tunable parameter is the minimum fractional direct illumination \( \Phi_{min}^* \), also called depth of shadow. Theoretically, it can be zero, i.e., a completely shadowed pixel receiving only diffuse solar illumination. However, a too low estimate close to zero will boost the surface reflectance, especially for channels in the 1.5 - 2.5 \( \mu m \) region (eq. 10.113), since the diffuse solar radiation term \( E_{dif} \) is very small. Therefore, small positive values of \( \Phi_{min}^* \) are recommended. The range of \( \Phi_{min}^* \) is typically from 0.05 to 0.1, with the default set at \( \Phi_{min}^* = 0.08 \). The third tunable parameter is \( \Phi_{max}^* \), providing the range of stretching of the unscaled shadow function into the scaled function. The default of \( \Phi_{max}^* \) is the location of the maximum of the histogram of \( \Phi \), but it could be set at a greater value if the corrected image is too dark in the expanded shadow regions, which indicates the histogram maximum does not represent fully illuminated areas.

The advantage of the presented method is its fast processing performance, because it relies exclusively on spectral calculations and avoids time-consuming geometric cloud/shadow pattern considerations. The drawback is that useful geometric information is neglected.

In some cases it is useful to have the de-shadowed digital number (DN) image in addition to the surface reflectance product. This facilitates a comparison with the originally recorded DN imagery. The conversion from reflectance to the corresponding at-sensor radiance is performed with eq. (10.1). Then eq. (10.6) is employed to compute the de-shadowed DN image for channel \( k \):

\[
DN(k) = \frac{L(k) - c_0(k)}{c_1(k)}
\]  

(10.116)

Figure 10.24 shows an example of de-shadowing. More images with the results of the de-shadowing method can be found on ATCOR’s web page, http://www.rese.ch or http://www.op.dlr.de/atcor .
Figure 10.24: De-shadowing of a HyMap scene.
Same image as Figure 10.23. Left: original scene, right: after de-shadowing.

10.6 Correction of BRDF effects

The bottom of atmosphere reflectance as retrieved after standard ATCOR atmospheric compensation is highly variable due to the influence of the bidirectional reflectance distribution function (BRDF, [56]). The observed reflectance value may deviate from the average spectral albedo by up to 30%, specifically for vegetation and man made surfaces.

ATCOR offers three different methods of correcting BRDF effects: The first method is mainly intended for flat terrain and normalizes the off-nadir reflectance values to the corresponding nadir reflectance values.

The second method is exclusively dedicated to rugged terrain imagery and corrects for BRDF effects due to the variability of the solar incidence angle. The reflectance values of areas with low local solar elevation angles, i.e., large local solar zenith angles, are often overcorrected by the assumption of isotropically reflecting surfaces. The method reduces these high, overcorrected values depending on the illumination and/or viewing angles.

The third method corrects the observation BRDF effects by fitting a physical BRDF model to a number of images and surface cover classes in order to obtain a generic BRDF correction function.
This function is used to calculate a per-pixel anisotropy factor which corrects for the deviation from an averaged spectral albedo.

In some cases of rugged terrain imagery it is useful to apply both, incidence and observation angle correction methods of BRDF correction.

10.6.1 Nadir normalization method

A simple algorithm was implemented as part of the ATCOR package to normalize the scan angle dependent brightness values to the nadir value. It is recommended to apply the method to imagery after atmospheric correction, i.e., to reflectance data. However, if only the across-track illumination gradients shall be removed without any further atmospheric correction, the algorithm can also be applied to radiance (DN) data. In this case, the brightness gradient may be caused by a combination of surface BRDF and atmospheric BRDF (left/right asymmetry in path radiance).

The algorithm is intended for large field-of-view sensors (minimum FOV=20°). It computes the column means with a certain angular sampling interval (1° or 3°). The input image may be geocoded, or not. If it is not geocoded the total field-of-view FOV corresponds to the number n of across-track image pixels per line. If geocoded, the scan angle for each pixel must be provided in a separate file (".sca"). It contains the scan angle in degree scaled with a factor of 100, and coded with 16 bits per pixel. This definition is taken from the airborne ATCOR/PARGE interface (Schläpfer and Richter 2002). Scan angles on the right hand side with respect to flight heading are defined as negative, those on the left side as positive, e.g., a value of -2930 represents a scan angle of 29.3° on the right side.

The nadir region is defined here as the ±3° scan angle range. Usually, a 3° angular sampling interval, from +3° to +FOV/2 on the left side and -3° to -FOV/2 on the right side, is adequate, except for geometries close to the hot spot geometry. In the latter case, a 1° sampling interval can be selected.

If \( \bar{b}_{nadir} \) denotes the averaged brightness value for the nadir region, i.e., reflectance or radiance, then the nadir normalized brightness value of a pixel with column number \( j \) is calculated as:

\[
b_{\text{norm}}(j) = \frac{b(j)}{\bar{b}_{nadir}} f_2(j)
\]

where the function \( f_2 \) is obtained with three processing steps:

- The first step is the averaging over each interval (3° or 1°). It yields a function \( f_1 \) with \( m+1 \) grid points for the \( m \) off-nadir intervals plus the nadir interval.

- Two cases are distinguished now: if the image is not geocoded, an interpolation from function \( f_1(m+1) \) to a function \( f_2(ncols) \) is performed where ncols is the number of column pixels of the image. If the image is geocoded, an interpolation from the 3° grid to the 1° grid is performed (no hot spot case).

- The third step is a filter with a moving average window applied to the \( f_2 \) function. The following cases are distinguished: if the image is not geocoded the window is 9 pixels (without hot spot) and 3 pixels (with hot spot option). If the image is geocoded, the moving window extends over a 5° angular interval (no hot spot) and over a 3° interval (with hot spot option).
Figure 10.25 shows part of a HyMap image (acquired 3 June 1999, Barrax, Spain, 12:09 UTC) containing the hot spot geometry. The solar azimuth was 181° and the sensor scan line azimuth was 179°, almost exactly pointing into the solar azimuth. The left image shows HyMap band 30 at 868 nm after atmospheric correction. The right image is the result after nadir normalization with a 1° sampling interval. In this example, the column means were calculated globally, i.e. surface cover independent. The algorithm also contains an option to compute the column means separately for 4 surface covers. It can currently only be selected if the input imagery is reflectance data and not geocoded. The processing time is much larger than for the global, cover-independent method. The four surface classes are:

- bright vegetation (ratio vegetation index NIR/RED > 10);
- medium/dark vegetation (6 < ratio vegetation index < 10);
- dry vegetation or mixed vegetation/soil (3 < vegetation index < 6);
- soil (vegetation index < 3).

The reflectance of off-nadir water pixels (criterion: near infrared reflectance < 5 %) is not modified.

![Figure 10.25: Nadir normalization of an image with hot-spot geometry.](image)

Left: reflectance image without BRDF correction. Right: after empirical BRDF correction.

### 10.6.2 Empirical incidence BRDF correction in rugged terrain

For many surface covers the reflectance increases with increasing solar zenith and/or viewing angle [45]. Scenes in mountainous regions often exhibit a large variation of terrain slopes, and thus bidirectional brightness variations for a certain surface cover, e.g., meadow or forest. This behavior cannot adequately be eliminated with the Lambertian assumption of equation (10.15). This equation leads to overcorrected reflectance values in faintly illuminated areas (having small values of $\cos\beta$).

**Background**

Several approaches have been pursued to solve this problem in the past:
• an empirical coefficient $C$ is calculated based on a regression of brightness values and the local illumination angle derived from the DEM. The coefficient depends on scene content and wavelength ([95], [58]).

• the sun-canopy-sensor (SCS) geometry is employed in forested terrain instead of the solely terrain-based geometry [29].

• the SCS method is coupled with the C-correction [93].

These approaches produced good results on sample scenes with uniform cover types presented in the above papers. When applying the methods to a wider range of areas, some of the practical problems are:

• mountainous scenes often contain a number of different covers, e.g., deciduous forest, coniferous forest, mixed forest, shrubs, meadow, rocks, etc.

• the computation of the $C$ coefficients for different surface covers would require a pre-classification.

• the correlation obtained for the $C$ coefficients is often less than 0.7, yielding unreliable results with this method.

These remarks are supported by reference [58]. These authors applied different correction approaches to a TM scene containing different cover types and noted that there is no optimum method for all cover types. A drawback of the Minnaert and empirical $C$-methods is that they do not distinguish between the direct and diffuse solar illumination as opposed to the physically based approach of ATCOR. Nevertheless, the latter approach also cannot avoid problems in faintly illuminated areas.

**Correction method**

The methods described in the above section are supplemented by an empirical method with three adjustable parameters ($\beta_T$, $b$, and $g$) as explained below. This approach was tested on different rugged terrain scenes with vegetated and arid landscapes and usually yields satisfactory results. It reduces overcorrected reflectance values starting at a threshold local solar zenith angle $\beta_T$ greater than the scene’s solar zenith angle $\Theta_s$. Equation (10.118) defines the implemented basic geometric correction function which depends on the local solar incidence angle (solar illumination $\beta_i$) and the threshold angle $\beta_T$. The exponent $b$ ( = 1/3, 1/2, 3/4, or 1) is the second parameter and can be selected by the user. Some guidelines on the choice of $b$ are are discussed below. The third adjustable parameter is the lower bound $g$ of the correction function, see Figure 10.26.

$$G = \left\{\frac{\cos\beta_i}{\cos\beta_T}\right\}^b \geq g$$ (10.118)

The threshold illumination angle $\beta_T$ should have some margin to the solar zenith angle to retain the original natural variation of pixels with illumination angles close to the solar zenith angle. The threshold angle can be specified by the user and the following empirical rules are recommended:

• $\beta_T = \theta_s + 20^\circ$ if $\theta_s < 45^\circ$

• If $45 \leq \theta_s \leq 55^\circ$ then $\beta_T = \theta_s + 15^\circ$

• If $\theta_s > 55^\circ$ then $\beta_T = \theta_s + 10^\circ$
These rules are automatically applied if \( \beta_T = 0 \), e.g., during batch processing.

The geometric function \( G \) needs a lower bound \( g \) to prevent a too strong reduction of reflectance values. Values of \( G \) greater than 1 are set to 1, and values less than the boundary \( g \) are reset to \( g \). This means the processing works in the geometric regime from \( \beta_T \) to \( 90^\circ \) and the updated reflectance is:

\[
\rho_g = \rho_L G, \quad (10.119)
\]

where \( \rho_L \) is the isotropic (Lambert) value.

Figure 10.26: Geometric functions for empirical BRDF correction. Left: Functions \( G \) eq. (10.118) for different values of the exponent \( b \). Right: Functions \( G \) of eq. (10.118) for \( b=1 \) and different start values of \( \beta_T \). The lower cut-off value is \( g=0.2 \).

Figure 10.26 shows a graphical presentation of equation (10.118). The left part displays the function \( G \) for different values of the exponent \( b \). For \( b=1 \) the decrease with \( \beta_i \) is strong with a constant gradient. For smaller values of \( b \) the decrease with \( \beta_i \) is moderate initially, but the gradient increases with larger \( \beta_i \). Currently, different functions \( G \) for soil/sand and vegetation can be selected in ATCOR (compare the graphical user interface of Figure 5.47). The function \( G \) for soil / sand is applied with a wavelength - independent exponent \( b \). After testing a large number of vegetated mountainous scenes two vegetation modes were finally selected because of their good performance:

1. \( b=0.75 \) for channels with \( \lambda < 720 \text{ nm} \) and \( b=0.33 \) for \( \lambda > 720 \text{ nm} \) (”weak” correction),
2. \( b=0.75 \) (\( \lambda < 720 \text{ nm} \)) and \( b=1 \) (\( \lambda > 720 \text{ nm} \)), (”strong” correction).

In most of the tested cases, the first mode was appropriate. A simple criterion (vegetation index \( \rho_{8650nm}/\rho_{660nm} > 3 \)) is used to distinguish soil/sand and vegetation.

The right part of Figure 10.26 shows the effect of shifting the threshold illumination angle \( \beta_T \). For larger values of \( \beta_T \) the decline of function \( G \) starts later with a larger gradient, and the lower bound \( g \) is met at slightly higher values of \( \beta_i \). In most cases, \( g=0.2 \) to 0.25 is adequate, in extreme cases of overcorrection \( g=0.1 \) should be applied.
CHAPTER 10. THEORETICAL BACKGROUND

Practical considerations

The angle $\beta_T$ can actually be calculated from the imagery as demonstrated by the following example. When processing the scene with ATCOR the map of local solar zenith angles is stored in a separate file "*ilu". If the output file after atmospheric / topographic correction contains bright overcorrected areas this file should be linked to the "*ilu" file using any available standard image processing software. The "*ilu" file contains the illumination map $\beta_i$ scaled as byte data:

\[ \text{ilu} = 100 \times \cos \beta_i \]
\[ \beta_i = \arccos(\text{ilu}/100) \]  \hspace{1cm} (10.120)
\[ \beta_i = \arccos(\text{ilu}/100) \]  \hspace{1cm} (10.121)

Let us assume an example:

A pixel in a dark area of the "*ilu" image has the value ilu=32, i.e., $\beta_i = 71^{\circ}$. The overcorrected reflectance value be $\rho_L = 80\%$ and this value shall be reduced to 40\%, a value typical for the flat-terrain neighborhood. Then the threshold angle has to be specified such that $\cos\beta_i/\cos\beta_T = 0.5$ (with exponent $b=1$ in equation (10.118), in this case $\beta_T = 50^{\circ}$). So, if the desired reflectance reduction factor is $G$ then the required threshold angle can be calculated from eq. (10.118 with $b=1$):

\[ \beta_T = \arccos\left(\frac{\cos \beta}{G}\right) = \arccos \frac{\text{ilu}}{100G} \]  \hspace{1cm} (10.122)

In many cases a separate treatment of BRDF effects for soil/rock and vegetation provides better results. For this purpose, two modes of BRDF correction are available for vegetation, compare the graphical user interface of Figure5.47. The first mode is superior in most cases.

Reference [71] contains a comparison of different topographic correction methods for several Landsat-TM, ETM+, and SPOT-5 scenes from different areas. The proposed empirical ATCOR approach performed best in most of these cases, but no method ranked first in all cases.

10.6.3 BRDF effect correction (BREFCOR)

A generic BRDF effects correction routine (BREFCOR, [86])) has been included in ATCOR. The idea is to apply a scaling of the volume scattering and the geometric scattering component within a well accepted BRDF model. A fuzzy surface cover index of the complete image image is used for this purpose, which covers all surface types from water to asphalt and concrete, soils, sparse vegetation and dense vegetation.

The Ross-Li-sparse reciprocal BRDF model has been selected as basis for the correction of reflectance anisotropy [33]. This model is mainly developed for vegetation, but we use it in a scaled way for all kind of surfaces. Literature mainly related to MODIS atmospheric correction routines showed the superior performance of this model if compared to others. However, for high spatial resolution instruments also other models may be applicable.

Selected BRDF kernels

The BRDF correction scheme is based on the Ross-Thick Li-Sparse Model (RTLS), potentially enhanced by the Hot-Spot function as proposed by Maignan et al. [50]. For the correction, a formulation of the model for the Bidirectional Reflectance Factor (BRF) is used. The BRF is well suited for correction of the HDRF, as both quantities are defined as 1.0 for a 100% reflecting target
at the same observation geometry, and as only the second dimension (observation direction) relative variation of the BRF is used for the correction. The generic RTLS-equation of the BRF for each pixel and spectral band is given as:

\[
\rho_{\text{BRF}} = \rho_{\text{iso}} + f_{\text{vol}} K_{\text{vol}} + f_{\text{geo}} K_{\text{geo}},
\]

(10.123)

where \(\rho_{\text{iso}}\) is the isotropic reflectance defined at nadir for both illumination and observation angle. The kernel factors \(f_{\text{vol}}\) and \(f_{\text{geo}}\) are weighting coefficients for the respective kernels. They depend on the ground coverage BRDF, whereas the kernels are fixed functions which define a fully bi-directional reflectance property. The kernels have been selected according to the findings of BRDF literature [96]. For the volume scattering, the Ross-Thick kernel is modified to include the hot-spot extension by Maignan, i.e.:

\[
K_{\text{vol}} = \frac{4}{3\pi} \frac{1}{\cos \theta_i + \cos \theta_r} \left[ \left( \frac{\pi}{2} - \zeta \right) \cos \zeta + \sin \zeta \right] - \frac{1}{3},
\]

(10.124)

where \(\zeta = \arccos(\cos \theta_i \cos \theta_r + \sin \theta_i \sin \theta_r \cos \phi)\).

The angle \(\theta_i\) is the incident solar zenith angle, \(\theta_r\) is the observation zenith angle, and \(\phi\) is the relative azimuth angle \(\phi = \phi_i - \phi_r\) (i.e., the difference between incidence and observation azimuth).

The extension of this volumetric kernel by Maignan is given as:

\[
K_{\text{vol}} = \left( K_{\text{vol}} + \frac{1}{3} \right) \left( 1 + \frac{1}{1 + \zeta/1.5} \right),
\]

(10.125)

The reciprocal Li-Sparse kernel is used for the geometric part. It is defined as:

\[
K_{\text{geo}} = \frac{1}{\pi} \left( t - \sin t \cos t \right) \left( \frac{1}{\cos \theta_i} + \frac{1}{\cos \theta_r} \right) - \left( \frac{1}{\cos \theta_i} + \frac{1}{\cos \theta_r} \right) + \frac{1 + \cos \zeta}{2 \cos \theta_i \cos \theta_r},
\]

(10.126)

where

\[
t = \arccos \left( \frac{\sqrt{\tan^2 \theta_i + \tan^2 \theta_r - 2 \tan \theta_i \tan \theta_r \cos \phi + (\tan \theta_i \tan \theta_r \cos \phi)^2}}{1 / \cos \theta_i + 1 / \cos \theta_r} \right).
\]

10.6.4 BRDF cover index

A continuous BRDF cover index (BCI) function is used for characterization of the surface. It is calculated on the HDRF of four standard bands: blue at 460nm, green at 550nm, red at 670 nm, and near infrared at 840nm. This reduced selection of spectral bands makes the index applicable for most current optical remote sensing systems. The BCI function characterizes the image based on intrinsic BRDF properties from strong forward scatterers (water) to neutral targets (asphalt) to backward scatterers (soils and vegetation types). The index implementation is using the normalized difference vegetation index (NDVI) as a first input for vegetation density quantification due to its known relation to the leave area index (LAI), which has a significant influence on the BRDF [48].

The NDVI is increased in Equation (10.127) by a value of up to 0.5 using the fact that dense agricultural vegetation shows higher green reflectance than dense forests; i.e., the NDVI is increased by \(C_{\text{forest}}\) for dense forests having a green reflectance in a range below 7%. In a further step, the BCI is decreased for soils by \(C_{\text{soils}}\) using the effect that soils show a relatively low blue at-sensor radiance. A last adaption \(C_{\text{water}}\) is made for water such that clear water areas are always set to a minimum value:

\[
BCI = (NDVI + C_{\text{forest}} - C_{\text{soils}} - C_{\text{water}}) > -1.2.
\]

(10.127)
Note: the “>”-sign denotes a maximum operator between the left and the right side of the term.

The three correction functions in Equation 10.127 are given as follows, first for forests using the absolute HDRF value in the green $\rho_{green}$:

$$C_{forest} = \frac{0.5}{0.04 \cdot 0.2} \cdot (0.07 - \rho_{green}^{(0.04)}) \cdot (NDVI - 0.55^{(0.20)}).$$ (10.128)

The upper and lower values at the square brackets indicate a truncation at these values. The upper values could be adapted for better representation of biome types. For surface covers having a BCI below 0.1 (i.e., mostly soils), a reduction factor is found from the relation between blue and red HDRF as:

$$C_{soils} = \frac{\rho_{blue}}{\rho_{red}} \left[1 - 10 \cdot (NDVI + C_{forest})^{(1.00)} \right].$$ (10.129)

This factor accounts for the variability of non-vegetated areas in the visible. Finally, a summand to account for water is added, starting with $BCI_{soil} = NDVI + C_{forest} - C_{soils}$. It takes into account the relatively higher reflectance of water in the green spectral band in relation to the blue for discrimination to other surface targets such as shadows and dark asphalt:

$$C_{water} = \left(\frac{\rho_{green}}{2 \rho_{blue}} - 0.8 \right) > 0 \cdot (-3 \cdot (BCI_{soil} + 0.5 < 0)).$$ (10.130)

The range of the final BCI function is defined between values of -1.20 and 1.50. The BCI-index calculated in each image pixel can then be used for BRDF model calibration and subsequently for image correction.

**Model calibration**

For the calibration of the model, the BCI is divided into a number of 4-6 discrete classes. The evaluation has shown that increasing the number of classes often leads to worse fitting results and less stable BRDF correction whereas keeping the number of classes small is more stable. A second outcome was that it is hardly feasible to define generic class limits for any kind of data acquisition and sensor. The calibration follows the scheme shown in Figure 10.27.

Differences in limits can be attributed to the fact that the higher resolution images allows for a more accurate and statistically more relevant calibration of the model whereas for lower resolution, the number of classes should be reduced. The classes can be denominated as: water, artificial materials, soils, sparse vegetation, grassland, and forests. For each of the classes, the optimum kernel weights are calculated and stored for each image of a campaign. All weights are then averaged while bad fitting classes (with relative errors greater than 10%) are excluded from averaging. No BRDF correction is applied for classes without any fitting parameters (i.e. if less than 3 bands out of 4 within the class could be calibrated). The averaged model is stored for later application to the imagery.

**Image correction**

Finally, the derived BRDF model calibration data are to be applied to the image data. For application on the imagery, the BCI has to be calculated from each image and is used to get a continuous correction function.

The image processing procedure is following the below steps (compare Figure 10.28):
• calculate the BCI from image,
• calculate the scene-specific angular kernels subsets,
• interpolate the calibration data from BCI levels to a continuous BRDF model,
• calculate an anisotropy map by scaling the kernels using the BCI, the scan angles (observation zenith and azimuth angle), and the interpolated BRDF model, and
• apply the anisotropy map on a per-pixel basis.

The anisotropy factor is derived as relation of the directional model for each pixel to the same model averaged over all angles (i.e., to a good approximation of the spectral albedo BHR).

\[
ANIF = \frac{\rho_{iso} + f_{geo}K_{geo,\theta_i,\phi} + f_{vol}K_{vol,\theta_i,\phi}}{\rho_{iso} + f_{geo}K_{geo} + f_{vol}K_{vol}} \approx \frac{\rho_{BRF}}{\rho_{BHR}}. \quad (10.131)
\]

The bihemispherical reflectance is described by the two hemispherical averages \(K_{geo}\) and \(K_{vol}\) weighted by the respective factors and added to the constant isotropic reflectance \(\rho_{iso}\). Alternatively, the anisotropy with respect to nadir BRF would be an option as done in earlier BRDF
Figure 10.28: Image correction scheme.

research [80]. This option is currently not supported in BREFCOR, as the BHR is the more generic spectral albedo definition for surface object characterization. The corrected bihemispherical reflectance is finally calculated as
\[ \rho_{BHR} = \frac{\rho_{HDRF}}{\text{ANIF}}, \]
where \( \rho_{HDRF} \) is the bottom of atmosphere (directional) reflectance after standard ATCOR-4 atmospheric compensation, as described above.

Sample result

The BREFCOR correction is most appropriate for wide FOV sensors, i.e., with FOV values larger than approx. 20 degrees, typically found for airborne imagery (compare validation samples in [85]). A sample result of ADS-80 is displayed in Figure 10.29. The upper image is a mosaic of two east-west flown flights, where strong across-track gradients are visible. The middle image is the continuous, surface-cover dependent anisotropy factor used for correction. The lower image is the correction result based on the calibrated Ross-Li-sparse BRDF model. Most BRDF effects can be removed by this method as long as the image statistics allow for a good model calibration. However, shaded areas and forests are still affected by visible differences due to the variable spatial structure if seen from varying angles.

10.7 Summary of atmospheric correction steps

Although the case of a flat terrain could be treated as a special case of a rugged terrain with the same elevation everywhere this is not an efficient solution, because the rugged terrain algorithm runs 3 to 4 times slower than the flat terrain code. Therefore, the coding is done in separate modules as discussed below.
10.7.1 Algorithm for flat terrain

The complete sequence of processing for sensors with water vapor bands and a short wave IR band (1.6 or 2.2 µm region) consists of the following steps:

- masking of haze, cloud, water, and clear pixels
- haze removal
- de-shadowing
- masking of reference pixels

- calculation of visibility or optical thickness for reference pixels. The optical thickness for the remaining pixels can be defined as the average of the reference pixels or a spatial (triangular) interpolation is employed to fill the gaps. Finally, a moving low pass window with a box size
of 1.5\( km \times 1.5\)\( km \) (or the minimum of ncols/2 and nlines/2, ncols=image columns, nlines=lines) is applied to smooth sensor noise and small scale variations of the spectral correlation coefficient for the DDV reference pixels. The resulting visibility index and AOT maps are stored as separate files. The visibility calculation based on the reference pixels has to account for the adjacency effect, because reference areas are embedded in non-reference areas, see the sketch below. Since the weighting fraction of reference to non-reference area within the adjacency range is not known for each pixel, the visibility calculation is performed with an average adjacency weighting factor of 0.5 \( q \):

![Figure 10.30: Weighting of \( q \) function for reference pixels.](image)

\[
L_{\text{adj.cor}} = c_0 + c_1DN + 0.5q(DN - DN_{\text{av.clear}}) \quad (10.132)
\]

\[
L(VIS) = L_p + \tau r_{ref}E_y/\pi = L_{\text{adj.cor}} \quad (10.133)
\]

Next the visibility is converted into the nearest visibility index \( vi \) (range 0-182), compare Fig. 10.14, to store the visibility index map as byte data. Spatial interpolation is performed to fill the gaps for non-reference pixels or the average \( vi \) value can be taken to close the gaps. A moving average window of 1.5 km \( \times \) 1.5 km is employed to reduce the influence of noise. The cloud / building shadow map is stored separately (“fshd.bsq” file, containing the fraction of direct solar irradiance per pixel, scaled with the factor 1000). The scaled value 1000 indicates full solar irradiance, smaller values a corresponding fractional value.

- an update of the path radiance in the blue-to-red spectral region is performed, if required, provided a blue spectral band exists.
- water vapor retrieval using the previously calculated visibility map. If the scene contains no reference areas the user has to specify a constant visibility that enters the water vapor calculation.
- reflectance spectrum retrieval with pixel-based water vapor and visibility map. Iterations for adjacency effect and spherical albedo are included. For the adjacency correction, the reflectance of cloud pixels is replaced with the scene average reflectance to avoid an overcorrection of the adjacency effect.
- temperature / emissivity retrieval if thermal bands exist.
10.7.2 Algorithm for rugged terrain

The algorithm for rugged terrain basically consists of the same processing step as in the flat terrain, but every step has to take into account some or all DEM information:

- During the calculation of the visibility index map the DEM information (elevation, slope, aspect, skyview factor) is taken into account.

- The retrieval of the water vapor map has to include the terrain elevation.

- The empirical BRDF correction is based on the local illumination map (local solar zenith angle) derived from the slope, aspect and shadow channels.

- The retrieval of the spectral reflectance cube consists of the steps
  1. three iterations for terrain reflectance,
  2. empirical BRDF correction depending on illumination map, if enabled,
  3. adjacency correction, including proper treatment of cloud areas,
  4. spherical albedo correction.

- The retrieval of surface temperature and emissivity includes the maps of visibility index, water vapor (if water vapor bands exist), elevation, and scan angle. No slope/aspect correction is performed in the thermal region.

10.8 Accuracy of the method

There is not a single figure that can be given to summarize the accuracy for all situations, because the radiometric accuracy of the method depends on several factors: the calibration accuracy of the sensor, the quality of geometric co-registration of the spectral bands, the algorithm for ortho-rectification relying on auxiliary information such as attitude and GPS/DGPS, the accuracy of the radiative transfer code (MODTRAN 5), the correct choice of atmospheric input parameters, the terrain type (flat or rugged), and the surface cover.

Solar region:

In the solar region (wavelength $<2.5\ \mu m$), assuming a flat terrain, and avoiding the specular and backscattering regions, an accuracy of the retrieved surface reflectance of $\pm 2\%$ (for reflectance $<10\%$) and $\pm 4\%$ reflectance units (for reflectance $>40\%$) can be achieved [66]. For rugged terrain, the most important parameter is an adequate spatial resolution of the DEM or DSM (digital surface model) and the exact ortho-rectification of the imagery. It would be desirable to have a DEM of a quarter of the sensor’s spatial resolution or at least the resolution of the sensor’s footprint which is seldom available [63]. Even in the latter case, errors in the matching of imagery and DEM can lead to large relative reflectance errors exceeding $100\%$ for critical geometries (principal plane, e.g. a mountain ridge with half a pixel offset between imagery and DEM [63]). Thus, the quality of the required DEM will limit the final accuracy of the geo-atmospheric image product in many cases. For a flat terrain and larger off-nadir view angles, BRDF effects may have to be accounted for, and the appropriate surface-cover dependent BRDF model will influence the accuracy.

Thermal region:

In the thermal wavelength region beyond $8\ \mu m$, the surface temperature retrieval additionally depends on the correct choice of the surface emissivity. In the ATCOR model the emissivity in one
thermal band is based on a classification of the reflective bands if the sensor collects co-registered reflective and thermal band data. Depending on the surface cover classification (vegetation, soil, sand, asphalt, water, etc.), a typical emissivity value is assigned to each class [79]. If the deviation of the true surface emissivity to the assumed emissivity is less than 0.02 (a typical error margin), then the temperatures will be accurate to about 1-1.5 K. A rule of thumb is a surface temperature error of about 0.5-0.8 K per 0.01 emissivity error if the surface temperature is much higher than the boundary layer air temperature [90]. An accuracy of 1-2 K can be achieved if the emissivity estimate is better than 2% [14].
Bibliography


References


[46] Levy, R.C. et al., "Algorithm for remote sensing of tropospheric aerosol over dark targets from MODIS: collections 005 and 051, Revision 2; Feb 2009" (2009).


References


Appendix A

Comparison of Solar Irradiance Spectra

The following two plots show the relative differences between two extraterrestrial solar irradiance sources:

- Kurucz 1997 (distributed with MODTRAN, Berk et al. 2008 [8]). The previous high-resolution ("monochromatic") databases of ATCOR were calculated with this spectrum.

- Fontenla 2011 (Fontenla et al., 2009, 2011 [20, 21]). The new ATCOR release uses the improved quiet sun spectrum of Fontenla and co-workers, also referred to as "low activity sun".

As explained in chapters 5.8.6, 5.8.7 the user can convert the database of atmospheric look-up tables from one solar irradiance source to another one, provided that the spectral range and sampling distance agrees with the template spectra in the "sun_irradiance" directory of ATCOR. Currently, irradiance spectra of Kurucz 1997, Kurucz 2005 (distributed with MODTRAN [8]), and Fontenla 2011 are offered.

The plots show the detailed information (line structure) contained in the Fontenla spectrum. The curves with 2.8 nm and 10 nm represent results based on a moving average of the 0.4 nm data over 7 and 25 spectral points, respectively.