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NEW APPROACHES IN AIRBORNE THERMAL IMAGE PROCESSING FOR LANDSCAPE ASSESSMENT

NOVÉ PŘÍSTUPY ZPRACOVÁNÍ LETECKÝCH OBRAZOVÝCH TERMÁLNÍCH DAT K HODNOCENÍ KRAJINY

DIZERTAČNÍ PRÁCE DOCTORAL THESIS

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Abstrakt

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Summary

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Klíčová slova

diaľkový prieskum Zeme, letecké termálne hyperspektrálne dáta, separácia teploty a emisivity

Keywords

remote sensing, airborne thermal hyperspectral data, temperature-emissivity separation

Image Processing for Landscape Assessment	I thesis New Approaches in Airborne Thermal on my own according to advice of my super-D., and using the sources listed in references.
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Contents

1	Theoretical Background on Thermal Radiation	13
2	Airborne Thermal Hyperspectral Data Properties 2.1 Instrument Technical Specifications	
3	Temperature and Emissivity Separation 3.1 Available approaches	25
	3.2 TES Algorithm Improvement	
4	Ground Measurements	33
	4.1 introduction	33
	4.2 Data Description	34
	4.3 Methods	36
	4.4 Data Properties	37
	4.5 Spectral Smoothing Algorithm	37
5	Application to UHI Detection	39
6	Conclusion	41
\mathbf{B}^{i}	ibliography	43

Contents

1

Theoretical Background on Thermal Radiation

This chapter describes fundamental principles and concepts of EM radiation called *thermal radiation*. Every object with temperature above 0 K emits thermal radiation. The amount of thermal radiation as a function of wavelength depends on object's temperature and its surface as is described in following text.

Black body

The concept of black body is very well described in the work of Howell [12], where black body is defined as perfect absorber for all incident radiation. Apart of perfect absorber, the black body is perfect emitter as well. Thus black body absorbs and reemits all energy incident upon it. Black body does not exist in nature but its concept is used for determination of real object's surface property called emissivity, which will be defined in following text.

Planck's law

Concerning black body at thermal equilibrium, the amount and spectral distribution of emitted energy is described by Planck's law [19]:

$$B(T,\lambda) = \frac{2hc^2}{\lambda^5} \frac{1}{e^{\frac{hc}{\lambda kT}} - 1},$$

where $B(T,\lambda)$ is spectral radiance (W m⁻² µm⁻¹ sr⁻¹) of black body at temperature T (K) and wavelength λ (µm); k is Boltzmann constant (1.3806488 · 10⁻²³ J K⁻¹), h is Planck constant (6.62606957 · 10⁻³⁴ J s) and c is speed of light (299792458 m s⁻¹). Example of the black body radiation at three different temperatures, as described by Planck's law, is depicted in figure 1.1a.

Emissivity

The emissivity is defined as ratio of radiance of real surface to that of black body at the same temperature:

$$\varepsilon(T,\lambda) = \frac{L(T,\lambda)}{B(T,\lambda)},$$

where $\varepsilon(T,\lambda)$ is spectral emissivity and $L(T,\lambda)$ is real surface spectral radiance. The emissivity can be understood as real surface emission effectiveness in comparison with radiation emitted by a black body of the same temperature in the same wavelength. Let us note that emissivity depends on the viewing angle apart temperature and wavelength, as is defined in Hollow [12]. In remote sensing an observed objects are of the temperature within 270 – 330 K and the observation angle is close to nadir (usually maximum off-nadir angle is less than 30°), which causes negligible changes in spectral emissivity of most of the natural surfaces. Thus, it can be further assumed that emissivity depends just on wavelength.

Quartz was chosen to demonstrate the principles of radiation of real object's surfaces. Its spectral emissivity was taken from ASTER spectral library [1] and it is shown in the figure 1.1b. Quartz heated to the temperature T has spectral radiance $L(T, \lambda) = \varepsilon(\lambda)B(T, \lambda)$ as is illustrated in figure 1.1c.

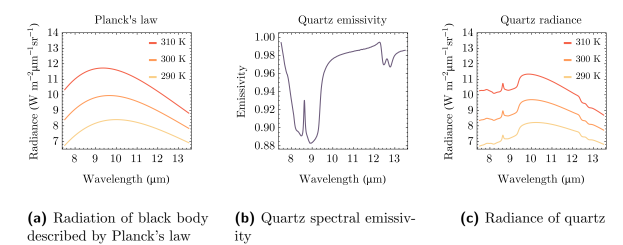


Figure 1.1: Principles of radiation of real surfaces.

Wien's displacement law

The peak of black body radiation at wavelength λ_{max} is described by Wien's displacement law [12]:

$$\lambda_{max} = \frac{b}{T},$$

where b is Wien's displacement constant (2.8977721 mK). As was mentioned before, the temperature of the most of the natural and artificial surfaces observed by airborne remote sensing ranges in 270 – 330 K. According to the Wien's displacement law, the peak of emitted radiation varies roughly from 8.8 µm to 10.7 µm. This range is in coincidence with the atmospheric window situated between 8 µm to 13 µm. The atmospheric transmittance in this atmospheric window is very high and thus it is relevant for acquisition of remotely sensed thermal data.

Kirchhoff's law of thermal radiation

Emitting and absorbing properties of an object at local thermodynamic equilibrium surrounded by isothermal environment are tied up by Kirchhoff's law of thermal radiation [14]. It states that object's surface absorptivity $\alpha(\lambda)$ at a given wavelength equals to object surface emissivity $\varepsilon(\lambda)$ at the same wavelength:

$$\alpha(\lambda) = \varepsilon(\lambda).$$

Energy conservation implies that energy incident to the object surface can be reflected, transmitted or absorbed. Considering the fractions of incident energy the following equation holds:

$$1 = \rho(\lambda) + \tau(\lambda) + \alpha(\lambda),$$

where $\rho(\lambda)$ is object surface spectral reflectivity, $\tau(\lambda)$ is object surface spectral transmissivity and $\alpha(\lambda)$ is object surface spectral absorptivity. Applying Kirchhoff's law to opaque material $(\tau(\lambda) = 0)$ results in following equation:

$$1 = \rho(\lambda) + \varepsilon(\lambda) \implies \rho(\lambda) = 1 - \varepsilon(\lambda).$$

All mentioned principles in this section will be further used in explanation of properties of airborne thermal hyperspectral data and its processing.

Airborne Thermal Hyperspectral Data Properties

This chapter provides insights into technical parameters of Thermal Airborne Spectrographic Imager (TASI) and processing chain of image data acquired by this sensor. Knowledge of the instrument parameters and processing chain gives important overview of the image data properties and their components. The result of the processing chain described in this chapter is georeferenced image data containing land-leaving radiance. Such an image data form input for further processing. Let us note that this chapter omits naming physical quantities dependent on wavelength as "spectral" for the sake of clarity. However, all quantities remain wavelength dependent.

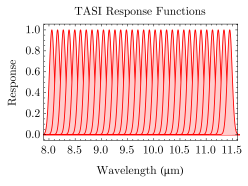
2.1. Instrument Technical Specifications

The TASI sensor is developed by Itres Ltd. (Calgary, Canada) and is one of the very few commercially available pushbroom hyperspectral TIR sensors equipped with mercury cadmium telluride array. Each of its 600 across-track imaging pixels contains 32 bands all of which are in the TIR region. Bands are situated in the 8 to 11.5 µm region and have a FWHM ≈ 0.11 µm with NE Δ T ≈ 0.1 K. The response functions of the TASI sensor are described by the Gaussian functions as depicted in Figure 2.6a.

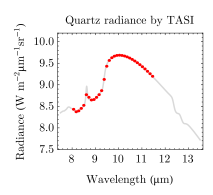
The shape of response functions implies that any quantity observed by TASI sensor is of finite spectral-bandwidth. Quantities needs to be transformed to band-effective quantities in order to relate them with certain wavelength. The band-effective quantities are obtained by using weighted average:

$$X_{i} = \frac{\int_{\lambda_{1}}^{\lambda_{2}} r_{i}(\lambda) X(\lambda) d\lambda}{\int_{\lambda_{1}}^{\lambda_{2}} r_{i}(\lambda) d\lambda},$$
(2.1)

where $r_i(\lambda)$ is response function of band i, λ_1 and λ_2 are lower and upper boundaries of band i and X can be substituted by any quantity. In Figure 2.6b is illustrated radiance of quartz (solid line) and band-effective values of radiance measured by the TASI sensor (red dots). Sensor of this type is available at Global Change Research Institute CAS (Brno, Czech Republic) and it is a part of Flying Laboratory of Imaging Systems (FLIS) [11].







(b) Radiance of quartz at 300 K measured by TASI sensor

Figure 2.1: TASI response functions.

2.2. Image data pre-processing

The main objective of image data pre-processing is transformation of acquired raw image data into the georeferenced radiance at-surface level. It is accomplished by three major steps: radiometric calibration, atmospheric corrections and geometric pre-processing. Radiometric calibration converts digital numbers (DN) into values of radiance at-sensor level. Atmospheric corrections compensates the influence of intervening atmosphere and produces land-leaving radiance. Finally, the geometric pre-processing compensates for image data distortions caused by aircraft movement and register image data into coordinate system.

Supportive field measurements of thermal radiance, temperature, emissivity and atmospheric parameters offers valuable data for calibration and validation purposes. Especially in cases of airborne image data for scientific purposes the high quality is strongly demanding. Thus it is necessary to perform supportive measurements in order to achieve precise results and determine the data quality.

It is important to emphasize, that currently does not exist any definitive standard preprocessing chain. It is caused mainly by huge number of sensors with various technical parameters and their different applications. Sensors usually have tailored pre-processing chains, which is the case of TASI sensor as well. Certain parts of processing chain are maintained by commercial tools. However, there are still parts of processing chain, which needs to be done by in-house tools. In Figure 2.2 is illustrated processing chain used in Global Change Research Institute CAS (Brno, Czech Republic) to pre-process image data acquired by TASI sensor. Individual parts of the diagram will be discussed in the following text.

2.2.1. Radiometric Calibration

Thermal radiation incident upon the sensor array originates from many additive components (e.g. observed scene, instrument enclosure, intervening atmosphere and others). Incident thermal radiation produces electrical signal, which is proportional to radiant intensity. Electrical signal is then amplified and converted into voltage and subsequently into DN values. Radiometric calibration consists of separating signal from viewed scene

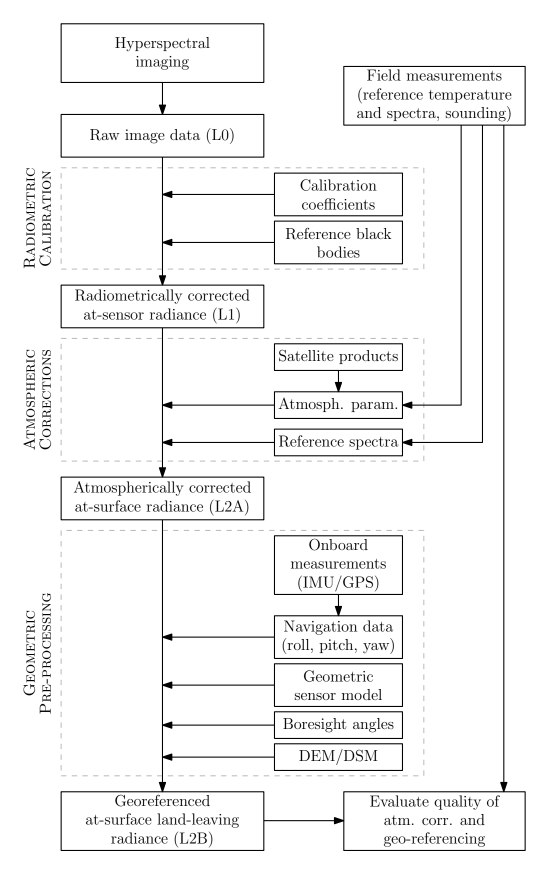


Figure 2.2: Processing chain applied to image data acquired by TASI sensor.

and converting it into physical units of radiance. Atmosphere influence is not accounted in this process and thus after radiometric calibration one gets radiance at-sensor level.

The relationship between DN and at-sensor radiance $L_{\rm m}$ is following:

$$DN = a + bL_{\rm m},$$

where a and b are calibration coefficients. The calibration coefficient a, also known as offset, represents radiation originating from instrument enclosure, sensor dark current and electronic offset. The calibration coefficient b, also called gain, determines sensor radiant sensitivity. Calibration coefficients are determined by imaging a set of reference black bodies of known temperature and emissivity. In this context, the term black body is ment to be a surface with emissivity very close to unity. These coefficients are usually determined applying one of two methods: 1) imaging two black bodies at different temperature directly before imaging, or 2) combining black body image data from laboratory and black body image data acquired before imaging.

In the first case are usually used two black bodies of different temperatures. Temperatures of these black bodies enclose temperatures expected to occur in the scene. Let us consider the radiance of cold black body $L(T_{\rm C})$ and the radiance of hot blackbody $L(T_{\rm H})$. The calibration coefficients can be obtained from:

$$a = \frac{DN_{H}L(T_{C}) - DN_{C}L(T_{H})}{L(T_{C}) - L(T_{H})}$$

$$b = \frac{DN_{C} - DN_{H}}{L(T_{C}) - L(T_{H})},$$

where DN_C and DN_H are digital numbers measured by sensor viewing cold black body and hot black body respectively. This procedure is commonly used in case of other instruments for measuring thermal radiation, such as $\mu FTIR$??.

The determination of calibration coefficients in the second case assumes that gain calibration coefficient b does not change under different conditions. Thus, it is sufficient to perform series of black body measurements at different temperatures in order to determine gain calibration coefficient b. These measurements can be performed in the laboratory once per season. However, offset calibration coefficient a does not remain stable and changes under different conditions. Hence, it is necessary to image a black body at known temperature directly before acquisition to account for variability of this coefficient.

Again, it is important to emphasize that all quantities and both calibration coefficients are wavelength dependent. Spectral calibrations are part of the radiometric calibrations. In the laboratory are determined band centers of every pixel using laser at different wavelengths. Determined positions does not change over time significantly. However, the spectral shift occurs under different conditions and thus it needs to be determined for every scene. Spectral shift estimation is usually based on the spectral features of the atmosphere or certain materials.

In case of TASI sensor are used commercial softwares delivered by Itres company (Calgary, Canada). SparCal software [56] is used to determine all parameters necessary for radiometric calibrations from laboratory measurements. RCX software [57] is used for additional estimation of calibration parameters and for processing raw image data. Both softwares are tailored for the TASI sensor. The resulting image data are made of radiance at-sensor level $L_{\rm m}$.

2.2.2. Atmospheric Corrections

Radiometric calibrations deliver image data containing radiation from the surface, attenuated by atmosphere, plus radiation from the atmosphere along the line of sight. Thus the measured radiance at-sensor level $(L_{\rm m})$ consists mainly of radiance emitted from the land surface, downwelling atmospheric radiance reflected by the surface $(L_{\rm atm}^{\downarrow})$ and the atmospheric upwelling radiance $(L_{\rm atm}^{\uparrow})$. The sum of all these components is expressed by a radiative transfer equation (RTE) as follows:

$$L_{\rm m} = \tau \varepsilon B(T_{\rm s}) + \tau (1 - \varepsilon) L_{\rm atm}^{\downarrow} + L_{\rm atm}^{\uparrow}, \tag{2.2}$$

where $B(T_s)$ is radiance of the surface at temperature T_s according to the Planck's law, ε is the surface's emissivity and τ is atmospheric transmittance. It is important to emphasize that all elements in the equation are wavelength dependent but notation for this is omitted for the sake of clarity. Since sensors are of finite bandwidth, quantities in (2.2) are replaced by band-effective equivalents according to the equation (2.1). Moreover, RTE can be used under the assumption of cloud-free atmosphere under local thermodynamic equilibrium. The meaning of the RTE is illustrated in the Figure 2.3, where ρ is reflectivity. Kirchhoff's law of thermal radiation implies that reflectivity ρ can be rewritten as $(1 - \varepsilon)$ for opaque materials.

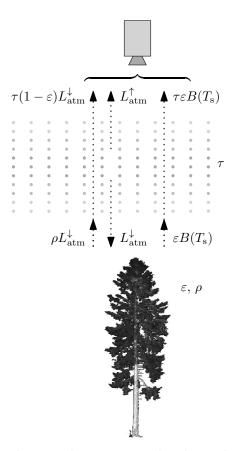


Figure 2.3: The radiance incident to the sensor in the thermal region originates mainly from three sources: 1) radiance $\tau \varepsilon B(T_s)$ emitted by object; 2) reflected downwelling atmospheric radiance $\tau(1-\varepsilon)L_{\text{atm}}^{\downarrow}$; 3) upwelling atmospheric radiance L_{atm}^{\uparrow} emitted by atmosphere itself.

The goal of the atmospheric corrections is to determine atmospheric transmittance, downwelling and upwelling atmospheric radiance and compensate for them. The quantifi-

cation of these quantities are usually based on radiative transfer models of the atmosphere. For this purpose is usually used MODerate resolution atmospheric TRANsmission (MOD-TRAN) model [3]. MODTRAN simulates atmospheric parameters such as atmospheric transmittance, downwelling and upwelling atmospheric radiance based on input parameters such as vertical profiles of water vapour content and temperature, CO₂ concentration, the choice of model atmosphere (if measured profiles are not available) and many others. In general, input parameters can be obtained in two ways: 1) by in-situ measurements; 2) by satellite-based products.

The most common in-situ measurement is radio sounding. Radiosonde is launched during the overflight and it is used to measure vertical temperature and water vapour profile of the atmosphere. Other in-situ instruments can be used as well, for example sun-photometer for obtaining water vapor content or different radiometers for measuring sky or surface radiance. Other source of water vapour and temperature profile is satellite-based products acquired close to the time of aircraft overflight. The most common is MOD07_L2 product [4] generated by Moderate Resolution Imaging Spectroradiometer (MODIS) instrument. Illustration of the transmittance, downwelling and upwelling atmospheric radiance generated by MODTRAN using MOD07_L2 products as input are depicted in Figure 2.4.

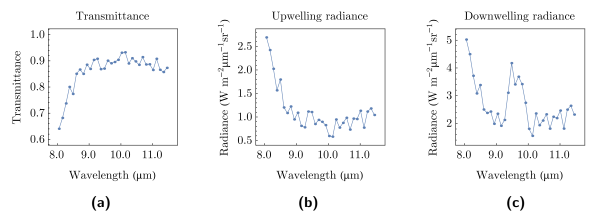


Figure 2.4: Example of atmospheric parameters used for atmospheric correction of TASI image data acquired at altitude of 2000 m during summer season.

In case of thermal hyperspectral images, various algorithms for estimating atmospheric effects based just on the image data itself were developed. Usually it is applied to one of the following: In-Scene Atmospheric Corrections (ISAC) introduced by Young et al. [28] and Autonomous Atmospheric Compensation (AAC) introduced by Gu et al. [10]. The advantage of using one of these algorithms is that no supporting data are necessary. The drawback of these methods remains in estimation of just atmospheric transmittance and upwelling atmospheric radiance.

Once all the atmospheric parameters are determined, it remains to compensate for them. Compensating for atmospheric transmittance and upwelling atmospheric radiance lead to land-leaving radiance:

$$L_{\rm LL} = \varepsilon B(T_{\rm s}) + (1 - \varepsilon) L_{\rm atm}^{\downarrow}. \tag{2.3}$$

The downwelling atmospheric radiance is not possible to separate without knowledge of emissivity. Hence, image data after atmospheric corrections are made of land-leaving radi-

ance $L_{\rm LL}$. Compensation for downwelling atmospheric radiance is part of the temperature and emissivity separation described in Chapter 3.

Atmospheric corrections for TASI sensor, as part of processing chain, are not performed by commercial products. However, there exists commercial tools that allows complex solution for atmospheric corrections. An example of such a tool is ATCOR [21] which is based on look-up tables generated by MODTRAN and takes into account terrain topography and sensor parameters. It offers basic temperature and emissivity separation algorithms as well. Apart of mentioned solution, atmospheric corrections rely on extracting data from in-situ measurements or satellite products, running radiative transfer models and applying derived atmospheric parameter on image data. Alternatively, algorithms for atmospheric parameters estimation from image data can be implemented. In both cases, atmospheric corrections involve creating in-house tools.

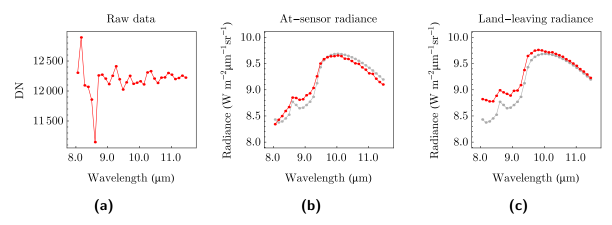


Figure 2.5: Example of data at various processing stages. Data simulates quartz at 300 K as would be acquired by TASI sensor at altitude of 2000 m under summer mid-latitude atmosphere. Case (a) shows raw data, case (b) shows data after radiometric calibration (in red) and case (c) shows data after corrections of atmospheric transmissivity and upwelling atmospheric radiance (in red). In cases (b) and (c) are shown pure quartz radiance in gray.

2.2.3. Geometric Pre-processing

Acquired image data are distorted during its acquisition and geometric pre-processing is accounting for all factors causing these distortions. During geometric pre-processing are taken into account aircraft motion, terrain variations and geometric sensor model in order to register image data into reference frame.

Ancillary data about aircraft position and movement, terrain structure and geometric sensor model are necessary. Aircraft needs to be equipped with IMU/GNSS systems for recording aircraft position (longitude, latitude and altitude) and orientation (roll, pitch and heading angles). Terrain structure is obtained form Digital Surface Model (DSM) or Digital Elevation Model (DEM). These data are derived from aerial laser scanning or from stereo images. Aerial laser scanning can be performed either simultaneously with image data acquisition or separately. Other sources of DEM/DSM are national services or satellite products (e.g. ASTER product AST14DEM). Geometric sensor model is usually delivered by sensor manufacturer.

The process applied on image data during geometric pre-processing is called geoothoreferencing. It consists of two successive steps: direct image data geocoding and resampling. Direct image data geocoding consists of geometric corrections and orthogonalization of the image data. These are further resampled into regular grid of the reference frame with the desired coordinate system (e.g. Universal Transverse Mercator coordinate system). Image data are resampled into desired spatial resolution applying nearest neighbor, bilinear or cubic interpolation. For the scientific purposes is commonly used nearest neighbor interpolation since it preserves spectral information and does not combine spectra from surrounding pixels.

Geometric pre-processing of the image data acquired by TASI sensor are performed by GeoCor software [58] delivered by Itres company (Calgary, Canada). Importance of image data geometric pre-processing is illustrated in the Figure 2.6.





(a) Distorted image data

(b) Georeferenced image data

Figure 2.6: Illustration of land-leaving radiance image data before and after geometric preprocessing.

Temperature and Emissivity Separation

Pre-processed thermal image data provide valuable information about the properties of the observed surfaces, most importantly their temperature and emissivity. But in order to determine temperature and emissivity from observed radiance one must solve system of RTEs. Data from multispectral and hyperspectral TIR sensors offer the opportunity to derive both the temperature as well as an emissivity spectrum, which can be used to characterize the material composition of surfaces. However, observing radiance in Nbands yields N RTEs but N+1 unknowns (N emissivities plus temperature), which results in the underdetermined system of equations. The estimation of temperature and emissivity from such a system of equations is usually addressed as temperature-emissivity separation. This chapter describes several approaches for separating temperature and emissivity. It firstly introduces few commonly used ones and then focuses on the most popular approach called Temperature and emissivity separation algorithm (TES). Last part of this chapter focuses on enhancing the accuracy and precision of the products generated by the TES algorithm. The main improvement is accomplished by replacing one of the TES modules with a newly designed one that takes advantage of a relationship between brightness temperature and emissivity. The improved TES algorithm is called Optimized Smoothing for Temperature Emissivity Separation (OSTES).

3.1. Available approaches

Many approaches have been developed to overcome the problem of having an underdetermined system of RTEs [15]. Methods used to overcome the problem of underdetermined system of RTEs are usually based on adding empirical or semiempricial constraints.

Algorithms for temperature and emissivity estimation depend on sensor architecture and acquisition context. Some algorithms require knowledge of Normalized Difference Vegetation Index (NDVI) [?sobrino-towards-2000?], surface type [?snyder-classification-1998?] or even emissivity [?jimenez-munoz-revisions-2009?]. Others are based on multitemporal [?wan-new-2000?] or multiangle [?soria-derived-2007?] acquisition. There are only a small number of algorithms for simultaneous temperature and emissivity retrievals from a single scene without ancillary surface information, that are robust enough to be applicable to data acquired by multispectral or hyperspectral sensors. The most common are: grey body emissivity method [2], linear emissivity constraint temperature emissivity separation method [27], spectral smoothing [?B07?] and the TES algorithm [9]. Principles of last four mentioned methods are described in the following text. The most attention is

paid to the TES algorithm, as it is the most popular and it is widely applied to many sensors.

Gray body emissivity method

Barducci and Pippi [2] proposed algorithm, which is based on assumption of flat spectral emissivity beyond 10 µm. To solve the system of RTEs it is enough to find at least two spectral bands with the same emissivity. This can be achieved in case of airborne thermal hyperspectral data. Drawback of this method is its sensitivity to instrument noise.

Linear emissivity constraint temperature emissivity separation method

As Wang et al. [27] describe, this method is based on idea of substituting spectral emissivity with piecewise linear function. The emissivity spectrum is divided into segments, in which spectral emissivities are assumed to be linearly dependent on wavelength. Thus, it is necessary for every segment to estimate gain and offset. It implies that the number of spectral bands has to be equal or greater than number of unknowns resulting from segmentation to piecewise linear functions.

Spectral smoothing

Spectral smoothing algorithm, also known as ARTEMISS (Automatic Retrieval of Temperature and EMIssivity using Spectral Smoothness), was reported by Borel at [5] and [6]. Algorithm is based on the assumption that spectra of solids are much more smoother than spectra of gases. Thus by smoothing spectra one removes spectral features introduced by atmosphere and obtains spectral emissivity. Moreover, current implementation described in [6] includes modified ISAC algorithm called ARTISAC, which estimates atmospheric transmissivity for further choice correct atmospheric model. Atmospheric models contains so called TUD (atmospheric Transmissivity, Upwelling and Downwelling atmospheric radiance) and are stored in look-up tables (LUT). Then temperature is varied until the spectral emissivity is the smoothest possible, where the smoothness criterion is standard deviation of measured radiance minus simulated radiance. Spectral smoothness method can be described briefly by following steps:

- 1. estimation atmospheric transmissivity using ARTISAC algorithm
- 2. determination few closest atmosphere models from TUD-LUT according to the estimated atmospheric transmissivity
- 3. use these atmosphere models as input to spectral smoothness algorithm for a few pixels chosen from the image and the atmosphere model which results in smoothest emissivity in the most of the cases is chosen as the correct one
- 4. use chosen atmosphere model for the whole image and estimate temperature and emissivity by applying spectral smoothing procedure

Temperature and emissivity separation algorithm (TES)

TES algorithm was originally developed for the Advanced Spaceborne Thermal Emission and Reflection Radiometer (ASTER) sensor [9]. ASTER was launched in December 1999 onboard NASA's Terra platform. TES has since then found widespread use with other multispectral and hyperspectral sensors.

Several studies have discussed the implementation of TES with Airborne Hyperspectral Scanner (AHS) data [24, 38]. Application of TES to data acquired by the TASI sensor is mentioned in a few studies as well [26, 18]. Apart from the mentioned sensors, the TES algorithm has been modified for the Digital Airborne Imaging Spectrometer (DAIS) sensor [39]. Concerning spaceborne sensors, the TES algorithm has also been suggested for Spinning Enhanced Visible and Infrared Imager (SEVIRI) [40], Moderate Resolution Imaging Spectroradiometer (MODIS) [41] and Multispectral Thermal Imager (MTI) [42] data processing. Moreover, the TES algorithm is being suggested for the future HyspIRI mission [?hulley-hyspiri-2011?].

The TES algorithm is based on a semi-empirical relationship between spectral contrast (i.e. difference between the highest and lowest values in the emissivity spectrum) and the minimum emissivity. The algorithm consists of three modules, namely the Normalization Emissivity Module (NEM) [7], the Ratio module and the Maximum-Minimum Difference (MMD) module [16]. The inputs of the algorithm are land-leaving radiance $L_{\rm LL}$ and downwelling radiance $L_{\rm atm}^{\downarrow}$. Let us remind the reader that land-leaving radiance is obtained from (2.2) by compensating for atmospheric transmissivity τ and atmospheric upwelling radiance $L_{\rm atm}^{\uparrow}$:

$$L_{\rm LL} = \varepsilon B(T_{\rm s}) + (1 - \varepsilon) L_{\rm atm}^{\downarrow}. \tag{3.1}$$

The NEM module performs an iterative process for estimating temperature and emissivity, and compensating for the downwelling radiance. The output of the NEM module is an initial estimation of temperature and emissivity. Then the ratio module normalizes the emissivities obtained by the NEM module by their arithmetic mean. Thus one obtains the so called β spectrum, which should be less sensitive to sensor noise. Finally, the maximum and minimum of the β spectrum are found and their difference (MMD) is used in following semi-empirical relationship:

$$\varepsilon_{\min} = 0.994 - 0.687 \times \text{MMD}^{0.737}.$$
 (3.2)

Derivation of (3.2) is explained in following paragraph. Ratioing the β spectrum back to an emissivity spectrum with knowledge of minimum emissivity results in a more precise estimation of the emissivity spectrum. The band with highest emissivity is then used for temperature estimation.

The relationship between spectral contrast and minimum emissivity, shown in (3.2), is a regression based on 86 laboratory spectra of rocks, soils, vegetation, snow and water chosen from the ASTER spectral library [1]. This relationship is depicted in the Figure 3.1. It is important to note that (3.2) is tailored for the ASTER sensor. To apply TES to a different sensor, the regression of ε_{\min} on MMD must be refined by using sensor specific response functions.

After ASTER was launched, [45] and [22] suggested to replace the power regression shown in (3.2) with linear regression. The replacement is connected with modification of the threshold for separating materials with low spectral contrast. The main advantage

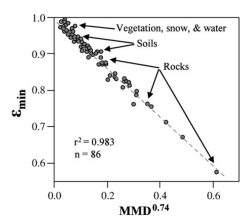


Figure 3.1: Semi-empirical relationship between emissivity contrast and minimum spectral emissivity as shown in study reported by Sabol et al. [22].

is elimination of artefacts in retrievals. However, the drawback is loss of accuracy in cases of materials with low spectral contrast [22]. The TES algorithm used for generation ASTER standard products [?bjorn-personal-communication?], as well as its modifications for other sensors [24, 38, 26, 39, 40, 41, 42, ?hulley-hyspiri-2011?], is based on the power law regression. Thus, in this work the TES algorithm is considered to be that using the power regression.

3.2. TES Algorithm Improvement

he algorithm described below brings a new approach for the TES algorithm by replacing the NEM module with a completely new module. The new module is based on the similarity between brightness temperature spectral features and emissivity spectral features. Brightness temperature is obtained from land-leaving radiance under the assumption of emissivity $\varepsilon = 1$ for every wavelength. Although land-leaving radiance includes some portion of reflected downwelling radiance, it still retains the spectral features arising from the emissivity of the surface materials, which is 0.6 or higher for natural materials [9]. Since the magnitude of downwelling radiance is usually much lower than the surface radiance the features contained in the brightness temperature spectra may be distorted but will not be completely hidden. The new module approximates this relation between brightness temperature $T_{\rm b}$ and emissivity.

In order to demonstrate the relationship, three emissivity samples with different spectral contrasts were chosen from the ASTER spectral library, namely green grass, fine sandy loam and altered volcanic tuff. Emissivities are depicted in Fig. 3.2 (solid lines) together with corresponding band-effective values for TASI sensor (empty symbols). These emissivities were applied to Plank's law at temperature 300 K and combined with downwelling radiance from standard mid-latitude summer atmosphere generated by MODerate resolution atmospheric TRANsmission (MODTRAN) [3]. The resulting radiances, were transformed to band-effective quantities with respect to the ASTER, AHS and TASI response functions. Brightness temperatures for every band of each sensor were obtained by applying inverse Planck's law on sample land-leaving radiances under the assumption of $\varepsilon = 1$. Fig. 3.2 also includes brightness temperatures (full symbols) in order to demonstrate spectral similarity with emissivity. Fig. 3.3 plots emissivity against bright-

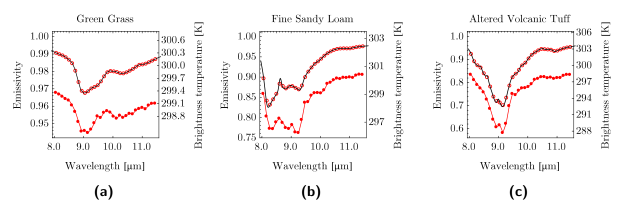


Figure 3.2: Emissivity spectra (black solid line) of three samples chosen from ASTER spectral library [1]. Symbols represent band-effective values of emissivity (empty symbols) and brightness temperature (full symbols) for ASTER (orange triangles), AHS (blue squares), and TASI (green dots) sensor.

ness temperature for chosen samples (empty symbols). These quantities clearly exhibit relationship with linear trend regardless of spectral contrast. Also displayed in Fig. 3.3 are lines that approximate this relationship, derived in the manner described later in the next. The algorithm description below uses band-effective values of quantities linked to i-th band by subscript index i.

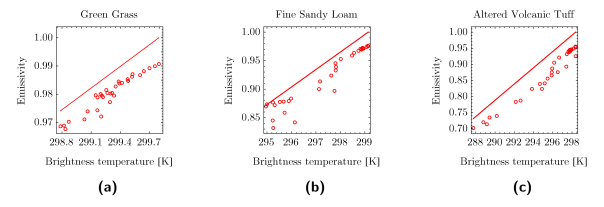


Figure 3.3: Symbols represents examples of the relationship between brightness temperature $T_{\rm b}$ and emissivity as would be observed by ASTER (orange triangles), AHS (blue squares), and TASI (green circles). Lines illustrate the approximations of the relationship between brightness temperature and emissivity for ASTER (orange dotted line), AHS (blue dashed line), and TASI (green full line) sensor. The procedure used for estimation of the brightness temperature and emissivity relationship is described in text.

Spectral features of brightness temperature will be further used for emissivity estimation. The dependence of emissivity ε_i on brightness temperature $T_{\mathbf{b}_i}$ will be approximated by following equation:

$$\varepsilon_i = pT_{\mathbf{b}_i} + q,\tag{3.3}$$

where p and q are empirical coefficients. These coefficients are determined by solving the system of two equations using two points, namely maximum brightness temperature

```
 \begin{split} & \textbf{Function SmoothingErr}(\varepsilon_{\min}, L_{\text{LL}}, L^{\downarrow}) \\ & 1. \quad T_{\text{b}_i} = B^{-1}(L_{\text{LL}_i}) \\ & 2. \quad \text{Find } p \text{ and } q \text{ by solving} \\ & 1 = p \max(T_{\text{b}_i}) + q \\ & \varepsilon_{\min} = p \min(T_{\text{b}_i}) + q \\ & 3. \quad \text{Estimate emissivity} \\ & \varepsilon_i = p \, T_{\text{b}_i} + q \\ & 4. \quad \text{Estimate spectrum} \\ & \quad L'_i = \frac{L_{\text{LL}_i} - (1 - \varepsilon_i) L_i^{\downarrow}}{\varepsilon_i} \\ & 5. \quad T_{\max} = \max(B^{-1}(L'_i)) \\ & 6. \quad \textbf{return} \\ & \quad \sum_i \left| \frac{B_i(T_{\max})}{||B(T_{\max})||_1} - \frac{L'_i}{||L'||_1} \right| \end{aligned}
```

Figure 3.4: Pseudocode of the function that is being minimized in order to estimate the value of ε_{\min} .

coupled with emissivity equal to 1 and minimum brightness temperature coupled with lowest emissivity ε_{\min} :

$$1 = p \max(T_{b_i}) + q,$$

$$\varepsilon_{\min} = p \min(T_{b_i}) + q.$$
(3.4)

The next step is estimation of the lowest emissivity ε_{\min} .

This is done by varying ε_{\min} over the range of possible emissivities for natural materials [0.6, 1], determining corresponding coefficients p and q by solving (3.4) and then approximating emissivity by (3.3) using brightness temperature for all spectral bands. The estimated emissivity is then used together with land-leaving radiance L_{LL} and downwelling radiance L^{\downarrow} in a computation that yields spectral radiance:

$$L_i' = \frac{L_{\mathrm{LL}_i} - (1 - \varepsilon_i)L_i^{\downarrow}}{\varepsilon_i}.$$
 (3.5)

The temperature in every spectral band is derived from spectral radiance L' applying inverse Plank's law. The highest one is chosen as the reference temperature T_{max} . Finally, the estimated spectral radiance L' and Planck's law at the reference temperature T_{max} are normalized and compared against each other as follows:

$$\sum_{i} \left| \frac{B_i(T_{\text{max}})}{||B(T_{\text{max}})||_1} - \frac{L_i'}{||L'||_1} \right|. \tag{3.6}$$

The value of ε_{\min} is considered final if its corresponding spectral radiance L' is the best fit to Plank's law.

The whole process of determining ε_{\min} can be understood as smoothing the spectrum by finding the optimal value of ε_{\min} . Pseudocode depicted in Fig. 3.4 summarizes the above described procedure as a function SMOOTHINGERR($\varepsilon_{\min}, L_{LL}, L^{\downarrow}$) evaluating the error between Planck's law and estimated spectral radiance. This function is minimized with respect to the variable ε_{\min} as follows:

$$\underset{\varepsilon_{\min} \in [0.6,1]}{\operatorname{arg\,min}} \operatorname{SMOOTHINGERR}(\varepsilon_{\min}, L_{\mathrm{LL}}, L^{\downarrow}). \tag{3.7}$$

Continuous curves in Fig. 3.3 show the optimal brightness temperature and emissivity relationship approximation. Let us emphasize that by applying emissivities obtained

from the approximated relationship between brightness temperature and emissivity to (3.5), one gets L' as the best fit to Planck's law. This means that $B^{-1}(L'_i)$ produces a temperature value for each band. These temperatures have minimum variability since they are derived from the best fit to Planck's law. Let us also remind the reader that maximum brightness temperature is coupled with emissivity equal to 1, which implies that it is part of the set of temperatures with smallest variability. It is important to note that maximum brightness temperature T_b computed from land-leaving radiance is usually smaller than surface temperature T_b computed from surface radiance. Land-leaving radiance is smaller than surface radiance since natural materials are of emissivity higher than 0.6 and the contribution from reflected downwelling radiance is usually much lower than surface radiance. By reason of maximum brightness temperature T_b being smaller than surface temperature T_b and by being part of the set of temperatures with smallest variability, it can be concluded that maximum temperature from the set of temperatures tends to be the closest to the surface temperature T_b and is therefore taken as the reference one.

Before passing emissivity to the Ratio and MMD modules, it is recomputed according to (3.8):

$$\varepsilon_i = \frac{L_{\mathrm{LL}_i} - L_i^{\downarrow}}{B_i(T) - L_i^{\downarrow}},\tag{3.8}$$

where T is the maximum temperature associated with optimal ε_{\min} . Equation (3.8) is derived from (3.1) and it is important for relating temperature and emissivity. This recomputation keeps temperature and emissivity consistent with each other (i.e. the same temperature can be derived from any emissivity band). The emissivity is then further processed with the Ratio and MMD modules, with minor changes to the original version of the TES algorithm as it is described in [9] and [8]. These changes include: 1) there is no refinement of ε_{\max} according to the emissivity spectral contrast, 2) the threshold T_1 for separation emissivities with small spectral contrast is not applied, and 3) the number of MMD iterations is set to one. Let us emphasize that before reporting algorithm outputs, emissivity is recomputed by (3.8) using the final value of temperature.

Ground Measurements

4.1. introduction

Post-mining sites represent areas of large-scale and intensive disturbance. They have serious impact on the surrounding landscape in many countries of the world. Original ecosystems are destroyed, and the restoration of ecosystem functions and services is necessary [1]. Afforestation is widely used reclamation method. Many studies demonstrate that post-mining sites have a large potential for carbon sequestration if afforestation has been applied [2-5]. This can contribute to mitigate the increase in atmospheric CO2 concentrations.

During opencast mining a large amount of substrate above the coal layer is removed and relocated in heaps covering extensive areas. These heaps consist of material often excavated from depths of several hundred meters. This material is called spoil substrate and it can vary in its physical and chemical properties. The heterogeneity is largely affected by geology and the method of mining and heaping. For this reason the substrates differ substantially from recent soils. They often have extreme pH and may contain high concentrations of heavy metals, polyphenols (i.e., products of coal decomposition) and salt content. Such properties can significantly impact a success and/or speed of vegetation development at post mining sites. Therefore a proper knowledge on spoil substrate properties and distribution is necessary in land rehabilitation.

Thermal remote sensing can provide beneficial tools for monitoring of post-mining areas. Particularly, land surface emissivity (LSE) can be used for spoil substrates classification. In addition, physical and chemical properties can be estimated by spectral analysis of LSE. Land surface temperature (LST) is closely connected to soil moisture which is important for establishment of new ecosystems.

LST is coupled with LSE and thus one quantity cannot be derived without knowledge of the second. These quantities cannot be explicitly derived from radiance measurement. The reason is that by observing radiance in N bands one gets N unknown emissivities plus one unknown temperature. Such a system of equations is underdetermined (i.e., more unknown than known variables). Several algorithms have been suggested to solve this problem [6]. These algorithms either require knowledge of LSE in advance, or an estimate LSE as a part of their output. A library of spectral emissivities can be utilized for: 1) determination of LST, 2) material classification, and 3) LSE validation of airborne and satellite thermal remote sensing data.

This work describes a spectral library of spoil substrate emissivities from brown coal mining sites in the Czech Republic near towns of Sokolov, Chodov, Bílina and Ustí nad

Labem (Figure 1). The spectral library contains emissivities, soil pH in water and in KCl, soil conductivity, content of water soluble Na and K, Al and Fe in KCl, loss on ignition and content of polyphenols. In addition to all measured physical and chemical parameters, sample's latitude and longitude are listed. The dataset consists of 24 spoil substrate samples, which were homogenized by mixing and sieving before any sample analysis. The toxicity test and measurement of chemical properties are discussed at length in [7]. Data collection for emissivity retrievals was performed outdoors in Petri dishes using a Fourier transform infrared (FTIR) spectrometer Model 102 (D&P Instruments, United States). The emissivity of each sample was estimated by a spectral smoothing algorithm [8].

Datasets containing LSE are rare in comparison with datasets containing LST measurements. The most well-known spectral libraries containing emissivities are the ASTER Spectral Library [9], Johns Hopkins University Spectral Library [10], Arizona State University Spectral Library [11], United States Geological Survey Spectral Library [12] and the Spectral Library of Urban Materials (SLUM) [13]. However, these spectral libraries do not include neither geographical coordinates of samples nor representatives of spoil substrates. One example of a spectral library of emissivities from calibration/validation sites containing coordinates for each sample is described in [14]. The dataset described in this paper is exceptional in its nature and location.

The data presented in this paper were used in a study focused on mapping of spoil substrates for site re-cultivation [15] as well as in a study discussing spoil substrates toxicity [7]. The mining site was also mapped with the Airborne Hyperspectral Scanner (AHS) in visible, near infrared, shortwave infrared and longwave infrared regions for mineral classification purposes [17]. Examples of emissivity spectra retrieved from AHS and their corresponding samples spectra extracted from the library are depicted in Figure 2. Samples spectra from the library were spectrally resampled with respect to AHS response functions using weighted averages [1]. Comparison of retrieved spectra in case of samples 11 and 19 shows good agreement in shape. Sample 12 exhibit deviations mainly between bands 3 and 4 (9.24 and 9.68 μ m). This can be explained by the fact that AHS pixel has 5x5 m pixel size and these pixels were not pure thus had more complex mineral composition than the collected samples. Discrepancies in magnitude can be addressed to imperfect atmospheric corrections or to different soil state during overflight.

Any activity involving remote sensing over these mining sites can benefit from publicly releasing the spectral library of spoil substrates emissivity. Apart from remote sensing application, data in spectral library can be further analyzed for identifying relationships between a sample's spectral emissivity and its chemical properties.

4.2. Data Description

The spectral library consists of 24 ASCII files. Each file describes one spoil substrate. Individual files are named according to the sample number. Files consist of a file header and spectral emissivities. Both file parts are described in the subsections below.

4.2.1. Header

The format of header is similar to format of ASTER Spectral Library header [9]. Each file contains 26 lines of header, which includes available sample information. The header is

divided into four sections separated by empty lines. First part contains 9 lines discussing sample classification, particle size and sample origin. Sample origin is expressed by latitude and longitude on the reference ellipsoid WGS84. This information is summarized in the following fields:

- 1. Name
- 2. Type
- 3. Class
- 4. Particle size
- 5. Sample No
- 6. Owner
- 7. Origin
- 8. Latitude
- 9. Longitude

A second section contains information about sample toxicity and chemical properties. The unit of each quantity is indicated in square brackets after quantity name. This header section contains following fields:

- 1. toxicity
- 2. pH in H20
- 3. pH in KCl
- 4. conductivity
- 5. water soluble Na
- 6. water soluble K
- 7. Al in KCl
- 8. Fe in KCl
- 9. loss on ignition
- 10. polyphenol content

A third section contains reference to [7], which discusses toxicity measurement and chemical analysis. Finally, the fourth header section contains the names of two columns, in which the following spectral emissivity data are aligned. Metadata in each header line contains an attribute name followed by a colon (ASCII Character 3A) and tab (ASCII Character 09) and then the corresponding value.

4.2.2. Spectral Emissivity

After the header part, the file continues on lines 27-213 with spectral emissivity data aligned in two columns. As header file indicates, the first column contains wavelength in micrometers and the second column contains corresponding emissivity value. Values in each row are separated by tab. The emissivity of each sample is provided in wavelengths interval from 8 μ m to 14 μ m. Sampling in this interval is non-linear.

4.3. Methods

The study area is situated around two post mining districts: 1) Sokolov – coal-mining district near towns of Sokolov and Chodov (North-West Czech Republic) and 2) North Czech coal mining district near towns of Bílina and Ustí nad Labem (North Czech Republic). Open-pit mines produce large areas of tailings where spoil material was sampled. Claylike tertiary sediments dominate in these districts.

Spoil substrates were sampled from bare soil without vegetation. Samples contained negligible amounts of organic matter. Extracted samples were further homogenized by mixing and sieving trough a 2 mm screen. Homogenized samples were divided into two groups, from which the first one was used for chemical analysis and the second one for toxicity testing. Samples set for chemical analysis were air dried and stored in a dark place at room temperature. Soil pH in water and in 1N KCl (which is 74.56 g of potassium chloride diluted in 1000 mL of water [17]) was measured using a pH meter with glass electrode in suspension. The suspension was prepared in 1:5 spoil to water ratio and 1:5 spoil to KCl ratio. Conductivity was measured in filtrated suspension using a conductometer. The suspension was prepared in 1:5 spoil to water ratio. Content of water soluble Na and K was also measured in filtrated water suspension (1:5 spoil to water ratio) using an ion selective electrode. Both suspensions were left to stay overnight. Al and Fe contents in 1N KCl eluate, (1:5 spoil to water ratio), were determined by spectrophotometer Spectra AA 640 (Varian, Australia). Loss on ignition was measured by burning spoil samples at 600 °C for 5 h. This process is called ashing. To determine the amount of polyphenols samples were kept in 80% ethanol (1:5 spoil to ethanol ratio) and stayed for 24 h. Samples were then filtrated and the polyphenol content was determined spectrophotometrically by Folin-Ciocalteau reagent at a wavelength of 765 nm [18]. Gallic acid was used as a standard for calibration. The polyphenol content was expressed as mg/100 g of soil. Toxicity was determined by enchytraeid toxicity test. The test is based on the population growth of pot worms in substrates. The details of the measurement are discussed in [7].

Spoil substrates emissivity measurements were collected with Designs and Prototypes Model 102 (United States) portable FTIR spectrometer. The measurement was performed outdoor under clear sky conditions during two consecutive days in summer season. The spectroradiometer was pre-heated to maximum expected ambient day temperature during the nights before both measurement days. Spectrometer orientation during measurements was south to avoid shadows. Fore-optics field-of-view was 4.8° and it was 60 cm away from the sample. Such a configuration resulted in spot size of approximately 5 cm. Samples were put in Petri dish of diameter 14 cm and they were let to be heated up naturally by sunlight. Samples temperatures ranged from 40 °C to 50 °C. Every sample was measured at three different spots. The measurement of one spot consists of ten measurements, which were averaged. The resulting emissivity of each sample is the average of all three mea-

surements. Sample temperature and emissivity were determined by a spectral smoothing algorithm, as described in [8].

During the measurements the instrument was calibrated using two blackbodies at different temperatures. A cold blackbody was set to ambient temperature (30 °C) and warm blackbody was set just above the sample temperature (40 – 50 °C). The calibration procedure during first four spoil samples was done between the changing of each sample. The calibration procedure during the rest of the measurements was done between every fourth sample. Before every sample a measurement was made of a diffuse gold reflectance plate (Infragold from Labsphere Inc.), to compensate for downwelling radiance, as suggested in [19]. The measurement of one sample along with instrument calibration and measurement of the diffuse gold reflectance plate took around 15 minutes. A description of the procedures for converting instrument response to radiance and compensating for downwelling radiance can be found in [8] and [20].

Some of the spoil substrate emissivity spectra are greater than one at certain wavelengths. This inaccuracy occurs at both ends of provided wavelengths interval (i.e. near 8 μ m and near 14 μ m). Data at these wavelengths are on the edge of atmospheric window and thus the cause of the inaccuracy is imperfect compensation for downwelling radiance. Samples with numbers 33, 34 and 38 are missing header information of latitude and longitude. Absent values are indicated by 'NA' string. In these cases the origin of the sample is specified with respect to closest town (either Bílina or Sokolov). We still find these data meaningful, since they can be used as spectral endmembers.

4.4. Data Properties

All samples are claylike sediments which imply that spectral features of clay can be found in every emissivity spectra. Figure 3 depicts three examples of spoil samples taken from the spectral library. Sample 02 (Figure 3a) is clay consisting mostly of kaolinite with significant peaks at 8.90, 9.50, and 10.00 μ m. Sample 06 (Figure 3b) is coal combined with sand and clay. The emissivity spectrum of this sample contains kaolinite features mixed with a quartz feature at 8.50, 8.90, 9.30 and 10.00 μ m. The sample 33 (Figure 3c) is bentonite rich for montmorillonite. Montmorillonite has typical dip in spectral emissivity at 9.43 μ m. The spectral emissivity library of spoil substrates includes also image providing a preview of all samples in library similar to images shown in Figure 3.

Thermal remote sensing can be used for classification of spoil substrates and for determination of their physical and chemical properties. The spectral library presented in this paper can ease and enhance all these activities. Obtained information together with LST are valuable for selection and monitoring of restoration process on post-mining sites.

4.5. Spectral Smoothing Algorithm

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Application to UHI Detection

Conclusion

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