Research paper

European spruce bark beetle (Ips typographus, L.) green attack affects foliar reflectance and biochemical properties

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1. Introduction

Bark beetles (Ips typographus, L., and Dendroctonus spp.) are important biotic disturbance agents in the coniferous forests of Europe and North America, respectively (Christiansen and Bakke, 1988; Fauhe and Heurich, 2011; Raffa et al., 2008; Seidl et al., 2011; Seidl et al., 2014). In the past decades, an increasing number of severe bark beetle outbreaks has led to an extensive economic loss in the forest industry (Goheen and Hansen, 1993; Waring et al., 2009). The economic impacts include a reduction in the commercial value of the infested trees and increased management costs (Schowalter, 2012). Besides causing economic losses, outbreaks of the bark beetle form an important factor in the ecological development of the forest landscape, in terms of forest structure and composition (Pfeifer et al., 2011; Schowalter, 2012; Simard et al., 2012; Zeppenfeld et al., 2015), as well as biodiversity and ecosystem services (Thom and Seidl, 2015). Bark beetles can affect forest ecosystems both directly and indirectly. Direct impacts include an increase in tree mortality rates and a reduction in forest stand densities (Bright et al., 2013; Eitel et al., 2011; Filchev, 2012; Hais and Kucera, 2008; Schowalter, 2012; Vanderhoof et al., 2013; Verbesselt et al.,...
2009). Other indirect impacts are; reduction in carbon uptake, changes in tree species’ distribution, as well as changes in erosion processes, fire frequency, and nutrient cycling (Beudert et al., 2015; Kurz et al., 2008; Lehnert et al., 2013; Lindenmayer and Franklin, 2002; Mikkelsen et al., 2013). Additionally, recent studies showed that the infestation of bark beetle leads to biodiversity enhancement by opening the canopy layers and altering microclimate condition in the forest. This alteration will provide the essential habitats and sources of energy for various organisms, and allow them to persist in the disturbed areas by bark beetle (Beudert et al., 2015; Lehnert et al., 2013; Müller et al., 2008).

The phenology of bark beetle and the associated host responses are well described in Wermelinger, (2004). The infested tree goes through three stages of attack (Coulson et al., 1985; Sprintsin et al., 2011; Wermelinger, 2004). These stages are termed green, red and grey attacks, respectively. During the green attack stage, the foliage remains green (hence the name green attack), and therefore, it is difficult to detect this stage by the human eye at leaf and canopy levels (Niemann and Viisintini, 2005; Wulder et al., 2006). However, the subsequent degradation of the needles can be noted by regular field observations, as, during the red attack stage, the needles of the tree turn from green to yellow and red-brown. Finally, the needles fall off, and only the grey bark will remain, hence the last stage being called a grey attack. This discolouration of the attacked trees is evident at canopy level (Coulson et al., 1985). It is expected that bark beetle infestation induces changes in the spectral response of the infested trees (Filchev, 2012; Meddens et al., 2013), as the biophysical characteristics of the entire tree, and very likely the biochemical features of the needles, change. Therefore, during the infestation period, the trees are subjected to increasing stress and face physiological change (Heath, 2001). This is due to the interruption of the water flow and the deterioration of chloroplasts as the beetle drills into the tree’s cambium tissue (Yamaoka et al., 1990). The fungi carried by the beetles penetrate the living phloem and xylem cells, hampering the translocation of water, sugar and other nutrients within the bole of the tree (Paine et al., 1997; Rohde et al., 1996; Safranyik et al., 2007; Wermelinger, 2004). This leads to a gradual change in biochemical and water content in the attacked tree, thus inducing alterations to its spectral characteristics over the course of the infestation (Deshayes et al., 2006; Lawrence and Labus, 2003; Marx and an der Havel, 2010; Reid, 1961; Sprintsin et al., 2011; Yamaoka et al., 1990).

A considerable increase in unplanned harvesting of European forests following bark beetle infestation has led to increased research interest in understanding the dynamics and improving the management of Ips typographus, L. outbreaks (Scidl et al., 2011). Furthermore, an increase in the frequency and severity of bark beetle outbreaks is expected due to global climate change (Bentz, 2010). Consequently, more attention is devoted to this topic. It has also been evoked to increase discussion regarding salvage logging in protected areas in Europe (Lehnert et al., 2013). Early detection of Ips typographus, L. outbreaks at the green attack stage may prove an important step, as management intervention to prevent further outbreaks may, therefore, involve the removal of infested trees before the new brood emerges and migrates (Wermelinger, 2004; Wulder et al., 2009). Traditionally, foresters have performed field surveys to identify infested trees; such surveys are very laborious, costly, and therefore it is inefficient and hard to apply for management purposes in large areas. Remote sensing has the potential to detect pest infestations over large areas in relatively short periods of time. Employing remotely sensed data allows monitoring of the changes in leaf and canopy properties before and after insect infestation (Bentz and Endreson, 2003; Deshayes et al., 2006; Dye et al., 2008; Jensen, 2009). To date, the utilisation of remote sensing for the monitoring and detection of bark beetles by forest managers has mainly focused on the last two attack stages (i.e., the red and grey stage) and has achieved high degrees of accuracy. During the last two stages of the attack, the changes in canopy colour effect the spectral reflectance signature which are mainly been used as an indicator to detect infestations (Carter et al., 1998; Franklin et al., 2003; Heurich et al., 2010b; Latifi et al., 2014; Meddens, 2012; Nikolov et al., 2014; Skakun et al., 2003; Wulder et al., 2006). However, detecting the infestation in the last two stages is not sufficient for appropriate management, as phenological research proved that during the red attack stage the newly developed beetles have already left their host trees and started to attack new trees. Therefore, the continuing of an outbreak can not be prevented by salvage logging during this stage. Consequently, the detection of the bark beetle at the green attack stage by means of remote sensing is necessary to have a meaningful effect on the spreading of the beetle, but is challenging due to the lack of apparent visual symptoms in needles.

Early detection of infestations by Dendroctonus spp in lodgepole pine trees has been investigated at canopy level (Gimbarzevsky et al., 1992; Heath, 2001; Klein, 1973; Murtha and Wiart, 1989; Murtha, 1972). Similarly, detection of a bark beetle green attack (ips typographus, L.) at canopy level in Norway spruce trees has been investigated by (Immitzer and Atzberger, 2014; Lausch et al., 2013; Marx an der Havel, 2010; Ortiz et al., 2013). However, these studies did not succeed in discriminating healthy from green attacked trees. More recently, Niemann et al. (2015) used LIDAR and hyperspectral data to examine the spectral properties of healthy trees and those under mountain pine beetle green attack and demonstrated that the most promising wavelengths, for the detection of mountain pine beetle at green attack stage, is located in the shortwave infrared region. Furthermore, Näsi et al. (2015) used a hyperspectral sensor (500–990 nm) mounted on an unmanned aerial vehicle (UAV) to map bark beetle damage at the tree level, by dividing tree stands into three different classes (healthy including trees with potential early infestation stage, red attack; and dead). They found that the healthy and dead trees can be classified with 90% over-all accuracy, however, when all classes were considered (healthy, red and dead), the overall accuracy dropped to 76%.

At the leaf level, there are very few studies, although with different beetle and tree species, that have examined the differences in spectral reflectance between healthy needles and those that have been under bark beetle green attack (Ahern, 1988; Cheng et al., 2010; Foster et al., 2017). In addition to a change in spectral reflectance properties, Cheng et al. (2010) observed that there were differences in water content of healthy and infested needles, particularly, between 1318 and 1322 nm. These results show that remote sensing has the potential to detect early stages of bark beetle attacks. However, these studies have mostly considered the effect of early infestation on reflectance spectra and not on the biochemical properties of the needles. Therefore, it is highly important to understand whether, in addition to the spectral reflectance, the biochemical properties are also affected at the bark beetle early infestation stage. Consequently, the topic warrants further investigation.

As mentioned earlier, it is expected that the infested tree will exhibit a change in terms of its biochemical and spectral properties, due to the beetle larva and blue stain fungi such as (Ophiostoma and Cerato-cystis species) carried by the beetles starts to penetrate the living phloem and xylem cells, hampering the translocation of water, sugar and other nutrients within the bole of the tree (Paine et al., 1997; Rohde et al., 1996; Safranyik et al., 2007; Wermelinger, 2004). Therefore, in this study, we aim to investigate the possible early detection of a bark beetle green attack by examining and comparing the foliar biochemical (chlorophyll and nitrogen) and spectral properties (400–2000 nm) of both healthy and green attacked trees. Chlorophyll and nitrogen are two important elements that have a key role to play in plant life and status and can be considered as indicators that reflect the status of plant growth and health (Heinze and Fiedler, 1976; van Maarschalkerweerd and Husted, 2015; Wang et al., 2015a; Wang et al., 2015b). As such changes, initiated by bark beetle attack cause stress during the infestation time, we hypothesise that chlorophyll and nitrogen concentrations are reduced during such an attack. These can thus provide suitable proxies for detecting the presence of Ips typographus L. during a
green attack stage. Specifically, there are three main objectives: (a) to investigate the impact of an *Ips typographus* L. green attack on foliar spectral reflectance; (b) to examine the changes in foliar biochemical properties due to the *Ips typographus* L. green attack; (c) to explore the impact of the *Ips typographus* L. green attack on the estimation accuracy of foliar biochemical properties utilizing hyperspectral measurements.

2. Material and methods

2.1. Study area

The study area is the Bavarian Forest National Park (BFNP) in southeast Germany, which covers an area of 240 km² between 13°12′9″ E (longitude) and 49°3′19″ N (latitude) (Fig. 1). The elevation in the BFNP ranges from 600 m to 1453 m. This region is characterised as having a temperate climate with a total annual precipitation along the gradient between 900 and 1800 mm as well as a mean annual temperature that varies between 3.5 and 7.2 °C (Bässler et al., 2008; Lehnert et al., 2013). The area is divided into three ecological zones: high elevations, hillsides, and valley bottoms. Around 90% of the tree stands in high elevations are Norway spruce (*Picea abies* (L.) Karst), 2% are beech (*Fagus sylvatica* L.), and the remaining 8% are covered by other broad-leaves trees, mainly Common Rowan (*Sorbus aucuparia*). While on the hillsides, around 58% is occupied by Norway spruce, and the rest is a mixture of European silver fir (*Abies alba*) and beech. In the valley bottoms, 83% of the trees are Norway spruce, and the rest is a mix of species (Cailleret et al., 2014; Heurich et al., 2016a). Multiple storm events in the 1980s and a series of hot summers in the 1990s have lead to an extensive mortality of canopy trees by bark beetles on about 8000 ha (Lausch et al., 2013).

2.2. Data acquisition

The adult bark beetle starts to attack new host trees during the European springtime (between the end of April and the beginning of May). This process strongly depends on the air temperature as the beetles start their swarming when the air temperature reaches 16.5 °C (Lobinger, 1994; Wermelinger, 2004). Therefore, the field data were collected in the early summer of 2015 within 15–30 days from the early stages of infestation. In total 120 trees were measured, of which 66 were healthy, and 54 were freshly infested trees. An extensive field survey was conducted to identify the trees under bark beetle green attack by spotting piles of dry, boring dust, which had been pushed out onto the bark surface (Fig. 2). An average of 2–3 branches were taken from each tree. Needle samples from each branch were collected separately. All the branches were taken from the upper part of the trees, which was exposed to the sunlight. The heights of the trees varied between approximately 25 and 30 m. A crossbow was used to shoot an arrow with a fishing line attached to a branch with sunlit leaves (Ali et al., 2016). The fishing line was used to feed a rope over the targeted branch. Once the rope was in place, the branch was pulled down gently until it broke off. Needles were immediately removed from the fallen branches and placed in a labelled plastic zip-locked bag, which was then covered with wet pulp paper and subsequently transported to the laboratory in a portable cooling box filled with frozen ice packs to keep the sample cool. The aim was to retard, as much as possible, any changes in the needles’ reflectance spectra and biochemical characteristics (Malenovský et al., 2006). Once the spectral measurements were completed, the same sample branch was transported to the laboratory to measure the fresh weight, leaf area and the biochemical properties of the foliar samples.
2.3. Chemical analyses

The concentration of Chlorophyll a and b in the collected fresh leaf samples was determined destructively by a spectrophotometer following the Lichtenthaler (1987) method using acetone (v 100%). In addition, to determine foliar nitrogen, the needles were dried for 72 h using an oven dryer at 60° C. The dried needles were properly grounded using mortar and pestle until the ground leaves became a soft powder and were passed through a 0.25 mm mesh screen. Subsequently, 2 mg of powdered leaves was transferred to a small aluminium capsule to measure the nitrogen content, using an organic elemental analyser (FLASH 2000).

2.4. Reflectance measurements

The leaf directional hemispherical reflectance from 350 to 2500 nm was measured for collected samples, using an ASD FieldSpec-3 Pro FR spectrometer equipped with an ASD RT3-3ZC integrating sphere (Analytical Spectral Devices, Inc., Boulder, Co, USA). The integrating sphere had a port diameter of 15 mm for a reflectance measurement. The Norway spruce needles did, however, not cover the entire sample port of the integrating sphere. Therefore, the method proposed by Daughtry et al. (1989) was applied, where a universal sample holder, designed by Malenovský et al. (2006), was used to accommodate all sizes and lengths of Norway spruce needles. For every measurement five to six needles were secured on the sample holder with cellophane tape, leaving a gap of one needle’s width between needles to avoid multiple reflectances of the adjusted needles (Ali et al., 2016; Daughtry et al., 1989). To minimise the effect of spectral signal noise, for every needle sample two hundred scans were averaged to present a single spectrum. The integrating sphere has an internal light source, and therefore, the sun angle and cloudiness does not affect the spectral measurement (Foster et al., 2017). However, to prevent possible light entering into the integrating sphere ports, covering the integrating sphere with a black piece of cloth is recommended by Mesarch et al. (1999).

The measured spectral reflectance from the sample holder contained the effects of the gap between the attached needles to the sample holder; therefore, the spectral measurements were corrected by calculation of the gap fraction. To do so, a black painted paper mask with a 15-mm diameter circular aperture was precisely superimposed onto the samples and photographs were taken using a 16.1 megapixel Panasonic LUMIX camera (Ali et al., 2016). Then, the images were imported into ImageJ software (Schneider et al., 2012), to compute the gap fraction, based on the illuminated area of the sample port. The following equation was adapted from Mesarch et al. (1999) for the Field ASD spectrometer and to compute the hemispherical reflectance of the sample needles:

\[
\text{Reflectance} = (\frac{p - R_d/I - R_d}{R_r})/(1 - GF);
\]

where: \( p \) is the measured reflectance from the sample holder; \( R_d \) is stray light (ambient light) inside the integrating sphere, measured as a radiation flux of the empty illuminated sample port in reflectance mode; \( R_r \) is reference of sample reflectance; and \( GF \) is the gap fraction of the sample.

2.5. Data preprocessing and statistical analysis

A series of pre-processing steps were performed on the measured reflectance data. First, the spectral reflectance was examined, and “noisy” bands (in the spectral ranges 350–399 nm and 2000–2500 nm) were excluded from the analysis. Secondly, to eliminate and reduce the sensor noise, a Savitzky – Golay smoothing filter with a frame size of 15 (2nd-degree polynomial) was applied to the reflectance spectra (Savitzky and Golay, 1964).

The significance of differences (\( p \leq 0.05 \)) in foliar reflectance as well as chlorophyll and nitrogen concentration between leaf samples from green attacked and healthy trees, was examined using Student \( t \)-tests, in order to determine whether there is spectral variation at any spectral band. Because the considered adjacent wavelengths are highly correlated, we corrected the p-value using Holm (1979) procedure, next the significant wavebands with mean reflectance spectra of both healthy and infested samples were plotted following the technique used by Schmidt and Skidmore (2003).

To investigate the relationship between the spectral reflectance of the healthy and infested samples and their biochemical parameters, the Pearson’s correlation coefficients were calculated between spectral reflectance (400 – 2000 nm) and chlorophyll as well as nitrogen concentration for both healthy and infested samples to identify the most sensitive wavebands in healthy and infested samples.

2.5.1. Partial least square regression model (PLSR)

Partial least squares regression (PLSR) was used to investigate the impact of the bark beetle green attack stage on the retrieval accuracy of the chlorophyll and nitrogen concentrations in needles. PLSR is a regression method that takes into account both the variance of the explanatory and the dependent variables. It specifies a linear relationship between a set of dependent (Y) variables and a set of predictor (X) variables (Wold et al., 2001). PLSR performs particularly well when the various X-variables have strong correlations, as is normally the case with hyperspectral data. (Carrascal et al., 2009; Nicolai et al., 2007). Further details on PLSR can be found in Geladi and Kowalski (1986).

PLSR was used to predict the foliar chlorophyll and nitrogen concentrations (dependent variables) of the healthy and infested needles from spectral reflectance (predictor variables). PLSR models were built independently for the healthy and infested samples. For the chlorophyll...
concentration, PLSR was fitted to the VNIR range (400–790 nm), as this spectral region mainly contributes to total chlorophyll estimation in plants (Curran, 1989; Yoder and Pettigrew-Crosby, 1995). However, as nitrogen is associated with many other leaf chemical compounds, the spectral wavebands corresponding to nitrogen are distributed over the whole spectrum (VIR, NIR, SWIR) (Curran, 1989; Peterson et al., 1988) and consequently, for the nitrogen concentration, PLSR was fitted to the entire spectrum (400–2000 nm). To determine the number of components needed to build the PLSR model, Monte Carlo cross-validation (MCCV) was applied (Xu and Liang, 2001). The procedure involves adding an extra component to the model when the value of the root mean square error of cross-validation (RMSEcv) is reduced by ≥2% (Darvishzadeh et al., 2008; Geladi and Kowalski, 1986). The performance of the PLSR models were assessed by computing the coefficient of determinations ($R^2$), the normalised root mean square error of predictions (nRMSEP) and normalised root mean square error of cross-validations (nRMSEcv) of predicted versus measured values.

We further examined whether the most informative wavebands for the estimation of chlorophyll and nitrogen concentrations in both healthy and infested samples correspond. To achieve this, the variable importance in the projection (VIP) was calculated for all wavebands from the PLSR models. VIP scores summarise the influence of individual X variables (reflectance) in the PLSR model. A variable with a VIP score close to, or greater than 1, can be considered significant in a given model (Chong and Jun, 2005; Tenenhaus, 1998; Wold et al., 1993). A detailed description of the calculation of VIP scores can be found in Farres et al. (2015); Wold et al. (1993). TOMCAT and libPLS toolbox was used within MATLAB to establish PLSR analysis and calculate VIP score, respectively (Daszykowski et al., 2007; Li et al., 2014).

3. Results

3.1. Spectral reflectance variation due to bark beetle green attack

The results demonstrated that the mean reflectance spectra of healthy and green attacked foliage were statistically different ($p \leq 0.05$) for 917 wavebands out of the 1600 wavebands used in this study (Fig. 3). As can be seen from Fig. 3, a clear distinction was observed in the mean reflectance between healthy and infested foliage in the visible and near-infrared regions. In the visible region, the mean reflectance of the infested foliage was higher than of the healthy leaves, in conjunction with chlorophyll degradation (Fig. 4). However, there was a larger difference ($p \leq 0.05$) between infested and healthy needles in the wavelength range 730–1370 nm, as the healthy needles had higher reflectance compared to mean infested spectra.

3.2. Impact of bark beetle green attacks on foliar biochemical properties

A Student $t$-test demonstrated a significant difference between total chlorophyll and foliar nitrogen concentrations of healthy and infested leaves ($p < 0.05$). In healthy foliage, the mean and ± of the total chlorophyll concentration were 0.766 ± 0.140 mg/g, respectively, whereas for the green attacked leaves they were 0.657 ± 1.62 mg/g, respectively. Furthermore, the concentration of nitrogen was 1.25 ± 0.21% within healthy leaves, while for the infested leaves they were 1.13 ± 0.18%, respectively (Fig. 4). The correlation between total chlorophyll and nitrogen concentration was higher in needles from healthy trees ($r = 0.68$, $p < 0.05$) than in needles from green attacked trees ($r = 0.57$, $p < 0.05$).

3.3. Effects of bark beetle green attack on the retrieval accuracy of leaf biochemical properties

The impact of bark beetle green attack on the retrieval accuracy of foliar chlorophyll and nitrogen concentrations was assessed using PLSR. In general, higher accuracies were observed when healthy samples were assessed (Table 1). For healthy foliage, the coefficients of determination for chlorophyll and nitrogen concentrations were $R^2 = 0.64$ and $R^2 = 0.76$, respectively. However, these coefficients decreased for models fitted to data from infested foliage, both for chlorophyll ($R^2 = 0.55$) and for nitrogen ($R^2 = 0.68$) (Table 1). Similar results were discerned with the Pearson correlation coefficient between chlorophyll and the reflectance data, as well as between nitrogen and the reflectance data. As can be observed from Fig. 6, significantly higher correlations were observed for healthy leaves than for infested leaves. In addition, the prediction error for chlorophyll in healthy leaves ($nRSMSEP = 0.20$) was much lower than for chlorophyll in infested leaves ($nRSMSEP = 0.62$) (Fig. 5). However, this variation in nRSMSEP for nitrogen content was much lower (0.12 and 0.22) in both healthy and infested samples respectively.

As shown in Figs. 6 and 7, the spectral regions, in particular, the green and the red edge (536–559 nm and 732–790 nm, respectively), demonstrated the highest VIP (VIP > 1) and a moderately negative correlation ($r = -0.41$ and $-0.32$, for the green and red edge, respectively; $P < 0.05$) between chlorophyll and the reflectance spectra of healthy foliage. The high VIP value indicated that the spectral feature was of major importance in estimating the chlorophyll concentration. A weaker negative correlation with a lower VIP score was observed for the infested samples ($r = -0.12$, $P < 0.05$). The spectral regions that were most sensitive to changes in nitrogen content were in the range of 980–1000 nm and 1448–1469 nm, where
the infested leaves had a lower VIP score (Fig. 6). Furthermore, the correlation coefficients between foliar nitrogen concentration and the reflectance spectra show a distinct variation between healthy and infested foliage, especially in the spectral region between 720–1400 nm. The healthy foliage showed a moderately negative correlation ($r = -0.30, P < 0.05$), while for the infested foliage a positive correlation was found (Fig. 6).

4. Discussion

It has long been believed that the detection of bark beetle green attack using remote sensing data is a challenging task (Wulder et al., 2009; Niemann and Visintini, 2005). However, recent studies have shown that remote sensing data obtained across a number of wavelengths have the potential for the detection of bark beetle green attack (Cheng et al., 2010; Foster et al., 2017; Niemann et al., 2015). In this study, we investigated whether bark beetle green attack has a concurrent effect on key foliar biochemical properties (i.e. chlorophyll and nitrogen concentration) as well as spectral responses (400–2000 nm).

Table 1

<table>
<thead>
<tr>
<th>Sample Condition</th>
<th>Variables</th>
<th>Nr. Of latent variables</th>
<th>nRMSEP</th>
<th>nRMSECV</th>
<th>$R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Healthy Chl</td>
<td>9</td>
<td>0.20</td>
<td>0.24</td>
<td>0.64</td>
<td></td>
</tr>
<tr>
<td>Healthy N</td>
<td>10</td>
<td>0.12</td>
<td>0.20</td>
<td>0.76</td>
<td></td>
</tr>
<tr>
<td>Infested Chl</td>
<td>10</td>
<td>0.62</td>
<td>0.75</td>
<td>0.55</td>
<td></td>
</tr>
<tr>
<td>Infested N</td>
<td>8</td>
<td>0.22</td>
<td>0.35</td>
<td>0.68</td>
<td></td>
</tr>
</tbody>
</table>

The healthy foliage showed a moderately negative correlation ($r = -0.30, P < 0.05$), while for the infested foliage a positive correlation was found (Fig. 6).
The results demonstrate that the foliar reflectance of needles from green attacked trees differs significantly from healthy needles (p ≤ 0.05), in particular between wavelengths of 730 and 1370 nm. Furthermore, the infestation significantly (p ≤ 0.05) affected the biochemical concentrations of total chlorophyll and foliar nitrogen and reduced their retrieval accuracy using PLSR (Figs. 3 and 4).

The measured reflectance spectra of healthy needles matched those of similar studies which measured needle reflectance of Norway Spruce (Ali et al., 2016; Atzberger and Werner, 1998; Ković et al., 2012). However, there were significant differences observed between the reflectance spectra of the healthy and the green attacked samples. This can be attributed to differences in their foliar biochemical properties, especially to their chlorophyll concentration which is known to be affected by vegetation stress. In our study, the reflectance increased in the visible region, and decreased in the NIR, for the green attack leaves. This is in line with the findings of Abern (1988), who studied the spectral range of 400-1100 nm and showed that the spectral band at the green peak increased for the lodgepole pine needles infested by similar beetle species (Mountain pine beetle) at the green attack stage. In the visible region, the concentration of pigments such as total chlorophyll is the main factor for determining leaf spectral variation and absorption peaks (Carter and Knapp, 2001; Demetriades-Shah et al., 1990; Feret et al., 2008; Zhang et al., 2008). The infested trees had significantly (p < 0.05) lower chlorophyll and foliar nitrogen concentrations than the healthy ones, resulting in lower absorption and higher scattering (Figs. 3 and 4). The observed behaviour of the reflectance spectra in the visible region in this study is a clear indication of stress caused by early infestation and is in agreement with the results of earlier studies focused on plant stress (Carter, 1993; Carter and Knapp, 2001; Zarco-Tejada and Sepulcre-Cantó, 2007). Furthermore, wavelength ranges between 730 and 790 nm that contained the most informative spectral region regarding variations in chlorophyll concentration, demonstrated the greatest amount of dissimilarity (58 wavebands) between the healthy and infested leaf samples (Fig. 7). This spectral region is often referred as red-edge, and it responds quickly to changes in foliar chlorophyll (Carter, 1993, Carter and Knapp, 2001; Smith et al., 2004). This is in good agreement with finding by Lottering et al. (2016), as they studied the utility of spectral vegetation indices derived from WorldView-2 data in detecting and mapping G. scutellatus induced vegetation defoliation. Their result showed that the combination between red-edge and NIR region has the potential to detect stress induced by G. scutellatus insect.

Moreover, as noted in Fig. 3, the more pronounced changes in the reflectance spectra were observed at the wavelength region between the NIR and SWIR regions (730-1370 nm), where the reflectance of the infested leaves was distinctively lower than that of the healthy ones. This can probably be explained by the changes in water content that occur due to the infestation, which leads to a degeneration of the internal leaf structure at the cellular level (Miller et al., 1991; Murtha, 1978; Paine et al., 1997; Slaton et al., 2001; Zhang et al., 2012). This result partially agrees with the findings of Niemann et al. (2015), who demonstrated that at the wavelength of 970 nm there was no separation between healthy trees and samples infested by mountain pine beetle, while at the wavelength of 1200 nm a significant difference existed between them. Additionally, Ismail and Mutanga (2010) showed the importance of the wavelength between 900 and 1110 nm and identified both RatioO275 and water index has the ability to assess water stress induced by S. noctilio in P. patula trees from an early stage of infestation when there is no sign of infestation at the canopy level.

In addition to the variations in VIS and NIR, the reflectance spectra

Fig. 6. Correlation between chlorophyll and nitrogen and the reflectance of individual wavebands. The bold lines represent the wavebands at which the reflectance spectra correlated significantly with the chlorophyll and nitrogen concentrations. The arrows show the wavebands in which recorded the highest correlation with chlorophyll content in both healthy and infested samples.

Fig. 7. Importance of wavelengths corresponding to the highest value of variable importance in the projection scores of partial least squares regression in healthy and infested samples, used for chlorophyll and nitrogen estimation.
of healthy and green attacked samples were also significantly different \((p < 0.05)\) in two short-wave infrared regions \((1430–1500 \text{ nm and } 1897–2000 \text{ nm})\) (Fig. 3). The most likely cause for this variation is the low water and nitrogen content in the infested samples \((\text{Curran et al., 1992; Mirzaie et al., 2014; Peñuelas and Filella, 1998})\). Physiological research has shown that the fungi carried by the beetles penetrate the living phloem, hampering the translocation of water, sugar and other nutrients within the bole of the tree \((\text{Paine et al., 1997; Rohde et al., 1996; Safranyik et al., 2007; Wermelinger, 2004})\). Not only are the fungi instrumental in the decline of trees, but they also provide a necessary food source for insects during the infestation period \((\text{Runesson, 1991})\). This leads to a gradual decrease in needle biochemical content and water content. \(\text{Foster et al. (2017) and Cheng et al. (2010)}\) showed that a number of wavebands in the shortwave infrared region have the potential for the detection of bark beetle infestation at the green attack stage which is in line with our observation. Furthermore, in our study, we have verified that the infested trees had a significantly \((p < 0.05)\) lower nitrogen concentration than that found in the healthy trees.

5. Implications for remote sensing applications

Our study confirms the importance of hyperspectral measurement as well as foliar biochemical properties \(\text{(i.e. chlorophyll and nitrogen)}\) for the detection of \textit{Ips typographus}, \text{L.} green attack. An extension of this finding would be to investigate the up-scaling of our findings to the canopy level. However, this forms a challenge as many factors such as logistical and technological aspects may limit the scaling up to the canopy level \((\text{Foster et al., 2017; Wulder et al., 2009})\). One of the critical logistical factors that affect a remotely sensed survey for green attacked trees is the timing of the optimal conditions for image acquisition. In addition, spatial and spectral resolutions play an important role in the studies of bark beetle green attack. However, this limitation may be overcome by utilising a UAV-based application and newly available multispectral satellite data, such as provided by Sentinel-2. In our study, the estimation accuracy of foliar chlorophyll and nitrogen concentration using hyperspectral measurements decreased when the tree was infested by bark beetle green attack. This might be used as an indicator for the efficient landscape-wide detection of bark beetle green attack. It is, however, important to note that retrieval accuracies for the total chlorophyll and foliar nitrogen concentrations at the leaf level...
would probably be different from those at the canopy level, because structural and external factors such as illumination and atmospheric conditions may affect the reflectance spectra at the canopy level. Additionally, we found that the wavelength region between 730 and 790 nm (red-edge) is the most informative spectral region regarding variations in chlorophyll concentration due to bark beetle green attack (Fig. 6). This portion of the spectrum can be found in new multispectral satellites such as Sentinel-2, World-View-2 & 3 RapidEye and therefore it might be possible to scale up our findings to the canopy level, particularly considering the high spectral and spatial resolution data. Further investigation will show how accurate green attack stages can be detected with different air and spaceborne sensors.

6. Conclusion
This study demonstrates that reflectance properties of healthy and green attacked Norway spruce trees are significantly different in 917 wavebands between 400 and 2000 nm. We also observed differences in the biochemical properties chlorophyll and nitrogen between healthy and green attacked needles by the bark beetle (Ips typographus, L.). Furthermore, the results demonstrated that the relationship between reflectance and both chlorophyll and nitrogen deteriorated as the leaves became stressed by a bark beetle green attack. Such an infestation affected the estimation accuracy of chlorophyll and nitrogen concentrations. The research findings indicate that hyperspectral measurements are promising, and present a powerful tool to determine the damage caused by bark beetle green attack at the leaf level. Further research is required to assess whether or not, the findings of the current study can be verified at the canopy level using different remote sensing data.

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