Spectral studies of ocean water using DOAS

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Abstract

Methods enabling the retrieval of oceanic parameter from the space borne instrumenta-
tion Scanning Imaging Absorption Spectrometer for Atmospheric ChartographY (SCIA-
MACHY) using Differential Optical Absorption Spectroscopy (DOAS) are presented.
SCIAMACHY onboard ENVISAT measures back scattered solar radiation at a spectral
resolution (0.2 to 1.5 nm). The DOAS method was used for the first time to fit mod-
elled Vibrational Raman Scattering (VRS) in liquid water and in situ measured phy-
toplankton absorption reference spectra to optical depths measured by SCIAMACHY.
Spectral structures of VRS and phytoplankton absorption were clearly found in these
optical depths. Both fitting approaches lead to consistent results. DOAS fits correlate
with estimates of chlorophyll concentrations: low fit factors for VRS retrievals corre-
spond to large chlorophyll concentrations and vice versa; large fit factors for phyto-
plankton absorption correspond with high chlorophyll concentrations and vice versa.
From these results a simple retrieval technique taking advantage of both measure-
ments is shown. First maps of global chlorophyll concentrations were compared to the
corresponding MODIS measurements with very promising results. In addition, results
from this study will be used to improve atmospheric trace gas DOAS-retrievals from
visible wavelengths by including these oceanographic signatures.

1 Introduction

Ocean color sensors on board earth orbiting satellites provide a long-term record of
remotely sensed aquatic parameters. In practice such parameters are retrieved using
appropriate radiance or reflectance ratios. Despite of their limitations these approaches
have been very successful leading to significantly improved knowledge of inherent and
apparent properties of sea water, chlorophyll concentration, and provided important
information about the carbon cycle.

In recent studies Vassilkov et al. (2002) and Joiner et al. (2004) showed an alterna-
tive approach using backscatter ultraviolet instrumentation utilizing Vibrational Raman Scattering (VRS) spectral signature of liquid water. They clearly demonstrated the capability to evaluate oceanic chlorophyll content using VRS at a spectral resolution from GOME (Global Ozone Monitoring Experiment). Vountas et al. (2003) confirmed the coherence between VRS and chlorophyll concentration using Differential Optical Absorption Spectroscopy (DOAS) analysis of GOME data.

In this study DOAS, which is a wide-spread technique in atmospheric trace gas retrieval, is applied to backscatter measurements from the spaceborne instrument SCIAMACHY (SCanning Imaging Absorption spectroMeter for Atmospheric CHartographY) to retrieve ocean parameters. SCIAMACHY is a follow-on mission of GOME with extended characteristics. It was launched on board ESA’s ENVironmental SATellite, ENVISAT, in 2002.

As for example shown in Vountas et al. (2003) VRS spectral signature can clearly be detected in oligotrophic waters due to larger penetration depths of light. Opposite behavior is shown in eutrophic waters such as large parts of the coastal zones. DOAS retrievals in such regions will not show significant spectral structure from VRS because of increased absorption by dissolved matter and phytoplankton.

As direct and scattered UV-Vis. light penetrates the ocean surface and therefore interacts with phytoplankton the phytoplankton absorption spectra are imprinted on the backscattered radiance spectrum in the visible also. The molecules of the photosynthetic pigments exhibit well structured absorption features in the visible wavelength range. Analogue to the VRS approach, a straightforward way to detect the phytoplankton biomass is therefore the DOAS evaluation of phytoplankton absorption.

The objective of this publication is to show the potential of the DOAS analysis to retrieve oceanic parameters (as fit factors) using two complementary methods taking into account i) VRS spectral imprint in the UV and ii) the spectral fine structure of phytoplankton absorption bands in the visible. Such evaluations require spectral resolution. Modern spaceborne spectrometers for analysis of atmospheric constituents, such as SCIAMACHY measure radiances at a spectral resolution around 0.2 nm (within the
spectral intervals used in this study). On the other hand the instrument provides only a rather coarse spatial resolution $30 \times 60 \text{ km}^2$ compared to established ocean color missions like for example MODIS or SeaWiFS. In order to resolve scales of variability in the ocean at least a spatial resolution of $1 \text{ km}^2$ is necessary. Therefore this study aims to show the feasibility to retrieve valuable oceanic information from hyperspectral data and provides the opportunity to further exploit spectral fine structure for other oceanic applications.

The publication is structured as follows: After giving some information about the SCIAMACHY instrument in Sect. 2.1 we describe briefly the DOAS method in Sect. 2.2. While phytoplankton absorption spectra can be measured, VRS spectra have to be modelled. Therefore Sect. 2.3 gives a description of a simple model to compute VRS spectra (which is a brief summary of Vountas et al., 2003). Finally both approaches are applied to SCIAMACHY backscatter measurements (Sect. 3) and the retrieval of chlorophyll concentration is demonstrated (Sect. 4).

2 Data sets and methods applied

2.1 SCIAMACHY

SCIAMACHY (Scanning Imaging Absorption Spectrometer for Atmospheric CHartographY) (Bovensmann et al., 1999) was launched on board ESA’s ENVironmental SATellite, ENVISAT, in 2002. The instrument is designed to measure a broad band of solar radiation, spanning from the UV to the near infrared. Apart from its broad spectral coverage, a unique feature of SCIAMACHY is the ability to detect sunlight that has been transmitted, scattered and reflected in the Earth’s atmosphere in different observation geometries. In particular, the spectrometer continuously alternates between limb and nadir modes, which allows the observation of the same volume of air under different viewing angles, facilitating the separation of stratospheric and tropospheric components of molecular absorbers. The instrument takes backscatter measurements
at high spectral resolution (in the UV-Vis.: 0.26–0.44 nm).

This study exclusively uses spectra measured in nadir-viewing. The main objective of SCIAMACHY, as well as the new Ozone Monitoring Instrument (OMI) on AURA and GOME-2 (Global Ozone Monitoring Experiment-2) on METEOSAT, is to determine the abundances of atmospheric trace gases. Although SCIAMACHY is primarily an atmospheric mission, part of the detected solar radiation penetrates the ocean surface and picks up absorption and backscattering signals from sea water.

2.2 Differential Optical Absorption Spectroscopy (DOAS)

The DOAS method (Perner and Platt, 1979) has proven to be a powerful, computationally fast and simple tool in atmospheric remote sensing. As already shown to be feasible for GOME (Vountas et al., 2003) data we will apply and further develop DOAS retrievals of ocean parameters using SCIAMACHY data.

The DOAS technique usually exploits the sharp spectral features in backscatter radiance spectra that are caused either by absorption due to atmospheric constituents or spectral re-distribution features as induced by VRS in ocean waters (Vassilkov et al., 2002; Vountas et al., 2003). Spectrally dynamic features are separated from the slowly varying attenuation due to scattering by subtracting a low-degree polynomial (typically of a degree of three) in a spectral fit procedure. However, depending on the size of the fitting window and the degree of the polynomial the algorithm is also able to retrieve broad spectral structures (Eisinger et al., 1996).

The DOAS retrieval involves the fitting and scaling of a set of spectra within a certain wavelength window, which can be composed of:

– atmospheric trace gas absorption cross-sections,
– a polynomial,
– so-called pseudo-absorber spectra for example a reference spectrum for VRS in liquid water and
– additional spectra for chlorophyll absorption
to the measured optical depth $\tau(\lambda, \theta) = \ln(I/I_0)$. With $I$ being the backscattered radiance, $I_0$ the extraterrestrial irradiance. $\tau$ is a function of the wavelength $\lambda$ and the solar zenith angle $\theta$ (dependence omitted in the following). The fitting is formalized as a least-squares minimization:

$$\left| \tau(\lambda) - \sum_{i=1}^{L} S_k \sigma_i(\lambda) - S_a a(\lambda) - S_v \nu(\lambda) - \sum_{j=1}^{M} S_j r_j(\lambda) - \sum_{k=0}^{N} x_k \lambda^k \right|^2 \rightarrow \text{min.} \quad (1)$$

Here the input parameters to DOAS are:

- $\sigma_i(\lambda)$ being the cross-sections for each atmospheric trace gas $i$ (total number of gases is $L$),
- $a(\lambda)$: is the spectrum for phytoplankton absorption and
- $\nu(\lambda)$: is a spectrum accounting for VRS (see below).
- $M$ pseudo-absorber spectra $r_j(\lambda)$. Taking into account other pseudo-absorbers is often necessary and depends on the wavelength range investigated.

The output of DOAS are the fit factors for each spectrum and coefficients for each polynomial variable (here the wavelength $\lambda$). Where $S_k$ is the fit factor of the $k$th atmospheric absorber and $S_j$ the fit factor of the $j$th pseudo absorber. $x_l$ is the $l$th polynomial coefficient (degree of the polynomial is $N$) accounting for spectral broadband features like Rayleigh and Mie scattering in atmosphere. The fit factor of chlorophyll absorption is $S_a$ and the one of VRS is $S_v$.

The DOAS fit factors $S_a$ and $S_v$ contain specific information about ocean optical characteristics and will therefore be the target quantities in the following.

It should be noted that the DOAS method will lead to erroneous results if the reference spectra used in the fit have spectral correlation, i.e. the fit algorithm will not be able to distinguish between similar spectral features.

As a scalar indicator of fit quality $\chi^2$ values are often used. The $\chi^2$ values are defined as the square of the wavelength-integrated fit residual weighted with the square of the measurement error.
2.3 Optical properties of ocean water

Before we can use DOAS to determine the fit factors $S_v$ and $S_a$ to derive further information from them all relevant input spectra have to be selected carefully. Of particular importance are $\nu(\lambda)$ and $a(\lambda)$.

**Determination of $\nu(\lambda)$:** In order to account for VRS in DOAS retrievals an adequate VRS spectrum has to be used in the analysis. Such a spectrum can be considered as a-priori information for the retrieval.

VRS is an inelastic scattering process which provides a mean wavenumber shift of 3357 cm$^{-1}$. It involves two fundamental OH stretch vibration modes of the water molecule that are further modified by hydrogen bonding and rotational fine structure. These interactions induce a broad band of emissions around the mean wavenumber shift so that water Raman emissions occur over a band of $\approx$30–50 nm. In order to account for this redistribution of photons a spectroscopic model of VRS has to be embedded in an adequate description of the interaction between light, water and atmosphere. This requires a coupled atmosphere-ocean radiative transport model.

As a first step we will have to define a reference spectrum $\nu$ accounting for VRS in liquid water at a wavelength $\lambda$ as:

$$\nu(\lambda) = \ln \frac{I^{+\text{VRS}}(\lambda)}{I^{-\text{VRS}}(\lambda)}.$$  

(2)

With $I^{+\text{VRS}}$ being the modeled radiance taking into account VRS and $I^{-\text{VRS}}$ neglecting it (Vountas et al., 2003). A prerequisite for taking into account VRS in DOAS-type retrievals is therefore a precise knowledge of the quantities $I^{+\text{VRS}}$ and $I^{-\text{VRS}}$.

The following paragraphs list the underlying spectra and assumptions necessary to create the reflectance being the central input for the determination of $I^{+\text{VRS}}$ and $I^{-\text{VRS}}$. The way how the reflectance is determined is shown elsewhere in detail (Vountas et al., 2003; Sathyendranath and Platt, 1998).

We adopt a reflectance model, originally proposed by Sathyendranath and Platt...
(1998) which is based on a modified Quasi-Single-Scattering Approximation. It takes into account elastic backscattering and photon redistribution by VRS at the sea surface. This model has been incorporated into an atmospheric model, SCIATRAN (Rozanov et al., 1997, 2005), by passing the reflectance function for a given chlorophyll concentration to the atmospheric model which computes the radiance or flux for a given measurement geometry and atmospheric scenario. This hybrid model version will henceforth be referred to as SCIATRAN-OC (OC: Ocean).

Apparent optical properties such as the reflectance require the knowledge of inherent optical properties (IOP) of the water body. With the exception of the VRS (back) scattering coefficient which is determined through a simple exponential expression both IOPs, the total absorption coefficient \( a_t \) and the total backscattering coefficient \( b_t \) will depend solely on chlorophyll concentration [mg/m\(^3\)], i.e. we consider case-1 waters (Morel and Prieur, 1977). It should however be noted that measurements and corresponding parametrizations of IOPs in the UV are rare and often unreliable (Vassilkov et al., 2002b). Accordingly, errors in the IOPs will translate into the determination of the VRS reference spectra.

For the determination of \( a_t \) two pure water absorption spectra from Buiteveld et al. (1994) and Quickenden and Irvin (1980) have been used. In the UV-overlap region both spectra were smoothly spline-interpolated.

The (specific) phytoplankton absorption is taken from Bracher and Tilzer (2001). These spectra were derived from in situ measurements of particulate absorption between 300 and 750 nm on water samples from different biogeochemical provinces of the South Atlantic Ocean which were afterwards corrected for detritus absorption and normalized to chlorophyll concentration according to the method developed by Yentsch (1962) and modified by Bricaud and Stramski (1990) using the \( \beta \)-correction of Mitchell and Kiefer (1988).

Current operational schemes for determining phytoplankton biomass from ocean color data use a single bio-optical model. For instance OC4V4 by O'Reilly et al. (2000) is currently used to generate global maps of chlorophyll-a from both SeaWiFS and
MODIS imagery. Global remote sensing algorithms, such as version four of the NASA algorithm (OC4) are designed for use at global scales and are found to be less accurate at local and regional scales. This is because the simplified parameterization of seawater composition in terms of chlorophyll-a concentration alone does not account for much of the optical variability observed in natural waters. This is in case-1 waters mainly caused by the variation of the specific phytoplankton absorption among species, but also within the same species depending on the photoacclimative state due to differences in packaging (package effect) and the composition of photosynthetic pigments (see e.g. Sathyendranath et al., 1987; Hoepffner and Sathyendranath, 1991; Bracher and Tilzer, 2001; Ciotti et al., 2002).

For the determination of $b_i$ the pure sea water scattering coefficients $b_w$ were taken from (Smith and Baker, 1981) and used within the bio-optical model for the elastic backscattering coefficient proposed by Morel (1988). The VRS backscattering coefficient is computed following the approach described in Vountas et al. (2003).

Finally, absorption from dissolved organic matter (DOM) is treated according to the model of Morel (1988). Here, the DOM absorption at 440 nm is 20% of the total absorption of pure seawater and particulate matter and an exponential function is used to describe the spectral variation. The spectral slope of DOM is set to the wide-spread value of 0.014 nm$^{-1}$. As can be seen in Fig. 1 DOM absorption changes smoothly with respect to the wavelength within a typical DOAS wavelength window of a size of 30–70 nm used here. Such spectral imprint is safely removed by the fitted polynomial.

For a wavelength range of 300–550 nm relevant scattering and absorption coefficients are shown in Fig. 1.

**Determination of $a(\lambda)$**: As in case of the determination of $\nu(\lambda)$ a phytoplankton absorption spectrum from Bracher and Tilzer (2001) has been used. We expect most reliable DOAS fits in regions where the selected spectrum has been measured (i.e. in the South Atlantic). Even though we will not restrict our DOAS retrievals to this region but perform global fits the results can give valuable and simple information and show the feasibility of the approach.
3 Applications

Prior to application of DOAS using a VRS spectrum and phytoplankton absorption spectrum, additional fit parameters have to be defined. Depending on the target quantity slightly different wavelength windows have been selected: the regular VRS retrieval is performed between 349.5 nm and 382 nm, where the spectral structure of VRS can easily be distinguished from other spectral components (Vountas et al., 2003). First tests of Sierk et al. (2004) could show that fitting of phytoplankton absorption spectra performs well in another wavelength window located at about 428–496 nm, where photosynthetic pigments of phytoplankton have generally their maximum absorption.

In the considered wavelength ranges are weak atmospheric absorbers potentially disturbing the DOAS fit if not accounted for such as ozone, bromine monoxide, nitrogendioxide and the oxygen dimer. These spectral features have been accounted for using the appropriate reference spectra in the DOAS fits.

Additionally (inelastic) rotational Raman scattering at air molecules can fill-in solar Fraunhofer and telluric lines and is able have large impact on DOAS retrievals if not properly accounted for. The effect became known as the Ring effect (after Grainger and Ring, 1962) and can be accounted for by fitting a modelled pseudo-absorber–also called Ring-spectrum, defined as in Vountas et al. (1998).

For the retrievals described in the following, a fixed setup for the VRS and Ring spectrum computations has been used. SCIATRAN-OC computed VRS (Vountas et al., 2003) and Ring spectra (Vountas et al., 1998) with a fixed chlorophyll concentration of 0.1 mg/m$^3$ at a solar zenith angle of 30°. A maritime aerosol setup with moderate visibility has been used. Trace gas and temperature profiles were taken from a climatology (Brühl and Crutzen, 1992$^1$) for 25° southern latitude.

Small wavelength shifts due to tiny differences in the wavelength scale used in the VRS, Ring, cross-sections, SCIAMACHY irradiance and radiance spectra are corrected

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by allowing a non-linear shift-and-squeeze of the wavelength axis for each spectrum relative to an arbitrary reference, here the SCIAMACHY radiance.

One important prerequisite is that the groundpixel has to be cloud-free. A very simple but effective approach has been used to identify clear sky pixels: the measured radiance is integrated over the wavelength window and normalized to the number of wavelengths. The resulting value is compared to an empirical threshold value determined beforehand. The underlying physical principle goes back to increased backscatter over cloudy scenes leading to significantly larger radiance values. This approach helps masking cloudy pixels fast and effective but is very sensitive to the threshold used. In this study we have fixed the threshold for all cases conservatively low in order to reject even partially cloudy pixels.

As the spatial resolution of SCIAMACHY is rather poor compared to specrally lower resolving imagers such as SeaWiFS, MERIS or MODIS larger periods of data have to be investigated. For the global maps shown here one month of cloud-free SCIAMACHY nadir data served as a basis. Depending on the data density on overcast situation shorter periods of about two weeks are possible.

3.1 Regional VRS-fitting

The above described retrieval technique has been applied to several thousand SCIAMACHY groundpixels but as a first step a study on a regional scale has been performed to identify the spectral signature of VRS, \( \nu(\lambda) \), in SCIAMACHY data. As an example orbit 12429 measured on 16 July 2004 has been selected. In Fig. 2 the fit factor \( S_\nu \) of a VRS reference spectrum modelled for a chlorophyll concentration of 0.1 mg/m\(^3\) is shown in a color-coded way.

An obvious feature is the water-land contrast: pixels over land are significantly down-scaled to values of around zero. This indicates, as expected, that the VRS spectral signature cannot be found in land pixels. For coastal zone pixels, where water and land has been in the field of view of the sensor the fit factor is \textit{diluted} to values around \(-0.3\). Interestingly some pure land pixels still have fit factors different from zero. We suspect
uncompensated spectral Ring effect structures to be the reason for this. This mismatch could be reduced when modelling the Ring reference spectrum for the adequate scene albedo. The sign of the fit factors is negative. This fact is a consequence of our DOAS implementation, defining ln(I/I_0)=τ and not like the physical law ln(I_0/I)=τ (compare with Eq. 1). Changing the sign of the resulting fit factor after retrieval is uncritical but has not been performed.

The consistent picture persists if the spectral fit results over land and open water are compared. A widespread approach testing the fit quality for a specific target species using DOAS retrievals is to plot the scaled reference spectrum and overplot the scaled reference spectrum with the added fit residual. This approach simplifies the spectral and quantitative comparison of the scaled reference and the residual. Figure 2 (upper right panel) shows the spectral fit results for an open water retrieval of the selected orbit in the Mediterranean sea. Obviously the VRS reference spectrum fits well and is clearly uncorrelated to the residual. Expected opposite behaviour is seen for results over land (lower right panel): the scaled VRS spectrum is almost zero which indicates that no spectral signature of VRS could be retrieved.

Yet an important issue is that over open water the VRS reference spectrum is not adequate for every groundpixel but has a fit factor different from (negative) unity. The reason for this can easily be understood if the relation between Fraunhofer line filling and chlorophyll concentration is recapitulated. Figure 3 shows the modelled dependence of the in-filling at 397 nm (a Calcium Fraunhofer line) and the chlorophyll concentration. With increasing chlorophyll concentration the filling in for this line decreases rapidly. The same mechanism can be seen in Fig. 2. Here, a fit factor larger than −1 (tending to zero) means that the modelled VRS reference spectrum provides more filling-in than necessary for the SCIAMACHY spectrum currently evaluated. As a consequence the reference has to be scaled down in order to fit to the effective situation during measurement. A lower net-in-filling corresponds to a scene with higher chlorophyll concentration, according to Fig. 3. The VRS spectrum has to be downscaled in practically all cases which indicates that the chlorophyll content for the particular ground
pixels is larger than the one related to the modelled VRS spectrum (therefore the actual chlorophyll concentration can be considered larger than \(0.1 \text{ mg/m}^3\)).

3.2 Global VRS-fitting

Applying the above described technique to one month of SCIAMACHY nadir data leads to a dense global map of VRS fit factors. July 2005 has been selected and 999 Orbits have been evaluated. As in the regional study a clear water-land contrast can be shown globally (for clarity land retrievals are not depicted).

Regions with practically no scaling for VRS (fit factors near \(-1\)) correspond to well known large fields of oligotrophic waters. The reason for this has been described above.

Low overall \(\chi^2\) values can be found and indicate good global fit quality. Figure 4 shows the map of VRS fit factors. The global fits have been performed with a solar zenith angle limit of 60° due to signal to noise issues. However, in the southern hemisphere this still seems to be insufficient because of very low fit factors for VRS over the whole range of longitudes. The reason for this is still not understood.

Not shown are the fit factors for the Ring reference. They show no systematic and significant scaling in oligotrophic waters. Therefore the Ring effect does not induce artificial spectral structures that are erroneously compensated by VRS. Interesting is that the Ring fit factor shows a clear latitudinal variation, arising from strong solar zenith angle dependence and is significantly smaller over mountain areas which is in full agreement with our model.

3.3 Regional phytoplankton-fitting

The next step has been to fit phytoplankton spectra, \(a(\lambda)\), to SCIAMACHY optical depths in the wavelength window of 428–496 nm taking into account atmospheric absorption and pseudo absorbers (VRS and Ring reference spectrum).

For our retrieval of phytoplankton distribution from SCIAMACHY data we used sev-
eral (23) specific absorption spectra derived from surface water samples (from Bracher and Tilzer, 2001) in the DOAS fit. As for the current operational chlorophyll-a algorithms used in case-1 waters by MODIS, SeaWIFS and MERIS, we chose one single specific absorption spectrum as standard for the DOAS-fit with SCIAMACHY data in order to obtain distributions of phytoplankton: Our investigations showed that global distributions of phytoplankton absorption retrieved from SCIAMACHY data compares very well to global distributions derived from MODIS operational chlorophyll-a level-2 data (see below) when a specific absorption spectrum measured at surface water samples from the Antarctic Circumpolar Current was used. Here, the phytoplankton was characterized by low biomass (<0.5 mg chl-a/m^3), high specific phytoplankton absorption, a mixed phytoplankton composition with almost all within the size class smaller than 20 µm and where diatoms, dinoflagellates, prymnesiophytes and chrysophytes contributed more or less equally to the overall biomass.

Figure 5 depicts fit factors of a phytoplankton spectrum over an upwelling region near the African West coast. The figure shows clear indication of increased chlorophyll concentration near the upwelling. Here, the phytoplankton absorption spectrum is strongly scaled (color-coded in green and red). Both plots on the right hand side show fit results (so-called differential optical depths) for phytoplankton absorption near the upwelling (upper right panel) and over the open oligotrophic ocean (lower right panel). As expected, low scaling can be observed for oligotrophic and strong for eutrophic waters.

3.4 Global phytoplankton-fitting

An example for a global fit of chlorophyll created for the whole month of July 2005 from SCIAMACHY nadir data is shown in Fig. 6.

The map shows that increased fit factors for the phytoplankton reference spectrum clearly correlate to high chlorophyll concentrations (as for example seen in the corresponding MODIS global chlorophyll concentration map in Fig. 7). Clearly this approach is not only valid on a regional scale, but shows consistent global results.

Improved DOAS fits can be expected when pre-defining ocean provinces and using
appropriate phytoplankton absorption spectra which will be performed in a successive study.

4 Quantitative assessment

4.1 $S_a$ vs. chlorophyll concentrations

We performed a large quantitative comparison between MODIS OC4V4-Aqua chlorophyll-a data ($\approx 1 \text{ km}^2$) with results derived from SCIAMACHY data using DOAS-fitting of phytoplankton absorption. Collocations were chosen with comparing the mean of all MODIS data measured within a SCIAMACHY pixel at the same day. In order to avoid artifacts, only comparisons were made when at least 10 MODIS pixel were collocated to a SCIAMACHY observation.

Figure 8 (left) shows a scatter plot for all collocations in July 2005. Results show a “gamma”-shaped ($\Gamma$) relation between the DOAS fit factor and the MODIS chlorophyll concentration. A simple exponential curve could be fitted according to the equation $f(\phi) = a_0 - a_1 \exp(-a_2 \cdot \phi)$, with $\phi$ being the chlorophyll concentration as obtained from MODIS. The scatter around the fitted curve is still not fully understood and might partially be attributed to spatial and temporal variations of chlorophyll-concentrations.

The curve shows a “saturation” of the SCIAMACHY fit factors for MODIS chlorophyll concentrations higher than 0.5 mg/m$^3$. Obviously the input radiance spectra from SCIAMACHY are not changing significantly with higher chlorophyll concentrations which is likely to be related to the selected wavelength interval where the penetration depth of light is not sufficient and should be adapted in further studies.

4.2 $S_v$ vs. chlorophyll concentrations

MODIS chlorophyll-concentrations have also been compared to VRS-fitting results from SCIAMACHY. The procedure was analogous to the one described above. A comparable but mirrored “gamma”-shape relation between the DOAS fit factor for VRS and...
the MODIS chlorophyll concentration could be found (see Fig. 8, right). The functional
dependence was again as for \( f(\phi) \) (see above), with \( \phi \) being the chlorophyll concentra-
tion as obtained from MODIS.

The absolute value of the VRS fit factor decreases with increasing MODIS chloro-
phyll concentrations, which is expected behavior. A similar “saturation” effect as in the
comparison of chlorophyll concentrations and chlorophyll fit factors can be observed.

4.3 Chlorophyll concentrations

Both approaches, fitting VRS or phytoplankton absorption spectra, exhibit expected
clear correlations to chlorophyll concentrations. Obviously both methods could be used
to retrieve this quantity. An optimized retrieval technique would take advantage of a
hybrid approach: VRS fits strongly in regions where low chlorophyll concentrations
prevail whereas direct fitting of phytoplankton behaves vice versa. One method could
potentially stabilize the other.

The fit factor \( S_a \) for phytoplankton (or to be precise for the specific absorption spec-
trum) is given in \([\text{mg/m}^2]\) which is a mass column. If the penetration depth \( \delta \) of light
for the wavelength window considered is known this column can be converted into a
concentration by the ratio:

\[
C = \frac{S_a}{\delta}
\]  

(3)

VRS is strongly related to \( \delta \) and serves therefore as a proxy: A single vibrational Ra-
man scattering event is always accompanied by an elastic scattering process. There-
fore, the fit factor of VRS, \( S_v \), is directly related to the same quantity for elastic scatter-
ing only. As described above (and in more detail in Vountas et al., 2003) a bio-optical
model from Morel (1988) has been used to describe the dependence of the elastic
backscattering coefficient \( b_b \).

The backscattering coefficient scaled with the same factor as the VRS spectrum
(\( S_v \)) can be understood as the true \( b_b \) for the real situation considered. As \( b_b^{-1} \) is the
modelled penetration depth, \( S_v \ast b_b^{-1} \) can be associated with the measured one. For
the whole retrievals of $S_a$ for July 2005 (as described above) all corresponding values of $S_v$ have been used within Eq. (3) to model a global map of chlorophyll concentrations $C$, which is displayed in Fig. 9. A first visual comparison with MODIS chlorophyll concentrations (Fig. 7) shows good agreement. Further studies will optimize this approach and will also include a thorough quantitative comparison. An important point of investigation must be the impact of the “saturation effects” (shown in Sect. 4.1) on the chlorophyll concentration conversions. The patchy appearance of the mapped chlorophyll concentrations is related to the fact that due to fitting failures, either in case of VRS, or phytoplankton fitting, not both corresponding quantities could be related.

However, there are alternative ways to use hyperspectral “imagery” data to retrieve chlorophyll concentrations. For instance, fitting a liquid water spectrum also has been tested with interesting results: first a stand-alone chlorophyll concentration retrieval from phytoplankton-fitting has been tested. One advantage of the DOAS approach retrieving phytoplankton absorption is that in the wavelength range considered the effective liquid water path can be determined. As the fit factor of the liquid water spectrum $S_{lw}$ represents the number of water molecules encountered by photons along the average propagation path the under-water light path, $L_{wp}$ can easily be estimated (assuming constant water density near surface) from:

$$L_{wp} = \frac{S_{lw}}{r_{lw}}$$  \hspace{1cm} (4)

with $r_{lw}$ being the density of liquid water. Assuming a constant chlorophyll concentration along the identical average light path yields:

$$r_{chl} = \frac{S_{chl}}{L_{wp}}$$  \hspace{1cm} (5)

First tests (Sierk et al., 2004) showed promising results but this approach has still to be used in global retrievals and must be validated.

Yet another promising stand-alone test has been performed for VRS-fitting. Here we have calibrated VRS fit factors ($S_v$) with MODIS chlorophyll concentrations ($C$), as
given in Fig. 8. Using the exponential dependence between $S_v$ and $C$ we applied this function to one month of SCIAMACHY VRS fit factors. The validation will be part of a subsequent study. First comparisons of SCIAMACHY’s and MODIS’ chlorophyll concentrations show good agreement. However, this approach is strongly dependent on the quality of MODIS data. Problems with MODIS’ chlorophyll concentrations significantly interfere with such a retrieval.

5 Conclusions

For the first time the DOAS retrieval method has been used fitting VRS reference spectra to optical depths measured from SCIAMACHY. We clearly have found spectral structures of VRS in SCIAMACHY measurements. Furthermore, a novel approach has been described to fit phytoplankton absorption spectra within DOAS.

Both fitting VRS or phytoplankton reference spectra lead to consistent results. VRS fit factors clearly correlate with chlorophyll concentrations. Low factors correspond to high chlorophyll concentration and vice versa. As expected opposite behavior is observed when fitting phytoplankton spectra: Here, low factors correspond to low chlorophyll concentration and vice versa. Both approaches are promising tools for retrievals of chlorophyll concentration. First results of a hybrid approach show good agreement with chlorophyll concentrations derived from MODIS and confirm the feasibility of the approach. However, more fine-tuning of the retrievals will be necessary. A more in-depth validation result will be shown in a consecutive study.

Despite the spatial resolution of SCIAMACHY being far lower than for the ocean color imagers such as MODIS, SeaWIFS, MERIS, MOS, etc. further developments of our method to identify phytoplankton absorption in satellite spectra are of great relevance for improving chlorophyll concentration determinations for ocean color imagers. So far it has become clear that remote sensing algorithms derived for ocean color imagers are designed for use at global scales but less accurate at regional and local scales (Sathyendranath et al., 2004). Phytoplankton and associated particulate and dissolved
material absorb and scatter light, and their collective influence dominates the optical variability of the water column, at least in the open ocean waters generally referred to as case-1 waters. Phytoplankton absorption varies between species and phytoplankton groups but also in dependence to their physiological states (see Sathyendranath et al., 1987; Mitchell and Kiefer, 1988; Babin et al., 1993; Bracher and Tilzer, 2001). Therefore, errors arise in the common ocean color retrievals because the corrected water-leaving radiance can vary independently of changes in chlorophyll concentrations, as in dependence to the specific phytoplankton absorption. In our study we sought to use the high resolution spectra of the satellite instrument SCIAMACHY in order to look for phytoplankton absorption in open ocean areas. These retrievals could be used to improve chlorophyll retrievals from common ocean color sensors by using the right regional phytoplankton spectrum, which will improve modelling primary production and food web dynamics in the global open oceans. In addition, also retrievals of atmospheric constituents from backscatter UV satellite measurements will improve when the ocean optical signals, such as phytoplankton absorption are also considered in radiative transfer models used in these retrievals.

Further analyses are planned to check whether discrimination between different phytoplankton groups dominating the chlorophyll concentration is possible since the DOAS method is highly sensitive to spectral structures. If this parameter can be retrieved from remote sensing data, further improvements for regional and global marine chlorophyll and marine primary production estimates are expected.

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Fig. 1. Absorption coefficients for phytoplankton (Bracher and Tilzer, 2001) (assuming a chlorophyll concentration of 0.1 mg/m$^3$), DOM (Morel, 1988), pure water (Buiteveld et al., 1994; Quickenden and Irvin, 1980) and total backscattering (Smith and Baker, 1981) in [1/m], as well as a VRS spectrum [without units] as defined in Eq. (2). Each of both dominant peaks in the VRS spectrum correspond to two strong CaII Fraunhofer lines in the solar spectrum. The values between 0.04 – 0.05 can be interpreted as filling-in of 4 – 5 % w.r.t to non-filled lines.
Fig. 2. Left: VRS fit factor for SCIAMACHY orbit 12429. Right: VRS fit for readout 2 of state 5 over open water (above) and readout 253 of state 5 over land (below).
Fig. 3. Chlorophyll concentration vs. VRS filling-in at 397 nm.
Fig. 4. Global VRS fit factors from SCIAMACHY data for July 2005.
Fig. 5. Left: Phytoplankton fit factor for SCIAMACHY orbit 17712. Right: Phytoplankton fit for readout 214 of state 7 over upwelling (above) and readout 6 of state 7 over oligotrophic water (below).
Fig. 6. Global chlorophyll fit factors from SCIAMACHY data for July 2005.
Fig. 7. Global chlorophyll concentration from MODIS for July 2005.
Fig. 8. Scatter plots, left: MODIS chlorophyll-a concentration vs. SCIAMACHY chlorophyll fitting factors for collocations in July 2005. Right: MODIS chlorophyll-a concentration vs. SCIAMACHY VRS fit factors for collocations in July 2005.
Fig. 9. Global chlorophyll concentration from SCIAMACHY for July 2005.