Inherent optical properties of dissolved and particulate matter in an Arctic fjord (Storfjorden, Svalbard) in early summer

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Abstract. There have been considerable efforts to understand the hydrography of Storfjorden (Svalbard). A recurring winter polynya with large sea ice production makes it an important region of dense water formation at the scale of the Arctic Ocean. In addition, this fjord is seasonally influenced by freshwater inputs from sea ice melt and the surrounding islands of the Svalbard archipelago, which impacts the hydrography. However, the understanding of factors controlling the optical properties of the waters in Storfjorden are lacking and are crucial for the development of more accurate regional bio-optical models. Here, we present results from the first detailed optical field survey of Storfjorden conducted in early summer of 2020. Our observations are based on spectrometric analysis of water samples and in situ vertical profiles with an absorption and attenuation meter, a fluorometer, and a conductivity, temperature, and depth (CTD) sensor. In addition to the expected seasonal contribution from phytoplankton, we find that in early summer waters in Storfjorden are optically complex with a significant contribution from coloured dissolved organic matter (CDOM, 33%–64% of the non-water absorption at 443 nm) despite relatively low CDOM concentrations and in the nearshore or near the seabed from non-algal particles (up to 61% of the non-water absorption at 550 nm). In surface waters, the spatial variability of light attenuation was mainly controlled by inorganic suspended matter originating from river runoff. A distinct subsurface maximum of light attenuation was largely driven by a subsurface phytoplankton bloom, controlled by stratification resulting from sea ice melt. Lastly, the cold dense bottom waters of Storfjorden from winter sea ice production periodically overflows the sill at the mouth of the fjord and can thus reach the Fram Strait. It contained elevated levels of both non-algal particles and dissolved organic matter, which are likely caused by the dense flows of the nepheloid layer interacting with the sea bed.

1 Introduction

Located in the northern Barents Sea, Storfjorden, the largest fjord in the Svalbard archipelago, is influenced by two major water masses (Skogseth et al., 2005b), namely warm Atlantic water (AW) from the Norwegian Atlantic Current and cold Arctic water (ArW), which is AW-cooled along the traverse around Svalbard, entering the fjord from the north and east with the East Spitsbergen Current (ESC). The recurring winter polynya in Storfjorden (Skogseth et al., 2005a) is considered an important “ice factory” and winter sea ice formation here supports the production of about 5%–10% of the dense water formed in the whole Arctic Ocean (Smedsrud et al., 2006). This is related to subsequent release of salty brine from sea ice to the underlying relatively shallow waters, which creates favourable conditions for the formation of cold, salty brine-enriched shelf waters (BSWs) of high density (Skogseth et al., 2005b). While these phenomena are well documented, mainly from hydrographical surveys, knowledge of the inherent optical properties (IOPs) of waters...
in Storfjorden is lacking, which holds back the development of regional bio-optical models.

At a larger scale, over the Arctic Ocean, the largest recent reduction in the extent of sea ice in winter has been found to take place in the Barents Sea (Onarheim et al., 2018). While typically the Barents Sea is completely ice-free in summer and early to late autumn, a reduced winter/spring ice cover can have a significant impact on the light climate and the productivity of associated ice algae and phytoplankton (Leu et al., 2011). Recent studies have shown a northward progression of the polar front with associated changes to the seasonal plankton bloom and the sea ice reduction in the Barents Sea (Neukermans et al., 2018; Oziel et al., 2017). As sea ice continues to retreat, understanding the evolution of the optical properties of the water column, and in particular IOPs, is crucial for a better understanding of the potential changes in phytoplankton dynamics.

The two complementary ways of getting IOP observations in oceanic waters are (i) passive multispectral satellite imagery in the visible, often referred to as “ocean colour” imagery and (ii) field-based measurements. While the former enables improved spatial and temporal coverage of the IOP dynamics, it is especially difficult to operate in polar regions (International Ocean Colour Coordinating Group (IOCCG), 2015). Despite their limited availability, field observations thus remain the main source of information regarding IOPs in the Arctic. They also constitute a unique way of capturing the complex vertical structure of the optical properties in Arctic waters, the knowledge of which is essential for the remote sensing of stratified waters (Lee et al., 2020). In addition, increasing the number of observations through new surveys is critical for strengthening the statistical characterization of the parameters used when building regional bio-optical models (Kostakis et al., 2020), both for calibration and validation of ocean colour satellite products (Orkney et al., 2020) and for getting an accurate representation of optical properties in numerical models.

While we are not aware of any existing optical surveys in Storfjorden, there have been a few studies examining the optical characteristics of waters in the Barents Sea proper (Aas and Berge, 1976; Aas and Høkedal, 1996; Falk-Petersen et al., 2000; Hancke et al., 2014; Hovland et al., 2014; Kostakis et al., 2020; Orkney et al., 2020). Further, a number of other studies have been conducted in the region of AW inflow west and north of Spitsbergen (Kowalczyk et al., 2019; Makarewicz et al., 2018; Pavlov et al., 2015, 2017). All these studies point toward a significant contribution of phytoplankton to light attenuation in the open Barents Sea both north and south of the polar front, since the concentration of coloured dissolved organic matter (CDOM) in water masses of AW origin is relatively low, compared to “true” polar waters with a distinct higher CDOM signal (Pavlov et al., 2015). The cold ArW entering the Barents Sea from the north (Loeng, 1991), and located north of the polar front, is also low-CDOM water (Hancke et al., 2014), likely modi-
nent sill (sill depth about 120 m while the bottom depth of the deepest part of the inner fjord is ~180 m). In addition, a section at the latitude of the sill was conducted across the fjord. Optical measurements and sampling were undertaken at the stations shown in Fig. 1, while extra hydrographical casts were conducted in addition to these stations (positions not shown). This study only focuses on the south–north transect for the sake of brevity, but the entire optical dataset is available (Petit et al., 2022).

2.2 In situ measurements and sampling procedures

Vertical profiles of IOPs (absorption $a$ and attenuation $c$), fluorescence, together with conductivity (salinity, $S_p$, in practical salinity scale), temperature, and depth (pressure) were measured at all stations from the surface down to close to the bottom using an instrument package consisting of a 25 cm pathlength attenuation and absorption meter (ac-s, WET Labs Inc., USA), a fluorometer measuring fluorescence by dissolved organic matter (WetStar FDOM, Wet Labs Inc., USA), and an SBE37 SIP conductivity–temperature–depth (CTD) probe (Sea-Bird Electronics, USA). All these instruments were calibrated in the manufacturer in 2018 and had not been used since then. The data from all three instruments were stored by a DH4 data logger (WET Labs Inc., USA). Since there was no real-time pressure reading from the optical instrument package, the maximum depth for each cast had to be estimated using wire length. Thus, the actual distance to the bottom (from echo depth from the ship’s echo sounder) varied from cast to cast. The battery failed at the two northernmost stations, and only one profile down to half the bottom depth could be achieved at station 513.

Total absorption ($a$) and attenuation ($c$) were obtained from the two tubes of the ac-s with a sampling frequency of 4 Hz. The processing steps involved (i) blank subtraction, (ii) salinity and temperature correction, and (iii) scattering correction of the absorption and attenuation channels. The blanks were measured by injecting ultra-pure water successively inside each channel using a peristaltic pump with constant flow of 0.6 L min$^{-1}$. This was done once before and three times during the cruise to track any potential drift of the instrument. No evidence for any drift was found but we experienced more variability of the blank performed during the cruise and thus decided to use the blanks measured in the laboratory before the cruise in all the data processing. Salinity and temperature values were taken from the CTD data, which had a sampling frequency of ~0.5 Hz, and the correction was applied following Van Zee et al. (2005). Scattering correction was performed using the flat$^2$ method proposed by Röttgers et al. (2013). This correction assumes spectral invariance of the volume scattering function (VSF). It makes use of an empirical law for estimating the true absorption at 715 nm, from the 715 nm a channel of the ac-s. This value is then used to correct the whole spectrum. The rationale behind the use of a flat method instead of a proportional one is that with the latter the spectral shapes of the corrected spectra were less realistic and very low absorption spectra resulted in some (unrealistic) negative values.

$\text{FDOM}$ was measured using a WET Labs WETStar fluorometer, with three excitation–emission pairs as follows: channel 1 (FDOM1, 310/450 nm), which represents marine ultraviolet humic-like and marine humic-like material; channel 2 (FDOM2, 280/450 nm), which represents terrestrial humic-like material; and channel 3 (FDOM3, 280/350 nm) representing protein-like tryptophan-type material resulting from the presence of phytoplankton (for details, see Makarewicz et al., 2018). Fluorescence intensities acquired from the WETStar fluorometer are reported here in background-corrected raw counts (RCs). The background values were estimated in a lab with a dark environment by creating a flow of ultra-pure water into the sensor tube with a peristaltic pump.

Water was collected on a separate cast using a rosette with seven 4 L Niskin bottles with an SBE 19plus CTD (Sea-Bird Electronics, USA) giving real-time information about depth, temperature, and salinity. Water samples were collected to determine oxygen isotope ratio ($\delta^{18}$O), CDOM, and particulate matter (total, algal, and non-algal) absorption. The samples were collected at 10, 20, and 50 m and near bottom depths.

Water samples for CDOM were collected by gravity filtration through pre-rinsed 0.2 µm Millipore Opticap XL filter capsules connected to the Niskin with silicon tubing as described in Makarewicz et al. (2018). The samples were stored in 20 mL pre-combusted amber glass vials in the dark at $+4^\circ$C until analysis. CDOM samples were analysed on board within days of collection.

Samples for particulate absorption measurements were collected on 25 mm filters (Whatman GF/F, nominal pore size 0.7 µm) as described by Röttgers and Gehnke (2012). Special care was taken to keep a low vacuum during filtration to avoid algae cell breakage. The filtered volume was adjusted systematically following two criteria: (i) colour of the filter and (ii) filtering speed for anticipating potential clogging. Samples were immediately flash-frozen at $-196^\circ$C and stored in liquid nitrogen during the cruise and were later stored at $-80^\circ$C until analysis.

Samples for $\delta^{18}$O were collected into 20 mL polyethylene bottles, filled completely to avoid any headspace, and closed carefully, and the caps were sealed with parafilm. Samples were shipped to the Jan Veizer Stable Isotope Laboratory (University of Ottawa, Canada) for determination of the oxygen isotope ratio on a Finnigan MAT Delta plus XP + Gasbench. A precise amount of 0.6 mL of water was pipetted into Exetainer vials. No catalyst was required. The vials were flushed and filled with a gas mixture of 2% CO$_2$ in helium. The flushed vials were left at room temperature for a minimum of 5 d. The CO$_2$ gas was analysed automatically in continuous flow. The results were normalized to VS-MOW (Vienna Standard Mean Ocean Water) standard using https://doi.org/10.5194/os-18-455-2022
three calibrated internal standards spanning most of the natural range. The precision of the analysis was ±0.15 ‰.

2.3 Laboratory methods

Spectral CDOM absorption coefficients were measured on board using a liquid waveguide capillary cell (LWCC) system following the IOCCG protocol (Mannino et al., 2019). This system included a DH-mini deuterium/tungsten source (Ocean Optics), a 1 m long LWCC, and a Flame-T spectrometer covering the UV–NIR (250–800 nm). The LWCC was cleaned with methanol at the beginning and end of each day. For each sample, intensity measurements in digital counts of the dark current \( I_{DC} \), reference purified water \( I_{ref} \), and sample water \( I_S \) were performed successively, and this was repeated three times. The apparent absorption coefficient (in m\(^{-1}\)) was then determined at each wavelength by averaging the three \( a_{app} \) spectra, each computed as follows:

\[
a_{app}(\lambda) = \frac{-\ln [(I_S(\lambda) - I_{DC}(\lambda)) / (I_{ref}(\lambda) - I_{DC}(\lambda))]}{l},
\]

with \( l = 1.00 \text{m} \) as the optical path length. The apparent absorption coefficient of a 100 mg L\(^{-1}\) pre-burnt HPLC-grade NaCl solution \( a_{NaCl} \) (in m\(^{-1}\)) was measured with the same protocol. The salinity correction applied was as follows:

\[
a(\lambda) = a_{app}(\lambda) - \frac{a_{NaCl}(\lambda)}{91} \cdot S_p,
\]

with \( S_p \) the practical salinity of the sample. It should be noted that the salinity-normalized apparent NaCl absorption was obtained by dividing \( a_{NaCl} \) by 91 instead of 100 for taking into account the fact that seawater comprises salts other than NaCl. This number has been determined empirically (personal communication with Rüdiger Röttgers, 2020).

The particulate absorption coefficient was quantified for each sample filter by two independent methods of the quantitative filter techniques (QFTs) using integrating spheres large enough for placing the filters inside them. The first one (Röttgers and Gehnke, 2012, hereafter QFT-Perkin) uses a commercial laboratory UV–Vis–NIR spectrophotometer (Lambda 950, Perkin Elmer, USA), while the latter (Röttgers et al., 2016, hereafter QFT-ICAM) uses a custom-made portable integrating cavity absorption meter. In both cases, the optical density of the sample filters, OD\(_S\), was measured against the optical density of a wet reference blank filter OD\(_{ref}\), taking a dry filter as the reference in both cases. The particulate absorption \( a_p \) was then determined as

\[
a_p(\lambda) = \frac{(OD_S(\lambda) - OD_{ref}(\lambda)) \cdot A \cdot \beta}{V},
\]

where \( A \) (m\(^2\)) is the filter patch area, \( \beta = 4.5 \) (Röttgers and Gehnke, 2012) is the path length amplification factor, and \( V \) (m\(^3\)) the volume of water filtered. The particulate absorption coefficient was measured between 350–750 and 390–850 nm with the QFT-Perkin and QFT-ICAM, respectively. The custom-made QFT-ICAM showed very good agreement.
with the QFT-Perkin method (Fig. 2). Non-algal particle (NAP) absorption $a_{\text{NAP}}$ (m$^{-1}$) was measured with the QFT-ICAM method using the same filters after bleaching them with a 1% NaOCl solution for 1–3 min. The bleach was removed by oxidation using a 10% H$_2$O$_2$ solution, with the filters stored and measured 1 d later. The phytoplankton absorption $a_{\text{phy}}$ (in m$^{-1}$) was computed as the difference between total and NAP absorption ($a_{\text{phy}}(\lambda) = a_p(\lambda) - a_{\text{NAP}}(\lambda)$).

2.4 Satellite data

Clear-sky conditions prevailed during the days of the cruise, and thus, as background information to this study, we included a qualitative analysis of a Sentinel-2 image taken on 22 June 2020 over Storfjorden. The Sentinel-2 mission, launched by the European Space Agency (ESA) under the Copernicus programme, is a constellation of two satellites launched in 2015 and 2017, respectively. Each of them encompasses a multispectral imager (MSI) capturing light in 13 spectral bands from 443 nm in the visible to 2190 nm in the short-wave infrared. Compared to existing satellite missions like Sentinel-3 OLCI, it has a higher spatial resolution (10–60 m depending on the band), which makes it suitable for catching the high spatial variability typically encountered in coastal environments.

The Sentinel-2 satellite data were downloaded from the Copernicus Open Access Hub via the sentinel2sat python package in geo-corrected top-of-the-atmosphere (TOA) spectral radiance (L1C product). An atmospheric correction algorithm based on Acolite (Vanhellemont and Rudnick, 2016) was used for converting the TOA data into a bottom-of-the-atmosphere (BOA) reflectance. The algorithm was implemented using the python package Py6S (Wilson, 2013). This solution uses the state-of-the-art 6S vector radiative transfer model (Vermote et al., 1997) for estimating and removing the contribution of atmospheric gases and aerosols to the remote-sensing signal. The gas (aerosol) model was set to subarctic summer (maritime), and a manual value of 0.1 was set for the aerosol optical thickness at 440 nm (aiming at BOA reflectance of around 0 in the near-infrared in areas with low turbidity). There was no swell and no wind at the time of acquisition as well as favourable solar-sensor geometry, and we thus did not have to apply any sunglint removal strategy. The algorithm of Nechad et al. (2010) was used for estimating the turbidity and enhancing the visual analysis of the surface water. It has been shown to be robust to various environmental conditions (Dogliotti et al., 2015). Considering the very low concentration of phytoplankton in surface waters, both noticeable from our observations (see Sect. 3.) and expected due to the stratification-related summer nutrient depletion of the upper layer (Tremblay et al., 2015), we did not apply any chlorophyll $a$ retrieval algorithm to the satellite data.

3 Results and discussion

3.1 Inputs from the shore

An assessment of the impact of terrestrial runoff on the optical properties of surface water on 22 June 2020 was done using Sentinel-2 bottom-of-the-atmosphere reflectance (Fig. 3a) and turbidity (Fig. 3b). As expected from the season (late June) and associated land snow and glacier melt, significant turbidity plumes were observed nearshore with turbidity $> 3.5$ FNU (red colour in Fig. 3b), while it was $< 0.8$ FNU (blue colour in Fig. 3b) in the central parts of the fjord where sampling effort was concentrated. Along the main south–north transect, only the stations 509, 511, and 513, all sampled on 26 June (4 d after the satellite acquisition), seem to be close enough for being potentially impacted by the coastal inputs of particles and coloured dissolved organic matter. This confirms the similar assertion made from the analysis of the $\delta^{18}$O data (see Sect. 3.2).
3.2 Hydrographic setting in Storfjorden in early summer

Section plots of salinity and temperature for the south–north transect are presented in panels a and b of Fig. 4. A surface layer of fresher ($S_p \sim 33$) and warmer ($T > 2^\circ C$) water was found down to about 20 m depth, followed by a strong density gradient between 20–30 m depth. These section plots also highlight very dense and cold ($S_p > 35.2$ and $T < -1.5^\circ C$) bottom waters for the stations north of the sill (at depths > 125 m) but also south of the sill at the bottom. At the southernmost station in Storfjordrenna there are warmer temperatures down to 125 m depth, indicative of warmer AW from the West Spitsbergen Current (WSC) entering the Storfjordrenna. In addition, dense, cold BSW was present at the bottom, likely a result of an overflow of BSW over the sill from Storfjorden proper. $\delta^{18}O$ values (Fig. 4c) were positive and ranged from 0.2‰–1‰, which indicates contribution from sea ice melt at the surface and limited impact from runoff along the main south–north transect in the middle of the fjord (e.g. station 535). Only a few northern stations (stations 509 and 511) with the lowest $\delta^{18}O$ values at the very surface compared to other stations show signs that local runoff could have reached the offshore location of the sampling stations.

This situation appears typical for the fjord in summer (Fossile et al., 2020), with a rather shallow fresher and warmer, well-stratified surface layer present over the entire fjord and near-freezing, salty, brine-enriched waters at the bottom of the fjord. The surface layer was shallower than later in summer (see Fossile et al., 2020), likely because the sea ice had just disappeared, which has limited the time for mixing and deepening of the surface layer. Below sill depth we find evidence for the presence of BSW in the fjord from winter sea ice production.

3.3 Observed inherent optical properties in relation to hydrography

In situ observations of the attenuation coefficient at 440 nm, $c(440)$ (Fig. 5b), which is the sum of absorption and scattering coefficients and varies with CDOM, phytoplankton, and non-algal particle concentration, show how IOPs are related to the stratification and water masses in the fjord and/or to
the inputs from land. The ratio between the scattering and absorption at 440 nm, \( b(440)/a(440) \) (Fig. 5a), gives some indication of the nature of the IOPs, with very high values corresponding to strong scatterers like mineral particles, and lower values obtained when the IOPs are dominated by CDOM or phytoplankton.

There are distinct subsurface maxima of attenuation (\( \sim 1.5 \text{ m}^{-1} \) at 440 nm) with a very high scattering-to-absorption ratio (\( > 11 \)) for the northernmost station 513. This could be linked to land inputs visible in the satellite data (Fig. 3). Indeed, particles coming from land are typically strong scatterers, especially the mineral ones due to their high refractive index. The presence of such particles in the water would then increase the scattering-to-absorption ratio observed in the areas of the fjord influenced by runoff. For most of the other stations we can see distinct subsurface maxima in \( c(440) \) between 25–50 m depth with \( b(440)/a(440) \) showing lower values (5–7). This may be linked to the presence of phytoplankton as phytoplankton cells, which contain a lot of water, are weaker scatterers compared to other types of particles present in seawater. AW in Storfjordrenna and waters at intermediate depth in the fjord show the lowest attenuation (\( \sim 0.25 \text{ m}^{-1} \) at 440 nm), and here also the contribution from scattering is lowest with a scattering-to-absorption ratio of \( \sim 5 \). We found elevated scattering relative to absorption at the very bottom at several stations, mostly next to the sill. Apart from station 513, we also observed widespread highly scattering particles in the surface layer (often with low concentration, as deduced from concomitant low \( c(440) \)). They could potentially result from very small (fine grain size) non-algal particles melted out from sea ice (Bélanger et al., 2013; Granskog et al., 2015) or, as suggested very recently in Davies et al. (2021), by some large non-mineral particles such as fish eggs or zooplankton. The ratio of the scattering to absorption coefficient (Fig. 5a) was generally very high in Storfjorden: there was up to 11 times more scattering than absorption at 440 nm (which corresponds to a single-scattering albedo \( b/c \) of 0.92). A similar increase in scattering was also observed west and north of Spitsbergen in surface layers affected by sea ice melt (Granskog et al., 2015).

### 3.4 Characteristics of CDOM and FDOM

The concentration of CDOM, here quantified in Fig. 6a by its absorption coefficient at 440 nm \( a_{\text{CDOM}}(440) \), shows values within the range 0.02–0.05 m\(^{-1}\), which are levels typical for AW in the WSC (\( a_{\text{CDOM}}(443): 0.016–0.51 \text{ m}^{-1} \); Kowalczuk et al., 2019) and in the Barents Sea proper (\( a_{\text{CDOM}}(443): 0.035–0.162 \text{ m}^{-1} \); Hancke et al., 2014). Although the resolution of water samples was low, some distinct patterns appear. The very surface layer seems to have the lowest CDOM, which is likely linked to dilution from recent melting of low-CDOM sea ice (Granskog et al., 2015; Mascarées and Zielinski, 2019; Kowalczuk et al., 2017). This reveals an opposite trend compared to what is usually found in estuar-
The fluorescence originating from three pairs of excitation–emission gave more insights about the origin of the dissolved organic matter and processes regulating distribution of different dissolved organic matter (DOM) fractions in the water column. As already observed for AW in the WSC (Makarewicz et al., 2018) we found overall low values of humic-like FDOM in Storfjorden (Fig. 7a, b). The FDOM1 and FDOM2 fluorescence intensities were \( \sim 1.5 \) higher at the core of Storfjorden compared to the southernmost part of the transect in the presence of AW. The surface, low-salinity layer was characterized by the lowest FDOM1 and FDOM2, linked to the same dilution-based phenomenon as observed for CDOM, while they showed the highest values near the bottom, especially north of the sill. These high values are likely linked to the accumulation of BSW with humic-like DOM fraction originating from brine release (Müller et al., 2013). On the other hand, the tryptophan-like compound (Fig. 7c) shows a higher dynamic range and distinct patterns with a subsurface maximum linked to the density gradient visible in Fig. 5 (white isolines). This is linked to the subsurface chlorophyll maxima (SCM) that we observed (see Sect. 3.), which are common in stratified Arctic waters (Ardyna et al., 2013). Fluorescence by tryptophan-like compounds was also found by Makarewicz et al. (2018) to be tightly related to chlorophyll concentration in the AW. The absorption line height at 676 nm \((a_{LH}(676), \text{Roesler and Barnard}, 2013)\) was computed from in situ ac-s absorption at 650, 676, and 715 nm as a proxy for phytoplankton biomass. It was computed as follows:

\[
a_{LH}(676) = a(676) - \frac{a(715) - a(650)}{715 - 650} \cdot (676 - 650) - a(650).
\]

The section plot of \(a_{LH}(676)\) (Fig. 8a) confirmed that the subsurface maxima in tryptophan-like FDOM, scattering, and absorption were linked to phytoplankton biomass. The absorption line height at 676 nm \((a_{LH}(676), \text{Roesler and Barnard}, 2013)\) was computed from in situ ac-s absorption at 650, 676, and 715 nm as a proxy for phytoplankton biomass. It was computed as follows:

\[
a_{LH}(676) = a(676) - \frac{a(715) - a(650)}{715 - 650} \cdot (676 - 650) - a(650).
\]
around 50 m depth at station 502) part of the fjord. This sub-
surface layer of increased phytoplankton abundance was in-
terfacing the frontal zone between inflowing AW and ArW.
This is likely due to nutrient limitation in surface waters and
light limitation in the deeper zone.

Sample-based values of $a_{phy}(440)$ and $a_{NAP}(440)$
(Fig. 8b, c) ranged between 0.025–0.2 and 0.008–0.55 m$^{-1}$
respectively. These values and their distribution with depth
were in a similar range as reported in WSC waters by Kowal-
czuk et al. (2017, 2019), further pointing to the presence
of AW. Although the depths of subsurface chlorophyll $a$
maxima were not sampled (since in situ data were not
available to guide the water sampling), the sample-based
phytoplankton absorption at 440 nm (Fig. 8b) indicates
increased absorption by phytoplankton in the subsurface and
at deeper depth in the southern fjord compared to the central
part of the fjord.

The surface waters of Storfjorden were characterized by
very low phytoplankton absorption. This is also clearly vis-
ible from the spectral absorption of particles shown for sta-
tions 501 and 508 (Fig. 9); here the typical absorption peaks
for chlorophyll are evident only at subsurface depths. Slight
increase in surface non-algal particle absorption was only ob-
erved for the northern station 509 and, to a lesser extent, at
station 511 (Fig. 8b). This implies overall limited influence
of absorbing particles from land runoff along the south–north
transect.

Non-algal particle absorption from the water samples
(Fig. 8c) showed elevated values for the bottommost samples
in the BSW (with nearly no phytoplankton absorption), sug-
gest that there are turbid, near-bottom flows dominated
by inorganic material, likely resuspension of bottom sedi-
ments. This increase in $a_{NAP}(440)$ is most likely due to min-
eral particles deposited on the seabed and resuspended due
to water flows typically encountered in the nepheloid layer.
This increase may also partly originate from both incomplete
mineralization of settled phytoplankton particles or be an ef-
fect of the adsorption of the CDOM fraction to resuspended
particles. Yamashita et al. (2021) has described the effect
of enrichment in bottom water flowing over the productive
shelf of the Sea of Okhotsk with allochthonous FDOM con-
tained within sediments pore waters. This effect could also
be present in Storfjorden, supported by our observation of
a significant decline in the local $a_{LH}(676)$ (phytoplankton
biomass proxy) maximum with depth (Fig. 7c).

3.6 Relative contribution of the optically active
constituents

Understanding the relative contribution of the optically ac-
tive constituents, namely phytoplankton, CDOM, and non-
algal particles, in determining the optical characteristics of
seawater is of particular importance for any radiative transfer
modelling or optical remote-sensing application of a specific
area. Considering the optical complexity found in Storfjor-
den, we decided to conduct a separate analysis for the sur-
face (0–15 m) water, subsurface (15–60 m) water, and dense
bottom (< 20 m from the bottom, stations 505, 508, and 509)
waters. In addition, we further separated the surface and sub-
surface into two groups: (i) northern stations 509, 511, and
513, which are potentially land-influenced, and (ii) southern
offshore stations 501, 502, 505, 535, and 508, which are less
influenced by land runoff. For each of the five resulting cases,
an absorption budget (excluding water itself) was calculated
based on the water sample data (Fig. 10) and an attenuation
budget (excluding water itself) was computed based on the
in situ ac-s data (Fig. 11). A corresponding table with aver-
ages of relevant optical properties for each of the five cases
is presented in Table 1.

The non-water IOPs in Storfjorden were driven by non-
algal particles and CDOM for the northern surface water,
by CDOM for the southern surface waters, by phytoplank-
ton in the subsurface waters, and by non-algal particles
and CDOM for the dense bottom water. The variability in attenu-
ation was driven by the variability of scattering (Fig. 11, light

![Figure 8. Section plots of (a) absorption line height at 676 nm ($a_{LH}(676)$) (data from the ac-s), (b) absorption by phytoplankton (data from the water samples), and (c) absorption by non-algal particles (NAPs) (data from the water samples) all at 440 nm. White points indicate sample locations. Plots are made with Ocean Data View (Schlitzer, Reiner, Ocean Data View, https://odv.awi.de/, last access: 30 January 2022).](https://doi.org/10.5194/os-18-455-2022)
Figure 9. Comparison of total particulate $a_p$ (dashed line) and phytoplankton $a_{phy}$ (solid line) absorption in Storfjordrenna at station 501 (a) and in the fjord north of the sill at station 508 (b) with depth.

Table 1. Average values of the main water IOPs for the five cases corresponding to northern surface (NSurf), southern surface (SSurf), northern subsurface (NSub), southern subsurface (SSub), and dense bottom water (Bott). The number of samples $n$ is given for the water-sample-based quantities and does not apply to $b_{440}/a_{440}$, which comes from the ac-s data.

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<th></th>
<th>$n$</th>
<th>$a_{cdom 440}$ (m$^{-1}$)</th>
<th>$S_{cdom 350 550}$ (nm$^{-1}$)</th>
<th>$a_{NAP 440}$ (m$^{-1}$)</th>
<th>$S_{NAP 350 550}$ (nm$^{-1}$)</th>
<th>$a_{phy 440}$ (m$^{-1}$)</th>
<th>$b_{440}/a_{440}$ (No unit)</th>
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<td>0.0148</td>
<td>0.0091</td>
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<td>0.0168</td>
<td>0.0369</td>
<td>0.0067</td>
<td>0.0061</td>
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The green curves confirm the evidence of a subsurface algae bloom with the highest phytoplankton contribution for the subsurface layer (max. 0.063 m$^{-1}$) and the lowest for the bottom water (0.006 m$^{-1}$). The CDOM had the most stable contribution across the five different categories, with average values of $a_{CDOM 440}$ ranging between 0.028–0.043 m$^{-1}$. Relative contributions of the three optically active constituents to the non-water absorption are presented in Table 2 for three specific wavelengths (443, 550, and 670 nm). For phytoplankton, a minimum (maximum) contribution of 6% (82%) was found for the bottom water at 550 nm (the southern subsurface water at 670 nm). The CDOM contribution ranged from 6% for southern subsurface water at 670 nm to 64% for southern surface water at 443 nm. Non-algal particle contribution to the non-water absorption had values ranging between 8% for the southern subsurface water at 443 nm to 70% for bottom water at 670 nm. As it could also be assessed qualitatively from Fig. 10, this also highlights the strong dependence of the absorption budget to the wavelength of the radiation, the CDOM contribution dramatically decreasing with increasing wavelength. The relative CDOM contribution was thus higher than what has been found in the AW in the WSC with an average CDOM contribution of 42% at 412 nm (Kowalczuk et al., 2019) as well as in AW north of Spitsbergen with a CDOM contribution of 43% at 443 nm (Kowalczuk et al., 2017). However, it remains significantly lower than what has been found in the central and eastern Arctic (Gonçalves-Araujo et al., 2018) with 85% of the non-water water absorption attributed to CDOM at 443 nm.

4 Conclusions

This first study of the inherent optical properties (IOPs) in Storfjorden (Svalbard) highlighted its optical complexity that we linked to the variable influence from sea ice melt (summer surface stratification) and formation (dense cold bottom waters), runoff from nearby land, and local phytoplankton productivity. As observed earlier in the Barents Sea proper
Table 2. Relative contributions (in %) of the phytoplankton, CDOM, and non-algal particles (NAPs) to the absorption budgets at 443, 550, and 670 nm for the five cases corresponding to northern surface (NSurf), southern surface (SSurf), northern subsurface (NSub), southern subsurface (SSub), and dense bottom water (Bott). Bold numbers indicate the constituent dominating the absorption for a given case and wavelength.

<table>
<thead>
<tr>
<th></th>
<th>443 nm</th>
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<td>NAP</td>
<td>Phyto</td>
<td>CDOM</td>
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<td>64</td>
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<tr>
<td>NSub</td>
<td>6</td>
<td>33</td>
<td>50</td>
<td>17</td>
<td>29</td>
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<tr>
<td>SSub</td>
<td>10</td>
<td>59</td>
<td>33</td>
<td>8</td>
<td>52</td>
</tr>
<tr>
<td>Bott</td>
<td>3</td>
<td>7</td>
<td>50</td>
<td>43</td>
<td>6</td>
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</tbody>
</table>

Figure 10. Absorption budget for five typical cases, corresponding to (a) northern surface (NSurf), (b) southern surface (SSurf), (c) northern subsurface (NSub), (d) southern subsurface (SSub), and (e) dense bottom water (Bott). Surface (subsurface) water is defined as < 15 m (15–60 m) water depth. The dense bottom water corresponds to the layer trapped by the sill at stations 505, 508, and 509.

Figure 11. Attenuation budget for five typical cases, corresponding to (a) northern surface (NSurf), (b) southern surface (SSurf), (c) northern subsurface (NSub), (d) southern subsurface (SSub), and (e) dense bottom water (Bott). Surface (subsurface) water is defined as < 15 m (15–60 m) water depth. The dense bottom water corresponds to the layer trapped by the sill at stations 505, 508, and 509. Dark blue corresponds to absorption and light blue to scattering.

and Atlantic waters (AWs) west and north of Spitsbergen, the contribution of CDOM to the absorption budget varied quite little. Also, the contribution from humic-like FDOM was low and invariable, which indicated limited terrestrial contribution to the DOM pool in the fjord. Conversely, tryptophan-like FDOM closely followed the spatial and vertical distributions of optical proxies for chlorophyll a concentration. Nevertheless, in surface waters not influenced by land runoff, CDOM was the main contributor to the non-water light absorption, despite its relatively low concentrations in AW, since stratification resulted in a subsurface phytoplankton bloom with a maximum magnitude of between 25–50 m depth. We surmise that in shallow nearshore waters with more direct input from land runoff, the contribution from turbid plumes and non-algal particles is much larger during spring freshet and was partly found in our dataset as an increased and non-negligible level of non-algal particle absorption as well as very high scattering-to-absorption ratios. The dense and cold bottom water, originating from winter sea ice formation and brine rejection, was found to contain higher levels of dissolved organic and non-algal particulate matter. Its transport across the sill needs further attention to better understand the potential implications regarding the bottom waters of the adjacent basins and material transports. The statistics obtained on the different IOPs and on their relative
contribution to the light absorption can be used, in conjunction with the ones from previous studies in nearby areas, for improved regional parameterizations of bio-optical models used in the field of remote sensing or climate modelling.

Appendix A: List and definitions of acronyms and abbreviations

<table>
<thead>
<tr>
<th>Acronym/abbreviation</th>
<th>Definition</th>
</tr>
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<tbody>
<tr>
<td>a</td>
<td>Light absorption in m$^{-1}$</td>
</tr>
<tr>
<td>ArW</td>
<td>Arctic water</td>
</tr>
<tr>
<td>AW</td>
<td>Atlantic water</td>
</tr>
<tr>
<td>b</td>
<td>Light scattering in m$^{-1}$</td>
</tr>
<tr>
<td>BSW</td>
<td>Brine-enriched shelf water</td>
</tr>
<tr>
<td>c</td>
<td>Light attenuation in m$^{-1}$</td>
</tr>
<tr>
<td>CDOM</td>
<td>Chromophoric dissolved organic matter</td>
</tr>
<tr>
<td>CTD</td>
<td>Conductivity, temperature and depth sensor</td>
</tr>
<tr>
<td>ESC</td>
<td>East Spitsbergen Current</td>
</tr>
<tr>
<td>FDOM</td>
<td>Fluorescent dissolved organic matter</td>
</tr>
<tr>
<td>IOP</td>
<td>Inherent optical properties</td>
</tr>
<tr>
<td>NAPs</td>
<td>Non-algal particles</td>
</tr>
<tr>
<td>OD</td>
<td>Optical density</td>
</tr>
<tr>
<td>VSF</td>
<td>Volume scattering function</td>
</tr>
<tr>
<td>WSC</td>
<td>West Spitsbergen Current</td>
</tr>
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</table>

Data availability. The hydrographic and optical data (Petit et al., 2022) are available at https://doi.org/10.21334/npolar.2022.e6974f73.

Author contributions. BH and MAG designed the study. HS and TP conducted the sample collection and cast profiling. RR and TP performed the sample analysis. TP, MAG, and RR conducted the data analysis. TP and MAG wrote a first draft of the paper. All authors contributed to editing the paper.

Competing interests. The contact author has declared that neither they nor their co-authors have any competing interests.

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