Underwater hyperspectral imaging as an *in situ* taxonomic tool for deep-sea megafauna

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Identification of benthic megafauna is commonly based on analysis of physical samples or imagery acquired by cameras mounted on underwater platforms. Physical collection of samples is difficult, particularly from the deep sea, and identification of taxonomic morphotypes from imagery depends on resolution and investigator experience. Here, we show how an Underwater Hyperspectral Imager (UHI) can be used as an alternative *in situ* taxonomic tool for benthic megafauna. A UHI provides a much higher spectral resolution than standard RGB imagery, allowing marine organisms to be identified based on specific optical fingerprints. A set of reference spectra from identified organisms is established and supervised classification performed to identify benthic megafauna semi-autonomously. The UHI data provide an increased detection rate for small megafauna difficult to resolve in standard RGB imagery. In addition, seafloor anomalies with distinct spectral signatures are also detectable. In the region investigated, sediment anomalies (spectral reflectance minimum at ~675 nm) unclear in RGB imagery were indicative of chlorophyll *a* on the seafloor. Underwater hyperspectral imaging therefore has a great potential in seafloor habitat mapping and monitoring, with areas of application ranging from shallow coastal areas to the deep sea.

Traditionally, assessment of the megafauna inhabiting a region of deep-sea seafloor has been determined either from direct, invasive physical sampling using box corer, trawl1–5 or Remotely Operated Vehicles (ROVs)3,4, or by remotely imaging the seafloor with video or still camera systems mounted on towed platforms6–8, ROVs3,9–11 or Autonomous Underwater Vehicles (AUVs)11–13. Over the last decade, alternative optical methods have been developed for identification and characterization of organisms. These methods are based on hyperspectral imaging, i.e., the acquisition of images integrating hundreds of contiguous colour bands (wavelengths), such that each image pixel contains a full spectrum, typically with 1 nm spectral resolution11. Each image pixel spectrum describes the percentage of light reflected by the material of that image pixel for each wavelength. The reflectance spectra represent so-called optical fingerprints that are specific to the object of interest (OOI).

To study biological organisms underwater, the visible range of the electromagnetic solar spectrum (400–700 nm) is typically used, due to the strong attenuation of light in the ultraviolet (<400 nm) and infrared (>700 nm) regions of the electromagnetic spectrum14–17. However, some studies also extended reflectance measurements to the near-infrared region (up to 900 nm)18–21. The high spectral resolution current hyperspectral imaging systems can achieve (up to 1 nm) and extended spectral range therefore provide a greater volume of colour information than conventional RGB imagery, facilitating the detection, and potentially aiding in the identification of organisms in seafloor images.

Previous studies have focused on the detection of pigmented organisms based on pigment-specific spectral characteristics, using a combination of pigment extraction from plants, algae, and fauna, and subsequent analysis by high performance liquid chromatography, followed by measurements of absorption and reflectance spectra16–22. As reflectance and absorption spectra are inversely related to each other, absorption spectra can be used...
to verify reflectance signatures from organisms. Applications have included *in vitro* studies of biofilms and *in vivo* analysis of sponges, cold-water corals, micro- and macroalgae, and coraline algae. These results have shown that hyperspectral imaging represents a useful bio-optical taxonomic tool for *in vivo* identification of biological organisms.

During the last eight years, hyperspectral studies of marine organisms and other seafloor OOI s have been conducted *in situ*, using diver-operated Underwater Hyperspectral Imagers (UHIs) in shallow waters, as well as by mounting UHIs on mechanical sledges on ROVs, and on AUVs. These studies indicate the promising potential for underwater hyperspectral imaging to greatly aid in the mapping and characterization of different seafloor OOI s *in situ*, including microphytobenthos, kelp forests, coralline red algae, and coral reefs, as well as identifying and characterizing seafloor mineral deposits and geological structures.

By using UHIs on ROVs instead of diver-operated or sledge-based UHIs, seafloor areas of up to 1000 m² may be surveyed in water depths of up to 6000 m. At such depths, sampling of animals for accurate taxonomic identification is generally difficult and time-consuming. Identification of fauna from these remote regions can therefore be very problematic, given the few available physical samples. Whilst standard RGB-based optical imaging methods may collect large datasets, interpretation of these data is not straightforward, with groups of experts potentially interpreting abundances in images differently. At present, there is a tendency to differentiate fauna to a fairly broad taxonomic ‘morphotype’, given the great disparity that can result from even multiple experts investigating a deep-sea fauna image dataset. Attempts at using machine learning algorithms to identify fauna vary in success with ecosystem and fauna.

By identifying fauna by their specific spectral responses, underwater hyperspectral imaging may provide *in situ* taxonomic identifications of benthic communities in a more standardized fashion than is currently achievable by human researchers or algorithms applied to human-identified training sets. This is particularly useful in areas where long-term monitoring of seafloor community structure may have important management applications, such as in the vicinity of oil and gas drilling, deep sea fishing, or deep sea mineral extraction sites.

In this study, we show that underwater hyperspectral imaging can be used as a non-invasive, *in situ* taxonomic tool for benthic megafauna in deep-water habitats. Megafauna may be identified based on their spectral characteristics in UHI image data without the need for physical sampling of animals. In addition, we show that the approach is also useful for detection and mapping of sediment anomalies, including potential seafloor biomass accumulations of phytodetritus, as indicated by elevated chlorophyll-a-like signatures.

**Results**

Seabed OOI s, i.e., megafauna individuals and sediment anomalies, were detected across all 11 ROV transects analysed, with all transects containing both megafauna and sediment anomalies. A total of 30 OOI s were identified based on the video data and previous sampling, and included 28 mobile and sessile megafauna across nine of the 11 transects. Up to four fauna categories with 1–2 individuals per category were observed on each of these transects, including sponges (Demospongiae and Hexactinellida; Fig. 1c.g.h), ophiuroids (Ophiuroidea; Fig. 1d), holothurians (*Paelopatides* sp. and *Synallactidae*; Fig. 1e), corals (Fig. 1f.i), polychaetes (*P. mirabilis*; Fig. 1j), dead salps (Fig. 1a), as well as a crinoid (*Crinoidea*; Fig. 1k) and an isopod (*Munnoopsidae*). One bony fish (*Ipnops* sp.; Fig. 1b) was also observed. Megafauna individuals detected in the video data measured about 2–15 cm in length (average 6 cm). Although more animals were visible in the video data, they were not clearly resolved due to their small sizes (<2 cm), and hence identification was not possible.

In addition, two anomalous sediment patches of a lighter colouration than the surrounding seafloor, likely representing recently exposed sediments from below the active surface sediment layer, were observed.

The 30 OOI s identified from the video data were clearly visible in the UHI images and exhibited spectral signatures that were distinct from the background spectra of sediment and manganese nodules, which were relatively stable throughout the transects. The OOI spectra also differed between the imaged fauna morphotypes, whilst the spectral signatures of different individuals of the same morphotypes were similar (Fig. 2). For example, the three crustaceans (*P. mirabilis*; Fig. 2e) were characterized by very similar spectral responses. Five dead salps, common seafloor ecosystem components during the research cruise, also showed similar spectral responses (Fig. 2f). In the case of corals, spectral signatures of surveyed green and yellow corals did not match those of three surveyed white corals (Fig. 2b). Similarly, two white ophiuroids exhibited a different spectral response than a red ophiuroid (Fig. 2d). Two observed holothurians, which were of different species (*Synallactidae* and *Paelopatides* sp.), also had different spectral signatures, differing most markedly in the red part of the spectrum (600–710 nm; Fig. 2c).

In addition to the 30 OOI s identified from the video data, 48 further OOI s were detected in the UHI images of each transect, with numbers ranging between two and six additional OOI s per transect. The calculated spectral contrast angle, which was used to compare the spectra of these 48 OOI s to the 30 reference spectra, varied between 0.63° and 30° in general and between 0.63° and 3.03° for the smallest angle (1.75° on average). Based on the smallest spectral contrast angle, 13 OOI s were associated with megafauna, in particular white corals and polychaetes (Table 1). The remaining 35 OOI s represented sediment anomalies. Six of these were reminiscent of the exposed deeper sediment patches, while the other 29 OOI s had spectral signatures characterized by a distinct intensity minimum around 668–680 nm (Fig. 3) that was not observed in the spectra belonging to the exposed deeper sediment patches. These OOI s were roughly circular spots with maximum diameters of 2 cm that appeared either green or white in the UHI images displayed in pseudo-RGB (R: 645 nm, G: 571 nm, B: 473 nm; Figs 4 and 5).

Along each transect, at least one of these anomalies was observed in the UHI images, while they were barely or not visible in the video data. Due to the characteristic intensity minimum, these OOI s were assigned to two new spectral categories termed “green spot” and “white spot” (Table 1). Spectral signatures were relatively similar for both the green and the white spots, although pseudo-reflectance intensities were slightly higher for the white spots, especially for wavelengths >650 nm (Fig. 3).
Using the average spectra from the 78 identified OOIs as training spectra, the SVM method revealed a total of 328 OOIs across the 11 transects (Table 1). In general, all of the known OOIs were detected and classified correctly. In some cases, the marginal pixels of organisms were misclassified as other fauna categories with similar spectra, e.g. sponge instead of white coral, but the bulk part of the organism surface was classified correctly. This allowed each OOI to be clearly assigned to one of the previously defined categories.

In addition to the 78 known OOIs, 250 new OOIs were detected and classified by the SVM method (Table 1). On all transects, the number of OOIs revealed by the SVM results (6–71 OOIs) exceeded that of the OOIs identified in the video data (0–6 OOIs), showing that more anomalies were detected using the spectral classification approach than from analysis of the visual image data. This distribution is illustrated in Fig. 6 for a section of the track shown in Fig. 5. In this section, a dead salp and a sponge (Fig. 1c) identified from the video data were well apparent in the UHI image (Fig. 6a). Additional OOIs observed in the UHI image were identified as white spots based on the spectral contrast angle of the associated spectra (Fig. 6b). By including the average spectra from the identified OOIs in the training data, the SVM method correctly classified the training OOIs and revealed additional OOIs belonging to the white spot and exposed sediment categories (Fig. 6c).

Besides increasing the number of detected OOIs, the SVM approach was also able to detect OOIs with sizes on a sub-cm scale (down to 0.8 cm) and generally showed an increased detection rate for small (<2 cm) organisms.

Figure 1. Images from HD video frames for different mobile and sessile megafauna observed in the study area. All scale bars measure 6.5 cm. (a) dead salp, (b) bony fish (Ipnops sp.), (c) Demosponge (round) on manganese nodule, (d) ophiuroid, (e) holothurian (Paeleopatides sp.), (f) holothurian (Synallactidae) with a sessile polychaete on the left side and a small coral on the right side, (g) Demosponge (disc), (h) Hexactinellid, (i) stalked sponge with a green coral attached to the stalk and an ophiuroid wrapped around it, (j) crustacean (Probeebei mirabilis), (k) stalked crinoid. Images: ROV KIEL 6000, ROV Team GEOMAR.
such as small corals and polychaetes compared to the video-based OOI detection (Table 1). For larger megafauna, such as crustaceans, salps, and holothurians (7–15 cm in length), the SVM method did not reveal any additional individuals besides those already identified from the video data.

In total, 328 OOIs, i.e., spectral anomalies that differed from the background sediment and manganese nodules, were identified across the 72.3 m² covered by the 11 survey tracks. Megafauna constituted 83 of these OOIs, the majority being white corals (39 individuals, or 0.54 m⁻²), polychaetes (10 individuals, or 0.14 m⁻²), and sponges (8 individuals, or 0.11 m⁻²). Of the 245 sediment anomalies, 187 were green or white spots, corresponding to 2.59 m⁻².

Discussion
Our results show that more megafauna individuals were detected and identified from the SVM classification results than from visual identifications in the video data. For example, only three white corals were observed by human observers in the video data, while 39 individuals were revealed by the spectral classification results. Similarly, only one polychaete was identified in the video data alone, whereas spectral classification revealed a total of 10 polychaetes across three different transects.

We note that it was not possible to verify the spectra-based identifications in all cases, due to the small sizes of most of the animals (<2 cm) and the associated difficulty in clearly resolving them in the video data. The overall classification accuracy of the SVM results for this dataset was estimated to >90%²⁶, suggesting that most OOIs, but not all, may have been classified correctly. This is a greater accuracy than is commonly produced from visual identification alone, given the inter-observer bias commonly reported from multi-observer studies⁷.
Despite this uncertainty associated with the SVM classification results, our results demonstrate that if a set of reference spectra from known megafauna exists, further individuals of the same fauna type can be identified in situ based on their spectral characteristics, or individual optical fingerprints. Although previous studies linked organism identifications based on spectral reflectance signatures to the pigment compositions of the organisms\(^\text{16,18,24}\), we show that identifications based on spectral characteristics are possible even if the pigment compositions are unknown. Consequently, additional sampling of marine organisms – a challenging procedure in the deep sea – and subsequent extraction of pigment information is not required. Our study thus extends the use of hyperspectral imaging as a taxonomic tool for marine organisms from in vivo applications\(^\text{16,21}\) to in situ surveys.

| Category                  | Taxonomy                          | Training spectra from video/samples | from spectral anomalies | total training spectra | SVM classification results |
|---------------------------|-----------------------------------|-------------------------------------|-------------------------|------------------------|---------------------------|
| Megafauna                 |                                   |                                     |                         |                        |                           |
| Sponge (round)            | Porifera, Demospongiae sp1        | 4                                   | 4                       | 8                      |                           |
| Sponge (disc)             | Porifera, Demospongiae sp2        | 1                                   | 1                       | 1                      |                           |
| Glass sponge              | Porifera, Hexactinellida          | 1                                   | 1                       | 1                      |                           |
| Coral (white)             | ?                                 | 3                                   | 8                       | 11                     | 39                        |
| Coral (yellow)            | ?                                 | 1                                   | 1                       | 1                      |                           |
| Coral (green)             | ?                                 | 1                                   | 1                       | 1                      |                           |
| Holothurian Synallactidae| Echinodermata, Holothuroidea, Synallactidae | 1                                    | 1                       | 1                      |                           |
| Holothurian Paeopatides   | Echinodermata, Holothuroidea, Paeopatides sp | 1                                    | 1                       | 1                      |                           |
| Ophiuroid (white)         | Echinodermata, Ophiuroidea sp1    | 2                                   | 2                       | 2                      |                           |
| Ophiuroid (red)           | Echinodermata Ophiuroidea sp2     | 1                                   | 1                       | 1                      |                           |
| Crustacean                | Arthropoda, Decapoda, Proteebei mirabilis | 3                                    | 3                       | 3                      |                           |
| Salp (dead)               | ?                                 | 5                                   | 5                       | 5                      |                           |
| Polychaete                | ?                                 | 1                                   | 3                       | 4                      | 10                        |
| Crinoid                   | Echinodermata, Crinoidea          | 1                                   | 1                       | 2                      | 4                         |
| Isopod                    | Arthropoda, Isopoda, Munnopsidea  | 1                                   | 1                       | 2                      | 4                         |
| Bony fish                 | Chordata, Actinopterygii, Ipnops sp. | 1                                    | 1                       | 1                      |                           |
| Other anomalies           |                                   |                                     |                         |                        |                           |
| White exposed sediment    |                                   | 2                                   | 6                       | 8                      | 58                        |
| Green spot                |                                   | 9                                   | 9                       | 9                      | 56                        |
| White spot                |                                   | 20                                  | 20                      | 131                    |                           |
| Identified/reference spectra |                               | 30                                  | 48                      | 78                     | 328                       |

Table 1. Overview of OOI categories and classification results. Training spectra for the SVM classification were obtained from OOI identified based on video data and samples, and from additional spectral anomalies in the UHI images. SVM – Support Vector Machine.

Figure 3. Examples of normalized pseudo-reflectance spectra of the anomalous green spots (a) and white spots (b). Green spots g1-g4 are shown in Fig. 4, white spots w1-w4 are marked in Fig. 5. Both green spots and white spots are characterized by a prominent intensity minimum around 675 nm, resembling reflectance dips of chlorophyll a or its degraded form, phaeophorbide a and/or phaeophytin a.

Despite this uncertainty associated with the SVM classification results, our results demonstrate that if a set of reference spectra from known megafauna exists, further individuals of the same fauna type can be identified in situ based on their spectral characteristics, or individual optical fingerprints. Although previous studies linked organism identifications based on spectral reflectance signatures to the pigment compositions of the organisms\(^\text{16,18,24}\), we show that identifications based on spectral characteristics are possible even if the pigment compositions are unknown. Consequently, additional sampling of marine organisms – a challenging procedure in the deep sea – and subsequent extraction of pigment information is not required. Our study thus extends the use of hyperspectral imaging as a taxonomic tool for marine organisms from in vivo applications\(^\text{16,21}\) to in situ surveys.
The high spatial resolution of the UHI image data (1 mm pixel size), combined with the spectral classification approach, is especially valuable for the detection of smaller fauna individuals (<2 cm in size). Examples include the tubed polychaetes, some of the sponges and less developed coral colonies in the current study, which were often difficult to detect or resolve in the lower-resolution video data (smallest resolvable size ca. 2 cm). By analyzing their spectral responses and applying spectral classification, the detection and identification rate for these organisms was increased considerably.

In contrast to smaller fauna, larger megafauna measuring about 4–15 cm in length, such as the adult crustaceans, ophiuroids, dead salps, and holothurians in the analysed dataset, were generally easy to detect both in the video data and in the UHI images, with the spectral approach not revealing any additional individuals to those already logged from the visual data by human observers. However, the UHI assessment, based on optical fingerprints rather than physical characteristics of the individual fauna, may well make the data more suitable for automated approaches, given the difficulty for semi-automated systems to be trained to recognize fauna of highly variable aspect or appearance7,28.

Even for the larger fauna, knowing the spectral responses of these may in some cases be used to indicate whether individuals belong to the same species. For example, in the case of the two holothurians observed, one was identified as belonging to the family Synallactidae based on the video data (Fig. 1f), while identification of the other holothurian was more difficult. This second individual (Fig. 1e) was interpreted as belonging to either Synallactidae or Paelopatides sp. Given that the spectral responses of these individuals differed markedly, especially in the red portion of the electromagnetic spectrum (Fig. 2c), they likely belonged to different species, and the second holothurian was therefore identified as Paelopatides sp.

Our results thus show that underwater hyperspectral imaging can be used as a non-invasive, in situ taxonomic tool for benthic megafauna. Even without knowledge of pigment composition and associated spectral absorption signatures, it is possible to differentiate marine organisms based on their unique spectral reflectance signatures in UHI images. Once a set of reference spectra based on identified organisms is established, spectral comparisons and application of supervised classification methods allow identification of benthic megafauna semi-autonomously without requiring additional sampling of animals. Especially in the deep sea, where sampling is difficult and time-consuming due to the high water depths, this approach is of great potential. For example, the assessment of the abundances of both small and larger megafauna may well be useful in gauging community recovery following a disturbance event in the deep sea. Aside from manganese nodule fields, deep-sea massive sulfide deposits and manganese crusts are economically attractive sources of deep-sea minerals32–35. Removal of these resources will wholly remove the colonized seafloor in these regions11,36, exposing new rock surfaces to the ocean. Monitoring the rate of post-exploitation colonization of these new surfaces by fauna will be an important component of management plans. The facility of underwater hyperspectral imaging to identify the early stages of colonization (i.e., the recently settled small corals, sponges, crinoids, etc.) could greatly aid in understanding faunal succession and relative success of any mitigation measures put in place in such areas36.

In addition to megafauna, the UHI is also able to detect and characterize seafloor anomalies with distinct spectral signatures. The distinct reflectance intensity minimum around 668–680 nm observed for the green and white spots is possible to differentiate marine organisms based on their unique spectral reflectance signatures in UHI images. Once a set of reference spectra based on identified organisms is established, spectral comparisons and application of supervised classification methods allow identification of benthic megafauna semi-autonomously without requiring additional sampling of animals. Especially in the deep sea, where sampling is difficult and time-consuming due to the high water depths, this approach is of great potential. For example, the assessment of the abundances of both small and larger megafauna may well be useful in gauging community recovery following a disturbance event in the deep sea. Aside from manganese nodule fields, deep-sea massive sulfide deposits and manganese crusts are economically attractive sources of deep-sea minerals. Removal of these resources will wholly remove the colonized seafloor in these regions, exposing new rock surfaces to the ocean. Monitoring the rate of post-exploitation colonization of these new surfaces by fauna will be an important component of management plans. The facility of underwater hyperspectral imaging to identify the early stages of colonization (i.e., the recently settled small corals, sponges, crinoids, etc.) could greatly aid in understanding faunal succession and relative success of any mitigation measures put in place in such areas.

In addition to megafauna, the UHI is also able to detect and characterize seafloor anomalies with distinct spectral signatures. The distinct reflectance intensity minimum around 668–680 nm observed for the green and white spots suggests a possible relationship between these sediment anomalies and chlorophyll a concentration, which has a characteristic in vivo absorption maximum of around 675 nm, or between 660–665 nm in in vitro studies. Given the water depth of nearly 4200 m, the presence of chlorophyll a and associated biomass at the seafloor may seem unlikely. However, concentrations of chlorophyll a and its degradation products...
(phaeophytin/phaeophorbide a) have been observed at the abyssal seafloor in both the equatorial Pacific and the North Atlantic40–43. Chlorophyll a concentrations in these areas have been associated with phytodetritus deposited on the seafloor after seasonal phytoplankton blooms in the shallow water column40–43, with deposits accumulating either as continuous greenish layers or as cm-scale aggregates42.

Chlorophyll a concentrations have also been measured in seafloor sediments in the nearby DISCOL area, 700 m northwest of the study area (Matthias Haeckel, personal communication), although direct evidence for phytodetritus was not observed during the study period. However, at the time of the survey, a large fall of dead salp, fauna which bloom in upper waters in response to algal blooms, was ongoing31. Elevated chlorophyll a concentrations may possibly be related to this event. As the spectral characteristics of the green and white spots (Fig. 3) resemble those of degraded chlorophyll a, such as phaeophorbide a and/or phaeophytin a44, we interpret the green and white spots as indications for the presence of chlorophyll a or associated breakdown products. However, physical samples from these anomalies would be required to confirm a relationship to chlorophyll a and its degradation products.

The difference between the green and the white spots is still unclear. One possibility is that the slightly different spectral responses reflect different chlorophyll a concentrations. Based on spectral reflectance studies in intertidal mud flats, Kromkamp et al.38 noted that an increase in biomass caused a decrease in reflectance, in particular around the chlorophyll a absorption maximum at 675 nm. As the green spots exhibit slightly lower spectral intensities than the white spots in this part of the electromagnetic spectrum (Fig. 3), the green spots may potentially be related to more biomass or higher chlorophyll a concentrations than the white spots. Chlorophyll a concentrations may be determined from reflectance spectra via a derived spectrometric index19, but this method could not be applied due to the availability of only pseudo-reflectance data, and hence chlorophyll a concentrations were not estimated.

Figure 5. A 2.9 m long UHI transect (in pseudo-RGB with R: 645 nm, G: 571 nm, B: 473 nm) across manganese nodules, a dead salp, a sponge and an infauna burrow trail. Annotations w1-w4 mark the white spots belonging to the spectral responses shown in Fig. 3b.
Alternatively, the difference in reflectance intensity between the green and white spots could be due to a difference in decay stage of the phytodetritus. Reflectance of organic material may vary over time, as shown by Dierssen et al.\textsuperscript{45} in a study on floating wrack of the genus \textit{Sargassum}. After an initial decrease of \textit{Sargassum} reflectance over three days, reflectance increased again, with the effect of varying reflectance being most prominent around the chlorophyll \textsubscript{a} absorption maximum.

Most of the green and white spots with distinct spectral signatures in the UHI data are not apparent in the HD video data and would thus have been overlooked if only standard RGB data had been used for seafloor imaging. Moreover, the limited spectral information provided by the three RGB bands would not have allowed establishing a potential link to chlorophyll \textsubscript{a} based on the spectral responses. Although sampling is still required to confirm the presence of chlorophyll \textsubscript{a} or degradation products at the green and white spots, our results indicate that hyperspectral image data are well suited for the detection and imaging of chlorophyll \textsubscript{a} distribution on the seafloor, including in the deep sea. The facility of the approach to detect chlorophyll \textsubscript{a} at depth, remotely, and over extended areas is highly attractive, and may well support investigations into carbon transport pathways, particularly in remote regions with a high seasonality\textsuperscript{46}. In addition to providing insight into the extent of seasonal phytodetritus deposition, mapping chlorophyll \textsubscript{a} distribution on the abyssal seafloor may also elucidate feeding habits of benthic megafauna, for which these surface deposits provide an additional food source\textsuperscript{40–42}.

Materials and Methods

Data acquisition and processing. The current study used the UHI data collected and presented in Dumke \textit{et al.}\textsuperscript{26} for location and identification of benthic megafauna and sediment anomalies. The data were acquired in a well characterized manganese nodule field in the Peru Basin (SW Pacific; 7°5′23″S, 88°26′46″W, 4195 m depth) during the RV SONNE cruise SO242/2, using a deep-sea UHI (UHI #4) by Ecotone AS (Trondheim, Norway) mounted on the KIEL6000 ROV (GEOMAR)\textsuperscript{26,31}.

The UHI is a push-broom scanner with beamwidths of 60° (transverse) and 0.4° (longitudinal). It was mounted on the ROV’s outstretched manipulator arm and looked vertically downwards to record lines of 1600 pixels perpendicular to the track direction\textsuperscript{26,31}. Seafloor illumination was provided by ten ROV light sources, which included an LED, halogen lamps (five Deep Multi-SeaLite lamps, two Sea Arc 5000 lamps), and two HMI lamps (SeaArc2). For each image pixel, the UHI recorded the intensities of the reflected light for 112 spectral bands between 378 nm and 805 nm with a spectral resolution of 4 nm.

UHI data were recorded throughout 11 ROV tracks across the seafloor, passing over various benthic fauna and manganese nodules (ROV dive SO242/2_191–193). The tracks were run at a constant speed (0.05 m s\textsuperscript{−1}) and heading, with track lengths ranging between 1.7 m and 20 m. The relatively constant altitude of 1–1.2 m resulted in a track width (field of view) of 1–1.2 m. In addition to the UHI data, HD and SD video data were simultaneously collected by the ROV cameras during each survey track.
As the spectral bands <400 nm and >710 nm were rather noisy, spectral subsetting was applied to the UHI data to reduce the data to the 83 bands between 400 nm and 710 nm. The raw UHI data contained several spectral components associated not only with the OOIs, but also with illumination, the inherent optical properties of the water column (scattering and absorption), vehicle motion, and imager properties. These external influences had to be corrected for in order to obtain reflectance data in which the spectral response is specific to the OOIs.

For this initial deep-sea deployment of the system, it was not possible to completely correct for all external influences. The raw UHI data were first calibrated to radiance data by applying radiometric correction using the Hypermap software tool (Ecotone AS). To process the radiance data to true reflectance data, knowledge of the illumination influence (the combined spectral characteristics of the ROV lights for each image pixel location) and the inherent optical properties of the water column were required. As these characteristics could not be determined, they were approximated by a reference spectrum calculated from the UHI image data collected during each track. By diving each pixel spectrum by its reference spectrum, followed by smoothing of the spectra with a moving average filter for a window of 11 bands (44 nm), most external influences could be eliminated (for further information on data processing, see Dumke et al.26). However, as the corrected data still contained some residual influences, mainly from the illumination source and geometry, they represented “pseudo-reflectance” data rather than true reflectance data.

The pseudo-reflectance data were of sufficient quality to allow differentiation between seafloor features and fauna based on the different optical fingerprints, but they may not be fully comparable to true reflectance data. The corrected UHI data were georeferenced (for further details, see Dumke et al.26) and output with a pixel size of 1 mm. The UHI data used in this study are available from PANGAEA (https://doi.pangaea.de/10.1594/PANGAEA.874408).

Data analysis. The analysis of the UHI data and the video data involved several steps. Firstly, the video data collected at the same time as the UHI data were analysed for the presence of OOIs, in this case megafauna individuals and any obvious sediment anomalies. These OOIs constituted the reference group. Animals were identified to the highest possible taxonomic level based on their appearance in the video data, and from previous physical sampling of megafauna in the area30,31,47. For each identified OOI, an average pixel spectrum was extracted from the UHI images. To account for any spectral variations or effects of bi-coloration across the OOI surface (such as overgrowth of portions of a surface by small encrusting sponges etc.), the investigation of which was outside the scope of this study, these pixel spectra were averaged over OOI surface areas of between 20 and 600 pixels, depending on the size of the OOI.

The UHI images were then analysed for additional OOIs that had not been identified from the video data and were associated with distinct spectral anomalies. Average pixel spectra from pixel areas of 20–200 pixels were extracted for these selected OOIs. To identify the OOIs, each OOI spectrum was compared against the 30 reference spectra from the OOIs identified from the video data by calculating the spectral contrast angle \( \theta \) (Equation (1))48, which has to date primarily been applied in the comparison of mass spectra48–51 but is also applicable for determining similarities between any kind of spectra6. For each unidentified OOI, \( \theta \) was calculated for each reference spectrum and the OOI was subsequently assigned to the category associated with the most similar spectrum (smallest \( \theta \)).

Following this step, spectral classification of the UHI images was performed using the ENVI software (v. 5.3; Exelis VIS). In spectral classification, each pixel spectrum is compared to training data, i.e., a set of known reference spectra from identified organisms or materials. The pixel spectra are then assigned to categories depending on which reference spectrum they match best23,25. Spectral classification was conducted using the Support Vector Machine (SVM) method, a standard classification technique often superior to other classification algorithms54,55 and known for its robustness with noisy and complex data6. The classification results were output as classification images showing the distribution of the different spectral categories.

This study used the same classification images presented in Dumke et al.26, but results and interpretations herein focus on benthic megafauna and sediment anomalies, features which were not discussed by Dumke et al.26. For spectral classification of the megafauna and sediment anomalies, the spectra of the previously identified OOIs were used as training data. The SVM classification results were then analysed to determine (1) if the training OOIs had been classified correctly, and (2) if any additional OOIs had been detected by the SVM method.

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Author Contributions
S.M.N. collected the data, I.D. performed the analyses, A.P. and Y.M. provided taxonomical identifications, and M.L., G.J. and F.S. organized funding for the work. I.D. wrote the manuscript and all authors contributed substantially to revisions.

Additional Information
Competing Interests: The authors declare no competing interests.

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