SPECTRAL CHARACTERISTICS OF SELECTED HERMATYPIC CORALS FROM GULF OF KACHCHH, INDIA

Nandini Ray Chaudhury

EHD/ABHG/EPSA, Space Applications Centre, ISRO, Ahmedabad – 380015 – nandinirc@sac.isro.gov.in

Commission VIII, WG VIII/6

KEY WORDS: Ecosystem, Hyper spectral, Multispectral, Image, Identification

ABSTRACT:

Hermatypic, scleractinian corals are the most important benthic substrates in a coral reef ecosystem. The existing, high (spatial) resolution, broad-band, multi-spectral, space-borne sensors have limited capability to spatially detect and spectrally discriminate coral substrates. In situ hyperspectral signatures of eight coral targets were collected with the help of Analytical Spectral Devices FieldSpec spectroradiometer from Paga and Laku Point reefs of Gulf of Kachchh, India to study the spectral behaviour of corals. The eight coral targets consisted of seven live corals representing four distinct colony morphologies and one bleached coral target. The coral spectra were studied over a continuous range of 350 to 1350 nm. The corals strongly reflected in the NIR and MIR regions with regional central maxima located at 820 and 1070 nm respectively. In the visible region the live coral spectra conformed to “brown mode” of coral reflectance with triple-peaked pattern at 575, 600 and 650 nm. All coral spectra are characterized with two distinct absorption features: chlorophyll absorption at 675 nm and water absorption at 975 nm. The live and the bleached corals get distinguished in the visible region over 400 to 600 nm region. Water column over the targets modifies the spectral shape and magnitude. First and second-order derivatives help in identifying spectral windows to distinguish live and bleached corals.

1. INTRODUCTION

Worldwide, hermatypic or reef-building corals occupy a keystone position in the overall functioning of reef ecosystem. Live coral cover of a reef habitat is one of the most widely accepted parameters of coral reef health (Green et al. 2000). Space-borne, optical remote sensing has sensor limitations in automated extraction of live coral cover parameter from space data. Digital delineation of this parameter has so far mostly used the expert knowledge of local reef systems in terms of specific geomorphological features and associated benthic classes (Hochberg and Atkinson, 2003). The existing, medium to high resolution, broadband, multi-spectral sensors have limited capabilities to provide synoptic data on live coral cover at community level. Poor penetration of sunlight and low signal to noise ratio of backscattered reflectance from deeper reef habitats are considered to be the major limiting factors to the performance of these sensors (Green et al. 2000). Therefore, live coral cover assessment through space-based imaging has been successfully confined to emergent, shallow, reef flat environments preferably at low tides. Relative abundance of reef biota and litho-substrates at different bathymetric depths with varying water column (i.e. depth, quality, etc.) add high degree of natural variability to coral reefs as optically complex, shallow water, remote sensing targets. This poses significant challenge in remote assessment of reef-scale biodiversity in terms of spatial detection and spectral characterization of individual corals at colony scale.

As an underwater remote sensing target, coral reefs appear as mosaics of diverse “substrates” or “bottom types” when viewed from the space (Hochberg et al. 2003). Spectral nature of the space-borne data has been recognized as the basic link between coral reef substrates and remotely sensed images (Hochberg and Atkinson, 2003). Substrate detection and quantification efforts have applied a deterministic, in situ data based, “reef-up” approach which often use proximal remote sensing of corals and other reef benthos. This approach calls for spectral cataloguing to identify distinct spectral features of a substrate. In situ reef-up approach has demonstrated the potential applications of field spectroscopy and hyperspectral analyses to characterize the reflectance properties of individual reef substrates which usually comprise an image (Goodman and Ustin, 2002). In situ hyperspectral signatures of corals and other reef benthos obtained with the help of portable field spectroradiometers have mostly been used for spectral discrimination of reef substrates (Hochberg and Atkinson, 2000, Hochberg et al. 2003, Hochberg and Atkinson, 2003, Kutser et al. 2003) and for health of corals (Holden and LeDrew, 1998; 1999; Clark et al. 2000). In situ reef spectra have been also used as input to simulate top-of-atmosphere spectral reflectance using radiative transfer models (Lubin et al. 2001).

In situ spectral reflectance of corals per se has been recognized as a fundamental parameter in coral reef remote sensing (Hochberg et al. 2004). Spectral characteristics of coral organisms at colony or at community level have been correlated primarily with coral pigments, fluorescence and colony morphology (Hedley and Mumby, 2002; Hochberg et al., 2004; Joyce and Phinn, 2002 and 2003).

Broad-band, multi-spectral signatures obtained from image-pixels of high (spatial) resolution Indian Remote Sensing (IRS) satellite
Indian coast is endowed with spatially limited but strategically located coral reef habitats which offer a myriad of marine biodiversity combined with unique regional characteristics. In India, major coral reefs occur in four distinct locations. Two of them occur in gulf settings: Gulf of Kachchh in Arabian Sea and Gulf of Mannar in Bay of Bengal while the other two are offshore island groups of Lakshadweep in Arabian Sea and Andaman and Nicobar in Bay of Bengal. Indian coral reefs share sixty genera of hermatypic, scleractinian corals out of the one hundred and eleven genera reported in the world (Venkataraman, 2003). Thus, Indian corals share 54% of global coral diversity at genera level.

The southern part of Gulf of Kachchh from Gujarat coast represents the sturdiest scleractinian coral species of India. Occurring in the northernmost limits of Indian reef regions (22°20’ to 22°40’ north latitudes and 68°30’ to 70°40’ east longitudes), these corals grow in a highly turbid and saline, macro-tidal environment marked with semi-diurnal desiccations due to fluctuating tidal exposures (Navalgund et al. 2010). Out of the sixty genera of scleractinian corals reported from India, only twenty genera are found in Gulf of Kachchh (Venkataraman, 2003) reef region which is declared and protected as a Marine Sanctuary since 1983.

2. STUDY AREA

Paga reef and Laku Point (near Poshitra) were chosen as two specific sites (Figure 1) for collection of in situ coral spectra from Gulf of Kachchh. These sites had been reported to have relatively high generic diversity of scleractinian corals within the region (Patel, 1978). Moreover the yearly, equinoctial spring tides (negative low tides) result in maximum exposure of inter-tidal and sub-tidal areas of these reefs facilitating in situ coral spectra collection with virtually no water column. Paga is an off-shore, patch reef where diverse coral colonies occur mostly in the reef slope, reef crest and outer reef flat areas while Laku point is a narrow fringing reef where coral colonies grow in shallow, rock-pools.

3. MATERIAL & METHODS

3.1 Multi-spectral Signatures of Reef Substrates from Resourcesat-1 LISS-IV Data

Linear Imaging and Self Scanning sensor: LISS-IV, onboard Resourcesat-1 (IRS-P6) and 2 (RS2) satellites have been the most preferred imaging sensor for Indian coral reefs for its high spatial resolution (5.8 m at nadir) complemented with three spectral channels (located in Green, Red and NIR regions) and 10 bit level of quantization. LISS-IV in multi-spectral mode has performed considerably well to characterize reef geomorphology of the smaller reefs of Central Indian Ocean (Navalgund et al. 2010). However, at an orbital altitude of 817 km, detection capability of this sensor gets spatially limited for coral colonies within a reef. Three discrete, broad-band, spectral channels, positioned at 530 to 590 nm, 620 to 680 nm and 770 to 860 nm usually fall short to spectrally resolve a “pure coral signature”. Coexistence of macro reef-benthos like corals and macro-algae along with underlying litho-substrates (sand, mud, etc.) under varying depths of water column make this task all the more difficult. Atmospheric interferences also alter the strength of the back-scattered signal through atmospheric absorption and scattering. The back-scattered signal from a reef for a single pixel can thus be a mixed representation of the natural heterogeneity present in the corresponding reef area. This problem has been demonstrated with the help of Figures 2 and 3 as a case study using a subset of an archived IRS- P6 LISSIVMX (multi-spectral) data acquired on 16th March, 2005 pertaining to Paga reef. The spectral behaviour of selected reef substrates have been analysed with respect to Top of the earth’s Atmosphere (TOA) spectral radiance. No atmospheric correction has been performed on this subset image for this study.

Figure 3 shows multi-spectral signatures (in terms of mean spectral radiance of randomly selected thirty pixels for each group) of five distinct reef categories obtained from the subset of LISS-IVMX image of Paga reef (Figure 2). This subset image was digitally enhanced by applying standard deviation stretch for better visual appreciation of the reef substrates. Four out of the selected five pixel-groups which represent four different reef substrates are clearly, visually distinguishable by their respective tones. In a standard LISS-IVMX False Colour Composite (FCC), white pixels representing pure, exposed sand get well distinguished from rest of the reef pixel classes, as a substrate giving the highest spectral response in all the three channels. For Paga reef the submerged sand pixels on reef flat (free of any kind of benthic substrates) appeared in light green tone as the sand is mixed with silt and clay.

Figure 1. Location of the study sites
The effect of water column in suppressing the magnitude of sandy substrates from reef is visible in Figure 3 if one compares the values of mean spectral radiance in all the three channels represented by grey and cyan triangles. Benthic green (chlorophyceae) and brown (phyaophyceae) macroalgae groups can be differentiated in terms of pixels appearing in orange and brown colours respectively. Spectrally, chlorophyceae group dominates the phyaophyceae in all the three channels as shown in Figure 3. The magnitude of difference in their spectral response is minimal in the red band (spectral channel 2, 620-680 nm) while in the green band (spectral channel 1, 530-590 nm) there is a slight increase in this difference. In NIR (spectral channel 3, 770-860 nm) this difference magnifies drastically which is well evident in Figure 3.

The fifth pixel group represents mixed pixels (appearing in different tones in the FCC) randomly selected from the ‘outer reef flat’ zone of Paga (Figure 2). This zone is naturally characterized by diverse benthic and litho-substrates including live coral colonies. The natural diversity allows this zone to appear as a ‘rough textured’ zone adjacent to ‘smooth textured’ chlorophyceae dominated areas. Interestingly, the position of mean spectral radiance of this mixed pixels lie very close to the centre of the vertical distance representing the magnitude difference in spectral response of sand on reef and benthic brown algae categories in green and red bands (red filled triangles vis-à-vis cyan and light green triangles). In NIR, sand on reef category is replaced by chlorophyceae representing the upper limit of this vertical distance as reef sand shows relatively less spectral response due to water column absorption.

Thus, in NIR the mean spectral radiance of mixed pixels lie within the vertical range defined by chlorophyceae and phyaophyceae. So it can be inferred that sand on reef and benthic macro-algae contribute to the backscattered signal of these mixed pixels. This fact is confirmed if one numerically calculates the spectral radiance of mixed pixels assuming that sand on reef and phyaophyceae contributes in equal proportion to a mixed pixel signal in green and red bands while chlorophyceae and phyaophyceae in NIR. This is demonstrated in figure 3 by the red outline triangles against the red filled/solid triangles. This reaffirms the fact, that even in high (spatial) resolution, broad-band, multi-spectral images, pixel-based spectral signature of coral colonies is dominated and obscured by other reef substrates.

Hence there is a definite need to explore the hyperspectral domain in remote sensing to understand the spectral behaviour of coral colonies at all possible scales and modes of acquisition: in situ, air-borne and space-borne.

### 3.2. In situ Spectral Measurements and Data Processing

*In situ* coral spectra were collected during the equinoctial spring tide (i.e. maximum negative tide = -0.09 m) of March, 2011 when low tide exposures of reefs coincided with early hours of local day time (i.e. 09:00 to 11:00 hrs) suitable for passive, proximal sensing of coral colonies with no or minimal water column. Coral reflectance spectra were collected with Analytical Spectral Devices (ASD) FieldSpec3 spectroradiometer having a spectral range of 350 to 2500 nm and spectral resolution of 3 nm (at 700 nm) and 10 nm (at 1400, 2100 nm). The sampling interval is 1.4 nm for 350-1000 nm wavelength region and 2 nm for 1000-2500 nm regions. The fibre optic probe has a Field of View (FOV) of 25° full conical angle. Since the objective was to study *in situ* spectral reflectance of diverse coral communities, a point sampling strategy was followed. The field spectroradiometer was calibrated with reference to a Spectralon white plate and thereafter multiple coral spectra were recorded from different sample stations. For each station, a minimum of thirty reflectance spectra was logged along with GPS coordinates, water depth and water transparency (visual). Spectral measurements were carried out for twenty two stations over three consecutive days during 09:00 to 10:30 hrs (to reduce illumination variations) when the live coral colonies were submerged in less than 10 cm of clear, water column and data logging was completed within 15-minute period for each station. The field spectra were subsequently processed with the help of ViewSpecPro software (version 5.6).

Twenty two hermatypic coral targets representing different taxonomic genera and colony morphologies (with varying levels of underwater polyp exposures) were sampled on field. Eight sample stations (representing seven live coral genera and one bleached coral, with least water depths) were later selected out of these twenty two stations as pure samples. The details of these eight coral targets are given in Table 1 and Figure 4 shows their field photographs.
Table 1. Details of the eight sampled coral targets

<table>
<thead>
<tr>
<th>Sr. No.</th>
<th>Coral Genera</th>
<th>Colony Morphology</th>
<th>Water column (in cm)</th>
<th>Field Site (Tidal Zone)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Favia</td>
<td>Massive</td>
<td>1.50</td>
<td>Paga (Intertidal)</td>
</tr>
<tr>
<td>2</td>
<td>Symphillia</td>
<td>Massive</td>
<td>8.00</td>
<td>Paga (Subtidal)</td>
</tr>
<tr>
<td>3</td>
<td>Goniopora</td>
<td>Sub-massive</td>
<td>1.50</td>
<td>Laku Point (Intertidal)</td>
</tr>
<tr>
<td>4</td>
<td>Porites</td>
<td>Sub-massive</td>
<td>2.50</td>
<td>Paga (Intertidal)</td>
</tr>
<tr>
<td>5</td>
<td>Goniastrea</td>
<td>Encrusting</td>
<td>1.50</td>
<td>Paga (Intertidal)</td>
</tr>
<tr>
<td>6</td>
<td>Platygyra</td>
<td>Massive</td>
<td>1.50</td>
<td>Laku Point (Intertidal)</td>
</tr>
<tr>
<td>7</td>
<td>Turbinaria</td>
<td>Foliose</td>
<td>4.00</td>
<td>Laku Point (Intertidal)</td>
</tr>
<tr>
<td>8</td>
<td>Turbinaria</td>
<td>Foliose</td>
<td>4.00</td>
<td>Laku Point (Intertidal)</td>
</tr>
</tbody>
</table>

Figure 5 shows the in situ reflectance spectra of these coral targets over the spectral range of 350 to 1350 nm. All corals have central maxima near 820 nm and another prominent peak at 1070 nm. The spectra are marked with two characteristic absorption features located at 675 nm (chlorophyll absorption) and at 975 nm (water absorption). The chlorophyll absorption at 675 nm is followed by an abrupt, steep rise in the NIR region. As per the spectral characteristics apparent in Figure 5 the eight coral targets can be grouped into three distinct groups: Group A consisting of Favia, Porites, Goniastrea and Platygyra; Group B comprising of Symphillia, Goniopora and Turbinaria (Live) and Group C: the bleached Turbinaria.

Group A corals represent massive, sub-massive and encrusting colonies with equal proportion of exposure of soft, live coral polyps and their calcium carbonate corallites. Two of the Group B corals (Symphillia and Goniopora) representing massive and sub-massive colonies had relatively more exposure of the soft, live coral polyps than the calcium carbonate corallites. Turbinaria (live) on the other hand, represents a foliose colony with more of a calcium carbonate structure with live coral polyps. Group C: the bleached Turbinaria represents the same with polyps largely devoid of endo-symbiotic zooxanthellae.

4. RESULTS & DISCUSSION

Favia was the most dominant scleractinian genus, ubiquitously occurring on the inter-tidal reef flat and also in the sub-tidal areas of Laku Point and Paga reefs. The second dominant genus was Porites followed closely by Goniastrea and Goniopora in both the locations. Symphillia was found only in the sub-tidal zones of Paga while Turbinaria were sampled from Laku Point. Platygyra was sampled from Paga reef.

Figure 4. Field photographs of the eight coral targets (numbered serially)
characteristic spike located near 760 to 762 nm. This spike is however a contribution from the atmospheric oxygen and brings out the limitation of the instrument to outweigh this spike while sensing and recording the target spectra. *Favia* spectra shows a small hump at 728 nm which is absent in rest of the Group A corals. The spectra show a rapid descent from 820 to 840 nm followed by a gradual descent over 840 to 910 nm region. Beyond 910 nm the spectra are characterized by a steep descent to the water absorption feature located at 975nm. The absorption band depth appears to be a function of the depth of overlying water column. In the MIR region, the spectra show similar symmetrical bell-shape architecture with central maxima located at 1070 nm. Beyond 1150 nm up to 1350 nm the spectra show a featureless straight line trend except *Platygyra* which show a plateau at 1270 nm.

In case of Group B corals, the spectral architecture as shown by the Group A corals get considerably modified in the NIR region. The triple-peaked pattern is quite pronounced in case of *Turbinaria* and *Goniopora* while gets subdued for *Symphyllia*. In NIR region these corals exhibit a hump at 725 nm which from Group A only *Favia* showed at 728 nm. Contrary to the Group A corals’ steady rise from 710 nm to 820 nm Group B corals show a peak and valley pattern in this region. Group B corals exhibit a pronounced descent from 820 to 840 nm deviating from the Group A trend of gradual descent over 840 to 910 nm. *Turbinaria* and *Goniopora* show a steep, stepped fall from 840 to 890 nm, 900 to 930 nm and 933 to 999 nm interrupted by small spikes at 898 nm and 933nm. This is due to the ripple effect of the water column present on the targets. The water absorption trough centered at 975 nm is present for both. *Symphyllia* though conforms to the trend of steady descent from 820 to 840 nm but deviates from the Group B trend in 840 to 950 nm region by showing a convex drop up to 935 nm. The water absorption feature of *Symphyllia* is absent from display in the 950 to 1000 nm region because of negative reflectance values indicating a strong water column absorption since this coral was submerged under 8 cm water column– the maximum recorded on field. In MIR region however, Group B corals conform to the trend of Group A corals.

Group C represented by the bleached coral spectrum of *Turbinaria* is characteristically different from the live corals in UV-Visible region with relatively much higher reflectance values, shooting up to its maximum at 590 nm (six times as that of its live counterpart). Right from 350 to 600 nm the bleached coral spectrum rises steadily with minor breaks of slopes. Between 590 and 650 nm, this spectrum again show a stepped pattern of descent with intermittent breaks of slopes. Thereafter, it plunges down to the chlorophyll absorption trough located at 675 nm. Unlike the live coral spectra, the bleached coral spectrum has a stepped rise up to 710 nm. Beyond 715 nm to 1350 nm this spectrum closely follows the trend of live *Turbinaria* with characteristic local shoulders and troughs getting vertically pronounced.

In order to exaggerate the spectral shapes and enhance the subtle features of the zero-order spectra, the first and second order derivatives (Figure 6 and 7 respectively) were numerically calculated over 4nm as finite band resolution. Figure 6 shows 700 and 760 nm as the locations corresponding to peaked coral reflectance for all the eight corals. However, the peak at 760 nm is not a contribution from the target. In the visible region two peaks out of that triple peak at 575 and 600 nm also become prominent in the first derivative spectra. The first order derivatives also bring out the characteristic absorption troughs located over 650 to 675 nm, 800 to 850 nm, 900 to 950 nm and 1100 to 1150 nm for all the eight corals. Moreover, the first derivative of averaged reflectance appears to provide a good separability between bleached and live corals over a large spectral region from 430 to 590 nm. At 631 and 647 nm the first derivative values of all the live corals are negative while for the bleached coral it is positive.

The second derivative of the averaged reflectance identifies 680 to 700 nm region as a window to separate live and bleached coral spectra. In this region all the live corals record a positive second derivative value while the bleached coral has a negative value. The second derivative plot also shows the enhanced reflectance and absorption features of *Symphyllia* which rather showed a subdued zero-order signal because of the maximum water column.

### 5. CONCLUSIONS

The results obtained in this preliminary study suggests that *in situ* hyperspectral signatures of corals can be used as an essential input to study the spectral behaviour of reef building corals. All seven live corals conform to the general belief of strong reflection
in the NIR region (Green et al. 2000) with central maxima at 820 nm and a secondary peak at 1070 nm in the MIR region. The live corals exhibited “brown mode” of coral reflectance with triple-peaked pattern in the visible region. All spectra have a characteristic spike located at 760 to 762 nm due to the presence of atmospheric oxygen. The coral spectra are characterized by two distinct absorption features located at 675 nm (chlorophyll absorption) and 975 nm (water absorption). The absorption band depth at 975 nm is a function of the depth of water column and water column does suppress the reflectance magnitude. Ripples on moving water column drastically change the spectral shape of corals at 840 nm. The live and the bleached coral spectra can be well distinguished in the 400 to 600 nm region. The relative exposure of coral polyps and corallite structure can be a major determinant in the spectral behaviour of the corals at colony scale. The first and second order derivatives can be employed to identify common reflectance and absorption regions of the coral targets along with potential region(s) of separation between live and bleached corals. Live and bleached corals get differentiated with first-order derivatives at 631 and 647 nm while 680 to 700 nm window show their clear separation with second-order derivatives. However, bleached coral target needs to be sampled more in order to strengthen these observations. Derivative analysis needs to be investigated in depth and with more number of samples to comment on the spectral separability of the coral targets.

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7. ACKNOWLEDGEMENTS

This study presents the preliminary results of the work being carried out under Optical Characterization of Coral Reef Diversity Project under Meteorology and Oceanography Programme of Space Applications Centre, ISRO. The author is thankful to Director, Space Applications Centre and Dr. J. S. Parihar, Deputy Director, EPSA for overall technical guidance and support for this study. The work has been carried out in collaboration with GEER Foundation, Gandhinagar. The author sincerely acknowledges the contribution of all the members of project team of GEER Foundation and Space Applications Centre, ISRO.