Environmental Monitoring in Offshore Areas

Sonali Kumbhare*, Dilip S. Ramteke and Pravin Charde

Sevadal Mahila Mahavidhyalaya, Nagpur, India
*Corresponding author

A B S T R A C T

The purpose of offshore environmental monitoring is to provide an overview of the environmental status and of trends over time seen in relation to offshore oil and gas activities. Monitoring is intended to indicate whether the environmental status is stable, deteriorating or improving, due to operator’s activities. There is lot of pollution in sea due to discharge of various chemicals. So, it is very essential to study the pollution impact on the environment. This paper is detailing about how the environmental monitoring in offshore areas is carried out. Monitoring of Marine water environment is essential to know the pollutants generated from the oil extraction activity.

Keywords: Environment, Monitoring, Pollution and Marine water.

Introduction

Environmental Quality monitoring is an essential component of any environmental impact assessment study conducted for developmental activities. While assessing the likely impacts due to development of oil / gas fields and subsequent proposed production in offshore region, monitoring of Marine water environment is essential as it serves as the ultimate sink for all the pollutants generated from the oil extraction activity. In the offshore region, the data pertaining to oceanography, meteorology, physico-chemical characteristics of sea water, sediments, benthos, fishery resources and primary/secondary productivity of biological species in required to be generated.

The water quality monitoring factors that influence are:

- Selection for Sampling locations
- Type of Samples
- Sampling Frequency
- Number of Samples
- Sampling Facilities
- Sampling Equipment’s for different parameters
- Selection of Parameters
- Samples collection frequency and size of sample
- Preservation of samples for different parameters
- Use of online analysers for in situ measurement/ on board analysis of field parameters (Non-conservative determinants which changes with time and could not be stabilized)
Selection of sampling location

In offshore region, sampling location in such areas where the maximum activities (oil and gas fields, processing platforms, fishing zone, tanker loading and unloading, ship routes etc.) are in existence. Generally complete zone is selected. The zone is further divided into different grids and sampling is carried out at the centre of the grid as per the latitude and longitude. In Bombay, offshore areas (west coast) there are different oil and gas fields and processing platforms. Offshore drilling and oil processing activities are in the progress in that particular area. These activities are polluting aquatic environment up to certain extent however the major pollution may occur due to petroleum hydrocarbons release mainly by ballast water and tank washing aboard ships during transportation (1). So there is necessity of regular water quality monitoring. Figure 4 shows the offshore production areas along the coast line in the form of grid pattern after selecting the site.

Sampling frequency

Water samples should be collected at intervals so that no change in quality could pass unnoticed. The quality of water in various water bodies is rarely if ever constant in time but is subjected to change. The larger the number of samples from which the mean is derived, the narrower will be the limits of the probable difference between observed and the true values. However, the sampling schedule is a compromise between accuracy and the funds, personnel available for the work (1, 2).

Sample collection

Sample from various depths can be collected by using any of the sampler, commercially available. A wide range of indigenous automatic sampling equipments are also available for continuous sampling or at fixed intervals. Any sampling technique may be used; however, adaptation of a particular technique depends upon what is being sampled and what constituents are to be determined (3).

Sample preservation

Between the time that a sample is collected in the field and it is actually analysed in the laboratory, physical changes and chemical/biochemical reactions, may take place in the sample container which will change the intrinsic quality of the sample. It is therefore necessary to preserve the samples to prevent or minimize their changes. This is done by various procedures such as keeping the samples in the dark, adding chemical preservatives (Table 1), lowering the temperature to related reactions by freezing of by a combination of the methods (4).

Selection of parameters

The selection of parameters is made according to the type of samples being collected in particular area. In general, the parameters are grouped in the following way:

Meteorological parameters: Temperature, Rainfall, Humidity

Physical parameter: Colour, Turbidity, dissolved Solids, conductivity

Chemical parameter: Inorganic and organic groups

Inorganic group

pH, alkalinity, chloride, sulphate, sulphide, salinity and heavy metals like, Fe, Mn, Cr, Cu, Pb, Zn and Cd.

Organic group

Dissolved oxygen, bio-chemical oxygen demand, chemical oxygen demand, hydrocarbons and oil and grease.
Nutrient parameters: Total nitrogen, nitrate nitrogen, total phosphate and orthophosphate

Biological parameters: Phytoplankton and zoo-plankton, water fowl, algae

Bacteriological parameter: total coliform, E. coli, Streptococci, Faecal coliform

A practical way to select suitable parameters for an individual water body is by means of a preliminary investigation programme, which provides an overview of the biological and chemical loads on the ecosystem.

**Marine abiotic environment**

Marine abiotic environment comprises two major components, viz. water column and sediments. It is important to study spatial and temporal variations in physico-chemical characteristics of water column for assessing the impacts due to discharges of drilling fluids, cuttings and produced water. The sediments act as major sink for accumulation of wastes hence it is required to study their characteristics for assessment of environmental impacts due to the drilling operations. Data on physico-chemical characteristics including levels of oxygen and nutrients, levels of heavy metals and hydrocarbons both in water column and sediments have therefore been required to be collated, critically reviewed and used for assessment of environmental quality (3).

**Water column**

Baseline environmental Quality of water column and sediments is mainly governed by hydrographical and meteorological features of the region which includes topographical features, winds, surface currents, water movements, waves, convergence and divergence zones and temperature. The water current and direction can be measured by lowering a water current meter DRDF from the vessel into sea at sampling point located at the centre of the grid under study and current speed and direction can be measured at various depths.

**Physico-chemical characteristics**

Seawater samples need to be collected at three depths (a) Surface, (b) Mid-depth of water column and (c) 2 m above the bottom of water column in each of the grid using Nishkin type sampler (Figure 5).

The field parameters can be estimated immediately onboard after sample collection whereas for other physico-chemical and biological parameters, ‘samples can be processed and preserved following the standard procedures. Standard analytical methods (5) as enlisted in Table 2 are practiced for analyzing the preserved samples. Following are the significant Parameters to be analysed for marine water quality, monitoring.

**Turbidity**

Turbidity results due to suspended substances such as silt, clay and/or planktonic organisms. It restricts the penetration of sunlight and hence reduces photosynthesis which in turn is related to productivity of biomass. Suspended particles causing turbidity may also adsorb considerable amount of nutrient, viz. phosphate, potassium and nitrogen in their ionic forms and hence are not available for plankton production. Turbidity caused by colloidal particles of different thermal properties also influences the temperature conditions of the biomass by restricting the penetration and scattering of sunlight. The turbidity levels during post-monsoon are higher than pre-monsoon. High turbidity values during post-monsoon can be attributed to discharges to suspended solids load in the
monsoon months coupled with resuspension of sediments caused by turbulence in the water column. Turbidity can be measured on board using a Hach or equivalent turbidimeter working on naphelometric principle.

**Salinity**

Salinity is a conservative parameter. It does not change with in-situ biochemical processes. Temporal and spatial variations in salinity are important to understand the dynamics of water column.

The salinity is estimated based on measurement of conductivity and chloride. Variation in surface salinity is caused by precipitation, evaporation and process mixing.

**pH**

pH of seawater varies between 7.0 and 8.5 indicating neutral to marginally alkaline conditions. Highly acidic or alkaline pH would be detrimental to biotic environment as well as biological processes occurring in the sea. The pH is measured on board using portable pH meter or by lowering a probe into water column at various depths for in situ measurements.

**Dissolved oxygen**

Dissolved oxygen in sea water is an essential requirement for the survival of aquatic organisms. The dissolved oxygen content from surface layers down to 50 m is more or less uniform. From 75 m downward there is a rapid decrease in the oxygen content and it attains the lowest value at a depth around 150m.

The DO is estimated on board either by Winkler method or by lowering the probe at depths and recording the DO value from portable DO meter.

**Nutrients**

In general, phosphate and nitrate show regular trend of distribution whereas nitrite shows an irregular pattern of distribution with depth. ‘It is assumed that the depthwise and regional distribution of nutrient components and hydrographic parameters appear to be more depended on dilution, general current pattern and source of water masses. Samples for ammonia, nitrite, nitrate need to be preserved on board and be analysed using spectrophotometric methods or using probes with ISE and portable ionmeter.

**Heavy metals**

Heavy metals discharged during oil/gas exploration and extraction activities have been reported to cause great impact on marine environment due to their refractory nature. It has been established (6) that metals such as barium, cadmium, chromium, copper, lead and zinc are present in substantial concentration in drilling discharges. Spatial and temporal distribution as well as chemical forms of these metals in sea would influence the bioavailability and biodesorption of these metals.

Samples for heavy metals are preserved with nitric acid immediately on board for total metals and for dissolved metals after filtration through 0.45 μm filter paper. The estimation is carried out by either AAS or ICP after preprocessing the samples.

**Hydrocarbons**

Transport of crude oil and petroleum products across Arabian Sea amounts to 576 X 106 tonnes/year. This is about 54% of the total oil transported through marine waters in the world. Out of this 340 X 106 tonnes are carried to the Far East and Japan. Oil spilled annually at sea has been estimated as 2.2 to 4.9 million tonnes (7 and 8).
Water samples should be collected in appropriate containers and analysed for hydrocarbons using carbon tetrachloride as solvent for extraction. Prior to analysis the extracts are subjected to treatment with silica gel for removal of organics other than hydrocarbons. Total petroleum hydrocarbons are then estimated using fluorescence spectroscopy (10). The reference standard is prepared using chrysene which represents petroleum hydrocarbons of natural origin as it cannot be biosynthesized. The photometric readings for extracts of samples are recorded at excitation and emission wavelengths of 318 nm and 362 nm respectively and the concentrations of hydrocarbons are determined using linear relationship of photometric data and standard hydrocarbon concentrations. The sequential procedure for hydrocarbon analysis of water sample is shown below.

**Sediments**

The marine environment is subjected to considerable stress through deliberate or accidental oil spills, ballast water discharges, dredging and infilling for coastal development, uncontrolled sewage and industrial wastewater discharges and oil and gas exploration activities. Most of the pollutants find their way to sediment which serve as sink. During drilling and development of oil and gas wells a wide variety of solids and liquids are produced on the platform, some of which are discharged directly into the ocean. Most of the inorganic elements potentially toxic to benthic biota will eventually reach the sediments. Continuous discharges of solid and liquid from platform will have significant impacts on meio and macro benthic biota (9, 10).

**Hydrocarbons**

The presence of hydrocarbons in sediments of offshore region can be considered as the indicator of pollution due to drilling and production activities that have undertaken. Additionally, hydrocarbons can also originate from urban and industrial effluents, accidental oil tanker discharges and discharges from ships sailing along the trade and tanker routes. Therefore identification of a specific source of hydrocarbon discharging into sea can be done only if data on composition of hydrocarbon from these sources are available and are correlated with the measured values.

In order to analyze hydrocarbons in sediments each preweighed sediment samples should be placed in soxhlet apparatus and refluxed for 8 hours using mixture of methanols and benzene (1:1 as solvent). The extracts obtained on refluxing should be subjected to preconcentration in a rotary evaporation and
individually passed through column of silica gel for removal of organics other than hydrocarbons and estimated using infrared spectroscopy at 2930 cm⁻¹ wavenumber (II). The sequential procedure for hydrocarbons estimation in sediment is given below:

**Marine biotic environment**

Biotic environment comprises micro-organism, phytoplankton, zooplankton, benthos and nekton. While phytoplankton are limited to the photic zone, other marine animals tend to move away from this zone. Further, geographic distribution of plankton depends on the habitat, water currents and the tolerance of organisms as water circulates in gyre systems.

Biological productivity of the ocean is influenced by factors, viz. light, nutrients, primary, secondary and tertiary production. Light penetration in the ocean determines the depth of the photosynthetic zone whereas the nutrients particularly nitrogen and phosphorus influence the promotion of productivity. Phytoplankton or the planktonic algae give the production at the primary stage of the food chain while zooplankton and fish are at the secondary and tertiary levels of the food chain.

**Fig.1 Sequential procedure for hydrocarbon analysis of water samples**

- **Water Sample**
  - Adjust pH at less than 2
  - Extraction with Hexane
    - Pass through anhydrous Na₂SO₄
  - Evaporate the Hexane at Room Temperature
  - Add Carbon Tetrachloride and Silica Gel
    - Stir in Magnetic Stirrer
      - Pass through Na₂SO₄
  - Make up Volume 25 ml with Carbon Tetrachloride
  - Analyse on Fluorescence spectrophotometer
Fig. 2 Sequential procedure for hydrocarbon estimation in sediments

1. Sediment
2. Freeze Drying
3. Soxhlet Extraction (1+1 Methanol, Benzene for 8 hours)
4. Rotating Evaporation under Vacuum (T=30°C)
5. Crude Residue
   - Extraction with 30 ml CCl₄
   - Solvent Extraction
     - Pass through Na₂SO₄
     - Solvent Extraction
       - Pass through Silica Gel Column
     - Hydrocarbon Extraction
6. Analyze on Fluorescence Spectrophotometer
Fig. 3 Sequential procedure for hydrocarbon estimation in fish tissues

100 gms Fish Tissues + 10 ml, 4 N NaOH
→ Reflux at 80°C
→ Suspension
→ Extract with CCl₄

Suspension
→ Solvent
→ Pass through Na₂SO₄

Collect in 100 ml Volumetric Flask Containing 10 gm Silica Gel

Clear Hydrocarbon Extract
→ Analyse on Fluorescence Spectrophotometer

Suspension
→ Extract with Fresh 20 ml CCl₄

Extract with Fresh 20 ml CCl₄
→ Solvent
→ Pass through Na₂SO₄

Extract with Another fresh 20 ml CCl₄
→ Solvent
→ Pass through Na₂SO₄

Solvent
→ Reject

Suspension
Fig. 4 Water quality monitoring stations
Fig. 5 Nisikin type water sampler with reversible thermometer

Fig. 6 Van Veen Grab sampler
**Fig.7** H. T. Net for plankton collection

**Table 1** Sampling containers and preservation

<table>
<thead>
<tr>
<th>Sr. No.</th>
<th>Determination</th>
<th>Container</th>
<th>Preservation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Temperature</td>
<td>Polyethylene or Glass bottle</td>
<td>Analyze immediately</td>
</tr>
<tr>
<td>2.</td>
<td>Turbidity</td>
<td>Polyethylene or Glass bottle</td>
<td>Analyze same day, store in dark up to 24 hrs</td>
</tr>
<tr>
<td>3.</td>
<td>Total Solid</td>
<td>Polyethylene or Glass bottle</td>
<td>Refrigerate</td>
</tr>
<tr>
<td>4.</td>
<td>Total Dissolved solid</td>
<td>Polyethylene or Glass bottle</td>
<td>Refrigerate</td>
</tr>
<tr>
<td>5.</td>
<td>pH</td>
<td>Polyethylene or Glass bottle</td>
<td>Refrigerate</td>
</tr>
<tr>
<td>6.</td>
<td>Alkalinity</td>
<td>Polyethylene or Glass bottle</td>
<td>Refrigerated</td>
</tr>
<tr>
<td>7.</td>
<td>Salinity</td>
<td>Glass bottle, wax seal</td>
<td>Analyze immediately</td>
</tr>
<tr>
<td>8.</td>
<td>Sulphide</td>
<td>Polyethylene or Glass bottle</td>
<td>Refrigerate, N Zinc acetate/ 100 ml</td>
</tr>
<tr>
<td>9.</td>
<td>Sulphate</td>
<td>Polyethylene or Glass bottle</td>
<td>Refrigerate</td>
</tr>
<tr>
<td>10.</td>
<td>Nitrate</td>
<td>Polyethylene or Glass bottle</td>
<td>Refrigerate, analyze soon</td>
</tr>
<tr>
<td>11.</td>
<td>Phosphate</td>
<td>Glass bottle (Rinsed with 1 + 1 HNO₂)</td>
<td>Refrigerate, analyze soon</td>
</tr>
<tr>
<td>12.</td>
<td>Dissolved oxygen</td>
<td>Glass, BOD bottles</td>
<td>Analyze immediately</td>
</tr>
<tr>
<td>13.</td>
<td>BOD</td>
<td>Polyethylene or Glass bottle</td>
<td>Refrigerate</td>
</tr>
<tr>
<td>14.</td>
<td>COD</td>
<td>Polyethylene or Glass bottle</td>
<td>Add H₂SO₄ to pH &lt;2</td>
</tr>
<tr>
<td>15.</td>
<td>Oil and grease</td>
<td>Glass, wide – mouth calibrated</td>
<td>Add H₂SO₄ to pH &lt;2</td>
</tr>
<tr>
<td>16.</td>
<td>Hydrocarbon</td>
<td>Glass bottle</td>
<td>Refrigerate</td>
</tr>
<tr>
<td>17.</td>
<td>Metals</td>
<td>Glass bottle</td>
<td>Add HNO₃ to pH &lt;2</td>
</tr>
</tbody>
</table>
Table 2 Physico-chemical analysis and methodology

<table>
<thead>
<tr>
<th>Sr. No.</th>
<th>Parameters</th>
<th>Method</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Alkalinity</td>
<td>Titrimetric Method/ Electrometric Method</td>
</tr>
<tr>
<td>2.</td>
<td>Chloride</td>
<td>Argentometric Titration Method</td>
</tr>
<tr>
<td>3.</td>
<td>Silica</td>
<td>Spectrophotometric method</td>
</tr>
<tr>
<td>4.</td>
<td>Sulphate</td>
<td>Turbidimetric Method</td>
</tr>
<tr>
<td>5.</td>
<td>Nitrate</td>
<td>Spectrophotometric method or UV Spectrophotometric method</td>
</tr>
<tr>
<td>6.</td>
<td>Phosphate</td>
<td>Ammonium Molydate Spectrophotometric method</td>
</tr>
<tr>
<td>7.</td>
<td>Heavy Metals</td>
<td>Atomic Absorption Spectrophotometric method or ICP</td>
</tr>
<tr>
<td>8.</td>
<td>Oil and Grease</td>
<td>Partition Gravematric Soxhiet Extraction Method</td>
</tr>
<tr>
<td>9.</td>
<td>Dissolved Oxygen</td>
<td>Winkler’s Modified Method</td>
</tr>
<tr>
<td>10.</td>
<td>Chemical Oxygen Demand</td>
<td>Open/ Closed Refluxing followed by Titration with FAS</td>
</tr>
<tr>
<td>11.</td>
<td>Bio-Chemical Oxygen Demand</td>
<td>5 days 20°C (Do estimation by winkler Method)</td>
</tr>
<tr>
<td>12.</td>
<td>Nitrogen (Total)</td>
<td>Digestion followed by distillation and Titrimetric Method</td>
</tr>
<tr>
<td>13.</td>
<td>Sulphide</td>
<td>Iodometric Method</td>
</tr>
<tr>
<td>14.</td>
<td>Hydrocarbons</td>
<td>Fluorescence Spectrophotometric or Infrared Spectrophotometric Method</td>
</tr>
</tbody>
</table>

**Phytoplankton**

Phytoplankton, mostly the unicellular organisms are either solitary or colonial. These autotrophs synthesize organic materials from inorganic substance in the presence of sunlight through the process of photosynthesis. Consequently, the depth of penetration of the light in seawater controls this process. Phytoplankton provide food for herbivores and hence form a major link in the food chain.

Samples for phytoplankton analysis should be collected from surface by vertically hauling the sample bucket. 750 ml of sample should be stored in 4% neutral formalin. The samples are then concentrated using centrifuge and further examined under microscope to identify the species and their number. This data is further used for determining Shannon Weaver Diversity Index of individual sampling location (12).

**Zoooplankton**

Zoooplankton form an important link in food chain and act as both primary consumer of marine aquatic ecosystem and a food source for many aquatic organisms including fishes. Unlike the freshwater zooplankton, the marine zooplanktons show considerable diversity as it is composed of members of every group from protozoa to chordata. Depending on seasons the plankton community show pronounced variation in its character and composition.

The zooplankton are indicators of the general fertility and water quality of a sea area. An imbalance in its population structure could bring about far reaching effects on the dependent fishery resources. The imbalance could be brought about by natural as well as man-made reasons. Fluctuations in the environmental conditions resulting in poor upwelling, rise in sea surface temperature,
underwater disturbances, altered monsoons and water currents from natural causes affect zooplankton productivity and species diversity while pollution especially due to oil spills represent one of the major man-made causes.

Samples should be collected using H.T. plankton net (300 microns) to represent all the available groups of zooplankton through a horizontal haul for 15 min duration at constant speed in each grid (Figure 6). Thesamples are to be fixed immediately with 5 percent formalin solution.

Benthos

The organisms which inhabit the bottom of aquatic body are known as benthos. Many of them are sessile, some creep over or burrow in mud and base of water body.

The quality and quantity of animals found at the bottom is not only related to the nature of substrata but also to the depth and type of aquatic plants present in such an environmental. Their number and distribution also depend upon physic-chemical characteristics of water and biological complexes such as food and other factors.

The bottom mud can be collected from various sampling points by Van Veen grab of 250 cm² size. For macrobenthos the sediment is sieved through 500 µ sieve and the organisms retained on the sieve are preserved immediately with 4 percent neutral formalin (without rose bengal).

A subsample of sediments is passes through 50 µ sieve for segregation of meiobenthos. The sediments along with meiobenthos retained on 50 µ sieve are preserved immediately for subsequent analyses in the laboratory (3).

Fisheries

India with 6100 km long coastline, continental shelf of 0.451 million sq. km and exclusive economic zone (EEZ) of 2.02 million sq.km has rich marine fishery potential. It is also established that coastal waters particularly off west coast are highly productive and act as spawning and nursery grounds for several commercially important fish species.

Existing fish species should be collected and identified by operating a bottom trawl net in” different locations by dividing the study area into uniform grid size. Fish catch should be weighed to estimate the yield of the individual species. The tissues viz. muscles, gonad, liver and gills are dissected out. Each of the tissues are weighed and preserved deep frozen for subsequent analysis (Heavy metals and hydrocarbons) in the laboratory (14). The sequential procedure for hydrocarbon estimation in fish tissues is given below:

Prevention of sample contamination

Field measurements should always be made on separate sub-sample which is to be discarded once the measurement have been made.

Sample bottles, new or used, must be cleaned according to the recommended methods.

Water sample bottle should be employed for water samples only. Laboratory bottles to store concentrated reagents should never be used.

Recommended preservation methods must be used. All preservatives must be of analytical grade.

The inner portion of the sample bottle and caps should not be touched with bare hands.
Sample bottle must be kept in a clean environment, away from dust, dirt, fumes and grime.

Petroleum products (gasoline, oil exhaust fumes) are prime sources of contamination. Spills or drippings must be removed immediately.

Filter units and related apparatus must remain sterile, until the sample is collected.

Specific conductance should never be measured in the sample water that was first pH measurement.

Samples must never be permitted to stand in the sun; they should be stored in cool place, ice chests are recommended.

Samples must be transferred to laboratory without delay.

Additional sampling precautions are required to be taken depending upon samples conditions at a particular location.

References

Reeve, R. N., (2002); “Introduction to Environmental Analysis” John Wiley and sons; England, PP 11-22


Kaul S.N. and Ashutosh Guatam; Daya Publishing House, New Delhi

Method of Soil analysis, Part – 11, American Society of Agronomy, ING publishers, Madison, Wisconsin USA (1965)


EPA (1985), “Assessment of Environmental Fate and Effects of Discharges from offshore oil and Gas operations” EPA 440/4-85/002

Monaghan et.al. (1980) “Marine Environmental Pollution” Elsevier Science Publication Vol. I PP 413-431


Law R.J. (1981); “Hydrocarbons concentration is waters and sediments from UK marine waters, Determined by Fluorescence Spectroscopy” Marine poll. Bull; 12(5); PP 153-157


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