Hydrogeochemistry and surface water quality assessment of IB valley coalfield area, India

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Abstract
Coal mining and ancillary activities have the potential to cause water pollution characterized by acid mine drainage, acid mine leachates, extreme pH conditions and heavy metal contaminations. In the present work, 33 water samples in premonsoon and 34 water samples in monsoon were collected from the surface water bodies of Ib Valley coalfield, India for hydrogeochemical analysis. In premonsoon, pH, TSS, Turbidity, DO, BOD, COD, Magnesium, Cadmium, Selenium, Nickel, Aluminum and in monsoon, pH, TSS, Turbidity, DO, BOD, COD, Iron, Cadmium, Selenium, Nickel and Aluminum were nonconforming to the permissible limit set by the Bureau of Indian Standards, World Health Organisation and Ministry of Environment, Forest and Climate Change, Government of India. The average BOD/COD ratio of less than 0.6 in both seasons indicated Ib valley coalfield water was not fairly biodegradable. The analysis of variance (ANOVA) revealed that significant seasonal variation \((p < 0.05)\) was observed in the hydro-chemical parameters viz. TSS, turbidity, redox potential, acidity, total hardness, bicarbonate alkalinity, chloride, sulfate, nitrate, sodium, calcium, magnesium, iron, cadmium, chromium and magnesium during the entire sampling period. Whereas, no significant seasonal variation \((p > 0.05)\) was observed in pH, EC, TDS, DO, BOD, residual chloride, COD, oil and grease, fluoride, potassium, zinc, copper, selenium, nickel, aluminum, boron, silica, temperature, salinity, cyanide and phenol. Water Quality Index revealed that 39.39% and 35.29% samples belong to poor water quality category in premonsoon and monsoon, respectively. As per Heavy Metal Pollution Index, Degree of Contamination (Cd) and Heavy metal evaluation index, medium degree of pollution were exhibited by 51.52%, 30.30% and 45.45% samples in premonsoon and 20.59%, 35.29% and 26.47% samples in monsoon. Whereas, 5.88%, 2.94% and 5.88% samples were having high degree of pollution with respect to Cd in premonsoon. However, EC, Na%, PI, SAR and RSC values suggested that the water can be used for irrigation. Water type of the region had been found to be Ca–Mg–Cl–SO₄ by Piper diagram.

Keywords Ib valley coalfield · Coal mining · Water quality index · Heavy metal pollution index · Degree of contamination · Heavy metal evaluation index

Introduction
Coal continues to be one of the major sources of non-renewable energy. To meet the increased demand for industrial sectors such as steel, power, cement there is a continuous increase in coal production in our country, and there is greater stress on opencast mines to achieve the target. Coal mining is carried out broadly by two methods, viz. opencast and underground mining. Opencast mining technique is the preferred method and contributed to 91.22% of India’s total coal production in 2013–2014 (Provisional Coal Statistics 2013–2014). The total production of coal for 2016–2017 was estimated at 724.71MT (Ministry of Coal 2016–2017). With increase in depth of mine, the workings in many opencast
mines extend below the water table. The mine excavations, therefore, are normally associated with a high degree of water influx due to the ingress of groundwater. A large quantity of water also accumulates in the deep part of the mines or mine sumps during monsoon. The mine waters in many mines is usually pumped and discharged outside the mine boundary, which end up in polluting the nearby water bodies. The water in Jharia, Raniganj and Northeastern Coalfields was found to be contaminated with high amount of sulfate (Singh 1988). Tiwary and Dhar (1994) observed that the key pollutants connected with coal mines were suspended solids, dissolved solids, acidity and iron compounds. Pathak and Banerjee (1992) studied water quality in Chapha incline of Umaria coalfield in eastern Madhya Pradesh and found that the mine water was severely polluted with turbidity, BOD and bacterial colonies. The acidity in acid mine drainage is caused by hydrolysis of dissolved Fe (II), Fe (III), Al, and Mn (II), as well as the precipitation of associated solid hydroxide substances. The pH and concentrations of alkalinity, acidity, and metals such as Fe, Mn, and Al, in mine discharge and receiving aquatic system usually determine possible environmental hazards, corrosiveness and water toxicity (Environmental Protection Agency 2000). Effluent from Mines if found suitable can be utilized for irrigation. However, the high salt content in mine water causes an increase in soil solution osmotic pressure. A high salt concentration (high EC) in water leads to the formation of saline soil, while high sodium concentration leads to the development of alkaline soil. The salts besides affecting the growth of the plants directly, also affect the soil structure, permeability and aeration, which indirectly affect the plant growth (Singh 2011). Therefore, proper scientific investigation is necessary to ensure mine water is safe for irrigation. To assess the water quality for irrigation purpose, TDS, Electrical conductivity, Sodium absorption ratio (SAR), Percent sodium (%Na), Residual sodium carbonate (RSC), Permeability index (PI), Kelley index (KI) and Magnesium hazard (MH) are required to be examined. The Odisha State Pollution Control Board has now made it mandatory for all the operating mines to follow “zero liquid discharge” concept. This means that the water has to be stored within the mine boundary and reused for different mining applications such as dust suppression, firefighting, plantation activities. However, if it is not feasible, then the water can be discharged outside only after proper treatment; and installation of the online real-time water quality monitoring system at the point of discharge. In monsoon, the rainfall occurs over the entire lease area. A part of it infiltrates downwards into the water table, a lesser amount evaporates to the atmosphere, and the rest contributes to the surface runoff. Unless specific arrangements have been made, the surface runoff of the region takes its usual course moving through the overburden (OB) dumps, coal stocks, workshops and railway sidings, etc., into the nearby water bodies. Untreated mine water and Leaching of the mine wastes can affect the hydrological system and disrupt the ecosystem. This research aims to assess the seasonal variations in concentrations of various water quality parameters in mines sumps, treatment plants, river and ponds to ascertain the suitability of water for drinking and irrigation.

**Study area**

The state of Odisha in India has two major coalfields, viz. Talcher and Ib Valley, which are under the administrative control of Mahanadi Coalfields Limited, a subsidiary of the public sector enterprise Coal India Ltd. It has a coal reserve of 22.3 billion tonnes, the third highest in the country. Ib Valley Coalfield is located between latitudes 21°41’N and 22°06’N and longitudes 83°30’E and 84°08’E with an area of 1,375 square kilometers (531 sq. mi). It covers part of Jharsuguda, and Sundargarh districts of the state Odisha, India (Fig. 1). Currently, there are five opencast and three underground mines in operation in the Ib valley area, and two opencast mines in Basundhra area in this coalfield. The Ib-valley coalfield is a part of the NE-SW chief basin belt of the Son-Mahanadi valley. The location map of study area was prepared using the ArcGIS 10.5. software.

**Geology**

Ib Valley Coalfield is a portion of large synclinal Gondwana basin of Raigarh-Hingir and Chattisgarh Coalfields. The stratigraphic progression (Table 1) begins with Precambrian rocks at the base followed by an unconformity. Gondwana supergroups such as Talchir, Karharabari, Barakar, lower Kamthi (Raniganj) of Permian time and upper Kamthi Formation of Triassic time lie over the unconformity and Precambrian rocks. At the top recent and subrecent, deposits are found (Goswami et al. 2006; Senapati 2015).

The chief coal-bearing formations in the Ib Valley coalfield area are Barakar and Karharbari of lower Permian age. Barakar formation has around twenty carbonaceous horizons. These horizons after being clubbed together are recognized into five seams, viz. Belpahar, Parkhani, Lajkura, Rampur, and Ib. Parkhani seam is very poor in quality and thickness and is usually not considered as mineable. Belpahar seam is situated in the deep side of Lakhapur OCP, and in this seam, mining operation is not expected in near future. Karharbari formation holds only thin coal bands below Ib river (Senapaty and Behera 2015). The two predominant seams being mined in this coalfield, therefore, are Lajkura and Rampur.
Drainage and hydrogeology

The study area is tropical and comes across three seasons, viz. summer, monsoon and winter. The average annual rainfall is around 1527 mm. The monsoon season usually lasts from June to September. Tropical rain forest type natural vegetation is predominant in this area (Odisha State Pollution Control Board 2010). The Mahanadi river system along with its tributary Ib and Hirakud reservoir is the chief aquatic body in Ib Valley and Jharsuguda area. They provide huge water supply and carry waste and effluent load from this area. Three streams namely Lilary, Bhenden and Kharkhari in the area usually get discharged water from various industries, mines, domestic and agricultural sector. Runoff during monsoon from various stockpiles such as coal, mineral, solid waste gets down to the stream network which ultimately drains into the Hirakud reservoir. The critical water quality parameters in the surface as well as ground-water are found to be fluoride, mercury, cadmium and BOD (Odisha State Pollution Control Board 2010). The hydrogeology of the area consists of Consolidated Formation, Semi-Consolidated Formation and Unconsolidated Formation.

Consolidated formation

About 60% of the area is underlain by consolidated formations comprising Precambrian metasediment of Sambalpur series, Iron ore and Gangpur series. These rocks
are very hard and compact, and the groundwater occurs mainly in secondary porosity. Groundwater occurs under semiconfined to confined conditions in fractured rocks. Granite Gneisses are the most predominant rock types usually occupying the underlying plains and topographic lows. These rocks are highly weathered, and the thickness of the weathered zone usually ranges from 4.57 to 12.03 mbgl and depth to water level from 3.91 to 9.92 and 1.36 to 6.69 mbgl in premonsoon and postmonsoon, respectively. The specific capacity of open wells varies from 8.30 to 76.32 m³/day/m. The quartzite of the iron ore group is bedded, joined but having little groundwater development prospects. The depth of open wells varies from 6.94 to 9.37 mbgl and depth to water level from 5.99 to 8.70 and 3.26 to 4.45 mbgl in premonsoon and postmonsoon, respectively. The specific capacity of the open well tested was found to be of the order of 8.30 m³/day/m. Mica schists have low permeability. The depth of the open wells ranges from 6.11 to 7.84 m, and depth to water level varies from 4.12 to 7.1 and 2.53 to 3.18 m in premonsoon and postmonsoon, respectively. The specific capacity of the open well tested was found to be of the order of 8.30 m³/day/m.

**Semi-consolidated formation**

Sandstones, shales, conglomerates grits, etc., belonging to Talcher, Barakar and Kamthis of lower Gondwanas constitute the semi-consolidated formations. The needle shales with a high frequency of intersecting joints form good aquifers. The depth of the open well varies from 6.6 to 12.5 mbgl. The premonsoon and postmonsoon water levels vary from 5.4 to 10.25 and 1.6 to 7.5 m, respectively. The specific capacity is found to be 10.29 m³/day/m.

**Unconsolidated formation**

Laterites and alluvium of sub-recent to recent age constitute the unconsolidated formations. Laterites that occur in patches as capping the older formations and form a very good shallow aquifer. The coarse grained sand with gravels and pebbles form repository of groundwater which occurs under water table conditions. Alluvium forms potential shallow aquifer to be developed through dug wells (Central Ground Water Board 2013).

**Materials and methods**

**Sampling**

The Ib Valley Coalfield area has been classified into two different areas, viz. Ib Valley area and Basundhara area. Ib Valley is further classified into Samaleswari area and Lakhanpur area. Samaleswari area has two opencast projects (OCP) such as Samaleswari and Lajkura. Pandenjhor and Bagmara nallah make their course to Ib river through this zone (Fig. 2). Similarly, Lakhanpur area has three opencast projects viz. Lakhanpur, Belpahar and Lilari. Lilari nallah and puljari nallah are flowing in this region (Fig. 2). Basundhara area has two opencast projects viz. Basundhara and Kulda. Basundhara river and Basundhara nallah flow in this area (Fig. 3). A total of 77 numbers of water samples, of which 33 samples were collected during premonsoon (April–May) and 34 samples were collected during monsoon (June–July) period in 2015 for analysis. These samples were collected from the mine sumps, inlet and outlet of effluent treatment plants (ETPs), inlet and outlet of mine discharge treatment plant (MDTP) from 7 different mines as mentioned earlier. Additionally, water
samples were also collected, from the upstream, downstream and confluence points of nallahs and rivers where mine water are discharged, in order to study the pollution load carried out by these drains.

The details of the water sampling sources are presented in Table 2. The maps of the water sampling locations of the area are presented in Figs. 4 and 5.

Fig. 2 Drainage pattern in Ib valley area

Fig. 3 Drainage pattern in Basundhara area

Analytical methodology

For water quality analysis, the sampling bottles were thoroughly cleaned with 1:3 nitric acid and distilled water solution followed by twice with double distilled water and finally rinsed with water sample to be collected. Grab
Table 2: Details of water sampling locations and sources of Ib Valley Coalfield

| Name of the coalfield | Name of the area       | Sample No | Sampling Point description for both Premonsoon and monsoon | Source         | Sample codes | Coordinates       |
|-----------------------|------------------------|-----------|-------------------------------------------------------------|----------------|--------------|-----------------|
| Ib Valley Coalfield   | Basundhara Area        | 1         | Basundhara Nallah Down-stream                               | Nallah/Stream  | BSND         | N 22°03’57.2”  |
|                       |                        |           |                                                             |                |              | E 83°42’12.4”   |
|                       |                        | 2         | Basundhara Nallah Upstream                                 | Nallah/Stream  | BSNU         | N 22°04’6.40”   |
|                       |                        |           |                                                             |                |              | E 83°42’16.8”   |
|                       |                        | 3         | Basundhara OCP West Sump                                   | Sump           | BSWS         | N 22°03’50.9”   |
|                       |                        |           |                                                             |                |              | E 83°42’29.8”   |
|                       |                        | 4         | Basundhara OCP East Sump                                   | Sump           | BSEC         | N 22°03’9.5”    |
|                       |                        |           |                                                             |                |              | E 83°43’56.1”   |
|                       |                        | 5         | Basundhara OCP Oil Trap Inlet                              | ETP            | BOTI         | N 22°03’34.2”   |
|                       |                        |           |                                                             |                |              | E 83°43’53.1”   |
|                       |                        | 6         | Basundhara OCP Oil Trap Outlet                             | ETP            | BOTO         | N 22°03’35.9”   |
|                       |                        |           |                                                             |                |              | E 83°43’53.0”   |
|                       |                        | 7         | Basundhara River Down-stream near Kulda OCP                | River          | KD           | N 22°02’25.4”   |
|                       |                        |           |                                                             |                |              | E 83°44’52.7”   |
|                       |                        | 8         | Kulda OCP MDTP Outlet                                      | Mine Discharge  | KMO         | N 22°02’11.4”   |
|                       |                        |           |                                                             |                |              | E 83°45’0.3”    |
|                       |                        | 9         | Kulda Mine Sump                                            | Sump           | KMS          | N 22°01’54.7”   |
|                       |                        |           |                                                             |                |              | E 83°44’43.2”   |
|                       | Samaleswari Area       | 10        | Samaleswari OCP Railway Siding Inlet                       | Railway Siding | SRSI         | N 21°49’16.4”   |
|                       | (Samaleswari area)     |           |                                                             |                |              | E 83°54’52.0”   |
|                       |                        | 11        | Samaleswari OCP Railway Siding Outlet                      | Railway Siding | SRSO         | N 21°49’15.2”   |
|                       |                        |           |                                                             |                |              | E 83°54’51.2”   |
|                       |                        | 12        | Samaleswari OCP Sump                                       | Sump           | SSS          | N 21°47’55.2”   |
|                       |                        |           |                                                             |                |              | E 83°53’28.8”   |
|                       |                        | 13        | Samaleswari OCP ETP Inlet                                  | ETP            | SEI          | N 21°48’3.0”    |
|                       |                        |           |                                                             |                |              | E 83°54’30.1”   |
|                       |                        | 14        | Samaleswari OCP ETP Outlet                                 | ETP            | SEO          | N 21°48’3.0”    |
|                       |                        |           |                                                             |                |              | E 83°54’30.1”   |
|                       |                        | 15        | Samaleswari OCP MDTP Inlet                                 | Mine Discharge  | SMI         | N 21°47’58.7”   |
|                       |                        |           |                                                             |                |              | E 083°54’45.1”  |
|                       |                        | 16        | Samaleswari OCP MDTP Outlet                                | Mine Discharge  | SMO          | N 21°47’54.2”   |
|                       |                        |           |                                                             |                |              | E 083°54’49.5”  |
|                       |                        | 17        | Samaleswari OCP MDTP extended Pond Outlet(500m from existing Outlet) | Mine Discharge | SMPI        | N 21°47’29.9”   |
|                       |                        |           |                                                             |                |              | E 083°55’19.7”  |
|                       | Lajkura OCP Sump       | 18        | Sump                                                        | LS             |             | N 21°49’30.8”   |
|                       |                        |           |                                                             |                |              | E 083°54’07.4”  |
|                       | Lajkura OCP ETP Inlet  | 19        | ETP                                                         | LJEI           |             | N 21°49’30.2”   |
|                       |                        |           |                                                             |                |              | E 083°54’50.5”  |
|                       | Lajkura OCP ETP Outlet | 20        | ETP                                                         | LJEQ           |             | N 21°49’30.9”   |
|                       |                        |           |                                                             |                |              | E 083°54’50.8”  |
|                       | Belpahar OCP ETP Inlet | 21        | ETP                                                         | BEI            |             | N 21°45’40.6”   |
|                       |                        |           |                                                             |                |              | E 083°51’44.2”  |
|                       | Belpahar OCP ETP Outlet| 22        | ETP                                                         | BEO            |             | N 21°45’40.9”   |
|                       |                        |           |                                                             |                |              | E 083°51’44.6”  |
|                       | Belpahar OCP MDTP Inlet| 23        | Mine Discharge                                              | BMI            |             | N 21°46’01.4”   |
|                       |                        |           |                                                             |                |              | E 083°51’42.7”  |
|                       | Belpahar OCP MDTP Outlet| 24        | Mine Discharge                                              | BMO            |             | N 21°46’06.3”   |
|                       |                        |           |                                                             |                |              | E 083°51’45.9”  |
|                       | Belpahar Mine Sump     | 25        | Sump                                                        | BMS            |             | N 21°44’17.8”   |
|                       |                        |           |                                                             |                |              | E 083°50’26.3”  |
water sampling method was followed to collect the water sample. About 5 L of sub-surface water was collected by using a shallow water sampler. In situ temperature was recorded using a Brannan thermometer. Samples were collected in high-density polyethylene (HDPE) containers for the estimation of general parameters (1 L) and Biochemical Oxygen Demand (BOD) (2.5 L). Chemical oxygen demand (COD) samples were collected in 100 mL pre-cleaned Nalgene bottles and preserved with 0.2 mL concentrated sulfuric acid to have Power of hydrogen (pH) ≤ 2. DO was fixed in situ with Winkler’s reagents. Samples collected from Effluent treatment plant (ETP), Mine discharge treatment plant (MDTP), drains for ammonia were preserved with concentrated sulfuric acid. Samples for heavy metal analysis were preserved with concentrated nitric acid on the field to bring the pH

| Name of the coalfield | Name of the area | Sample No | Sampling Point description for both Premonsoon and monsoon | Source | Sample codes | Coordinates |
|-----------------------|-----------------|------------|-------------------------------------------------------------|--------|--------------|-------------|
| 26                    | Lilari Mine Sump | Sump       | LIS                                                         | N 21° 45’ 23.1" E 083° 52’ 53.16” |
| 27                    | Lilari Mine ETP outlet | ETP       | LIEO                                                       | N 21° 46’ 40.2” E 083° 51’ 24.6” |
| 28                    | Lilari Nallah Downstream near Belpahar OCP | Nallah/Stream | LIDB                                                       | N 21° 46’ 09.0” E 083° 51’ 49.5” |
| 29                    | Lilari Nallah Downstream near lakhanpur OCP | Nallah/Stream | LIDL                                                       | N 21° 46’ 09.8” E 083° 51’ 49.4” |
| 30                    | Lilari Nallah Upstream | Nallah/Stream | LIUP                                                       | N 21° 46’ 09.7” E 083° 51’ 49.9” |
| 31                    | Lakhanpur OCP Central sump | Sump       | LKCS                                                       | N 21° 45’ 28.3” E 083° 49’ 18.0” |
| 32                    | Lakhanpur MDTP Inlet Mine Discharge | Mine Discharge | LKMI                                                       | N 21° 46’ 51’ 10.3” E 083° 51’ 02.0” |
| 33                    | Lakhanpur MDTP Outlet Mine Discharge | Mine Discharge | LKMO                                                       | N 21° 46’ 12.9” E 083° 50’ 10.0” |
| 34                    | Puljari Nallah upstream (Only for Monsoon) | Nallah/Stream | PNUS                                                       | N 21° 45’ 47.2” E 083° 49’ 23.9” |

![Fig. 4](image-url) Map showing Ib valley area with location of sampling points
down to two in order to avoid metal precipitation. Samples for oil and grease were collected in 1 Litre amber bottle and preserved with concentrated hydrochloric acid in the field. All the samples were preserved at 4 °C until analysis. Before analysis, all the glasswares were cleansed with 1:3 nitric acid and distilled water solution followed by twice with double distilled water and finally rinsed with corresponding water sample. All the reagents used were from Merck, Germany, and were of analytical grade quality. Highly purified water prepared by a Millipore water purification system were used for the experimentation. Analysis of all samples was completed within 2 days after sampling, except for BOD. The analysis of water quality parameters was carried out following the standard methods prescribed in American Public Health Association (APHA 2012). Some of the parameters viz. pH, Oxidation reduction potential (ORP), turbidity, Total dissolve solid (TDS), Dissolve oxygen (DO), electrical conductivity (EC), salinity were measured on the site itself with the help of a multi-parameter water quality analyzer (Model: Horiba U52G). Furthermore, DO, pH and EC were also measured in the laboratory using Azide Modification of Winkler Method, Systronics pH meter (Model 361) and Systronics Conductivity meter (Model 308), respectively. Chloride was measured using the Argentometric method. Fluoride was measured using 2-(parasulfophenylazo)-1,8-dihydroxy-3,6-naphthalene-disulphonate(SPADNS) method. Nitrate was analyzed by Orion™ Ion selective electrode method. Total hardness was measured by titration with Ethylenediamine tetraacetic acid (EDTA) using Erichrome black-T indicator. Gravimetric method was used to determine Total suspended solid (TSS) and TDS. Acidity and alkalinity were measured by Titration method. Sodium and Potassium were determined with the help of Systronics Flame photrometer (Model-128). Silica was measured by Molybdosilicate method using Ultra-Violet and Visible (UV–VIS) double-beam Spectrophotometer (Model: Electronics India 2375). Sulfate was measured using Turbidimetric Method. Ammonia–Nitrogen was measured using Phenate method for fresh water sample. Ammonia–Nitrogen for wastewater sample was estimated by distillation and followed by titration with 0.02 N sulfuric acid. Cyanide was measured by Silver nitrate titration method. Oil and grease were determined by extraction and gravimetric method. The results were cross-checked with National Institute of Standard Technology (NIST) certified oil and grease standards. BOD was measured by Bioassay test. The samples collected from fresh water for BOD were collected in 300 mL BOD bottles. Whereas, the samples collected from the wastewater were diluted with aerated water, added a known volume of micronutrients, and collected in 300 mL BOD bottles. The time zero samples were immediately fixed with Winkler’s reagents for the measurement of control. Another set of samples were maintained at 27 °C temperature in the BOD incubator for 72 h before final DO analysis. The change in DO concentrations between first day and final day was measured as BOD. COD analysis was performed by open reflux method and cross-checked with NIST certified COD standards. Phenol analysis was performed by distillation and spectrophotometric method. Heavy metals such as Calcium (Ca²⁺), Magnesium (Mg²⁺), Manganese (Mn), Iron (Fe), Zinc (Zn), Copper (Cu), Cadmium (Cd), Lead (Pb), Cobalt (Co), Selenium (Se), Arsenic (As), Mercury (Hg), Chromium (Cr), Manganese (Mn) and Aluminum (Al) were analyzed using an atomic absorption spectrophotometer (PerkinElmer AAnalyst 200).
**Results of the experimental investigation**

The results derived by the above methods are presented in Table 3 along with their descriptive statistics. The results are subjected to One-Way ANOVA, Correlation Analysis and Principal Component Analysis.

**One-Way ANOVA**

The results of one-way ANOVA revealed significant seasonal variation \((p < 0.05)\) in the hydro-chemical parameters viz. TSS, Turbidity, Redox Potential, Acidity, Total Hardness, Bicarbonate Alkalinity, Chloride, Sulfate, Nitrate, Sodium, Calcium, Magnesium, Iron, Cadmium, Manganese, Chromium and Magnesium during the entire sampling period. Whereas, no significant seasonal variation \((p > 0.05)\) was showed in hydro-chemical parameters such as pH, EC, TDS, DO, BOD, Residual chlorine, COD, Oil and grease, Fluoride, Potassium, Zinc, Copper, Selenium, Nickel, Aluminum, Boron, Silica, Temperature, Salinity, Cyanide and Phenol.

**Correlation analysis**

In this study, the correlation among various parameters had been established using Pearson ‘r’ correlation coefficient at 95% confidence interval level. In premonsoon, strong negative correlation was found between pH with Zinc and Copper. TDS exhibited strong positive correlation with chloride, calcium, magnesium and moderate positive correlation with iron. EC showed strong positive correlation with copper and nickel. BOD exhibited strong negative correlation with DO. Chloride displayed strong and moderate positive correlation with calcium and magnesium. Calcium showed strong and moderate positive correlation with magnesium and iron. Magnesium exhibited moderate positive correlation with Iron, selenium and arsenic. Zinc was having high positive correlation with copper and silica. In monsoon, strong negative correlation was found between pH-zinc and pH-nickel. EC exhibited moderate positive correlation with TDS. TDS exhibited moderate positive correlation with potassium, magnesium, zinc, copper, cadmium and nickel. BOD exhibited strong negative correlation with DO. Nitrate displayed positive correlation with calcium and magnesium. Calcium was having moderate positive correlation with magnesium. Zinc showed moderate positive correlation with calcium and cadmium. Cadmium and selenium exhibited moderate positive correlation with nickel.

**Principal component analysis**

Principal component analysis (PCA) is a significant statistical technique that was used to assess the interrelations between the water quality parameters in Ib valley coalfield. PCA was performed by considering the Rotated Varimax variables with Kaiser normalization on all the water quality parameters (Table 4). On the basis of Eigen values > 1.0, seven factors were extracted with the cumulative loadings showing a total variability of 89.88%. Factor 1 with eigen value 10.08 accounted for 33.59% of the total variance. Factor 1 showed strong positive loadings for TDS, residual chlorine, bicarbonate alkalinity, chloride, calcium, magnesium, iron and cadmium and strong negative loadings for manganese and aluminum. Factor 2 with eigen value 5.07 accounted for 16.88% of the total variance. Factor 2 showed strong positive loadings for EC, redox potential, acidity, copper, silica, salinity and strong negative loading for pH, TSS and Total hardness. Factor 3 with eigen value 3.91 accounted for 13.04% of the total variance. Factor 3 showed strong positive loadings for EC, redox potential, acidity, copper, silica, salinity and strong negative loading for pH, TSS and Total hardness. Factor 3 with eigen value 3.91 accounted for 13.04% of the total variance. Factor 4 with eigen value 3.13 accounted for 10.45% of the total variance. Factor 4 showed strong positive loadings for acidity, nitrate, sodium and selenium. Factor 5 with eigen value 1.96 accounted for 6.54% of the total variance. Factor 5 showed strong positive loadings for TSS, sulfate and temperature. Factor 6 with eigen value 1.54 accounted for 5.15% of the total variance. Factor 6 showed strong positive loadings for turbidity and potassium. Factor 7 with eigen value 1.27 accounted for 4.24% of the total variance. Factor 7 exhibited moderate positive loadings for iron and strong negative loading for fluoride.

**Data validation**

The analytical precision for the measurement of ions was determined by calculating Charge-Balance Error (CBE). CBE is defined as the fractional difference between the total cations and total anions (Edmond et al. 1995)

\[
\text{CBE} = \frac{\sum \text{Cation} - |\sum \text{Anion}|}{\sum \text{Cation} + |\sum \text{Anion}|} \times 100
\]

The CBE can be positive or negative. A positive CBE shows that the water has a higher grouping of cations than anions. On the other hand, a negative CBE demonstrates that anions are more. CBE less than \(\pm 5\%\) are considered acceptable (Huh et al. 1998). The CBE for all the water samples during premonsoon (Fig. 6) and monsoon season (Fig. 7) were found to be less than \(\pm 5\%\), indicating the acceptability of the results.
Table 3 Descriptive statistics for concentrations of chemical constituents in premonsoon and postmonsoon samples

| Parameters              | Premonsoon (Minimum–Maximum, Average ± SD) | Monsoon (Minimum–Maximum, Average ± SD) | BIS (2012) | WHO (2011) | MoEF &CC (2015) |
|-------------------------|--------------------------------------------|------------------------------------------|------------|------------|-----------------|
| pH                      | 3.77–7.64 (6.43 ± 0.94)                    | 3.05–7.85 (6.78 ± 1.12)                 | 6.5 to 8.5 | NA         | NA              |
| EC (µS/cm)              | 58–1580 (430.82 ± 370.38)                 | 32–2120 (406.62 ± 472.94)               | NA         | NA         | NA              |
| TDS (mg/L)              | 291–345 (316.55 ± 13.3)                   | 210–505 (300.53 ± 58.8)                | 500        | NA         | NA              |
| TSS (mg/L)              | 6–22.99 (13.64 ± 4.57)                    | 21.97–1304.54 (353.12 ± 264.33)        | NA         | NA         | < 10            |
| Turbidity (NTU)         | 1.2–14 (5.7 ± 3.44)                       | 0.8–1000 (188.03 ± 274.69)             | 1          | NA         | NA              |
| DO (mg/L)               | 0.19–6.95 (3.08 ± 1.99)                   | 0.2–6 (3.1 ± 2)                        | NA         | NA         | 6 mg/l or more for drinking water (Class A) |
|                        |                                           |                                          |            | NA         | 5 mg/l or more for bathing water (Class B)   |
|                        |                                           |                                          |            | NA         | 2 mg/l or less for drinking water (Class A) |
|                        |                                           |                                          |            | NA         | 3 mg/l or less for bathing water (Class B)   |
| BOD (mg/L)              | 10.1–73 (33.74 ± 20.83)                   | 3.6–67.6 (25.9 ± 22.1)                 | NA         | NA         | NA              |
| Redox Potential (mV)    | 110–379 (193.82 ± 56)                     | 181–384 (315.59 ± 51)                 | NA         | NA         | NA              |
| Acidity (mg/L)          | 4–15 (8.48 ± 2.87)                        | 3–146 (31.12 ± 31.43)                 | NA         | NA         | NA              |
| Total Hardness (mg/L)   | 60–133 (102.79 ± 18.7)                    | 63–130 (93.82 ± 16.85)                | 300        | NA         | NA              |
| Residual chlorine (mg/L)| BDL–0.26 (0.1 ± 0.07)                     | BDL–0.26 (0.09 ± 0.08)                 | 0.2        | NA         | NA              |
| COD (mg/L)              | 25.4–168.7 (78.14 ± 52.44)               | 19.87–141.19 (63.56 ± 49.61)          | NA         | NA         | < 10            |
| Oil and grease (mg/L)   | BDL–3.4 (1.59 ± 0.98)                     | 0.18–4.3 (1.62 ± 1.12)                | 10         | NA         | NA              |
| Bicarbonate Alkalinity  | 30–80.3 (61.01 ± 13.71)                   | 13.49–72.6 (39.54 ± 15.6)             | 200        | NA         | NA              |
| Chloride (mg/L)         | 12–42.8 (29.06 ± 9.79)                   | 17.66–33.11 (24.71 ± 4.75)            | 250        | 200–300    | NA              |
| Fluoride (mg/L)         | 0.1–0.54 (0.26 ± 0.13)                    | 0.007–0.79 (0.4 ± 0.22)               | 1          | 1.5        | NA              |
| Sulfate (mg/L)          | 10.82–177.97 (91.56 ± 44.08)             | 52.11–493.23 (155.47 ± 87.15)         | 200        | NA         | NA              |
| Nitrate (mg/L)          | 8.9–22.2 (15.71 ± 3.11)                   | 3.7–19.9 (12.49 ± 3.71)               | 45         | 50         | NA              |
| Sodium (mg/L)           | 5–29 (14.4 ± 6.48)                        | 6.96–47.45 (19.41 ± 9.38)             | NA         | NA         | NA              |
| Potassium (mg/L)        | 4.83–35.67 (18.35 ± 7.82)                | 5.5–45.32 (18.82 ± 11.08)             | NA         | NA         | NA              |
| Calcium (mg/L)          | 15.1–37.8 (24.02 ± 5.64)                 | 14.3–58.43 (30.73 ± 9.4)              | 75         | NA         | NA              |
| Magnesium (mg/L)        | 18.8–46.9 (31.81 ± 6.93)                 | 15.31–47.61 (28.35 ± 7.13)            | 30         | NA         | NA              |
| Iron (mg/L)             | BDL–0.17 (0.0455 ± 0.0401)               | 0.01–1.87 (0.5221 ± 0.6581)           | 0.3        | NA         | NA              |
| Zinc (mg/L)             | BDL–0.32 (0.0355 ± 0.0864)               | BDL–0.84 (0.0904 ± 0.1926)            | 5          | NA         | NA              |
| Copper (mg/L)           | 0.005–0.243 (0.0438 ± 0.0668)            | BDL–1.21 (0.044 ± 0.2065)             | 0.05       | 2.0        | NA              |
| Cadmium (mg/L)          | 0.001–0.01 (0.0049 ± 0.0016)             | BDL–0.009 (0.0033 ± 0.0023)           | 0.003      | 0.003      | NA              |
| Selenium (mg/L)         | 0.027–0.099 (0.0612 ± 0.0185)            | 0.011–0.579 (0.0596 ± 0.0933)         | 0.01       | 0.01       | NA              |
| Arsenic (mg/L)          | BDL–0.018 (0.0023 ± 0.0005)             | BDL–0.015 (0.0021 ± 0.0004)           | 0.01       | 0.01       | NA              |
| Mercury (mg/L)          | BDL–BDL (NA ± NA)                        | BDL–BDL (NA ± NA)                     | 0.001      | 0.006      | NA              |
| Chromium (mg/L)         | BDL–0.01 (0.0009 ± 0.0023)               | BDL–0.032 (0.0025 ± 0.0077)           | 0.05       | 0.05       | NA              |
| Nickel (mg/L)           | BDL–0.346 (0.0545 ± 0.0993)             | BDL–0.71 (0.0938 ± 0.179)             | 0.02       | 0.07       | NA              |
Index approach

Quality indices are suitable for getting a composite impact of all parameters into pollution. Additionally, it makes the evaluations into a reproducible form and compiles all the pollution parameters into some easy approach (Balakrishnan and Ramu 2016). The water quality index (WQI), heavy metal pollution index (HPI), Degree of contamination (Cd or C deg) and heavy metal evaluation index (HEI) are some of the indices that represent the quality of water at any locations, and these have been discussed here.

Water quality index

Figuring of water quality index is to transform complex water quality data into information that is understandable and usable by the general public. Water quality index is one of the most commonly used tools to ascertain water quality. There are a lot of water quality indices used nowadays. The most reliable among them is weighted arithmetic water quality index (WA-WQI). The calculation of WQI was performed by using the following equation (Yogendra and Puttaiah 2008):

$$ WQI = \frac{\sum_{i=1}^{n} Q_i W_i}{\sum_{i=1}^{n} W_i} $$

The Quality rating scale $Q_i$ is estimated by the expression:

$$ Q_i = 100 \left[ \frac{(V_i - V_0)}{(S_i - V_0)} \right] $$

where, $V_i$ is calculated concentration of $i$th parameter in the sample water, $V_0$ is the ideal value of this parameter in pure water $V_0 = 0$ (except pH = 7.0 and DO = 14.6 mg/L), $S_i$ is recommended standard value (Bureau of Indian Standards, 2012) of $i$th parameter, is the unit weight.

$$ W_i = \frac{K}{S_i} $$

where, $K$ = proportionality constant and is calculated as

$$ K = \frac{1}{\sum \left( \frac{1}{S_i} \right) } $$

The quality index of all the samples was determined using the above relationship, and the results are presented in Table 5.

| Parameters          | Premonsoon (Minimum–Maximum, Average ± SD) | Monsoon (Minimum–Maximum, Average ± SD) | BIS (2012) | WHO (2011) | MoEF &CC (2015) |
|---------------------|--------------------------------------------|------------------------------------------|------------|------------|-----------------|
| Lead (mg/L)         | BDL–BDL (NA ± NA)                          | BDL–0.013 (0.0004±0.00022)              | 0.01       | 0.01       | NA              |
| Manganese (mg/L)    | 0.06–0.01 (0.03±0.01)                      | 0.06–0.01 (0.03±0.01)                   | 0.1        | 0.4        | NA              |
| Aluminum (mg/L)     | 0.08–0.84 (0.37±0.17)                      | 0.08–0.84 (0.37±0.17)                   | 0.03       | NA         | NA              |
| Boron (mg/L)        | BDL–1.671 (0.1068 ± 0.3563)                | BDL–0.037 (0.0054 ± 0.0104)             | 0.5        | 0.5        | NA              |
| Silica (mg/L)       | 0.829–16.23 (4.38 ± 3.53)                  | 0.56–15.46 (3.79 ± 3.2)                 | NA         | NA         | NA              |
| Temperature (°C)    | 30.9–32.4 (31.7 ± 0.49)                    | 31.17–33.86 (31.65 ± 0.58)              | NA         | NA         | NA              |
| Salinity (ppt)      | BDL–0.8 (0.2061 ± 0.1983)                 | BDL–1.1 (0.2118 ± 0.2306)               | NA         | NA         | NA              |
| Cyanide (mg/L)      | BDL–0.02 (0.0016 ± 0.0042)                | BDL–0.017 (0.0012 ± 0.0037)             | 0.05       | 0.07       | NA              |
| Phenol (mg/L)       | BDL–0.006 (0.0004 ± 0.0012)               | BDL–0.003 (0.0002 ± 0.0006)             | 0.001      | NA         | NA              |
| Na%                 | 12.1–33.22 (22.6 ± 5.68)                  | 12.37–45.19 (25.09 ± 7.89)              | Refer Table 9 |
| SAR                 | 0.35–0.9 (0.57 ± 0.14)                     | 0.37–1.11 (0.65 ± 0.16)                 |            |            |                 |
| RSC                 | −3.54–0.82 (−2.02 ± 0.7)                  | −6.43–0.86 (−2.86 ± 1.13)               |            |            |                 |
| MH                  | 49.79–75.89 (68.45 ± 5.19)                | 46.31–75.36 (60.48 ± 7.11)              |            |            |                 |
Heavy metal pollution index (HPI)

HPI is a rating technique that represents the total quality of water with respect to their heavy metals content. The HPI model is calculated (Tiwari et al. 2016) as:

$$HPI = \frac{\sum_{i=1}^{n} W_i Q_i}{\sum_{i=1}^{n} W_i}$$

where $W_i = \frac{1}{S_i}$ (Prasad and Bose 2001). $Q_i$ is the sub-index of the $i$th parameter, $W_i$ is the unit weight of the $i$th parameter and $n$ is the total number of parameters taken into consideration. The sub-index $Q_i$ of the parameter is determined by the following equation

$$Q_i = \frac{\sum_{i=1}^{n} M_i(-)L_i}{(S_i - L_i)}$$

### Table 4
Rotated varimax principal component analysis of the water quality parameter

| Component loading |
|-------------------|
| Factor 1 | Factor 2 | Factor 3 | Factor 4 | Factor 5 | Factor 6 | Factor 7 |
| pH | 0.23 | −0.86 | −0.06 | −0.18 | −0.08 | 0.15 | −0.05 |
| EC (µS/cm) | 0.38 | 0.88 | 0.01 | 0.05 | −0.06 | 0.03 | −0.05 |
| TSS (mg/L) | 0.17 | −0.51 | −0.19 | 0.15 | 0.60 | 0.04 | −0.31 |
| TDS (mg/L) | 0.87 | 0.20 | 0.25 | 0.03 | 0.32 | 0.03 | −0.02 |
| Turbidity (NTU) | 0.06 | −0.34 | 0.02 | −0.28 | −0.11 | 0.69 | 0.04 |
| DO (mg/L) | −0.02 | −0.19 | −0.96 | −0.08 | −0.17 | 0.00 | 0.01 |
| BOD (mg/L) | 0.10 | −0.01 | 0.97 | 0.11 | 0.00 | 0.03 | −0.06 |
| Redox Potential (mV) | 0.26 | 0.88 | 0.06 | −0.15 | 0.17 | 0.12 | 0.16 |
| Acidity (mg/L) | −0.03 | 0.58 | −0.21 | 0.69 | 0.02 | −0.14 | 0.05 |
| Total Hardness (mg/L) | −0.03 | −0.63 | −0.27 | 0.25 | 0.44 | 0.29 | −0.05 |
| Residual chlorine (mg/L) | 0.85 | 0.13 | −0.07 | −0.21 | −0.11 | −0.02 | 0.10 |
| COD (mg/L) | 0.19 | 0.14 | 0.95 | 0.07 | −0.05 | 0.02 | 0.11 |
| Bicarbonate Alkalinity (mg/L) | 0.76 | 0.13 | 0.11 | 0.36 | 0.01 | 0.32 | −0.25 |
| Chloride (mg/L) | 0.89 | 0.30 | 0.13 | −0.03 | 0.28 | 0.05 | −0.06 |
| Fluoride (mg/L) | 0.08 | −0.41 | −0.03 | 0.05 | −0.07 | −0.25 | −0.77 |
| Sulfate (mg/L) | 0.33 | −0.11 | 0.31 | −0.07 | 0.72 | 0.00 | 0.19 |
| Nitrate (mg/L) | 0.21 | 0.01 | 0.15 | 0.84 | −0.29 | 0.14 | −0.10 |
| Sodium (mg/L) | −0.18 | −0.15 | 0.12 | 0.82 | 0.25 | −0.26 | −0.29 |
| Potassium (mg/L) | 0.21 | 0.28 | 0.03 | −0.09 | 0.29 | 0.82 | 0.19 |
| Calcium (mg/L) | 0.94 | 0.20 | 0.08 | 0.06 | 0.20 | 0.06 | −0.03 |
| Magnesium (mg/L) | 0.72 | 0.08 | 0.32 | 0.30 | 0.39 | 0.02 | 0.03 |
| Iron (mg/L) | 0.59 | −0.05 | 0.10 | 0.47 | −0.22 | −0.43 | 0.37 |
| Copper (mg/L) | 0.28 | 0.92 | 0.07 | 0.05 | −0.04 | 0.13 | 0.11 |
| Cadmium (mg/L) | 0.73 | 0.02 | 0.07 | 0.17 | 0.06 | −0.14 | 0.44 |
| Selenium (mg/L) | 0.16 | −0.04 | 0.08 | 0.82 | 0.11 | −0.29 | 0.34 |
| Manganese (mg/L) | −0.77 | −0.20 | −0.10 | −0.01 | 0.24 | −0.39 | 0.21 |
| Aluminum (mg/L) | −0.58 | 0.00 | −0.66 | 0.44 | 0.01 | 0.03 | −0.11 |
| Silica (mg/L) | 0.18 | 0.94 | 0.10 | −0.07 | −0.11 | 0.07 | 0.12 |
| Temperature (°C) | 0.59 | 0.30 | −0.03 | −0.15 | 0.58 | 0.33 | 0.10 |
| Salinity (ppt) | 0.39 | 0.85 | 0.06 | 0.08 | −0.10 | −0.01 | −0.02 |
| % of variance | 33.59 | 16.88 | 13.04 | 10.45 | 6.54 | 5.15 | 4.24 |
| % of cumulative variance | 33.59 | 50.47 | 63.51 | 73.96 | 80.49 | 85.64 | 89.88 |

Extraction Method: Principal Component Analysis
Rotation Method: Varimax with Kaiser Normalization

Rotation converged in 12 iterations
where $M_i$ is the estimated value of heavy metal of $i$th parameter, $L_i$ is the ideal value of the $i$th parameter and $S_i$ is the prescribed standard value of the $i$th parameter in ppb. The sign ($-$) shows the numerical difference of the two values, ignoring the algebraic sign; that is the absolute value.

Usually, the critical pollution index of HPI value for drinking water is 100. Any water having HPI > 100 is not suitable for drinking (Prasad and Bose 2001). The heavy metal pollution indices for all the samples collected during premonsoon and monsoon period are also presented in Table 5.

### Degree of contamination (Cd or C deg)

The degree of contamination (Cd or C deg) of metal explains the magnitude of contamination of metal. It describes the collective effect of heavy or trace metals, which is detrimental to drinking water (Loska et al. 2004). The Cd value is categorized into three classes: low (Cd < 1), medium (1 < Cd < 3) and high (Cd > 3). The degree of contamination is measured as:

$$C_d = \sum_{i=1}^{n} C_f$$

$$C_f = \frac{C_{Ai}}{C_{Ni}} - 1$$

where, $C_d$ is the degree of contamination, $C_i$ is the contamination factor, $C_{Ai}$ is the analytical value of the metal, $C_{Ni}$ is the upper permissible limit of metal of $i$th parameter.

The degree of contamination along with mean and percentage deviation of all the water samples is presented in Table 6.

### Heavy metal evaluation index (HEI)

Heavy metal evaluation index (HEI) explains overall water quality of a specific area in regard to heavy metals. The HEI is measured by the following equation (Edet and Offiong 2002).

$$HEI = \sum_{i=1}^{n} \frac{H_c}{H_{mac}}$$

where, $H_c$ is the analytical concentration of the $i$th parameter and $H_{mac}$ is the maximum permissible concentration of the $i$th parameter.

The HEI along with mean and percentage deviation for all the water samples is also presented in Table 6.

The Water Quality Rating for the WA-WQI method is presented in Table 7. When compared with the WQI values for all the water samples (Table 5), it could be seen that in premonsoon, 48.48% samples were good, 39.39% poor, and 12.12% very poor. Similarly, in monsoon, 41.18% water samples were good, 35.29% poor, 17.65% very poor, and 5.88% unfit for drinking. In general, it was found that the water quality of sump water is better compared to other samples. This may be due to the fact that coal is a natural filter and adsorbs a variety of pollutants.

The mean-based reclassification method as proposed by Edet and Offiong (2002) incorporating HPI, Cd and HEI is presented in Table 8. The samples for which the values of HPI, Cd, and HEI fall below the mean value or have negative percentage deviation, indicates comparatively healthier water quality (Prasad and Bose 2001; Edet and Offiong 2002). The water quality is assigned as “low” for values...
less than the mean value, medium for those values which remain between mean and two times of the mean value and, as high for those values which are more than two times of the mean. It could be seen that a total of 48.84% of the sample were under the mean of HPI in premonsoon and 73.53% in monsoon. Correspondingly, 54.55%, 66.67% samples had Cd and HPI indices below their mean values in premonsoon and 54.55% and 61.76% in monsoon.

**Graphical representation of water quality data**

**Hydrochemical facies**

Piper trilinear plot was made using Origin Pro 2016 to determine the major cations and anions of analyzed water samples (Figs. 8, 9). The dominant anions were found to be sulfate.
| Sample No | Premonsoon Season | Monsoon Season | Premonsoon Season | Monsoon Season |
|-----------|-------------------|----------------|-------------------|----------------|
|           | Cd                | Mean deviation | % deviation       | Cd             | Mean deviation | % deviation       |
| 1         | 11.34             | −1.13          | 90.94             | 6.19           | −7.71          | 44.51             |
| 2         | 7.81              | −4.66          | 62.66             | 3.23           | −10.67         | 23.21             |
| 3         | 14.65             | −5.13          | 117.51            | −19.03         | −36.91         | 80.94             |
| 4         | 13.77             | 1.30           | 110.45            | 15.07          | 1.17           | 104.84            |
| 5         | 4.96              | −7.51          | 39.78             | 88.85          | 74.95          | 639.23            |
| 6         | −1.18             | −13.65         | −9.49             | −2.42          | −16.32         | −17.43            |
| 7         | 9.19              | −3.28          | 73.67             | 14.30          | 0.40           | 102.88            |
| 8         | 12.29             | −0.18          | 98.58             | 27.46          | 13.56          | 197.54            |
| 9         | 9.70              | −2.77          | 77.79             | 18.81          | 4.91           | 135.31            |
| 10        | 28.19             | 15.72          | 226.07            | 20.31          | 6.41           | 146.13            |
| 11        | 25.93             | 13.46          | 207.98            | 66.13          | 52.23          | 475.79            |
| 12        | 26.32             | 13.85          | 211.05            | 17.85          | 3.95           | 128.42            |
| 13        | 13.92             | 1.45           | 111.60            | 5.60           | −8.30          | 40.29             |
| 14        | 9.98              | −2.49          | 80.01             | 8.53           | −5.37          | 61.39             |
| 15        | 14.73             | 2.26           | 118.12            | 7.93           | −5.97          | 57.07             |
| 16        | 13.78             | 1.31           | 110.51            | 2.30           | −11.60         | 16.55             |
| 17        | 14.66             | 2.19           | 117.54            | 7.13           | −6.77          | 51.32             |
| 18        | 30.26             | 17.79          | 242.66            | 9.34           | −4.56          | 67.19             |
| 19        | 3.78              | −8.69          | 30.33             | 0.88           | −13.02         | 6.33              |
| 20        | 1.20              | −11.27         | 9.64              | 3.17           | −10.73         | 22.81             |
| 21        | 12.44             | −0.03          | 99.79             | 7.24           | −6.66          | 52.11             |
| 22        | 7.40              | −5.07          | 59.34             | 4.33           | −9.57          | 31.13             |
| 23        | 22.15             | 9.68           | 177.63            | 18.45          | 4.55           | 132.75            |
| 24        | 20.36             | 7.89           | 163.27            | 13.21          | −0.69          | 95.02             |
| 25        | 25.64             | 13.17          | 205.64            | 21.23          | 7.33           | 152.76            |
| 26        | 14.53             | 2.06           | 116.55            | 25.58          | 11.68          | 184.04            |
| 27        | 3.23              | −9.24          | 25.93             | 2.02           | −11.88         | 14.51             |
| 28        | 2.68              | −9.79          | 21.46             | 6.95           | −6.95          | 49.98             |
| 29        | 1.97              | −10.50         | 15.77             | 7.47           | −6.43          | 53.72             |
| 30        | 2.83              | −9.64          | 22.67             | 7.47           | −6.43          | 53.72             |
| 31        | 16.06             | 3.59           | 128.76            | 14.69          | 0.79           | 105.71            |
| 32        | 9.19              | −3.28          | 73.70             | 10.59          | −3.31          | 76.18             |
| 33        | 7.63              | −4.84          | 61.16             | 9.94           | −3.96          | 71.50             |
| 34        | Nil               | Nil            | Nil               | 8.00           | −5.90          | 57.58             |
and chloride because of the oxidation of sulfides and disintegration of sandstone and shale. The major cations were calcium and magnesium. These may be produced due to weathering and leaching of carbonate and silicate minerals such as crystalline dolomitic limestone, calcium-magnesium silicate and biotite. The two major hydrogeochemical facies were found to be Calcium–Magnesium–Chloride and Calcium–Magnesium–Sulfate in the study area. The later was predominant in water contaminated areas due to mining activities.

**Concentration contour mapping**

Contour mapping is one of the ideal approaches to visualize spatial differences in water quality. The Geographical Information System (GIS)-based analysis of the concentration contour maps of the study area was made using ArcGIS 10.5 software with the kriging interpolation technique. Kriging is a statistical model that uses autocorrelation. In this technique, the distance or direction between the sample points is assumed that reflects a spatial correlation to explain variation in the surface. Kriging usually incorporates statistical analysis of the data, variogram model, creating surface and optionally exploring a variance surface. The concentration contour maps for some of the important parameters and indices viz. pH, Fe, Se, WQI, HPI and MH are presented in Figs. 10, 11, 12, 13, 14 and 15.

**Suitability for irrigation**

Most of the opencast and underground mines extend below the water table and hence change the hydraulic gradient. This lead to change in the surface as well as groundwater flow regime which affect their quality and quantity. Due to lack of water in the mining area, the mine discharge water can be utilized for irrigation after a proper scientific investigation. The most important water quality parameters to be considered for irrigation purpose are TDS, EC, Salinity, Percent Sodium (Na %), Sodium absorption ratio (SAR), Residual Sodium Carbonate (RSC) and Magnesium Hazard (MH) (Mondal et al. 2012). These parameters were determined for all the collected water samples, and the summary statistics of the results are presented in Table 9.

\[
SAR = \frac{\text{Sodium}}{\sqrt{\left(\frac{\text{Calcium} + \text{Magnesium}}{2}\right)}}
\]

\[
\text{Na}\% = \frac{\text{Sodium} + \text{Potassium}}{\left(\text{Calcium} + \text{Magnesium} + \text{Sodium} + \text{Potassium}\right)} \times 100
\]
Discussion

The following important observations were observed from the analysis in both premonsoon and monsoon season:

\[
\text{RSC} = (\text{Bicarbonate} + \text{Carbonate}) - (\text{Calcium} + \text{Magnesium})
\]

\[
\text{PI} = \left(\frac{\text{Sodium} + \sqrt{\text{Bicarbonate}}}{\text{Calcium} + \text{Magnesium} + \text{Sodium}}\right) \times 100
\]

\[
\text{MH} = \left(\frac{\text{Magnesium}}{\text{Calcium} + \text{Magnesium}}\right) \times 100
\]
**Fig. 9** Piper plot of water samples for monsoon season. The piper diagram can be separated in hydrogeochemical facies. Legend: A: Calcium type; B: No dominant type; C: Magnesium Type; D: Sodium and Potassium type; E: Bicarbonate type; F: Sulfate type; G: Chloride type; H: No dominant type; 1: Alkaline earths exceed alkalis; 2: Alkalies exceed Alkaline earths; 3: Weak acids exceed strong acids; 4: Strong acids exceed weak acids; 5: Magnesium Bicarbonate type; 6: Calcium chloride type; 7: Sodium Chloride type; 8: Sodium Bicarbonate type; 9: Mixed type (1:5 + 9 + 6, 2:8 + 9 + 7, 3:5 + 9 + 8 and 4:6 + 9 + 7)

**pH**

The mine water was acidic to neutral to slightly alkaline in nature. The minimum values of pH were recorded in Samaleswari OCP railway siding inlet (3.77) during premonsoon and south sump (3.05) in monsoon, making it highly acidic. The low pH values were probably due to the presence of sulfur in pyrite form in some coal patches. Acidic water might also have been caused due to the presence of CO₂ and oxidation of organic compounds. The acidity in mine water may also arise due to the presence of mineral acids formed by hydrolysis of salts of some heavy metals such as ferric chloride, aluminum sulfate, manganese oxide (Environmental Protection Agency 2000). Both Sulfate and Mn concentrations were found to have a significant difference between premonsoon and monsoon seasons. This might be the reason for significant difference of acidity between premonsoon and monsoon. In some locations, the mine water was observed to be slightly alkaline, which may be due to chemical buffering and release of bicarbonate and carbonate ions or salts (Verma et al. 2012). At some other locations, the mine water was neutral in nature which could be due to the effect of calcite in coal strata and in the aquifer, precipitation of iron, and the chemical buffering effect of the water (Liu et al. 1991).
Fig. 11 Contour map for Iron concentration in water samples during premonsoon and monsoon season

Fig. 12 Contour map for Se concentration in water samples during premonsoon and monsoon season

EC
The EC values in the range of 0–250, 251–750, and 751–2250 were categorized as low saline, medium saline and high saline, respectively (Islam et al. 2016). In Ib Valley coalfield, the EC values had been found in the range between 58 to 1580 in premonsoon and 32 to 2120 in monsoon, which indicated that the water quality of the region was low to highly saline. This might be due to the ionization of salts of various compounds.

Turbidity

Considerably turbid water at most of the mine locations had been recorded above the permissible limit. Turbidity of the water samples was found to be above the permissible limit with a maximum of 14 NTU in premonsoon and 1000 NTU in monsoon. Particulate substances mostly clay and silt, fine organic and inorganic compounds, soluble colored organic matter, algae and other microscopic particles in mine water might be the reason for the water to be found murky and turbid.

DO, BOD and COD

The Dissolve oxygen (DO) concentration is vital for aquatic lives in surface waters. DO concentration is governed by several factors like temperature, organic matter degradation, primary production and respiration (Paerl 2006). The concentrations of DO in the Ib valley coalfield were found to be in range of 0.19–6.95 (3.08 ± 1.99) mg/L in premonsoon and 0.2–6 (3.1 ± 2) mg/L in monsoon. The average DO in

**Fig. 13** Contour map of WQI in water samples during premonsoon and monsoon season
both the seasons was found to be noncompliant with standards (Table 3). BOD and COD are important indicators of organic pollution in the water bodies. BOD is the amount of oxygen necessary for the metabolic activities of aerobic microorganisms during the degradation of biodegradable organic matter (Chatanga et al. 2019). The BOD concentrations were observed in the range of 10.1–73 (33.74 ± 20.83) mg/L in premonsoon and 3.6–67.6 (25.9 ± 22.1) mg/L in monsoon. The average BOD concentrations were found to be very high irrespective of seasons. Chemical oxygen demand is a measure of chemical decomposition of organic and inorganic matter, dissolved or suspended substances in water. The COD indicates the contents of organics mostly, nitrites, sulfides, ferrous salts, etc. COD is expressed in terms of mass of oxygen consumed per liter of solution (Li and Liu 2019). The COD concentrations were recorded in the range of 25.4–168.7 (78.14 ± 52.44) mg/L in premonsoon and 19.87–141.19 (63.56 ± 49.61) mg/L in monsoon. The average COD concentrations were found to be high across both seasons. The low DO, high BOD and COD in Ib valley coalfield might be due to coal mine and domestic wastewater discharges without proper treatment. The BOD/COD ratio is found to be varied from 0.3 to 0.58 with an average of 0.45 ± 0.06 in premonsoon and 0.12 to 0.63 with an average of 0.39 ± 0.12 in monsoon. If the BOD/COD is less than 0.6, the wastewater is not fairly biodegradable and cannot be effectively treated biologically (Abdalla and Hammam 2014). The average BOD/COD was less than 0.6 in both seasons.

**TSS**

TSS was found in the range of 6 to 22.99 mg/L with an average of 13.64 ± 4.57 mg/L in premonsoon and 21.97 to 1304.54 mg/L with an average of 353.12 ± 264.33 mg/L in monsoon. Substantial quantities of suspended solids were recorded during monsoon. Suspended solids in coal mines originate from the mining activities mostly during excavation, surface runoff, dust and dirt from open mining areas. During monsoon, the kinetic energy of raindrops and subsequent runoff on exposed soil without vegetation cover causes
a substantial amount of soil to be eroded and become suspended in the runoff water. A large quantity of loose particles are generated due to mining activities, which are easily carried off by the runoff water. As the vegetation covers are taken off during mining, the exposure of bare soil by erosion and carriage of suspended solids via runoff becomes easier. It was also noted that in monsoon season, TSS concentrations in downstream samples were substantially higher compared to the upstream, demonstrating the direct impact of mining.

**Chloride**

Chloride was recorded in the range of 12–42.8 mg/L with an average of 29.06 ± 9.79 mg/L in premonsoon and 17.66–33.11 mg/L with an average of 24.71 ± 4.75 mg/L in monsoon. Chloride gets released into the aquatic bodies due to mineral disintegration, evaporation and evapotranspiration. Chloride is a minor constituent of sedimentary rocks, for example, limestone, sandstone, and shale, and in addition granitic rocks. Halite and other chloride salts being extremely soluble can easily get into the water bodies through surface runoff from mines. Warm temperature and strong light too reduce the chloride concentration in water (Sahoo et al. 2016).

**Fluoride**

The presence of fluoride was detected at all the sampling locations. Fluoride was found in the range of 0.1–0.54 (0.26 ± 0.13) mg/L in premonsoon and 0.007–0.79 (0.4 ± 0.22) mg/L in monsoon. The presence of fluoride in high concentration in water had been related to dental fluorosis (Brindha and Elango 2011) and even cancer (Peckham and Awofeso 2014), but its absence roots osteoporosis, wherein degeneration of bones and teeth happen. Fluoride pollution of water can be linked with fluoride-bearing minerals in rocks from volcanic and fumarolic actions, from industrial discharges, and from precipitation as downpour or dry drop out. The disintegration of fluoride, apatite, and topaz from zonal bedrocks prompts high fluoride concentration in surface water and groundwater.

**Sulfate**

Sulfate was observed in the range of 10.82–177.97 mg/L with an average of 91.56 ± 44.08 mg/L in premonsoon and 52.11–493.23 mg/L with an average of 155.47 ± 87.15 mg/L in monsoon. Overburden dumps generally have some low-grade coal with high sulfide mineral content such as pyrite, sphalerite, galena, chalcopryite and pyrrhotite. Sulfate from these minerals might have been leached out with surface runoff. Moreover, Jharsuguda town is an industry hub, located in close proximity to the coalfield, and SO\textsubscript{X} in the
atmosphere in the form of rainwater might have contributed to the sulfate concentration in water in monsoon.

Nitrate

Nitrate concentration was observed varied from 8.9 to 22.2 mg/L with an average of 15.71 ± 3.11 mg/L in premonsoon and 3.7 to 19.9 mg/L with an average of 12.49 ± 3.71 mg/L in monsoon. Nitrate in the coal mine water samples and nearby water bodies might be found due to the use of explosives for blasting in the mines and use of NPK fertilizers for plantation and agriculture in Ib valley coalfield area.

Heavy metals

Coal contains almost all the elements, whose concentration varies from parts per trillion (ppt) to greater than 50 wt% (Finkelman 1993). Chemical analysis of coal mine water samples revealed that selenium was present in all the samples in premonsoon as well as in monsoon. Both organic and inorganic selenium naturally occur in coal and the overburden. Selenite is found in the oxidized coal. Pyrite contains sulfidic form of selenium. Selenium in selendic form as Clausthalite (PbSe) is also found in coal (Yudovich and Ketris 2005). On contact with water, these species of selenium are leached out in a huge amount from the waste rock into the aquatic environment. Selenium in higher concentration may cause endocrine dysfunction, chiefly synthesis of thyroid hormones, metabolism of growth hormone and insulin-like growth factor-1, damage of natural killer cells action, hepatotoxicity and gastrointestinal issues. Dermatologic impacts include nail and hair loss and dermatitis, neurotoxicity effect such as deterioration of motor neurons may result in amyotrophic lateral sclerosis, and selenium compounds show a bivalent influence in cancer, either growing or declining (Vinceti et al. 2001). Arsenic was found to be present in 24.24% of the sample in premonsoon and 26.47% in monsoon. Arsenic has a strong affinity toward sulfide minerals in coal (Coleman and Bragg 1990). Combustion of fossil fuel leads to release of arsenic into the environment. Copper occurred in all water sample in premonsoon and 50% of the water samples in monsoon. Among all the heavy metals, only copper exhibited a significant difference between both the seasons. Earlier researchers had found high concentration of copper in Indian coals (Mukherjee 1982). Discharge from mine tailing and flyash also contributes to water pollution with copper (Krishnan 1995). Sorption of copper on suspended material increases in high pH condition (Aller et al. 1987). Coal ash contains a high amount of nickel (Mukherjee et al. 1982). In this study, Ni was found in 57.57% and 70% of the sample in premonsoon and monsoon, respectively. Nickel gets mobilized in the environment due to the burning of fossil fuel (Dara 1993). Nickel concentration in water increases under reducing condition in the presence of organic matter (Lu and Chen 1977). Iron was present in 93.93% of the samples in premonsoon and 100% in monsoon. Iron contamination in mine water may be attributed to oxidation of pyrite. Excess iron may cause cirrhosis of the liver and heart failure in human beings. Zinc is mostly found in coal waste and gets into the water bodies through surface runoff from the waste dump. The major geological source of cadmium is coal and overburden. Cadmium was present in all the samples in premonsoon and 88.23% of the sample in monsoon and was found mostly above the permissible limit. Mercury was not found in any of the collected samples during both seasons.

As per Heavy Metal Pollution Index (HPI), Degree of Contamination (Cd) and Heavy metal evaluation index (HEI), medium degree of pollution were exhibited by 51.52%, 30.30% and 45.45% samples in premonsoon and 20.59%, 35.29% and 26.47% samples in monsoon. Whereas, 5.88%, 2.94% and 5.88% samples caused high degree of pollution in monsoon and 15.15% samples caused high degree of pollution with respect to Cd in premonsoon. Water Quality Index (WQI) revealed that 39.39% and 35.29% samples belonged to poor water quality category in premonsoon and monsoon, respectively. Basundhara nallah upstream and downstream water (Fig. 3) exhibited a water quality index of 43.66 and 58.91 in premonsoon season and 38.30 and 56.33 in monsoon season, respectively. Further, Basundhara nallah downstream was observed to have higher HPI of 113.03 than the upstream HPI of 98.43 in premonsoon. This clearly indicated that mine drainage from Basundhara OCP contributed to increasing of pollution load in the Basundhara nallah. Basundhara river downstream near Kulda OCP (Fig. 3) had higher WQI than basundhara nallah upstream. Although heavy metals are present in the water; the HPI, Cd and HEI values indicated low heavy metal pollution in Basundhara river. Lilari nallah upstream (Fig. 2) point had WQI, HPI, HEI and Cd values lower than the downstream near Belpahar OCP and Lakhanpur OCP (Fig. 2). This might be due to mine discharge from both Belpahar and Lakhanpur OCPs. Water sample from Puljari nallah could not be taken during the premonsoon season as it was dry. The upstream sample was only collected in the monsoon as the downstream was inaccessible. The WQI for this sample was found to be 44.44, which showed it to be good. Moreover, the HPI, Cd and HEI values exhibited low pollution due to heavy metals.

Irrigation potential of mine water

The analysis results of irrigation potential of mine water are presented in Table 9. Sodium can react with soil to decrease its
degree of permeability (Todd 1980). Generally, little difficulties arise when SP values are less than 20%. When Na% more than 20%, permeability will reduce. The finer the soil texture is and the more the organic matter amounts are and the bigger

Fig. 16 Ib valley Coalfield map showing locations not suitable for Irrigation

Table 9 Assessment of irrigation potential of mine water based on analysis of Na%, SAR, RSC, EC, MH and PI

| Parameter | Premonsoon | Monsoon | Standards for Irrigation IS 11624 (1986) |
|-----------|------------|---------|----------------------------------------|
| Na%       | 12.1−33.22 (22.6 ± 5.68) | 12.37−45.19 (25.09 ± 7.89) | < 20 | 20–40 | 40–60 | 60–80 | > 80 |
| SAR       | 0.17−1.01 (0.46 ± 0.22) | 0.19−1.44 (0.61 ± 0.29) | < 10 | 10–18 | 18–26 | > 26 | > 26 |
| RSC       | −3.54−0.82 (−2.02 ± 0.7) | −6.43−0.86 (−2.86 ± 1.13) | < 1.25 | 1.25−2.0 | 2.0−2.5 | 2.5−3.0 | > 3.0 |
| EC (µS/cm)| 58−1580 (430.82 ± 370.38) | 32−2120 (406.62 ± 472.94) | < 250 | 250−750 | 750−2250 | 2250−4000 | > 4000 |
| MH        | 49.79−75.89 (68.45 ± 5.19) | 46.31−75.36 (60.48 ± 7.11) | < 50 | – | – | – | – |
| PI (%)    | 31.28−73.44 (54.44 ± 11.84) | 16.66−73.39 (40.26 ± 14.17) | > 75% maximum permeability (Class 1) | 75–25% maximum permeability (Class 2) | 25% maximum permeability (Class 3) |
the effect of sodium on infiltration of water and aeration (Joshi 2009). The Na% was found to be in the range of 12.1–33.22 with an average of 22.6 ± 5.68 in premonsoon and 12.37–45.19 with an average of 25.09 ± 7.89 in monsoon. This showed the water quality was good with respect to Na% irrigation standards IS 11624 (1986). The SAR was observed in the range of 0.17–1.01 with an average of 0.46 ± 0.22 in premonsoon and 0.19 to 1.44 with an average of 0.61 ± 0.29 in monsoon. This revealed the water was excellent with respect to SAR irrigation standards IS 11624 (1986). The water bodies having high RSC are highly alkaline due to high pH. Soil irrigated with such water becomes infertile due to deposition of sodium carbonate. RSC < 1.25 is considered to be excellent for irrigation. The average RSC was found as −2.02 and −2.86 in premonsoon and monsoon which was below 1.25 and therefore excellent for irrigation IS 11624 (1986). The water was good as far as EC was concerned in both seasons. The more the EC, the less water plant can take up. Usually, in water bodies, calcium and magnesium tend to maintain a steady state of equilibrium. However, more magnesium in aquatic system will reduce the plant productivity as the soil solution becomes further alkaline. The MH was found to be in the range of 49.79–75.89 with an average of 68.45 ± 5.19 in premonsoon and 46.31–75.36 with an average of 60.48 ± 7.11 in monsoon. The average MH was non-conforming to irrigation standards of less than 50 IS 11624 (1986) in both seasons. The average PI was recorded to be 54.44 ± 11.84% in premonsoon and 40.26 ± 14.17% in monsoon. Thus, the water belonged to the Class 2 (75–25%) type which was good for irrigation. The mine water was good with respect to Na%, EC and PI, excellent with SAR and RSC, but poor with MH. Overall, the mine water can be used for irrigation with very little or no treatment except for six locations such as Samaleswari OCP railway siding inlet and outlet, Basundhara OCP oil trap inlet, Samaleswari OCP ETP Inlet, Lajkura OCP ETP Inlet and Belpahar OCP ETP Inlet.

Conclusion

The water of the Ib valley coalfield is slightly acidic to neutral to slightly alkaline in nature except for Samaleswari OCP railway siding inlet (pH-3.77 in premonsoon) and outlet (pH-4.55 in premonsoon and 3.05 in monsoon) and Basundhara OCP oil trap inlet (pH-3.6 in monsoon).

In premonsoon, parameters which are found noncompliant to the standards are pH, TSS, Turbidity, DO, BOD, COD, Magnesium, Cadmium, Selenium, Nickel and Aluminum. In monsoon, noncompliant parameters include pH, TSS, Turbidity, DO, BOD, COD, Iron, Cadmium, Selenium, Nickel and Aluminum. Iron is found above the standards in monsoon season only. Magnesium is observed above the standard in premonsoon season only.

The average BOD/COD of less than 0.6 in both seasons indicates Ib valley coalfield water is not fairly biodegradable.

The major ions present in water are found to be Calcium–Magnesium–Sulfate–Chloride.

Water Quality Index (WQI) revealed that 39.39% and 35.29% samples belong to poor water quality category in premonsoon and monsoon. Moderate degree of pollution has been recorded in water bodies of Ib valley coalfield as per Heavy Metal Pollution Index (HPI), Degree of Contamination (Cd) and Heavy metal evaluation index (HEI) and is not suitable for drinking.

The mine water is good with respect to Na%, EC and PI, excellent with SAR and RSC, but poor with MH. Overall, the mine water can be used for irrigation with very little or no treatment except for six locations such as Samaleswari OCP railway siding inlet and outlet, Basundhara OCP oil trap inlet, Samaleswari OCP ETP Inlet, Lajkura OCP ETP Inlet and Belpahar OCP ETP Inlet.

Mine drainage from Basundhara OCP contributes to increasing of pollution load in the Basundhara nallah. Basundhara river downstream near Kulda OCP has higher WQI than basundhara nallah upstream. Although heavy metals are present in the water, the HPI, Cd and HEI values indicate low heavy metal pollution in Basundhara nallah and river.

Lilari nallah upstream point has WQI, HPI, HEI and Cd values lower than the downstream near Belpahar OCP and Lakhanpur OCP. This may be due to mine discharge from both Belpahar and Lakhanpur OCPs.

Mining activities, surface runoff, coal burning, atmospheric precipitation, weathering, leaching, industrial activities and anthropogenic activities could be mostly controlling the water chemistry of the Ib valley Coalfield.

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The authors have no affiliation with any organization with a direct or indirect financial interest in the subject matter discussed in the manuscript. On behalf of all authors, the corresponding author states that there is no conflict of interest.

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The text appears to be a page from a scientific journal, possibly discussing water quality and pollution. It includes references to various studies and guidelines, highlighting the importance of environmental protection and management. The page also contains a section on references, indicating the scholarly nature of the content. The presence of scientific terms and methodologies suggests that this document is intended for readers with a background in environmental science or a related field.
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