Distributions of Radionuclides (U & Th) and Pedogenic Characteristics as Indicators of Wet and Warm Climate during the Holocene in the Western Part of the Upper Gangetic Plain, India

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Abstract

Distribution of radionuclides in the soil samples, Infra-red stimulated luminescence dating techniques, electrical conductivity, pH measurements and grain size analysis of soils of the region between the Ganga and Yamuna Rivers (in the Upper Gangetic plain) have been studied. Soil characteristics are highly sensitive to climate changes and the degree of soil development indicated by higher thicknesses of A-Horizons, solum and clay accumulation in b-horizon are higher during the periods 1.7 - 3.6 ka and 6.5 - 9.6 ka, marked by wet and warm climates (inferred from earlier studies), the former period being marked by higher degree of soil development than the later. Radionuclides are significantly in higher amounts in soils developed during the period 1.7 - 3.6 ka, thus indicating that this was the wettest and warmest period, so these radionuclides could be released by weathering of primary rocks and be preserved in sedimentary rocks deposited during that period.

Keywords: Climate, Radionuclides, Gangetic Plain, Pedogenic Processes and India

1. Introduction

Reconstruction of paleoclimates in the Gangetic Plains has been attempted by several researchers using various techniques like study of clay minerals (Srivastava et al., 2003; Kumar et al., 1996; Pal et al., 1989; Srivastava et al., 1998), calcretes (Singh, 2001; Andrews et al., 1998), oxygen isotopes and organic remains (Sharma et al., 2004; Singh et al., 1999, Singh et al., 1974), lacustrine deposits (Sinha et al., 2006) and studies of the vertical sections exposed along the rivers (Sinha et al., 2005). This is the first attempt to reconstruct the past climates by the use of elemental distribution of radionuclides (Uranium and Thorium) and soil morphological aspects like solum thickness and texture of soils like relative clay accumulation in B-horizon, electrical conductivities and pH values of soils of different ages. The study area chosen is the Ganga-Yamuna Interflue (locally called as Doab) in the Upper Gangetic plains (Figure 1).

2. Methodology

Indian Remote Sensing Satellite (IRS-1D): Wide Field Sensors (WiFS, resolution: 188 m) and Linear Image Scanning Sensor (LISS-III, resolution 23.5 m) digital data were used to identify different soil-geomorphic units in the laboratory using the elements of image interpretation as described by Gupta (2003). 33 soil-geomorphic units were identified and mapped.

For ground truthing the laboratory generated data and to collect samples for various laboratory analyses detailed fieldwork was carried out. At least two soil profiles from each soil-geomorphic unit were studied in detail (Figure 2). Samples from C- or BC-horizons of soil profiles were taken for dating. The reasons for such type of sampling are 1) C-horizons are close representatives of the parent materials, 2) C-horizons are least affected by the pedogenic and organic processes and 3) contamination through clay translocation and leaching is minimal.
in this horizon. Also, samples from different horizons of one profile in a unit were collected for finding variation of radionuclides in an entire soil profile. In the laboratory samples from C-horizons were dated using Optically Stimulated Luminescence Technique (OSL). Concentration of Uranium, Thorium and Potassium in C-horizons, and electrical conductivity (EC) and Hydrogen ion concentration (pH) of samples from different horizons of soil profiles were determined and grain size analysis were carried out.

2.1. Optically Stimulated Dating of Soil Samples

OSL dating of the samples were carried out to know the ages of the last exposure of the sediments to day light before present. Soil samples were collected in 5 cm (diameter) X 15 cm (length) metal cylinders.

All laboratory procedures were carried out under subdued red light conditions. 1 inch of the exposed sample from both ends of the metal cylinder was removed. Details of the procedures of the dating are given by Aitken (1985, 1998). In the present study, optically stimulated luminescence (infrared stimulated luminescence, IRSL) was used, as it uses charges, which are removed rapidly on exposure to sun (Godfrey-Smith et al., 1988, Hutt et al., 1988). The sediments being dated are fluvial in nature that could have been exposed to the Sunlight for short periods during their transport. Feasibility study of OSL dating of Gangetic alluvium was carried out by Rao et al. (1997).

Samples were treated with 1 N hydrochloric acid, hydrogen peroxide (6% - 30%) and 0.01 N sodium oxalate, in order to remove carbonates, organic matter and to disperse individual grains, respectively. The fine-grain fraction (4-11 μm) was collected using Stokes’ law in acetone medium and it was deposited on 40 aluminum discs. Equivalent dose was determined by additive dose method (Aitken, 1998). Measurements were carried out on Daybreak version 1100 TL/OSL reader with infra-red (880 ± 80 nm) light stimulation and luminescence detection was through 7 mm Schott BG-39 and Corning 7-59 filters. The aliquots were short-shined to determine the normalization values. All the aliquots are divided in five groups with four aliquots in each group. From one group, the natural OSL signal is measured. The other groups are given known amount of increased radiation doses on top of their natural dose, using a calibrated source in the laboratory. The aliquots were pre-heated to 220°C for 0.5 seconds to remove unstable signals and given a range of beta doses. Main signals were read after shining source on a particular aliquot for 99 seconds to reduce the signal to its residual value. Signals read were normalized with short-shine values to remove disc-to-disc scatter. Growth curves were plotted by fitting linear line to luminescence vs. applied beta doses. Extrapolation of these curves to
residual value gave the equivalent dose. A plot of equivalent dose vs. shine-down time representing exposure energy was derived from 'shine plateau'. This representative plateau value is used for calculating the paleodoses. Rough estimates of water content were made by taking weight a sample with moisture and without moisture. The relative alpha efficiency for IRSL is taken as 0.07.

The concentrations of uranium (**238**U) and thorium (**232**Th) were measured using thick source alpha counters. The sample was crushed to about 10 µm grain size and then evenly spread in a thick layer on a ZnS screen placed at the base of a circular perpex container. Equal counts were assumed for **238**U . **232**Th series and the series were also assumed to be in radioactive equilibrium. All the measurements were done using Daybreak 583 alpha counters. Background was measured before measuring each sample and background values were then subtracted from the signal from the sample (Table 1).

For estimating potassium (**40**K) 0.5 mg of sample was taken, powdered and digested using the standard procedures of soil sample digestion. **40**K concentration was determined by using Atomic Absorption Spectrometry (AAS) technique.

### 2.2. Neutron Activation Analysis (NAA)

A general procedure for elemental analysis of soil samples involves several distinct stages, viz. sampling, sample preparation and analytical procedure. We have adapted two methodologies NAA and AAS (especially to determine the percentage of potassium in the samples). In case of NAA, it involves packing, irradiation, activity measurements, and calculations of results and data interpretation. On the other hand AAS involves sample dissolution, instrumental measurements followed by data analysis.

Samples in NAA or AAS cannot be analyzed on site as in case of multi channel portable Gamma Spectrometry. Hence their storage is also essential though it may be for a short period. In general, sample must not be contaminated by any chemical treatment or destabilized

<table>
<thead>
<tr>
<th>Sample No.</th>
<th>b-Horizon Thickness (cm)</th>
<th>Clay Accumulation Index</th>
<th>Pedogenic Clay Concentration (%)</th>
<th><strong>238</strong>U Concentration (ppm)</th>
<th><strong>232</strong>Th Concentration (ppm)</th>
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during sample handling and storage. Soil samples were thoroughly mixed and small clusters if present are broken manually; roots, organic remains, calcretes and Fe-Mn nodules, if present, are taken out completely. Samples were then air dried or dried in oven and small part of it is taken, finely crushed to less than 2 µm size and then the sample is stored for further processing.

Perhaps the simplest quality assurance procedure in NAA is to include two or more Standard/Certified Reference Material (SRM) in each batch of unknown samples for short or long irradiation. Das (1990) has recommended the simultaneous analysis of two or more SRMs preferably of similar matrix but from different agencies and known analysis by different techniques. In all our studies, we have used at least three different SRMs all having similar matrix from different agencies (Table 2).

### 2.2.1. NAA Methodology

All the powdered samples and standards each of 50 mg were accurately weighed and packed in alkathene (pure form of polypropylene) for long-term irradiation of 6 hrs. Long (6h) irradiation at thermal neutron flux of 1012 n cm⁻² s⁻¹ in APSARA reactor at the Bhabha Atomic Research Centre (BARC), Mumbai, India. Activities due to $^{153}$Sm, $^{121}$Eu, $^{82}$Br, $^{117}$Lu, $^{239}$U, $^{233}$Th, $^{149}$La, $^{24}$Na, $^{51}$Cr, $^{181}$Hf, $^{137}$Cs, $^{46}$Sc, $^{59}$Fe, $^{40}$K and $^{42}$K were measured using an 80 cm³ coaxial High Precision Germanium (HPGe) detector (EG & G ORTEC) and 8k MCA at the reactor site and later at the Radiochemistry Laboratories of BARC. All irradiations were carried out in APSARA/Dhruva reactors at the BARC, Mumbai. The irradiated samples were then brought to our laboratories in Roorkee and $\gamma$-activity was measured using a NaI (Sodium Iodide) detector having 2.2keV resolution at 1332keV of $^{60}$Co and 8k MCA with GENIE-2000 software (Canberra, Australia). Counting was followed after a delay of 10 days for 1, 2, 6 and 12 hours at different intervals up to 2 months. A typical $\gamma$-ray spectrum showing various photopeaks is shown in Figure 3. Peak areas were noted and background was subtracted to obtain net counts. Care was taken to obtain concentrations of the maximum number of elements from more than one counting and thus reproducibility of data was checked. Elemental contents were calculated by comparing with more than one reference materials.

The spectra of irradiated samples of natural matrices are usually complex with closely-lying multiple peaks. The peaks show Gaussian distribution as the first contribution to the photo peak in a semiconductor detector is due to statistical fluctuations in sharing the absorbed energy between ionization and heating the crystal network. The continuum under the peak is due to Compton effect from the $\gamma$-rays of higher energies and the back

### Table 2. List of standard reference materials.

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<tr>
<th>Geological Standard</th>
<th>Identification Number</th>
<th>Source</th>
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<td>IAEA, Vienna</td>
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<tr>
<td>Pond Sediment</td>
<td>No.2</td>
<td>NIES, Japan</td>
</tr>
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<td>Soil</td>
<td>Soil-5</td>
<td>IAEA, Vienna</td>
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<tr>
<td>Basalts</td>
<td>W-1 and BCR-1</td>
<td>USGS, USA</td>
</tr>
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</table>

### Figure 3 Typical $\gamma$-ray spectrum showing various photopeaks.

### 2.3. EC and pH Measurements

EC measurements of soil samples were carried out by preparing soil:water (1:2) mixture (Jackson, 1967), using distilled water. EC of the mixture was determined by digital conductivity meter (R-314, Raina Instrument, Delhi).

To determine pH a mixture of soil and distilled water in the ratio of 1:2 was prepared by shaking it intermittently for an hour (Jackson, 1967). pH reading for the mixture was taken by using Lutron PH-201 digital pH meter.

### 2.4. Grain Size Analysis

40 gm of sample was taken air or oven dried at low temperatures. Samples were prepared by following the standard method given by Galehouse (1971). Coarse (> 50 µ) and fine (< 50 µ, silt and clay) were separated by wet sieving through American Society for Testing and Mate-
Pedogenic clay content was determined by subtracting the percentages of clay of A or C-horizons from that of the B-horizons (Birkeland, 1984). Clay accumulation index (sum of increase in clay content over that of A or C-horizon multiplied by thickness of B-sub-horizon/s, C.A.I. index) proposed by Levine and Ciolkosz (1983) was also calculated for typical pedons. Where trenches were excavated up to C-Horizons, the clay percentages were calculated taking the clay abundances of C-horizon into account as C-horizon is least affected by pedogenic processes.

3. Results

From ages of C-horizons, soils have been categorized into five members of a morphostratigraphic sequence with ages < 1.7 ka (QGMS-I), 1.7 - 3.6 ka (QGMS-II), 3.6 - 6.5 ka (QGMS-III), 6.5 - 9.6 ka (QGMS-IV) and > 9.6 ka (QGMS-V).

B-horizon thickness of different pedons are higher in the QGMS-II and IV member soils as compared to soils of other members, as shown in bar diagrams for various pedons (Figure 4(a)). Pedogenic clay content in QGMS-II and QGMS-IV is higher as compared to the other members (Figure 4(b)).

The C.A.I. calculated for the soil samples of QGMS-II and IV show high values ranging from 1377.3.85 and 2233.55 as compared to soils of other members, where C.A.I. ranges from 153.85 to 475.95 (Figure 4(c)).

Bar diagrams (Figure 5(a), (b) and Table 1) showing concentration of Uranium and Thorium in different samples from five morphostratigraphic sequence members bring out that the soils of the QGMS-II members show higher concentrations as compared to the soils of other members. Also, U and Th values remain almost unchanged from horizon to horizon within a soil profile from QGMS-II member.

4. Discussion

The monsoonal variability during the last 18 ka inferred by Overpeck et al. (1996), Prell and Kutzbach (1987, 1992) and Goodbread (2003) during the last 150 ka show that the period at the beginning of the Holocene (about 10 to 11 ka) was a wet period. The periods 1.7 to 3.6 ka (QGMS-II) (Sharma et al., 2004; Andrews et al., 1998; Singh et al., 1974; Srivastava et al., 2003) and 6.5 to 9.6 ka (QGMS-IV) (Sharma et al., 2004; Srivastava et al., 2003) were marked by wet, warm climate.

Our data relating to soil morphology, texture and distribution of U and Th for different soil-morphostratigraphic sequence members can be explained in terms of climatic changes during the Holocene Period.

Studies on relation between clay content accumulation and climate suggests that both have a linear relationship i.e., rates of clay production increases with the increasing moisture content and temperature and the rate is highest in wet and warm climates (Jenny, 1985, McFadden, 1982 and Dan et al., 1982). Thus in wet, warm periods we expect highest pedogenic contents, high solum thickness and A-horizon thickness (indicating higher degree of soil development). Indeed that is the case in the present study, pedogenic clay content in B-horizons, clay accumulation index and solum thickness of soils are the highest during the wet and warm periods of QGMS-II and IV as compared to rest of the Holocene deposits in the study area. The QGMS-II soils are relatively more developed as compared to QGMS-IV soils. Thickness of A-horizons of soils of member QGMS-II (12 to 52 cm) is higher than soils of other members (0 to 45 cm). All these features thus indicate that probably periods of formation of QGMS-II soils showing higher degree of soil development was warmer and wetter than that of QGM_IV soils. The soils belonging to the QGMS-II and IV members are acidic in nature. An electrical conductivity measurement of these soils shows high conductivity values ranging from 0.041 - 0.60 mmhos/cm, which are higher than the soils of other ages (0.013 to 0.040 mmhos/cm). Electrical conductivities are the measure of the total dissolved solids in any medium so high values means that the period was marked by wetter and warmer climates, which are necessary for high rates of chemical weathering of minerals, leading to release to large amounts of soluble products. It will explain acidic nature and higher electrical conductivity of soils of the QGMS-II and IV deposited during warm and wet periods.

As Uranium and Thorium precipitation increases when the climatic conditions are wet and warm (Ramli et al., 2005; Vargas et al., 1997; Miah et al., 1998; Echevarria et al., 2001; Martinez-Aguirre et al., 1995). Distribution of Uranium is highly controlled by the insitu soil moisture and pH values of the deposition medium (water in this case) as envisaged by Langmuir (1997), which are high in warm and wet periods. High values of U, Th and K in parent materials for QGMS-II are as expected. Their low values during the period 6.5 - 9.6 ka indicates that though this period was marked by warm, wet climate, but it was subdued as compared to 1.7 - 3.6 ka period, so that smaller amounts of these elements were released by weathering of primary rocks. It supports the conclusions reached by degree of soil development, as discussed...
Figure 4. Variation in (a) B-horizon thickness, (b) Percentages of pedogenic clays and (c) Clay accumulation index (CAI) in soil profiles of different members.
Figures 5. Bar Diagram showing variations in concentration of (a) Uranium and (b) Thorium in different samples.

above.

Minor variation of U and Th within a profile of QGMS-II unit (warmest and wettest) suggests that pedogenesis does not redistribute these elements significantly and values from C-horizons are representative of the parent material.

5. Conclusions

Following conclusions can be drawn from the studies:

1) Pedogenic processes are highly sensitive to climatic changes and warmer and wetter periods of the Early Holocene period (6.5 - 9.6 ka) and late Holocene period (1.7 - 3.6 ka) are marked by higher thickness of A-horizons and solum and higher clay accumulations in B-horizons than the other periods.

2) Concentration of Uranium and Thorium show significantly high values in sediments deposited during relatively highly wet and warm period (1.7 - 3.6 ka), though it is not possible to provide exact climatic conditions. This is also supported by the degree of development of soils.

6. Acknowledgements

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7. References


