Comparison and correction of element measurements using qualitative and quantitative X-ray fluorescence in lacustrine sediments: A case study of Lake Hurleg

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Abstract: Using an X-ray Fluorescence (XRF) core scanner with nondestructive and successive, the chemistry features of lacustrine sediment can be measured directly. This method of XRF core scanner measurements has been widely applied to core sediment analysis but uncertain of the precision and accuracy. Comparison of intensities obtained by XRF core scanning and the concentration measured by conventional X-ray Fluorescence, indicates effects of physical properties varied from different elements on elemental intensities in the lacustrine sediments of a core from Lake Hurleg in the northeastern Tibetan Plateau. Correlation among elements Ti and Sr between the two measurement methods of the XRF and the conventional XRF is high. Using the intensity of Cl as an indicator of water content, the element intensities of Ti and Sr in the core samples is corrected. But the correlation coefficients of Ti and Sr is litter raised. The results show that XRF core scanning is a very useful tool for measuring element concentration in sediments particularly for high intensities elements.

1 Introduction

Lakes are very important for ecosystem, such as climate, tectonic human activities and so on. The important terrestrial records of environmental change over a range of timescale, recording both catchment-scale dynamics and regional climatic variability are preserved in lake sediments [1]. And lake sediments can provide long, continuous and high-resolution archives comparable to those from oceans and ice cores and the sedimentary records are sensitive to past variability in regional climate. And in most cases the changes of source from land can be transmitted to lake sediments. On the other hand, due to the Tibetan Plateau’s relatively stable sedimentary environment and the relative ease with which, in such a comparatively pristine environment can be identified[2], the deep-water lacustrine sediments of this region are likely to have faithfully recorded, and are also able to serve as a global indicator of the presence of elements within the environment[3-5]. Therefore, a multi-proxy approach should be required to settle the various factors, which are influence sediment composition and thus environmental interpretation. 

X-ray fluorescence (XRF) is a well-established analytical technique for estimating the composition of sediments. In this first study of the elements of Lake Hurleg, samples from core KLK02 were investigated. In particular, we measured the element concentrations sedimentary layer in an attempt to analyze them and to evaluate whether or not they were veracity. A comparison of the analyzed data from both XRF core scanning and conventional quantitative XRF measurements aided geochemical and environmental reconstructions.

2 Sample and methods

2.1 Study area

Lake Hurleg (37°17′N, 96°54′E; 2817 m above sea level) a freshwater lake and surrounded mostly by Quaternary lacustrine and alluvial deposits is located in the Qaidam Basin on the northeastern Tibetan Plateau[6]. The Qaidam Basin, surrounded by the Kunlun Mountains to the south, the Altun Mountains to the west and the Qilian Mountains to the north and east, is an area of 12000 km². The average elevation of the basin surrounding mountains (an elevation of >5000 m) is 2800 m., while the mean annual precipitation at nearby Delingha meteorological station (at 2982 m a.s.l.) is about 160 mm and is highly variable. The summer months has the most moisture falls as rain, and precipitation increases at about 45 mm per 100 m increase in elevation on the basis of seven stations in the region. So the actual precipitation at Lake Hurleg is likely <100 mm. Lake Hurleg is mainly fed by the Bayin River, the Balegen River and groundwater from the mountains to the north, and drains through Lianhu River into Lake Toson. The entire
catchment area is about 12,600 km², and the area is about 56.7 km² of Lake Hurleg. Meteorological data indicate a ~4°C of mean annual temperature, a ~160 mm of mean annual precipitation mostly during the summer, and ~2,000 mm of a mean annual potential evaporation during the period 1956-2004 from Delingha station (Fig.1).

![Figure 1. The map showing the catchment and location of core KLK02 in Lake Hurleg.](image)

2.2 Sample collection

The sedimentary environment of Lake Hurleg was reconstructed using data derived from sediment core KLK02 (Fig.1). The core was collected using a Hammer gravity coring system. The 850-mm-long sediment core (KLK02) close to the center of the lake was taken at Lake Hurleg in September 2014 from 7.2 m water depth using a custom-designed piston corer (6 cm in diameter). The sediment cores were transported back to the laboratory in PVC pipes and described. And shortly thereafter, the cores were split in two halves lengthwise. We examined the visual macroscopic features of the core in detail in order to determine the different sedimentary facies. The core shows successive with homogeneous lithology.

2.3 Analytical methods

Using an X-ray Fluorescence (XRF) core scanner, chemistry features of sediment core can be measured directly which does not destruct sediment surface nor split sediment cores. This method has been potentially and widely applied to core sediment analysis and collected of high-resolution chemical data from core profiles [7-9]. The high-resolution chemical data from core profiles can be rapidly collected from X-ray fluorescence (XRF) core scanning. The depth of surface sediment analyze by XRF core scanning generally does not exceed a few tens of microns. The step of analysis is set up to 1 mm in order to acquire a continuous record of elemental compositions along the sediment cores. The measurement time is 20s for each individual step analysis. And then the core was subsampled at 1.5-centimeter intervals. Dried sediment samples was crushed in an agate mortar and pestle, and analyzed for the concentration measured by conventional X-ray Fluorescence using powdered. Overall, about fifty-eight samples were analyzed by Conventional X-ray Fluorescence from core.

3 Results and analysis

The conventional quantitative XRF analysis of elements within the selected Lake Hurleg sedimentary core samples (n=58). The mean contents of the elements Ti and Sr in Lake Hurleg’s sediments were the highest and the mean concentrations of Ti and Sr in the sedimentary core samples were 1806ppm and 1687ppm respectively. Scanning X-ray fluorescence for major element chemical analysis was carried out on archive-core halves for KLK02 core from Lake Hurleg. The X-ray fluorescence (XRF) core scanner can provide a high-resolution record (every 1 mm) of geochemical variation and the XRF core scanner results were expressed hereafter as element peak intensities by counts per second (cps). Element concentrations were not directly available from the XRF measurements and the processing software. The areas of the element peaks provided an indication of the relative concentrations of each element [10].

The data derived from the conventional XRF were correlated with XRF core scanning results. The correlation coefficient of Ti and Sr was 0.5 and 0.66 respectively (Fig.2). During the XRF core scanning process, any measurements would relatively heavily influenced by the energy levels of any X-ray source, as well as by the physical and chemical properties of the sediment [11-13]. The XRF core scanning results from studies of marine sediments would necessarily have been affected by the moisture content of the core, as well as the particular core’s carbonate content, particle composition and any changes in the density and porosity of its mineral composition. Of these drawbacks, high water content could be seen as the most negative influence on the accuracy of any results, as a high water content would inevitably reduce the density of any elements scanned using the XRF method [14-16]. Nonetheless, the accuracy of such XRF scanning can be improved by making scientifically-based water content corrections. The strong chemical activity, and hence density, of the element Cl plays an important part in any sedimentary composition, meaning that this intensity can be used as an alternative indicator of the extent of any changes in moisture content in a sedimentary core. The higher the water content, the higher the Cl density. Measurements resulting from XRF scanning can therefore potentially be calibrated using the elemental intensity of Cl[17,18]. Out results showed a clear correlation between XRF scanning and conventional XRF data, and that the correlation coefficient between the calculated results for each element and the conventional XRF improved after water/Cl correction. The correlation coefficient of Ti and Sr was 0.55 and 0.72 (Fig.2). It demonstrated that water content exerted little impact upon the XRF scanning results of element Ti and Sr in Lake Hurleg.
In the sections extracted from core KLK02, the trends and ranges exhibited by element Ti and Sr contents appeared close to those found when statistical regression was applied to the conventional XRF results. And it showed the higher comparisons between the concentrations derived from conventional XRF measurements and those after water/Cl correction using XRF core scanning (Fig.3). This was especially true when looking at the accuracy of such ranges in element content. This combined methodology can therefore go some way to satisfying the need for an accurate, high-resolution analysis of both the paleoclimate and the paleoenvironment.

**Figure 3.** Comparisons between the concentrations derived from conventional XRF measurements and those calculated using XRF core scanning

### 4 Conclusion

When comparing this method with qualitative XRF core scanning analysis, it became clear that the results gained using these two different analytical methods were significantly comparable, especially when water content correction techniques were applied to the XRF scanning results. Correlation analysis of the two datasets XRF scanning results and another from conventional XRF measurements indicated that the element Ti and Sr fairly well and water content had little influence based on Cl-calibrated. This would suggest that interference of water content in core samples needed to be calibrated depending on the element. This indicated that, combined with the conventional XRF method, the XRF core scanning method can be a useful tool for establishing the contents of Ti and Sr in lacustrine sediments.

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