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Geochemical evolution and mechanisms controlling groundwater chemistry in the transboundary Komadugu–Yobe Basin, Lake Chad region: An integrated approach of chemometric analysis and geochemical modeling

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ABSTRACT

Study region: The study is conducted in the transboundary Komadugu–Yobe Basin, Lake Chad region, a critical groundwater resource shared across national boundaries of Nigeria and Niger Republic.

Study focus: The research investigates geochemical evolution and sources of chemical constituents in groundwater through an integrated methodology that integrates geochemical modeling, molar ratios, bivariate plots, and chemometric analysis. Groundwater samples (n=240) were collected during wet and dry seasons to identify seasonal variations and the impact of geogenic and anthropogenic processes on groundwater quality.

New hydrological insights: The findings revealed that Cl⁻ and NO₃⁻ are associated with anthropogenic pollution. The Principal Component Analysis identified three main components associated with geogenic and anthropogenic processes, agricultural pollution, and mineral weathering. Hierarchical Cluster Analysis highlighted geogenic, anthropogenic, and evaporative influences. Groundwater in the basin is predominantly of Ca–HCO₃ and Na–Cl types and is significantly undersaturated with calcite, dolomite, and fluorite, though seasonal variations show saturation in some samples. Elevated partial pressures of CO₂ (pCO₂) above atmospheric pCO₂ in nearly all samples suggests active biogeochemical processes. Moreover, Gibbs plots, molar ratios, and bivariate plots, along with chloroalkaline indices (CAI-I & CAI-II) confirms influence of mineral weathering and ion exchange reactions within the aquifer system. Few locations show evaporation during the dry season. This study provides valuable insights for sustainable management of groundwater resources in semi-arid and arid regions.

1. Introduction

Geochemical modeling and chemometric analysis of groundwater quality is paramount for groundwater sustainability. The

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demand for potable freshwater has significantly increased worldwide due to population growth, intensive agriculture, and pollution of surface and groundwater from both geogenic and anthropogenic activities (Awaleh et al., 2024; Elumalai et al., 2020; Mohammed et al., 2022; Samtio et al., 2023; Sikakwe and Eyong, 2022; Yang et al., 2021; Zhang et al., 2020). Groundwater provides drinking water supply for over one-third of the world's population (Ismail et al., 2020; Sheng et al., 2022). As a result, groundwater is overexploited globally, exposing it to significant risk of pollution and quality/quantity degradation (Elumalai et al., 2022; Ha et al., 2022; Mohammed et al., 2023; Sarti et al., 2021; Sheng et al., 2022; Yang et al., 2021). Consequently, groundwater quality is impacted by both natural and anthropogenic processes (Jabbo et al., 2022; Mohammed et al., 2023; Samtio et al., 2023; Subba Rao and Chaudhary, 2019) including factors such as leachate from dump sites, overexploitation of wells, accidental oil spillages, effluent from industries, residential wastewater, sewage, and intensive use of synthetic fertilizer (Awaleh et al., 2024; Shuaibu et al., 2024; Zhang et al., 2023). Natural processes that influence groundwater chemistry include rock—water reactions, oxidation—reduction, precipitation and dissolution of aquifer materials, and groundwater recharge and discharge processes (Elumalai et al., 2022; Jehan et al., 2019; Sikakwe and Eyong, 2022; Wali et al., 2019).

Groundwater is essential in arid and semi–arid regions (Elumalai et al., 2019; Loh et al., 2020; Mohammed et al., 2022). Aquifers in Komadugu–Yobe basin (KYB) are at risk of depletion and degradation due to overexploitation, climate change impacts, and pollution from geogenic and anthropogenic activities such as urbanization, industrialization, leachate from dumpsites, and indiscriminate discharge of solid wastes and wastewaters to the environment (Goni et al., 2019; Jagaba et al., 2020; Wali et al., 2020). Groundwater overexploitation in this region for domestic and irrigation water supply makes the aquifers vulnerable to depletion and quality degradation. The primary occupation of the residents in KYB is agricultural, including rain–fed and irrigation farming, contributing significantly to the food requirements in Nigeria (Adeyeri et al., 2020; Ahmed et al., 2018; Descloitres et al., 2013). It is crucial to know if groundwater in the basin meets the necessary quality standards for various uses. Several studies have recently been conducted in Komadugu–Yobe basin to evaluate groundwater quality (Abubakar et al., 2018; Bura et al., 2018; Garba et al., 2018; Goni et al., 2023; Hamidu et al., 2017; Ibrahim et al., 2021; Jagaba et al., 2020; Shuaibu et al., 2024; Suleiman et al., 2022). In contrast, geochemical modeling and chemometric analysis of groundwater at the regional scale remain challenging and had not yet been thoroughly investigated in the basin.

Geochemical modelling and chemometric analysis were employed in various studies worldwide to determine geochemical evolution and mechanisms influencing groundwater chemistry (Banda et al., 2024; Ding et al., 2024; Elumalai et al., 2019; Ha et al., 2022; Mohamed et al., 2022; Sikakwe et al., 2020; Spellman et al., 2024; Wang et al., 2024; Yang et al., 2021; Zhang et al., 2023). Chemometric analysis involves investigation of various water quality variables in a large hydrochemical data set simultaneously, compared to the univariate method, which involves evaluation of each variable in hydrochemical data set individually/using graphical methods (Liu et al., 2020; Rezaei et al., 2020; Sikakwe et al., 2020). The most effective way to examine hydrogeochemical datasets is by categorizing them into geospatial clusters with similar characteristics using chemometric analysis (Liu et al., 2020). It facilitates understanding the origin of dissolved ions in aquifer systems (Abdelaziz et al., 2020; Asomaning et al., 2023; Gautam et al., 2022; Liu et al., 2023; Sarti et al., 2021; Ullah et al., 2022), Graphical methods for assessing hydrochemical data are enhanced by employing multivariate statistical analysis (Gautam et al., 2022; Sikakwe et al., 2020; Singh et al., 2017), PCA is useful in reduction of large datasets into components and in assessing the interrelationship between different hydrochemical characteristics of analyzed water samples to understand sources and extent of pollution (Jehan et al., 2019; Wali et al., 2019), HCA is useful in identifying groups with similar characteristics in hydrochemical datasets (Elumalai et al., 2019; Kumar et al., 2018; Zhou et al., 2024). Groundwater geochemistry is usually influenced by geogenic processes and anthropogenic activities. Various factors, including aquifer lithology, travel time of groundwater, geological formations, sewage, effluent, leachates, and agrochemical applications, makes geochemical characterization of groundwater complex (Liu et al., 2020). However, the chemical reactions between water and minerals, as well as nature and characteristics of water that recharges the aquifer influences the hydrochemical characteristics of groundwater (Chen et al., 2020; Elumalai et al., 2020; Gautam et al., 2022; Kalin, 1996; Singh et al., 2017). Therefore, the interaction between water and mineral components of aquifer defines the general geochemistry of groundwater, providing valuable information on rock-water interaction process (Bradai et al., 2022; Kalin and Long, 1993).

The transboundary Komadugu-Yobe basin, located in the Lake Chad region, is characterized by intensive rainfed and irrigated agriculture, along with various industrial and domestic activities. The main source of water supply in this region is groundwater. Consequently, its significant exploitation results in a wide range of regional environmental problems, including degradation of groundwater quality/quantity. Although several studies have investigated groundwater quality in the transboundary Komadugu-Yobe basin, there has been limited research using integrated geochemical modelling and chemometric analysis on a regional scale. Previous studies in the basin focused on localized assessments of groundwater contamination, overlooking broader hydrogeochemical processes and various groundwater contamination sources. Therefore, this study has uniquely applied a combination of geochemical modelling, bivariate plots, molar ratios, and chemometric analysis to understand both geogenic and anthropogenic influences on groundwater chemistry at a basin-wide scale. It offers new insights into the impact of mineral weathering, ion exchange, and pollution from domestic and agricultural activities on groundwater quality, providing a comprehensive understanding that is critical for sustainable groundwater management. A conceptual model for groundwater evolution and distribution of geochemical processes controlling groundwater chemistry in KYB was established. The findings from this study are expected to support stakeholders and decision—makers in developing new insights into regional-scale groundwater management strategies in transboundary Komadugu—Yobe basin and the wider Lake Chad region.

2. The study area

The study area is the transboundary Komadugu–Yobe basin, a sub–basin of greater Lake Chad basin (Fig. 1a). The basin covers approximately 150,000 km². The topographic elevation of the basin varies from 294 m in Yobe to 1750 m in Jos. The Komadugu Gana and Komadugu Yobe river systems flow through Yobe river and drains into Lake Chad (Adeyeri et al., 2019; Gana et al., 2018). The Hadejia Nguru wetlands and its associated river systems support different economic activities such as agriculture, fish production, pastoralism, and trading to over 20 million people residing in the basin (Adeyeri et al., 2020; Ahmed et al., 2018). KYB has national and international value due to its various internationally shared water resources and wetlands, which boosts local, national, and international economies and requires transboundary integrated water resource management among several countries in the Lake Chad region (Adeyeri et al., 2020). The climate in the basin is arid to semi–arid, characterized by frequent droughts, significant rainfall variability, and intense evaporation (Adeyeri et al., 2019; Goes, 1999). The basin has an average annual temperature of 12 °C in December and January and 40 °C in the months of March and April. The basin has a relative humidity of about 40 % per year. A mean annual precipitation of 1360 mm is estimated in Jos, whereas an estimate of 400–600 mm were recorded in Nguru and Hadejia, ocurring in April to September. Mean annual evapotranspiration rate of 203 mm/year was estimated in the basin (Adeyeri et al., 2017; Shuaibu et al., 2023). The basin is characterized by arable land, and the vegetation in the basin is dominated by a variety of shrubs, dense grasses, and grasslands with scattered trees (Adeyeri et al., 2019).

2.1. General geology and hydrogeology

Geologically, the basin mainly consists of Precambrian basement complex rocks, sedimentary quarternary formations and Jurrasic younger granites (Fig. 1b). The Precambrian basement complex comprises crystalline rocks of Pan–African orogeny that have been

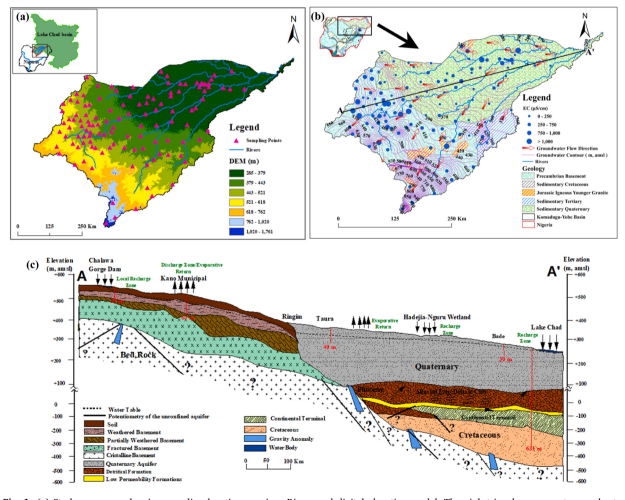


Fig. 1. (a) Study area map showing sampling locations, various Rivers and digital elevation model. The pink triangles represent groundwater sampling points (b) Generalised geological map showing groundwater flow direction and various geologic types (c) Hydrogeological cross–section A-A' of Komadugu–Yobe basin showing different lithologies, groundwater recharge and discharge zones and the groundwater table.

exposed and overlain by younger rocks (Schuster et al., 2009). These rocks consists of migmatite–gneiss, schists, and Pan–African granites that are mostly igneous and metamorphic rocks characterized by shallow weathered basement of low permaliblity. The mineral composition of the Precambrian basement complex are plagioclase, quartz, and biotite. The Precambrian basement is overlain by paleozoic to quaternary sediment deposits in the north-east (Fig. 1c). Jurrasic younger granites comprises syenites, biotite granites, gabbros, ignimbrites, and rhyolites form ring dyke structures around Ningi, Dutse, Shira, Birnin–Kudu and Riruwai. The mineralogical composition of the younger granites are pyroxene, olivine, amphibole, quartz, biotite and plagioclase. The stratigraphical sequence of sediment accumulation overlying the basement complex is the Palaeozoic, lower Cretaceous, Middle Cretaceous, Continental Hamadien, upper Cretaceous, and Continental Terminal (Maduabuchi et al., 2006; Obaje et al., 2004). The sedimentary quaternary consists of a fine to coarse grained sand with an alternation of sandy Aeolian deposits. The valley consists of clayed to sandy fluvial sediments. The sedimentary basin pattern formed a complex subsurface deposit intercalated with sandy to clayey layers (Descloitres et al., 2013; Le Coz et al., 2011). The Chad formation has been continuously sedimented from the Late Miocene to the present, resulting in the deposition of Aeolian sand and clay elements (Shuaibu et al., 2022; Wali et al., 2020). Sand dunes and various alluvial deposits coccurs in the Sedimentary Quaterary parts of the basin as a superficial deposits as parallel ridges extending several killometers with a depth of 15–20 m which influence the river system around Kafin–Hausa, Miga, Jahun, and Auyo in the basin.

The primary source of groundwater in the basin is weathered basement, fracture basement and Plio–Pleostocene argillaceous sequence of Chad Formation with minor arenaceous horizons and recent Quaternary sediments (Fig. 1c). The basin has three distinct aquifer zones in the north-eastern parts: upper aquifer, middle aquifer, and lower aquifer (Goni, 2006). The upper and middle aquifer are accessible for exploitation. However, alluvial deposits on the river floodplains along Hadejia River and Hadejia–Nguru wetlands provides groundwater at shallow depth through tube wells (Tukur et al., 2018). The upper unconfined aquifer zone consists of quaternary deposits, including sands from the lake edge and alluvial fans /deltaic sediments of varied sizes. The aquifer comprises three distinct units: an upper A unit, which is below the water table, and two other units, namely the upper B and C units, ranging from semi–confined to confined (Bura et al., 2018; Goni, 2006). The thickness of the upper aquifer varies from 15 to 100 m with depth to water table of around 20 m. This aquifer is recharged through rainfall runoff and has a transmissivity varying from 0.6 to 8.3 m²/day (Maduabuchi et al., 2006). The middle aquifer is composed of sand beds 10–40 m thick interbedded with clay and diatomites, as well as sand fractions of moderately coarse to finer quartz, feldspar, mica, and feoxides. The average transmissivity of the middle aquifer is about 360 m²/day whereas the lower aquifer has a transmissivity value ranging from 33 to 105 m²/day.

The Komadugu–Yobe River exhibits a seasonal flow pattern mainly between the months of June and December (Descloitres et al., 2013). Groundwater availability is dominant in the southern parts while the northeastern parts have some surface water resources. Groundwater flow is directed from southern parts of the basin to the northeastern parts towards the Lake Chad (Fig. 1b). The flow of groundwater in basement complex aquifer system is highly localized. Recharge likely occurs in the Southern part around Chalawa gorge dam while the discharge and evaporative return occurs around Kano towards Ringim. Moreover, recharge in the North-eastern parts is predominant around the Hadejia-Nguru wetlands through Komadugu-Yobe valley and Lake Chad region. Groundwater recharge is focused at wetlands and Komadugu–Yobe valley and through seepage from river channels and as infiltration of floodwater and runoff along the Yobe floodplain (Carter and Alkali, 1996; Le Coz et al., 2011; Maduabuchi et al., 2006).

3. Materials and methods

3.1. Groundwater sampling and laboratory analysis

In this study, 240 groundwater samples were collected in 50 mL polyethylene bottles from 120 boreholes and hand-dug wells over two sampling campaigns in August and September 2021 (wet season) and March and April 2022 (dry season) in KYB. Wells were pumped for a minimum of 5 minutes before sampling at each location to ensure the collection of samples free from contamination by borehole pipes and stagnant water. Prior to sampling, each sample bottle was rinsed twice with groundwater from the respective well. Groundwater samples were then filtered through a 0.45 μ m acetate cellulose syringe filter into two 50 mL polyethylene bottles: one bottle was acidified with 0.4 mL of nitric acid for cation analysis, while the other bottle, reserved for anion analysis, was left unacidified. Groundwater samples were sealed with watertight lids and kept below 4 °C in icebox coolers until laboratory analysis in Civil and Environmental Engineering Laboratory, University of Strathclyde, Glasgow, UK. Location of each sampled well were marked using a handheld portable global positioning system (GPS) and plotted on Fig. 1a. Prior to water samples collection, the pH, electrical conductivity (EC) were measured in–situ at each sampling location with hand–held digital electrical conductivity meter (Model 99720 pH/Conductivity meter). The equipment was calibrated using standard solutions before measurement. Total alkalinity was measured in–situ using a HACH digital alkalinity titrator (Model 16900, HACH International, Loveland, CO, USA). The major cations (Na⁺, K⁺, Ca²⁺, Mg²⁺ and Fe) were analysed with inductively coupled plasma optical emission spectrometry (ICP–OES, iCAP 6200, Thermo Fisher Scientific), while analysis of the anions (Cl⁻, F⁻, SO²₄, NO³₃) was achieved using ion chromatography instrument (Metrohm 850 Professional IC).

3.2. Quality control and assurance

The quality of the fieldwork followed standard procedures to ensure the integrity of groundwater samples and in–situ measurements. The field equipment was calibrated before the start of field measurement using standard solutions and procedures. The preventive course of action and standard operating procedures (SOP) were adhered to in the field, throughout transportation, preservation, and in laboratory analysis to avoid sample aging and cross–contamination as recommended by APHA (2012). Laboratory

analysis accuracy and precision were confirmed by running standards and blanks. Finally, the reliability and accuracy of hydrochemical data were ensured by assessing ionic balance error, which was within the acceptable threshold of \pm 10 % with significant percentage of wet and dry season samples below \pm 5 %.

3.3. Multivariate statistical analysis (Chemometric methods)

Various hydrogeochemical studies conducted globally employed chemometric approaches, including correlation matrix analysis, PCA, and HCA (Kumar et al., 2018; Liu et al., 2023; Samtio et al., 2023; Singh et al., 2017; Subba Rao and Chaudhary, 2019; Ullah et al., 2022; Yang et al., 2021). These methodologies are employed to evaluate overall water quality by identifying pollution sources and to effectively disseminate water quality information (Rezaei et al., 2020). The Pearson's correlation analysis, PCA, and HCA were performed in this study using Origin Pro 2023b to determine the interrelationship of hydrochemical parameters and pollution source distribution of groundwater in Komadugu–Yobe basin.

3.3.1. Pearson's correlation analysis

Pearson's correlation analysis was employed to classify groundwater quality variables and ascertain their interrelationships (Ullah et al., 2022). Perason's correlation analysis was applied to identify anthropogenic activities and underlying rock properties that affect the groundwater chemistry. Major ions, such as nitrate, sulfate, sodium, and chloride, get into groundwater aquifers through municipal waste, the application of synthetic fertilizer in agricultural lands, and organic and inorganic wastes (Wali et al., 2019; Yang et al., 2021).

3.3.2. Principal component analysis (PCA)

PCA was used to elucidate the relationship between large number of groundwater quality parameters (Kumar et al., 2018; Liu et al., 2023). It was applied in this study to identify potential sources of pollution, and statistical independent source tracers were chosen using Varimax rotation method with Kaiser normalization (Kumar et al., 2018; Yadav et al., 2020). The PCA was carried out on 13 groundwater quality variables, namely: pH, EC, TH, Na⁺, K⁺, Ca²⁺, Mg²⁺, Cl⁻, HCO $_3$, SO $_4^2$, NO $_3$, F⁻, and Fe. The basic steps followed in the PCA are the standardization of data and extraction of principal components (PCs) (Wali et al., 2019). Principal components (PCs) with eigen value > 1 were extracted from the scree plot. The variable in the respective components with higher loading were considered of greater significance regardless of sign. PCs are categorised into three classes: weak (0.30–0.50), moderate (0.50–0.750) and strong (>0.75) (Kumar et al., 2018).

3.3.3. Hierarchical clustering analysis (HCA)

This study adopted HCA to classify the groundwater into groups that are similar to each other (Lima et al., 2019). Moreover, the study employed the R and Q-mode HCA using Ward's approach (minimal variance) to find the best clusters (groups) and for comprehensive result interpretation. Distance between samples were measured by squared Euclidean distance (Eq. (1)). Dendrograms are widely used to show hierarchical clustering or grouping together with the associated linkage distances (Subba Rao and Chaudhary, 2019). The cohesiveness and correlation between the hydrochemical parameters and groundwater samples were observed by constructing a dendrogram.

$$d_{xy} = \sum_{i=1}^{p} (x_i - y_i)$$
 (1)

where dxy represents squared Euclidean distance between two points, x and y, in p-dimensional space. j is used to define each individual parameter (Kumar et al., 2018).

3.4. Geochemical modeling

Groundwater chemistry is mainly influenced by factors including structure and composition of mineral, and rock—water equilibrium (Elumalai et al., 2022; Kumar et al., 2018). Saturation index (SI) elucidates the equilibrium between water and minerals (Eq. (2)). The groundwater saturation indices for KYB were determined using Geochemist's WorkBench GWB software 17.0. SI value of zero shows that mineral saturation is in equilibrium in the aquifer, whereas positive value of SI signifies over–saturation of minerals in water and negative SI value signifies under–saturation of minerals in water.

$$SI = \log \left[\frac{IAP}{K_{SP}} \right] \tag{2}$$

IAP represents the ion activity product in the solution, while K_{SP} represents the solubility product or equilibrium constant of the reaction.

3.5. Chloro-alkaline indices (CAI)

The chloroalkaline indices (Eqs. (3) and (4)) were used to study ion exchange reactions occurring within aquifers of the study area. These indices were used in various studies to better understand general groundwater chemistry, ion exchange mechanisms, and

rock-water interactions (Chen et al., 2020; Elumalai et al., 2022; Liu et al., 2023; Mohamed et al., 2022; Sikakwe and Eyong, 2022).

$$CAI - I = \frac{Cl^{-} - (Na^{+} + K^{+})}{Cl^{-}}$$
 (3)

$$CAI - II = \frac{CI^{-} - (Na^{+} + K^{+})}{SO_{4}^{2-} + HCO_{3}^{-} + CO_{3}^{2-} + NO_{3}^{-}}$$
(4)

Where all ions are measured in meq/L.

4. Results and discussion

4.1. Hydrochemical characteristics

The hydrochemical characteristics of the groundwater in KYB are presented in Table 1. The concentration of hydrochemical parameters in wet and dry seasons groundwater samples displayed a wide variation. pH values varied from 5.52 to 8.24 with a mean of 7.2 in wet season and those of dry season varies from 4.81 to 8.30 with a mean of 6.4 indicating acidic to weak alkaline water in respective seasons. EC ranges from 15 to 2746 µS/cm, with an average of 462 µS/cm and 54–3560 µS/cm, with a mean of 538 µS/cm in wet and dry season respectively. A significant number of samples in both season have EC values < 1500 μS/cm, while some samples have EC value between 1500 to 3000 μS/cm and a few of dry season samples have EC > 3000 μS/cm. Therefore the groundwater in this region is largely fresh to brackish water in both seasons. The total hardness (TH) values of wet season samples varied from 0.8 to 704 mg/L as CaCO₃, with mean of 138 mg/L as CaCO₃. Most of the groundwater samples in both season appeared to be soft. However, few samples ranges from hard to very hard waters in respective seasons. The major ions dominance occurs in an order of $\mathrm{Ca}^{2+}>\mathrm{Na}^+>$ $K^{+} > Mg^{2+} > Fe$ and $HCO_{3}^{-} > Cl^{-} > NO_{3}^{-} > SO_{4}^{2-} > F^{-}$ for cation and anion and $Ca^{2+} > Na^{+} > Mg^{2+} > K^{+} > Fe$ and $NO_{3}^{-} > HCO_{3}^{-} > Cl^{-}$ > SO₄²⁻⁷ > F⁻⁷ for cation and anion in wet and dry seasons respectively. This variation in concentration is likely due to various anthropogenic and geogenic processes taking place within the study area. The hydrogeochemical analysis suggests that the wet season groundwater is mainly characterized by Ca²⁺, HCO₃ and Na⁺, Cl⁻ ions and those of dry season were dorminantly characterized by Ca²⁺, HCO₃ and Na⁺, NO₃ ions. This resonates with the findings of Wang et al. (2024) that the main cation and anion in the groundwater of Hutuo River alluvial-pluvial fan in China were Ca²⁺ and HCO₃. Similarly, Samtio et al. (2023) identified Ca²⁺, Na⁺, HCO₃ and Cl⁻ as the main ions in the groundwater of chachro sub-district in Pakistan. The dominance of these ions in the groundwater of the study area could possibly reflectes the influence of rock-water interaction and dissolution processes of plagioclase, amphibole, pyroxene, orthoclase and biotite rock minerals as well as contamination from irrigation return flows, synthetic fertilizers, potassium compost, domestic and industrial discharges, leachates and nitrification from pit latrines (Shuaibu et al., 2024; Subba Rao et al., 2022; Yu et al., 2024). It is worth noting that, the high nitrate concentrations shown in Table 1 during the wet and dry seasons were measured in open dug wells and boreholes, where significant pollution is attributed to agricultural activities and leachate from unimproved pit latrines, particularly in the downstream areas of the basin. This corroborates the findings of Goni et al. (2023) in parts of the Hadejia-Jama'are-Komadugu-Yobe Basin, where a high nitrate concentration of approximately 1003 mg/L was measured in a dug well affected by agricultural pollution.

4.2. Chemometric analysis

4.2.1. Pearson's correlation analysis

Correlation analysis provides an understanding of the relationship between variables for assessing their communal origin and/or

Table 1Descriptive statistics of hydrochemical parameters of wet and dry seasons groundwater.

Parameters		Wet Season			Dry Season	
	Range	Mean	Std Dev.	Range	Mean	Std Dev.
pН	5.52-8.24	7.2	0.6	4.81-8.30	6.41	0.5
EC	15-2746	462	470	54-3560	538	590
TH	0.8-704	138	125	5-1280	139	180
Na ⁺	2-285	36	45	1.8-247	33	40
K^+	0.1-96	10	20	0.7-173	9	20
Ca ²⁺	0.2 - 227	39	35	1.6-399	40	55
Mg^{2+}	0.9-58	9.9	10	0.2-69	9.2	10
Cl–	0.7-372	48	70	1.6-645	58	90
HCO ₃	1.5-379	120	85	0-220	64	45
SO ₄ ²⁻	0.1-133	15	20	0.1-226	16	30
NO_3^-	BDL-314	42	55	BDL-927	65	140
F-	BDL-2.3	0.3	0.5	BDL-2.4	0.7	0.5
Fe	BDL-19	0.98	3.0	BDL-12	0.66	2.0

Note: All units are in mg/L except for dimensionless pH, TH in mg/L as $CaCO_3$, and EC in μ S/cm. Std Dev.: Standard Deviation. BDL: Below Detection Limit.

sources (Singh et al., 2017; Wali et al., 2022). Fig. 2a and b presented the result of Pearson's correlation analysis of the hydrochemical parameters of wet and dry seasons. EC strongly correlated with TH ($r^2 = 0.82$), Na⁺ ($r^2 = 0.81$), Ca²⁺ ($r^2 = 0.81$), Mg²⁺ ($r^2 = 0.73$), Cl⁻ $(r^2 = 0.83)$, and SO_4^2 $(r^2 = 0.73)$ and moderately correlated with HCO_3 $(r^2 = 0.61)$ in wet season samples, whereas TH $(r^2 = 0.70)$, Na^+ $(r^2 = 0.72)$, Ca^{2+} $(r^2 = 0.71)$, $Cl^ (r^2 = 0.74)$, and Mg^{2+} $(r^2 = 0.60)$, SO_4^{2-} $(r^2 = 0.67)$, NO_3 $(r^2 = 0.60)$ correlates with EC in dry season groundwater samples (Table SM1 & 2, Fig. 2a &b). There was a weak correlation between EC and K^+ ($r^2 = 0.41$), and NO_3^- ($r^2 = 0.37$) in wet season groundwaters, and K^+ ($r^2 = 0.47$) and HCO_3^- ($r^2 = 0.29$) in the dry season groundwater samples (Table SM2). pH does not correlate with K^+ ($r^2 = -0.04$), and NO_3^- ($r^2 = -0.12$), both of which show a very weak correlation between other variables in wet season (Table SM1). Fluoride shows moderate correlation with HCO_3 and SO_4^{2-} and weak correlation with all other variables in the wet season, and Fe shows no correlation with all parameters in both seasons. Cl- significantly correlates with all parameters in wet season except pH, K⁺, NO₃, F⁻ and Fe. Unlike the wet season, it is positively correlated with all water quality parameters except pH, HCO₃, F⁻, and Fe, which implies anthropogenic sources, NO₃ and Cl⁻ displayed strong positive correlation during dry season, suggesting they were derived from both human and animal waste possibly from heavy application of chemical fertilizers, sewage and industrial effluents (Beshir et al., 2024; Dasari et al., 2024; Khan et al., 2023; Memon et al., 2023). Again, nitrate displayed a significant positive correlation with all variables in dry season except HCO₃, F⁻, and Fe, contrary to the wet season where the association is very weak except for F and Fe which shows no association. This could be attributed to the intensive use of synthetic fertilizer in irrigated areas, specifically in Kano Irrigation Project (KIP), Hadejia Valley Project (HVP), and other irrigation projects along Hadejia-Nguru parts of the basin (Shuaibu et al., 2024). This is inline with the findings of Bijay-Singh and Craswell (2021) that fertilizer N consumption in East and South Asia has adversely contributes to nitrate pollution of groundwater and surface water bodies. The correlation between Mg²⁺ and Ca^{2+} ($r^2 = 0.83$, wet season, and $r^2 = 0.89$, dry season) is an indication that magnesium in the groundwater is derived from either dissolution of carbonate minerals or weathering of silicate minerals such as biotite, pyroxene and amphibole in the study area. Likewise, SO_4^{2-} and Ca^{2+} displayed a moderate and strong positive correlation in the wet and dry season, signifying a probable dissolution of evaporate in dry season.

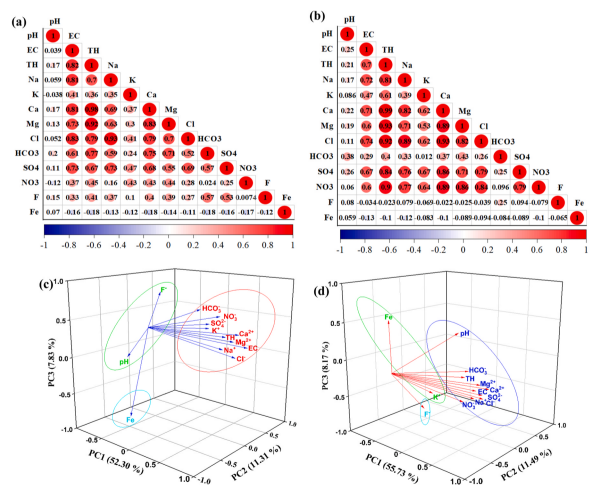


Fig. 2. Pearson's correlation analysis results (a) wet season (b) dry season. Loading plot of three PCs (PC1, PC2 and PC3) in 3D (c) wet season (d) dry season.

4.2.2. Principal component analysis (PCA)

Principal Component Analysis (PCA) was conducted on 13 groundwater variables, namely pH, EC, TH, Na⁺, K⁺, Ca²⁺, Mg²⁺, Cl⁻, HCO₃, SO₄², NO₃, F, and Fe in wet and dry season samples from the Komadugu-Yobe basin. This analysis aimed to understand the hydrogeochemical processes and identify sources of hydrochemical constituents within the groundwater of the basin. Three principal components (PCs) with eigenvalues greater than 1 were extracted for each season, as shown in Table 2. These components explain approximately 71.4 % and 75.4 % of the total variance for the wet and dry seasons, respectively. The loadings of the three PCs for the respective seasons are depicted in Figs. 2c and 2d, illustrating the relationship between variables. Notably, the first component exhibits a significant correlation among variables than the subsequent components in both seasons, indicating its primary influence on hydrochemical variations. In the wet season, the first principal component (PC1) accounts for 52.3 % of the variance in the groundwater dataset, with variables such as EC, TH, Na⁺, K⁺, Ca²⁺, Mg²⁺, Cl⁻, HCO₃, and SO₄²⁻ displaying substantial positive loadings (Table 2). This suggests that PC1 is heavily influenced by geogenic processes, including mineral weathering of minerals like plagioclase, biotite, amphibole, and orthoclase, as well as rock-water interactions. Additionally, anthropogenic contributions from domestic discharges and agricultural activities play a role in influencing groundwater chemistry. In this season, all variables exhibit positive loadings in PC1, except Fe, which shows a weak correlation with other variables. The second principal component (PC2), explaining 11.3 % of the variance, is marked by a strong negative loading on pH and significant positive loadings on K⁺ and NO₃. This pattern indicates non-point-source pollution from agricultural runoff and domestic waste. Elevated NO3 levels likely result from nitrification processes associated with the intensive use of nitrogenous fertilizers and manure in rain-fed farming. Furthermore, the widespread use of unimproved pit latrines and septic tanks in recharge areas can lead to nitrate leachate in groundwater (Chen et al., 2024; Dasari et al., 2024; Nyambar and Mohan Viswanathan, 2024). The third principal component (PC3) in the wet season, explaining 7.8 % of the variance, has significant positive loading on F and Fe, likely reflecting the influence of mineral weathering and dissolution, coupled with variations in groundwater flow and sediment transport. These results corroborates with those obtained by Ait Said et al. (2023) in South-East of Morocco.

In the dry season, PC1 explains 55.7 % of the variance, with EC, TH, Na $^+$, Ca 2 +, Mg 2 +, Cl $^-$, SO 2 -, and NO $^-$ 3 showing strong positive loadings. This component, like in the wet season, is influenced by mineral weathering and rock–water interactions, as well as anthropogenic inputs from agriculture and domestic sources (Bijay-Singh and Craswell, 2021; Dasari et al., 2024; Nyambar and Mohan Viswanathan, 2024). All variables, except F and Fe, exhibit positive loadings in PC1. The second component (PC2) in the dry season, accounting for 11.5 % of the variance, features significant loadings on pH, HCO $^-$ 3, and F, indicating a carbonate buffering effect that helps regulate groundwater pH. This buffering action suggests a distinct chemical environment compared to the wet season, with less contribution from agricultural runoff. Finally, PC3 in the dry season, which explains 8.2 % of the variance, again shows strong loading on F and notable loading on Fe. This component reflects the continued influence of mineral weathering and dissolution processes, particularly from biotite, pyroxene, and amphibole minerals (Dhaoui et al., 2023; Liu et al., 2023; Zhao et al., 2024).

4.2.3. Hierarchical clustering analysis (HCA)

R-mode and Q-mode HCA were performed to 13 physicochemical parameters, to identify interrelationship between water quality parameters and samples with similar chemical composition using Ward's method for wet and dry season samples. The R-mode HCA produces three cluster groups for both seasons (Fig. 3a and b, Table SM3). The first cluster reveals a strong correlation between EC, TH, Na^+ , Ca^{2+} , Mg^{2+} , CI^-

Table 2Component matrics of groundwater quality parameters, eigenvalues, % total variance and % cumulative total variance.

Parameters	Wet Season			Dry Season		
	PC1	PC2	PC3	PC1	PC2	PC3
рН	0.21	-0.56	-0.15	0.12	0.74	0.26
EC	0.89	0.16	0.04	0.76	0.20	-0.04
TH	0.94	0.06	0.12	0.96	0.19	-0.01
Na ⁺	0.86	0.01	0.01	0.84	0.20	-0.12
K^+	0.45	0.51	0.004	0.71	-0.17	0.06
Ca ²⁺	0.92	0.06	0.12	0.96	0.17	-0.01
Mg^{2+}	0.86	0.06	0.10	0.88	0.20	-0.02
Cl ⁻	0.89	0.14	-0.07	0.95	0.06	-0.03
HCO ₃	0.76	-0.33	0.32	0.22	0.79	-0.18
SO_4^{2-}	0.80	0.01	0.17	0.87	0.17	-0.03
NO_3^-	0.35	0.72	0.05	0.94	-0.09	-0.002
F-	0.47	-0.44	0.45	-0.11	0.47	-0.53
Fe	-0.06	-0.24	-0.88	-0.12	0.13	0.83
Eigenvalue	6.80	1.47	1.02	7.25	1.49	1.06
Total variance (%)	52.30	11.31	7.83	55.73	11.49	8.17
Cummulative total variance (%)	52.30	63.61	71.44	55.73	67.22	75.39

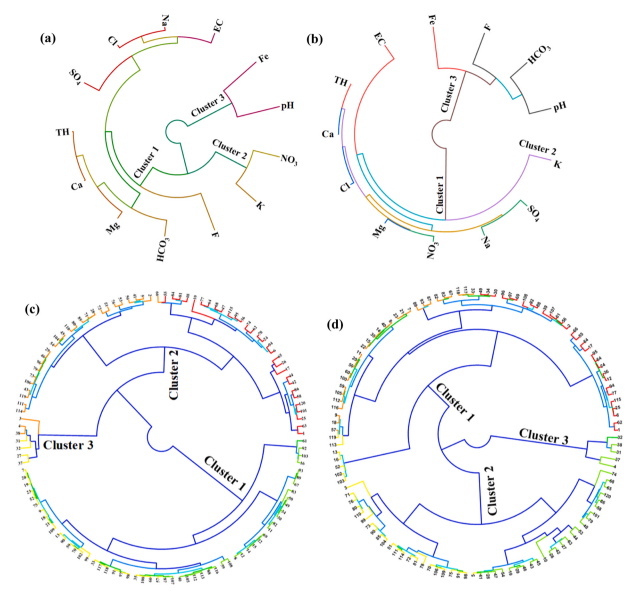


Fig. 3. Dendogram of groundwater quality parameters showing different clusters (a) Wet season (b) Dry season. Dendogram of sampling locations showing different clusters (c) Wet season (d) Dry season.

probable incongruent weathering of feldspar mineral (Bijay-Singh and Craswell, 2021). It is worth noting that cluster 3 is dominated by similar variables such as pH and Fe in wet season and pH, HCO₃, F⁻, and Fe in dry season indicating slight variation in groundwater chemistry which could be due to interactions between various geochemical processes (Karmakar et al., 2023; Liu et al., 2020).

Like R-mode HCA, Q-mode HCA performed on groundwater sampling points produced 3 cluster groups in both seasons (Fig. 3c and d, Table SM4). The first cluster comprises 56 (46.7 %) groundwater samples for wet season and 67 (55.8 %) for dry season sampling points. The value of the EC in this cluster varies from 15 to 438 μ S/cm with mean of 175 μ S/cm and 54–1266 μ S/cm with mean of 324 μ S/cm in the respective seasons dataset, which indicates less mineralized water which is evident in the concentrations of all the groundwater quality parameters (Elumalai et al., 2022; Spellman et al., 2024; Tziritis et al., 2024). Cluster 2 contains 57 (47.5 %) and 48 (40 %) of the respective wet and dry seasons sampling points, with an EC range of 188–2746 μ S/cm with a mean value of 593 μ S/cm and 70–3560 μ S/cm with a mean value of 635 μ S/cm. This signifies moderately mineralized water influenced by anthropogenic inputs. Furthermore, cluster 3 accommodates fewer sampling points for both seasons, with only 7 (5.8 %) in wet season and 5 (4.2 %) in dry season. Groundwater in these sampling points have an EC value ranging from 960 to 2503 μ S/cm with mean value of 1732 μ S/cm and 1846 to 3320 μ S/cm with mean value of 2465 μ S/cm, signifying highly mineralized/low brackishwater influenced by geogenic (plagioclase, biotite and pyroxine weathering and dissolution), salinization sources, and evaporation processes (Elumalai et al., 2022; Tziritis et al., 2024). Fig. 4a and b presents the spatial distribution of groundwater cluster groups related to geology and electrical conductivities in the study area. It is obvious that the Precambrian basement parts of the basin were dominated by moderately and

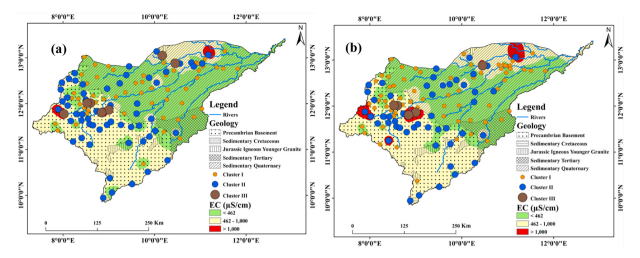


Fig. 4. Spatial distribution of groundwater clusters related with geology and electrical conductivities (a) wet season (b) dry season.

highly mineralized waters. This suggests weathering and dissolution of rock forming minerals (plagioclase, biotite, amphibole and pyroxine) in this region. However, less mineralized water predominates sedimentary quarternary parts of the study area.

4.3. Hydrochemical facies

The Piper (1944) diagram for hydrochemical classification is presented in Fig. 5a and b. Groundwater samples of wet season were plotted mainly in A, C, and D zones of lower–left triangle. This suggests the samples are characterized by sodium–type, calcium–type, and no dominant water type for cations. Moreover, the lower right triangle shows that groundwater samples of wet season were mainly plotted in G and F zones, signifying bicarbonate and chloride dominance. It is very clear that no samples are scattered in zones B and E, suggesting that Mg^{2+} and SO_4^2 are not among the major ions in wet season groundwater samples. Groundwater samples of wet season were projected onto zones 1, 2, and 4 of the central diamond–shaped plot (Fig. 5a). This suggests that wet season groundwater samples are mainly $Ca - HCO_3$, Na - Cl, and mixed water types. In the same vein, the dry season groundwater samples were plotted in D and C zones of lower–left triangle, and a few plotted in zone A, indicating Na^+ and Ca^{2+} as the predominant cations in dry season groundwater samples (Fig. 5b). Furthermore, significant percentage of groundwater samples for dry season were plotted in G and F zones of lower–right triangle (Fig. 5b) indicating the presence of HCO_3^2 and Cl^- anions (Hu et al., 2024; Yang et al., 2024; Zhang et al., 2024). It is obvious that dry season samples exhibited a wide range of hydrochemical compositions, and the vast majority of the samples were scattered in 1, 3, and 2 zones of central diamond–shaped plot (Fig. 5b). This indicates that dry season groundwater samples were predominantly $Ca - Na - HCO_3$, Na - Cl, and mixed water types. In overall, groundwater of the study area in the

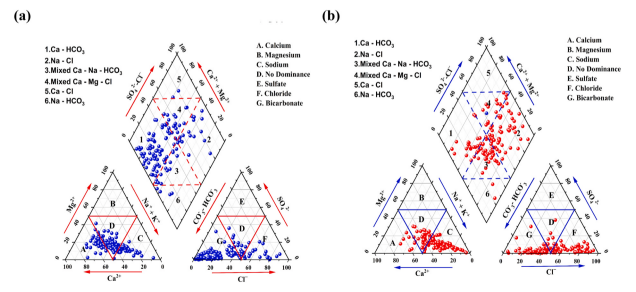


Fig. 5. Piper diagram for groundwater samples in Komadugu-Yobe basin(a) wet season (b) dry season.

respective seasons is greatly influenced by weathering of calcium and sodium feldspar minerals as well as anthropogenic contamination of surface water by irrigation return flows, drainage wastes and industrial discharges. Moreover, the dominance of Ca - HCO₃ water indicates dissolution of carbonates driven by soil CO₂ (Abu et al., 2024a and 2024b; Dasari et al., 2024; Elumalai et al., 2022; Liu et al., 2023; Subba Rao et al., 2022).

4.4. Geochemical modelling

Geochemical modeling of aqueous solutions using Geochemist's WorkBench GWB software 17.0 was employed to ascertain rock—water—mineral interactions using groundwater quality datasets. These modeling techniques have been extensively used in assessing mineral wethering of silicate and carbonate minerals dissolution (Bradai et al., 2022; Elumalai et al., 2022; Ganyaglo et al., 2024; Liu et al., 2015; Trabelsi and Zouari, 2019).

4.4.1. Saturation indices

In this study, Geochemist's WorkBench GWB software 17.0 was employed to determine saturation indices of various minerals in wet and dry seasons groundwater samples in Komadugu-Yobe basin (Table 3). It was observed that significant percentage of the samples shows undersaturation with respect to dolomite, calcite, and fluorite (Fig. 6) with some, 41 (34.2 %) and 4 (3.3 %), 27 (22.5 %) and 1 (0.83 %), and 1 (0.83 %) of the samples indicated near saturation with respect to dolomite, calcite, and fluorite in wet and dry seasons, respectively. Fig. 6 presents relationship between some selected saturation indices and electrical conductivities (EC). Few samples with high EC displayed over saturation with respect to dolomitic carbonate and calcite. It is noteworthy that dolomitesaturation is a proxy indicating high magnesium ions in the groundwater samples. This could be due to incongruent weathering of feldspar minerals. Therefore, the occurrence of dolomite in the study area is highly unlikely. Calcite mineral may be the main source of high Ca²⁺ and HCO₃ in the groundwater samples which results from congruent weathering and dissolution of calcite (Eq. (5)) (Elumalai et al., 2022; Ganyaglo et al., 2024). Moreover, carbonate minerals could be due to various rock minerals in the study area viz plagioclase, biotite, pyroxine and amphibole minerals, Fluorite saturation could be due to a common ion effect with carbonates together with weathering and dissolution of fluoride bearing minerals such as granite gneisses, fluorite and biotite minerals as a results of rock-water interaction under high alkaline condition which could be the main source of F in the aquifers of the study area (Abu et al., 2024a and 2024b; Sunkari et al., 2025), Kumar et al. (2018) posit that fluoride concentration in aguifer system results from dissolution of biotite minerals (Eq. (6)). Fluoride concentration in groundwater could also results from dissolution of fluorite mineral (Eq. (7)).

$$CaCO_3 + CO_2 + H_2O \leftrightarrow Ca^{2+} + 2HCO_3^{5}$$

$$(5)$$

$$KMg_3(AlSi_3O_{10})F_2 + 2OH^- \rightarrow KMg_3(AlSi_3O_{10})(OH)_2 + 2F^-$$
 (6)

$$CaF_2 + 2HCO_3^{-} \rightarrow CaCO_3 + H_2O + CO_2 + 2F^{-}$$

$$(7)$$

Figs. SM1 and SM2 shows spatial distribution of saturation indices of groundwater samples related to geology and electrical conductivities (EC) for wet and dry seasons. The figures revealed that saturation indices above the mean values are largely concentrated in Precambrian basement parts of the study area. Furthermore, most of the oversaturation in both seasons for dolomite, calcite, and fluorite dominated in Precambrian basement complex region. This could be due to the presence of plagioclase, biotite, pyroxene and amphibole rock minerals in this region. However, few samples shows oversaturation with respect to dolomite, calcite, and fluorite in the sedimentary quaternary formation and sedimentary tertiary regions of the study area in the respective seasons.

4.4.2. Partial pressures of carbon dioxide (pCO₂)

Geochemical processes of groundwater and reaction between groundwater and carbonate minerals are greatly influenced by partial pressure of carbon dioxide (pCO₂) which acts as a source of acid (H⁺) for mineral weathering reactions (Trabelsi and Zouari, 2019). The partial pressures of CO_2 of wet and dry season groundwater samples varied from $10^{-3.86}$ to 10^{-1} atm with a mean value of $10^{-2.36}$ atm and $10^{-4.2}$ to $10^{-1.34}$ atm with an average value of $10^{-2.08}$ atm, respectively. It was observed that about 96.7 % and 99.2 % of groundwater samples in wet and dry seasons have pCO₂ above atmospheric pCO₂, which is about $10^{-3.5}$ atm. This indicates the presence of CO_2 in the groundwater system due to biological activity such as respiration of vegetation roots and decomposition of soil organic matters. The partial pressure of CO_2 is observed to decrease as pH of wet and dry–season groundwater samples increases (Fig. SM3). This corresponds to the findings of (Adams et al., (2001); Elumalai et al., (2022); Liu et al., (2015) and Rajmohan et al., (2021)). Negative correlation of -0.61 and -0.21 were observed in groundwater samples of wet and dry seasons which might be

Table 3Descriptive statistics of mineral saturation indices.

Minerals	Wet Season				•	Dry Season			
	Min	Max	Mean	Std	Min	Max	Mean	Std	
Calcite	-6.65	1.15	-0.88	1.24	-6.78	0.65	-2.17	1.31	
Dolomite	-12.87	3.03	-1.27	2.52	-13.26	1.43	-3.85	2.66	
Fluorite	-6.89	0.08	-2.64	1.38	-2.70	0.09	-1.43	0.53	

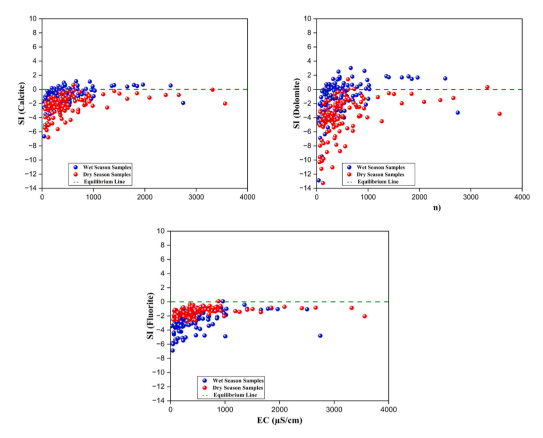


Fig. 6. Saturation indices of some selected minerals against electrical conductivity of groundwater samples of wet and dry seasons.

attributed to longer recidence time, rock—water interaction in aquifer systems and biogenic activities that produces CO_2 . According to Liu et al. (2015), negative correlation between pCO_2 and pH signifies dissolution of feldspar along groundwater flow path, and following reaction could be possible in the aquifer formations (Eq. (8)):

$$Na_2Al_2Si_6O_{16} + 2H_2O + CO_2 \rightarrow Na_2CO_3 + H_2Al_2Si_2O_8 + H_2O + 4SiO_2$$
(8)

This reaction resulted in the consumption of CO_2 and increase in concentration of Na^+ and HCO_3^- leading to pH increase and partial pressure of CO_2 decrease. It is established that the decay of organic matter and roots respiration reseases CO_2 which is the main source of HCO_3^- in groundwater (Eqs. (9) and (10)).

$$CO_2 + H_2O \rightarrow H_2CO_3 \tag{9}$$

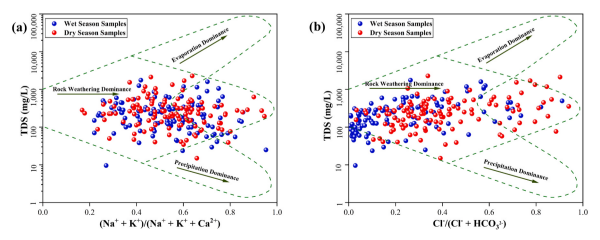


Fig. 7. Gibbs plots of geochemical processes governing groundwater chemistry (a) cations (b) anions.

$$H_2CO_3 \rightarrow H^+ + HCO_3^-$$
 (10)

4.5. Hydrogeochemical processes

Gibbs plots, molar ratios, and bivariate plots were employed in this study to estimate various hydrogeochemical processes that

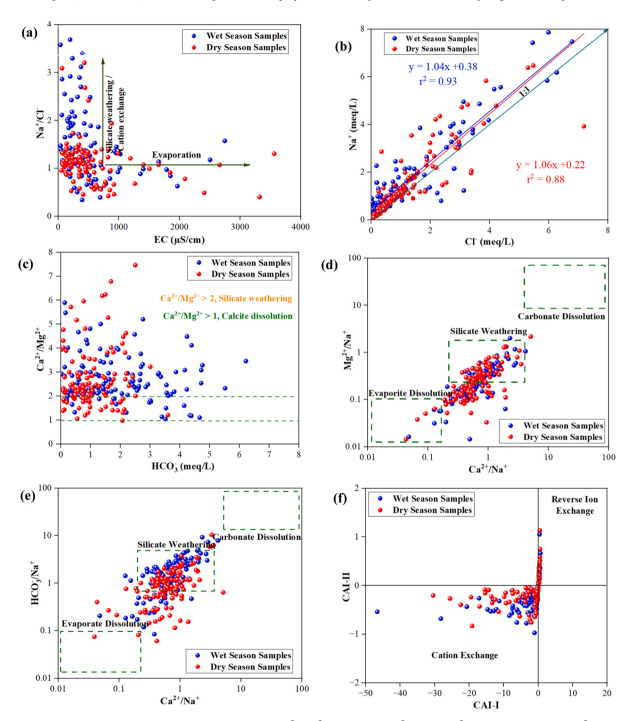


Fig. 8. Bivariate plots of: (a) Na $^+$ /Cl $^-$ vs. EC; (b) Na $^+$ vs. Cl $^-$ (c) Ca $^{2+}$ /Mg $^{2+}$ vs. HCO $_3^-$ (d) Mg $^{2+}$ /Na $^+$ vs Ca $^{2+}$ /Na $^+$ (e) HCO $_3^-$ /Na $^+$ vs Ca $^{2+}$ /Na $^+$ (f) CAI–II vs. CAI–I.

influence groundwater chemistry in the study area.

4.5.1. Gibbs plots

Gibbs plots (Gibbs, 1970) were used in this study to identify the major geochemical mechanisms influencing the groundwater chemistry of the study area (Fig. 7a and b). The figures show that majority of groundwater samples of wet and dry seasons were plotted in the region characterized by rock weathering/rock—water interaction dominance (Chen et al., 2024; Hu et al., 2024). A few samples with high TDS concentration during dry season were observed to trend towards evaporation dominance zone. Therefore, rock weathering is likely the predominant geochemical process controlling the chemistry of groundwater in the study area, except in a few areas associated with evaporation dominance during dry season.

4.5.2. Evaporation

Evaporation process has potential to increase concentration level of all ions present in groundwater in semi– arid areas due to climate change impacts. Na⁺/Cl⁻ vs EC plot was employed to determine influence of evaporation processes on chemistry of groundwater in the study area (Fig. 8a). According to Jankowski and Ian Acworth (1997), when evaporation has significant influence on the chemistry of groundwater, Na⁺/Cl⁻ vs EC plot will remain consistent as EC increases. Fig. 8a reveals that only few samples during wet season follows evaporation trend line, which suggests that evaporation does not play a vital role in influencing the groundwater chemistry during wet season. However, an appreciable number of samples during dry season follow evaporation trend line, which signifies that evaporation could be among the factors influencing the groundwater chemistry during dry season. Na⁺/Cl⁻ ratios shows significant correlation in wet and dry seasons. Na⁺/Cl⁻ ratios enrichment and depletion was observed, which is an indicative of the influence of ion exchange and silicate dissolution in groundwater chemistry in respective seasons. However, most of the groundwater samples in the respective seasons deviated from 1:1 section. This suggests that Ca²⁺ is being exchanged for Na⁺ from clay minerals into water thereby increasing its salinity levels (Elumalai et al., 2022; Hu et al., 2024).

4.5.3. Carbonate dissolution

Carbonate minerals dissolution in groundwater produces Ca^{2+} , Mg^{2+} , and HCO_3 ions (Elumalai et al., 2022; Rajesh et al., 2012; Rajmohan et al., 2017; Rajmohan and Elango, 2004). Molar ratio of Ca^{2+}/Mg^{2+} serves as a reliable indicator for calcite and dolomite dissolution processes within groundwater system. Dissolution of dolomite will maintain Ca^{2+}/Mg^{2+} ratio of 1, but Ca^{2+}/Mg^{2+} ratio > 1 indicates dissolution of calcite, and the process of silicate weathering is associated with Ca^{2+}/Mg^{2+} ratios > 2 (Rajesh et al., 2012; Rajmohan et al., 2017). It is noteworthy that about 81.67 % and 73.33 % of the respective seasons groundwater have Ca^{2+}/Mg^{2+} ratio > 2 (Fig. 8c). This implies that the groundwater samples are dominated by silicate weathering processes. Some portions of samples were observed to cluster between values of 1 and 2 on the Ca^{2+}/Mg^{2+} ratio scale, suggesting the dissolution of calcite which was described in Eq. (5). Moreover, the samples exhibited spatial distribution, and few samples were closely aligned with the $Ca^{2+}/Mg^{2+} = 1$ line. This implies occurrence of high magnesium carbonate dissolution, which is expressed by the following equation (Eq. (11)):

$$CaMg(CO_3)_2 + 2H_2O + 2CO_2 \rightarrow Ca^{2+} + Mg^{2+} + 4HCO_3^-$$
 (11)

4.5.4. Silicate weathering

High Na^+ in groundwater could be related to silicate rock weathering. According to Jankowski and Ian Acworth, (1997), $\mathrm{Na}^+/\mathrm{Cl}^-$ ratio remains constant if evaporation is the prevailing mechanism in the absence of mineral precipitation. The dissolution of halite can result in $\mathrm{Na}^+/\mathrm{Cl}^-$ ratio = 1 if present, however, $\mathrm{Na}^+/\mathrm{Cl}^-$ ratio > 1 increases Na^+ concentration in groundwater system due to silcate weathering and cation exchange processes. Fig. 8a presents $\mathrm{Na}^+/\mathrm{Cl}^-$ vs EC for wet and dry seasons in Komadugu–Yobe basin. A significant increase in $\mathrm{Na}^+/\mathrm{Cl}^-$ ratio is observed when electrical conductivity (EC) value is below 500, particularly in wet season groundwater samples. It was observed that 78.33 % and 71.67 % of respective wet and dry season samples exhibited $\mathrm{Na}^+/\mathrm{Cl}^- > 1$. This high Na^+ ion concentration could potentially originates from silicate weathering and/or cation exchange process (Fig. 8a). Furthermore, $\mathrm{Na}^+/\mathrm{Cl}^-$ and EC plots shows significant number of samples above evaporation line, particularly in wet season. However, few dry season samples were observed in evaporation zone. Therefore, silicate weathering with ion exchange is likely the primary processes that controls the chemistry of groundwater during both seasons (Abu et al., 2024a and 2024b; Yang et al., 2024).

According to Rogers, (1989), if sodium is likely from silicate weathering by soil CO_2 , groundwater would have bicarbonate (HCO_3) as the predominant anion. The release of HCO_3 is attributed to the reaction between feldspar crystals and carbonic acid in water. HCO_3 is the prevailing anionic species found in groundwater of the study area (Table 1). Bivariate ratio plots indicate that silicate weathering is the primary geochemical process influencing groundwater chemistry in the basin, rather than carbonate and evaporite dissolution (Fig. 8d & e). Geochemical evolution of groundwater is characterized by silicate weathering, carbonate dissolution, cation exchange, and evaporative dissolution (Adimalla and Taloor, 2020; Banda et al., 2020; Elumalai et al., 2020, 2022; Ganyaglo et al., 2024; Liu et al., 2023; Rajesh et al., 2012; Rajmohan et al., 2017; Rajmohan and Elango, 2004; Sikakwe and Eyong, 2022; Singh et al., 2017; Yang et al., 2024). Impact of evaporite dissolution appears to be limited in the basin. However, it is visible in few locations during the dry season. This observation is indicative of prevalence of silicate source rocks and limited presence of evaporites, such as halite and gypsum. The latter are commonly related to deposits characterized by poorer permeability, inadequate flushing, and the presence of shale or marl (Banda et al., 2020; Yuan et al., 2024). The process of silicate weathering can be explained using the following weathering reaction (Eq. (12)):

(12)

$$2\text{NaAlSi}_3\text{O}_8 + 2\text{H}_2\text{CO}_3 + 9\text{H}_2\text{O} \rightarrow \text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4 + 2\text{Na}^+ + 4\text{H}_4\text{SiO}_4 + 2\text{HCO}_3^-$$

4.5.5. Ion exchange reactions

Ion exchange and different weathering processes are best described using Na⁺/Cl⁻ vs. EC plot (Fig. 8a). The figure shows an increase in Na+ ion by ion exchange processes in water samples of both seasons. Therefore, cation exchange process is the predorminant mechanisms that control the groundwater chemistry in the respective wet and dry seasons compared to reverse ion exchange. Ion exchange processes in aquifer system have been explained in several studies using chloroalkaline indices (CAI–I and CAI–II) (Eq. (3) & (4)). Positive values of chloroalkaline indices indicate the occurrence of reverse ion exchange, whereas negative values indicate cation exchange reaction (Abu et al., 2024a and 2024b; Elumalai et al., 2022; Mgbenu and Egbueri, 2019). High Cl⁻ over Na⁺ and K⁺ results in positive chloroalkaline indices, suggesting reverse ion exchange reactions. Conversely, high Na⁺ and K⁺ compared to Cl⁻ result in negative values, indicating cation exchange processes (Ganyaglo et al., 2024). When the values of chloroalkaline indices are positive, Mg²⁺ and Ca²⁺ ions are exchanged with Na⁺ and K⁺ ions in water. Conversely, when chloroalkaline indices have negative values, it suggest that there is exchange of Mg²⁺ or Ca²⁺ ions in groundwater with Na⁺ or K⁺ ions in host rocks. Fig. 8 f shows positive correlation between CAI–I and CAI–II. 86.67 % of the samples had negative chloroalkaline index values during the wet season, whereas 80 % had negative values during the dry season. However, 13.33 % and 20 % have positive values during the respective seasons confirming that cation exchange reaction is predominant in the groundwater system compared to reverse ion exchange process. Cation exchange can be explained in the following reactions (Eqs. (13) and (14)):

$$2Na^{+} + CaX_{2} = Ca^{2+} + 2NaX \tag{13}$$

$$Ca^{2+} + 2NaX = 2Na^{+} + CaX_{2}$$
 (14)

Where X is the cation exchange sites.

4.6. Conceptual model for hydrogeochemical processes controlling the groundwater chemistry in Komadugu-Yobe basin

Fig. 9 presents a conceptual model for geochemical processes governing groundwater chemistry of Komadugu-Yobe basin. The study has established that the chemistry of groundwater in the basin is influenced by various geogenic and anthropogenic processes which indicates a subtle transformation as groundwater and surface water moves from upstream section passing through recharge zones in Hadejia Nguru wetlands to downstream parts along Komadugu Yobe river to Lake Chad. The study area is mainly characterized by local gravity driven flow due to varying topography from upstream to downstream. The aquifer in the upstream parts of the basin consists of a system of three aquifer layers resting on the Precambriam basement formation. The weathered basement which is semi-permeable in the vadose zone which hosted the water table. The partially weathered basement and the fractured basements provides groundwater in fractures, fault breccias, and joints which moves from upstream to downstream direction following natural slope of the bedrock. Localized infiltration from Chalawa gorge dam, river flows, industrial discharges, irrigation return flows and

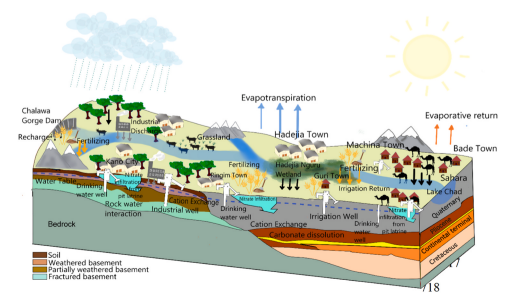


Fig. 9. Conceptual model for geochemical evolution and hydrogeochemical processes controlling the groundwater chemistry in Komadugu-Yobe basin.

domestic waste discharges as well as rock-water intraction influenced the groundwater chemistry and its evolution in upstream parts of the basin. The hydrochemical evolution shows that Ca-HCO₃ and Na-Cl water types predominated the region due to weathering/dissolution of Ca and Na bearing silicate minerals and calcite coupled with various anthropogenic inputs. Nitrate in the irrigation and residential areas may come from nitrogenous fertilizing and unimproved pit laterines respectively. However, in the downstream part of the study area, sedimentary quaternary aquifer provides groundwater mostly in unconfined aquifers. Groundwater occurs at shallow depth around Hadejia, Ringim and Guri town while it is at deeper depth in Machina town and its surroundings. Fig. 9 shows that groundwater in this region is recharged by infiltrating water in the Hadejia-Nguru wetland (HNW), Komadugu-Yobe valley and areas around Lake Chad. However, evapotranspiration occurs around HNW and farmlands as well as evaporation losses at the verge of downstream parts towards Lake Chad. The groundwater chemistry is highly impacted by anthropogenic activities rather than geogenic processes. Nitrate infiltrates into the subsurface from irrigation return flows, domestic waste discharges and unimproved pit latrines in the region.

5. Implications to groundwater sustainability

This study has successfully used a robust approach of geochemical modeling, bivariate plots, ionic ratios, and chemometric analysis and established the sources of chemical constituent influencing the general chemistry and geochemical evolution of groundwater in the transboundary Komadugu–Yobe basin, Lake Chad region. This information is crucial in developing groundwater monitoring and sustainable strategies for effective use and management of the basin's groundwater resources. Insights on the sources of chemical constituents in groundwater of the study area provides an avenue for understanding complex interplay between geogenic and anthropogenic factors influencing groundwater chemistry as well as groundwater evolution and hydrogeochemical characteristics for policy implementation as follows:

- The insights identified in this study regarding the sources of chemical constituents in groundwater are fundamental in providing strategic guides in developing and implementing effective policies for sustainable use and management of water resources. This would offer a more protective measures in ensuring healthy groundwater quality, particularly in areas prone to water contamination which is in line with SDG6 targets.
- Knowledge of various groundwater types in the study area and their distinct characteristics coupled with hydrochemical characteristics might influence decision on groundwater development projects in the region. Stringent policies should be implemented in areas at risk of contamination due to overexploitation or pollution from natural and anthropogenic sources.
- The Government of Nigeria should actively participate in the regional developmental projects being conducted by Lake Chad Basin
 Commission (LCBC) within Lake Chad region in restoring and safeguarding Lake Chad that provides a substantial groundwater
 recharge zones to the region's aquifers. These projects are essential in ensuring sustainable groundwater management and safeguarding water resources for the present population and the unborn generation.
- A significant percentage of potassium concentration due to incongruent weathering of potassium feldspar in the Precambrian basement region and nitrate contamination by irrigation return flows and pit latrines in irrigated and residential areas, as illustrated in Fig. 9 necessitates the development of detailed irrigation master plan and pollution model specifically addressing nitrate contamination from intensive synthetic fertilizing, pit latrines and sewerage systems. These activities would enable a more detailed analysis of nitrate-related contamination and support the establishment of stringent policies for proper land use planning to mitigate the potential health risk posed by nitrate in drinking groundwater.
- This study has undoubtedly provided an avenue for future research that would focus on trace element contamination and its
 associated health risk as well as the application isotope hydrology to understand groundwater recharge source/origin and possibly
 tracing various groundwater pollution sources.

6. Conclusion

This study employed an integrated approach of geochemical modeling, bivariate plots, ionic ratios, and chemometric analysis to explore geochemical evolution and mechanisms controlling the groundwater chemistry as well as origin/sources of chemical constituents in groundwater of the transboundary KYB. The groundwater in the basin is subjected to increased exploitation coupled with contamination from geogenic and anthropogenic inputs. The following conclusion can be drawn from the study:

- The Pearson's correlation analysis showed a positive relationship between EC and major ions, except with pH and Fe in the wet season and pH and F– in the dry season.
- Nitrate significantly correlated with all variables in the dry season, possibly due to excessive use of synthetic fertilizer during irrigation and nitrification from pit latrines.
- PCA results indicate that groundwater samples in PC1 were influenced by geogenic and anthropogenic sources, PC2 indicated the
 influence of agricultural and domestic waste inputs, while PC3 suggests fluoride enrichment due to mineral weathering and industrial activities.
- The R-mode HCA identified three cluster groups influenced by: both geogenic and anthropogenic factors (Cluster I); synthetic fertilizers and nitrification from pit latrines (Cluster II); and interactions among various geochemical processes (Cluster III).
- Q-mode HCA identified three water types with increasing mineralization levels influenced by geogenic, anthropogenic, and evaporation processes.

- Piper diagrams indicated Ca-HCO3, Na-Cl, and mixed water types, suggesting that groundwater is influenced by mineral weathering, ion exchange, and evaporation processes.
- Gibbs plots, bivariate plots, molar ratios, and chloroalkaline indices (CAI-I, CAI-II) confirms that the groundwater chemistry is
 influenced by geochemical processes like mineral weathering, evaporation, and ion exchange processes.
- Saturation indices revealed that most samples were undersaturated with respect to dolomite, calcite, and fluorite, as a result of
 carbonate precipitation.
- A significant percentage of groundwater samples (96.7 % in wet and 99.2 % in dry season) had partial pressure of CO₂ above atmospheric pCO₂ levels, as a source of acid for mineral dissolution.
- The study proves that geochemical modeling and chemometric analysis are effective techniques for assessing geochemical mechanisms and various chemical constituents within aquifer systems characterized by diverse contamination sources.
- Recommendations from the study include detailed trace element analysis and their associated health risk assessments, as well as
 isotope hydrology studies to identify groundwater recharge sources and contamination, supporting efficient groundwater management to mitigate significant risks to human health.

CRediT authorship contribution statement

Abdulrahman Shuaibu: Writing – review & editing, Writing – original draft, Validation, Software, Resources, Project administration, Methodology, Investigation, Funding acquisition, Formal analysis, Data curation, Conceptualization. **Robert M Kalin:** Writing – review & editing, Validation, Supervision, Funding acquisition, Conceptualization. **Vernon Phoenix:** Writing – review & editing, Validation, Supervision, Funding acquisition. **Ibrahim Mohammed Lawal:** Writing – review & editing, Visualization, Validation, Software.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.ejrh.2024.102098.

Data availability

Data will be made available on request.

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