

CLASSIFICATION AND GEOLOGICAL ENVIRONMENT IDENTIFICATION OF PRODUCED WATER IN OILFIELDS

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Abstract

Formation water or saline water known as produced water is generated in great volumes in onshore and offshore oil and gas production. It is important to determine its physical and chemical properties because of they are representing as a function of different factors such as depositional environment, water types and water classification. This study highlights the nature of the formation water and its characterizations in the light of different parameters and factors, in two oilfields namely Sarir and Nafora for oil producing wells, and in five gathering centers in Sarir oilfield throughout the chemical analysis of sixty water samples to determine the physiochemical characterizations. The data obtained from the chemical analysis were used to classify formation water and to determine the environment of deposition using Sulin's system. The results revealed that the type of formation water in both two fields are belonging to CaCl_2 type and deep marine environment. The Stiff diagrams of the studied formation water of Sarir field were constructed for five gathering centers and compared with different patterns of Stiff diagrams. Most of the studied water exhibits a similarity of these patterns, but some of them show slightly deviation, this may be attributed to the type of formation and the source of brine.

Index Terms: Oilfield, formation water, produced water, salinity, cations, anions, concentration.

المخلص

يُعرف الماء المُنتج المصاحب للنفط بالماء الطبقي أو الماء الأجاج، في كلٍ من حقول النفط والغاز البرية والبحرية المنتجة على حدٍ سواءٍ. ومن الأهمية تعيين الخصائص الفيزيائية والكيميائية لهذه المياه وذلك لأنها بمثابة دالة لعوامل عدة مثل البيئة الترسيبية، وتحديد نوع المياه ووضع تصنيف لها. هذه الدراسة تسلط الضوء على المياه المنتجة في حقلين نفطيين هما حقل السرير وحقل النافورة المنتجين للمياه المصاحبة للنفط بغرض تبيان مواصفات وخصائص هذه المياه. من جانبٍ آخر فقد تم دراسة خمسة مراكز تجميع بحقل السرير حيث تضمنت الدراسة تحليل 60 عينة من المياه لتعيين المواصفات الفيزيوكيميائية. وقد أُستخدمت النتائج المتحصل عليها من التحليل الكيميائي في تصنيف الماء الطبقي وتحديد البيئة الترسيبية باستخدام نظام سولين *Sulin's system*. حيث أظهرت النتائج أن نوع الماء الطبقي في كلا الحقلين يندرج ضمن النوع كلوريد الكالسيوم $CaCl_2$ وأن البيئة الترسيبية التي تشكل فيها هي بيئة بحرية عميقة. من جهةٍ أخرى تم دراسة الخصائص باستخدام مخططات ستيف للماء الطبقي بحقل السرير وتمت مقارنتها بأنماط مخططات ستيف لمراكز التجميع الخمسة، حيث أسفرت معظم المياه المدروسة عن وجود تماثل مع هذه المخططات عدا القليل منها الذي أظهر حيوداً طفيفاً والذي يعزو بطبيعة الحال إلى نوع التكوينات الصخرية ومصدر الماء الطبقي. المصطلحات الدالة: حقول النفط، الماء الطبقي، الماء المنتج، الملوحة، الكاتيونات، الأنيونات، التركيز.

1. Introduction

A basic knowledge of the physics and chemistry of subsurface waters and petroleum is essential for petroleum engineers because many problems associated with exploration, formation damage or production problems, enhanced oil recovery, wettability, and others are directly associated with the physical and chemical behavior of subsurface waters and petroleum as a whole, or as groups of constituents.

Water produced during oil and gas extraction operations may be called formation water, oilfield water and brine, and constitutes the industry's most important waste stream on the basis of volume. Today, nearly 115 billion barrels per year (bbl/y) of water are produced worldwide as a by-product of oil and gas [1,2]. In average, for every barrel of oil, three barrels of water are produced from oil wells. As the well ages, this ratio dramatically increases, sometimes rising as high as 50 barrels of water per barrel of oil produced [3]. The water varies greatly in quality and quantity and in some cases the water can be a useful by-product or even a salable commodity. reduced water handling methodology depends on the composition of produced water, location, quantity and the availability of resources [4].

2. Location of Study

The Sarir "C" field, which is 35 miles (56 km) long and 25 miles (40 km) wide covering approximately 146 square miles (378 km²), is located in a tectonically relaxed region of South Eastern Libya. Nafora oilfield was located in the south of Libya at Oasis area (Figure 1).

3. Constituents of Produced Waters from Oil and Gas Fields

Organic constituents are normally dispersed or dissolved in produced water and include oil and grease and a number of dissolved compounds. All formation waters contain dissolved solids, primarily sodium chloride.[5]. Generally, oilfield waters contain much higher concentrations of solids than seawater does. Formation waters have been reported with total solid concentrations ranging from as little as 200 ppm to saturation, which is approximately 300,000 ppm. Seawater contains about 35,000 ppm total solids. The dissolved cations commonly found in oilfield waters are Na⁺ Ca⁺⁺, and Mg⁺⁺ occasionally K⁺, Ba⁺⁺, Li⁺, Fe⁺⁺, and Sr⁺⁺ are present. The most common anions are Cl⁻, SO₄⁻ and HCO₃. Also CO₃, NO₃, Br, I, B₃O₃, and S are often present [6].

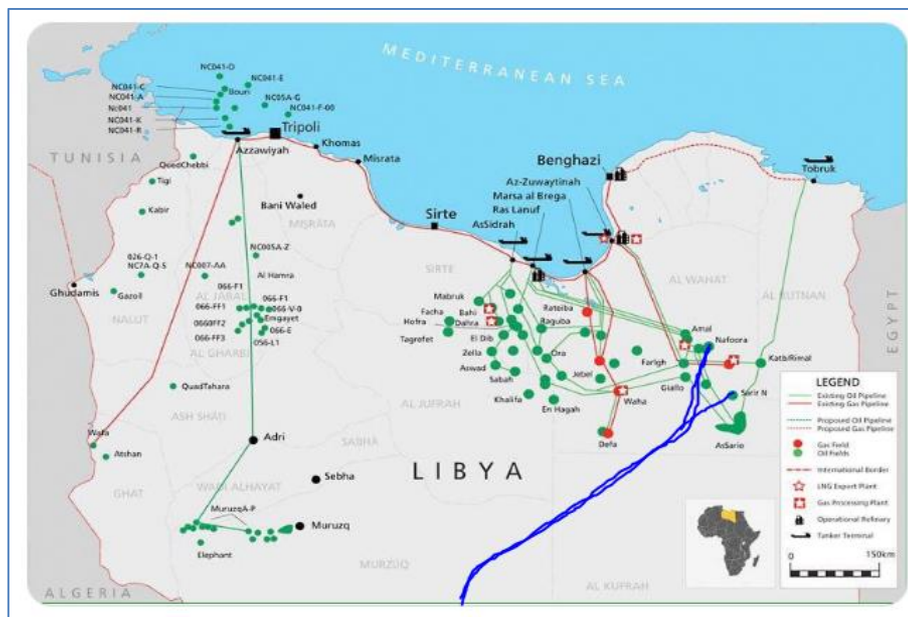


Fig. 1 A map showing wells locations of Sarir and Nafora oilfields [7]

4. Materials and Methods

4.1. Sampling

Sixty water samples were collected from the investigated wells and gathering centers. All the collected samples were preserved in accordance with guidelines and International Standards and analyzed in the field laboratory using XRF analytical technique.

4.2. Study Objectives

The main objective of this study is to recognize the large amounts of produced water that associated to the petroleum industry and investigate the characterization properties and classification of formation water produced in both Sarir and Nafora oilfields.

5. Oilfield Waters

The genesis of petroleum is intimately associated with shallow marine environments; hence, it is not surprising that water found associated with oil generally contains dissolved salts. Petroleum source rocks originally formed in lakes or streams, and the porous sediments that became today's petroleum reservoirs could have

acquired saline waters by later exposure to marine waters. After the original deposition, however, the oilfield sedimentary formations have histories of subsidence, uplift, reburial, erosion, etc. Therefore, the chemistry of the original water may have been altered by meteoric water, marine water infiltration at a later time, changes of salt types [8].

The common order of evaporative deposition from seawater in a closed basin is: calcium carbonate (limestone) > calcium-magnesium carbonate (dolomite) > calcium sulfate (gypsum) > sodium chloride (halite) > potassium chloride (sylvite). Dolomite begins to precipitate when the removal of calcium from solution increases the Mg/Ca ratio. The residual brines (containing unprecipitated salts at any period) may migrate away from the basin and leave the evaporites behind, or they may become the interstitial water of sediments that are rapidly filling the basin [9]. In accumulating marine clastic sediments, aerobic bacteria consume the free oxygen in the interstitial waters and create an anaerobic environment in which the anaerobes become active and attack the sulfate ion, which is the second-most important anion in seawater. The sulfate is reduced by the bacteria to sulfide, which is liberated as hydrogen sulfide (marsh gas) [9]. Thus, the composition of saline oilfield waters, or brines, is quite different from the average composition of seawater (Table 1). The ions in the brine are enriched with respect to seawater. Several mechanisms of enrichment are possible: (1) the original seawater may have evaporated if it was trapped in a closed basin; (2) movement of the waters through beds of clay may have concentrated cations by acting like a semipermeable membrane allowing water to pass through, but excluding or retarding the passage of dissolved salts; and (3) mixing with other subsurface waters containing high salt concentrations.

There are many reactions between the ions that can occur as the environmental conditions change with burial. Commonly, the salinity (total amount of dissolved salts, or TDS) of petroleum-associated waters increases with depth (there are a few exceptions to this). The principal anions change in a characteristic manner as depth increases: (1) sulfate is the major anion in the near-surface

waters; (2) below about 500 meters, bicarbonate may become the principal anion; and (3) in brines from deeper formations, chloride is the principal anion. The Ca/Na ratio increases, whereas the Mg/Na ratio decreases [9].

Table 1: Oilfield brine, average composition of seawater compared to Sarir, Nafora and Smackover, Arkansas [9]

Constituent	Seawater (mg/L)	Sarir (Libya) (mg/L)	Nafora (Libya) (mg/L)	Smackover brine (mg/L)
Lithium	0.2	-	-	174
Sodium	10,600	17000	897	67,000
Potassium	380			2,800
Calcium	400	1813	500	35,000
Magnesium	1,300	1387	280	3,500
Strontium	8	-	-	1,900
Barium	0.03	-	-	23
Boron	5	-	-	130
Copper	0.003	-	-	1
Iron	0.01	-	-	41
Manganese	0.002	-	-	30
Chloride	19,000	15953	1367	172,000
Bromide	65	-	-	3,100
Iodide	0.05	-	-	25
Sulphate	2,690	365	210	45

The concentrations of salts in formation waters are expressed as weight percent (wt%): milligrams per liter (mg/L) or parts per million (ppm). The quantities are related as follows: 1% = 10,000 ppm and mg/L = ppm/density .

Where ionic reactions are involved, the contents of ions are expressed as milliequivalents per liter (meq/L). One meq of a cation reacts quantitatively with exactly one meq of an anion:

$$meq/L = (mg/L) \times (valence/molecular\ weight)$$

The calcium and magnesium cation concentrations of subsurface waters are probably functions of the origin of the specific oilfield water as well as its history of contact with infiltrating waters. These cations undergo reactions forming dolomite and enter into ion

exchange reactions; consequently, they are normally found in lower concentrations than sodium cations. Other cations are present in concentrations less than 100 mg/L [10].

Oilfield waters are frequently referred to as connate or interstitial water, which is found in small pores and between fine grains in water-wet rocks [11]. “Connate” implies that the water is the original fossil water present in the rocks from the time of original deposition. One cannot be certain of this because the original water may have been displaced or mixed with other waters during the geologic history of the sedimentary formation.

6. Chemical Composition and Classification of Formation Water

6.1. Chemical Composition

Because of long-time contact with the reservoir rocks and crude oil, the formation water contains a considerable deal of metallic salts, such as sodium salt, potassium salt, calcium salt, and magnesium salt, and for this reason, the formation water is also called brine. Formation water has competitively higher salt concentration that makes the formation water distinguished from the land-surface water. The total salt concentration in formation water is called salinity.

Positive ions commonly encountered in formation water include Na^+ , K^+ , Ca^{2+} , and Mg^{2+} , and some other positive ions such as Ba^{2+} , Fe^{2+} , Sr^{2+} , and Li^+ also take a little share; the commonly encountered negative ions include Cl^- , SO_4^{2-} , HCO_3^- , CO_3^{2-} , NO_3^- , and Br^- , as well as some trace ions.

The microorganisms cause the corrosion of casings and build-up in pores of the formation. However, the origin of them is still rather vague for us. It is guessed that they may originally exist in the trapped and closed reservoirs or, perhaps, are brought to the underground formations during drilling operations.

6.2. Salinity and the milligram-equivalent concentration of the ions

Salinity is the total concentration of the mineral salts in water and is expressed with the unit mg/L or ppm (one millionth). It is the total

concentration of both the positive and negative ions in it. Salinity varies within a wide range from several thousand to hundreds of thousand (unit mg/L) in different oil field. A high salinity, which sometimes can be up to 3×10^5 ml/L, is a typical value for formation water [12].

The positive and negative ions react with each other in equivalent amounts, the salt concentration in formation water is usually expressed by milligram-equivalent concentration which equals the ratio of a certain ion's concentration to its equivalent value. The compounded equivalent values of some common ions in oil formations are shown in Table 2. For example, the concentration of chloride ion (Cl^-) is 7896 mg/L, and the compounded equivalent value is 35.3; then, the milligram-equivalent concentration of the chloride ion can be obtained as follows: $7896/35.3 = 225.6$ mg/L.

Table 2: Compounded equivalent values of common ions in oil formations

No.	Positive ion	Compounded equivalent values	No.	Negative ion	Compounded equivalent values
1	Ba^{2+}	68.7	1	CO_3^{2-}	30.0
2	Ca^{2+}	20.0	2	HCO_3^-	61.0
3	H^+	1.0	3	Cl^-	35.3
4	Fe^{3+}	27.9	4	OH^-	17.0
5	Fe^{2+}	18.6	5	O^{2-}	8.0
6	Mg^{2+}	12.2	6	SO_4^{2-}	48.0
7	Na^+	23	7	S^{2-}	16.0
8	Si^{2+}	43.8	8	SO_3^{2-}	40.0

6.3. Water hardness

Hardness of water is a term used to indicate its content of calcium and magnesium ions. In the chemical flooding (e.g., injection of polymers or active agents), high water hardness tends to worsen the effect of the stimulations by causing precipitation of the chemical agents.

In addition to the formation water's total salinity and water hardness, and to know quantitatively the species and individual

concentration of the ions, upon which the water type and the depositional environment it represents can be determined [12].

7. The Classification of Formation Water

Roughly, the total concentration of the formation water can be known in accordance with its total salinity. For further grasp of the individual concentration of every ion, either positive or negative, and their corresponding proportions, the formation water is classified to different types.

7.1. Classification of Water—Sulin's System

According to the chemical components in water or, in other words, their milligram-equivalent concentration:

$$\frac{Na^+}{Cl^-}, \frac{Na^+ - Cl^-}{SO_4^{2-}}, \frac{Cl^- - Na^+}{Mg^{2+}}, \frac{SO_4^{2-}}{Cl^-} \text{ and } \frac{Ca^{2+}}{Mg^{2+}}$$

and water is classified to four types.

- (1) *Sulfate-sodium (Na_2SO_4) type*: It represents an environment of continental washing.
- (2) *Bicarbonate-sodium ($NaHCO_3$) type*: It represents an environment of continental deposit.
- (3) *Chloride-magnesium ($MgCl_2$) type*: It represents an environment of marine deposit.
- (4) *Chloride-calcium ($CaCl_2$) type*: It represents an environment of deep enclosed construction.

This method discussed above is called Sulin's water classification system. [12].

7.2. Determination of Water Types

As illustrated in Figure 2, offers the principle on which the water types are determined:

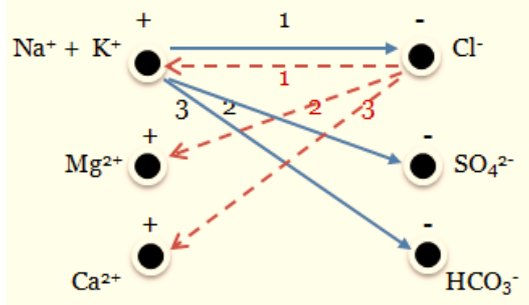


Fig. 2 Diagram of the order in which the ions combine with each other [12]

(1) When $Na^+/Cl^- > 1$, the redundant Na^+ will combine with the SO_4^{2-} or HCO_3^- ions and forms Na_2SO_4 type or $NaHCO_3$ type. Further judgment can be continued as follows:

(a) When $\frac{Na^+ - Cl^-}{SO_4^{2-}} < 1$, it suggests that after balancing the Cl^- , the redundant Na^+ will combine with the SO_4^{2-} , As a result, when the whole combination reaction ends is Na_2SO_4 the water belongs to the Na_2SO_4 type.

(b) When $Na \frac{Cl^- - Na^+}{Mg^{2+}} > 1$, it suggests that after balancing the Cl^- and SO_4^{2-} , the redundant Na^+ will combine with the HCO_3^- , upon the water is named $NaHCO_3$ type.

(2) When $Na^+/Cl^- < 1$, the redundant Cl^- will combine with the Mg^{2+} or ions and forms $MgCl_2$ type or $CaCl_2$ type. Further judgment is as follows:

(a) When $\frac{Cl^- - Na^+}{Mg^{2+}} < 1$, it suggests that after balancing the Na^+ , the redundant Cl^- will combine with the Mg^{2+} , As a result, the $MgCl_2$ type is formed.

(b) When $\frac{Cl^- - Na^+}{Mg^{2+}} < 1$, it suggests that after balancing the Na^+ and Mg^{2+} , the redundant Cl^- will combine with the Ca^{2+} , the water is named $CaCl_2$ type.

The procedure for water-type determination is summarized in Table 3.

Table 3: Sulin's system of water classification [12]

Equivalent Proportion	Parameter for judgment	Water type	Environment
$\frac{Na^+}{Cl^-} > 1$	$\frac{Na^+ - Cl^-}{SO_4^{2-}} < 1$	Na ₂ SO ₄ type	Continental washing (land, surface water)
	$\frac{Na^+ - Cl^-}{SO_4^{2-}} > 1$	NaHCO ₃ type	Continental deposit (water in oil and gas field)
$\frac{Na^+}{Cl^-} < 1$	$\frac{Cl^- - Na^+}{Mg^{2+}} < 1$	MgCl ₂ type	Marine deposit (water in oil and gas field)
	$\frac{Cl^- - Na^+}{Mg^{2+}} > 1$	CaCl ₂ type	Deep enclosed construction (gas field water)

8. Results and Discussion

The obtained results of chemical analysis were presented in Table 4.

Table 4: Chemical composition and water type of the formation water in oil fields [12]

Oil field	Total salinity (mg/L)	Na ⁺ (K ⁺)	Mg ²⁺	Ca ²⁺	Cl ⁻	SO ₄ ²⁻	HCO ₃	CO ₃ ²⁻	Water type
Daqang (China)	16,316	5917	11	95	7896	18	2334	45	CaCl ₂
Gudao (China)	3,228	1038	13	25	1036	0	1116	0	NaHCO ₃
Shengli (China)	17,960	4952	836	620	10,402	961	187	0	CaCl ₂
Renqiu (China)	178	21	9	20	43	18	67	0	CaCl ₂
East Texas (America)	64,725	23,029	536	1360	39,000	216	578	0	CaCl ₂
Gachsaran (Iran)	95,313	33,600	30	1470	55,000	4920	-	293	CaCl ₂
Furrial-Musipan (Venezuela)	5643	1739	59	54	1780	-	2001	-	MaCl ₂
Sarir (Libya)	26325	17000	1387	1813	15953	365	268.4	260	CaCl ₂
Nafora (Libya)	29250	897	280	500	1367	210	320	315	CaCl ₂

8.1. Formation Water Type

To determine the investigated formation water type, we use the chemical composition of the formation water of oil fields as following procedure.

1. Judge the equivalent proportion of the ions whose valence is 1:

$\frac{Na^{+} + K^{+}}{Cl^{-}} = < 1$, As initial judgment, the water can be $MgCl^{2-}$ or $CaCl^{2-}$ typed.

2. Because that $\frac{Cl^{-} - Na^{+}}{Mg^{2+}} = > 1$, the given water belongs to the $CaCl^{2-}$ type.

Table 4 shows the chemical compounds and types of the formation water exist in some oil fields comparing with Sarir and Nafora brine. The obtained results revealed that the investigated formation water of both Sarir and Nafora oil fields were belonging to the $CaCl_2$ type. This is clearly similar with that compared oilfields as shown in Table 4.

On the other hand, Figures 3 & 4 are depicted a comprehensive comparison among the formation waters characteristics; whereas, this comparison revealed the variance of total salinity, cations and anions in the studied formation waters in these oilfields.

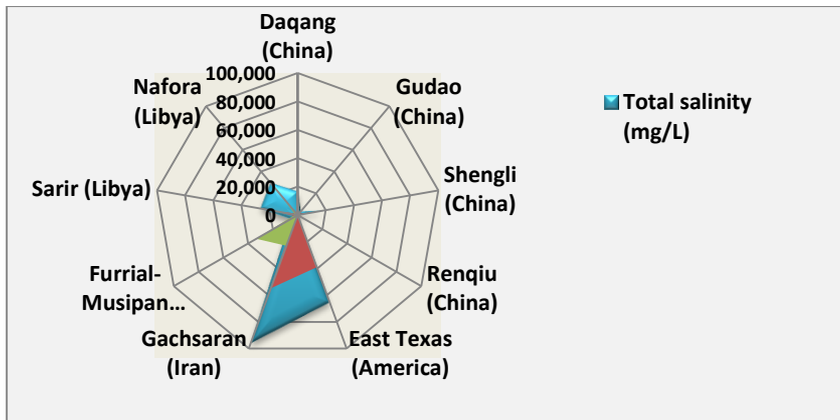


Fig. 3 Radar charts comparing the total salinity and Cl^{-} of formation waters

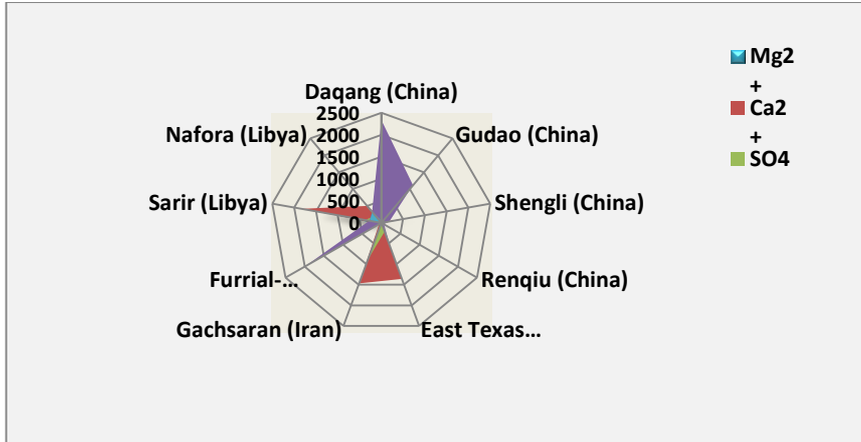


Fig. 4 Radar charts comparing the cations and anions of formation waters

8.2. Stiff Diagrams

The chemical analysis of formation water involved a range from 6 to 10 of numbers that representing the different composition of ions. So, the best way to represent the data was developed in patterns, these patterns called Stiff diagrams (Figure 5) .

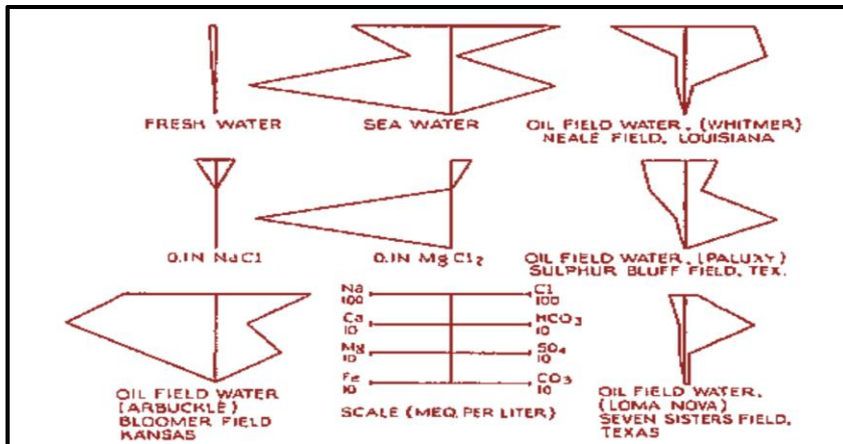


Fig. 5 Stiff patterns of formation water [12]

To construct Stiff diagrams for formation water, five gathering centers were investigated (Table 5), and we must be applied the following procedure:

Table 5: Determination of brine density

Gathering center No.	Unit	GC1	GC2	GC3	GC4	GC5
Parameter						
Total dissolved solids, TDS	ppm	68200	11800	220718	163545	24600
Total dissolved solids, TDS	%	6.82	1.18	22.07	16.4	2.5
Brine density	lb/cu ft	65.20	62.50	72.40	69.70	63.30
ρ_w	g/cc	1.044	1.001	1.159	1.116	1.014

8.3. Conversion the Concentrations from ppm to mg/l

1. Conversion the total dissolved solids, TDS from ppm into percentage (%) by dividing to 10^6 .
2. Brine density (lb/ft³) can be determine from the chart of density using the obtained TDS in percentage (%).
3. Conversion the brine density (lb/ft³) into g/cc.
4. Multiply the concentrations of cations and anions of brine in ppm by brine density to obtain the concentrations in mg/l.
5. The results obtained are presented in Table 6.

Table 6: Conversion the concentrations to mg/L

Chemical properties	Concentration (mg/L)				
	GC1	GC2	GC3	GC4	GC5
Calcium	28188	1892	17582	38455	43605
Magnesium	12047	1448	34572.9	932	10748
Salinity, NaCl	214020	27483	240017	206181	17238
Chloride, Cl ⁻	179568	16655	145446	125118	10444
Total iron, Fe	91	15.0	80.8	217.6	76.6
Sulphate, SO ₄	302	125	423	595.0	266.7
Alkalinity, HCO ₃	146.2	268.7	85.5	100.4	152

8.4. Conversion the Concentrations from mg/L into meq/L

To convert from mg/L to meq/L for cations and anions follow the following steps:

The atomic weight of Ca is 40.08 g/g mole, valence is 2 eq wt/g mole, then the equivalent weight is:

$$\text{For Ca, } \frac{40.08 \frac{g}{g \text{ mole}}}{2 \frac{eq \text{ wt}}{g \text{ mole}}} = 20.04 \text{ g/eq wt} = 20.04 \text{ mg/meq.}$$

and the milliequivalent is:

$$\frac{28188 \frac{mg}{l}}{20.04 \frac{mg}{meq}} = 1406.6 \text{ meq/L}$$

$$\text{For Na, } \frac{22.99 \frac{g}{g \text{ mole}}}{1 \frac{eq \text{ wt}}{g \text{ mole}}} = 22.99 \text{ g/eq wt.} = 22.99 \text{ mg/meq.}$$

$$\frac{214020 \frac{mg/liter}{g \text{ mole}}}{22.99 \frac{mg}{mg/meq}} = 9309.3 \text{ meq/L}$$

$$\text{For Mg, } \frac{24 \frac{g}{g \text{ mole}}}{2 \frac{eq \text{ wt}}{g \text{ mole}}} = 12 \text{ g/eq wt.} = 12 \text{ mg/meq.}$$

$$\frac{12047 \frac{mg/liter}{g \text{ mole}}}{12 \frac{mg}{mg/meq}} = 1003.9 \text{ meq/L}$$

$$\text{For Cl}^-, \frac{35.5 \frac{g}{g \text{ mole}}}{1 \frac{eq \text{ wt}}{g \text{ mole}}} = 35.5 \text{ g/eq wt.} = 35.5 \text{ mg/meq.}$$

$$\frac{179568 \frac{mg/liter}{g \text{ mole}}}{35.5 \frac{mg}{mg/meq}} = 5058.3 \text{ meq/L}$$

$$\text{For Fe, } \frac{56 \frac{g}{g \text{ mole}}}{3 \frac{eq \text{ wt}}{g \text{ mole}}} = 18.7 \text{ g/eq wt.} = 18.7 \text{ mg/meq.}$$

$$\frac{91 \frac{mg/liter}{g \text{ mole}}}{18.7 \frac{mg}{mg/meq}} = 4.87 \text{ meq/L}$$

$$\text{For SO}_4, \frac{96 \frac{g}{g \text{ mole}}}{2 \frac{eq \text{ wt}}{g \text{ mole}}} = 48 \text{ g/eq wt.} = 48 \text{ mg/meq.}$$

$$\frac{302 \frac{mg/liter}{g \text{ mole}}}{48 \frac{mg}{mg/meq}} = 6.29 \text{ meq/L}$$

$$\text{For HCO}_3, \frac{61 \frac{g}{g \text{ mole}}}{2 \frac{eq \text{ wt}}{g \text{ mole}}} = 30.5 \text{ g/eq wt.} = 30.5 \text{ mg/meq.}$$

$$\frac{1462 \frac{mg/liter}{g \text{ mole}}}{30.5 \frac{mg}{mg/meq}} = 47.9 \text{ meq/L}$$

The other calculations can be carried out as the above procedure. The results are presented in Table 7.

Table 7: Conversion from mg/l to meq/l for cations and anions

Chemical properties (Cations and anions)	Ion concentration (meq/L)				
	GC1	GC2	GC3	GC4	GC5
Calcium, Ca ⁺⁺	1406.6	94.41	877.4	1918.9	2175.9
Magnesium, Mg ⁺⁺	1003.9	120.7	2896.1	77.7	895.7
Sodium, Na ⁺	9309.3	1200.1	10481.1	9003.5	752.8
Chloride, Cl ⁻	5058.3	469.2	4097.1	3524.5	294.2
Total iron, Fe	4.87	0.80	4.3	11.6	4.10
Sulphate, SO ₄	6.29	2.60	8.81	12.40	556
Alkalinity, HCO ₃	47.9	8.8	2.8	3.29	4.98
Carbonate, CO ₃	0.0	0.0	0.0	0.0	0.0

The Stiff diagrams of the studied formation water were constructed for the different gathering centers as shown in Figures 6 through Figure 10 and compared with different patterns of Stiff diagrams. Most of the studied water exhibit a similarity of these patterns, but some of them show slightly deviation, this may be attributed to the type of formation and the source of brine.

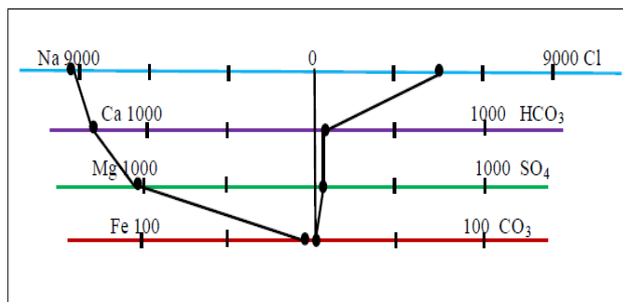


Fig. 6 Stiff diagram for GC1

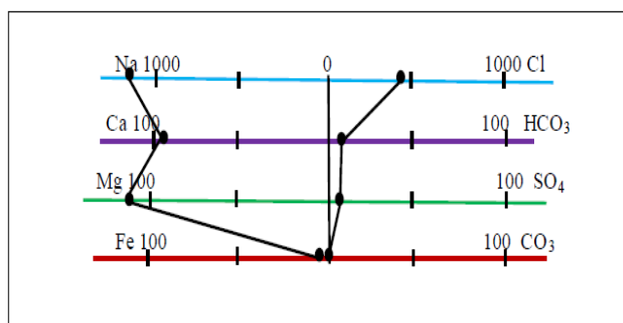


Fig. 7 Stiff diagram for GC2

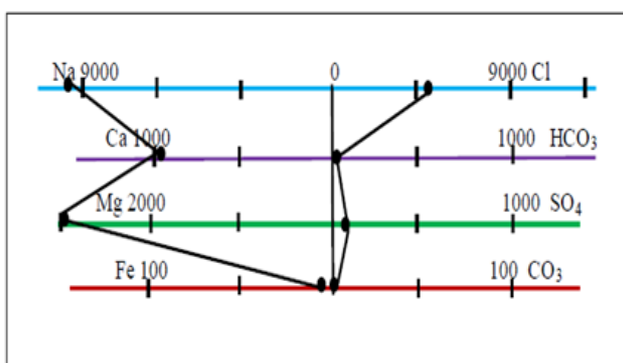


Fig. 8 Stiff diagram for GC3

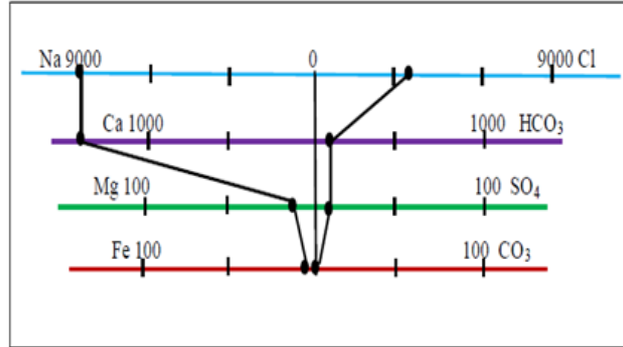


Fig. 9 Stiff diagram for GC4

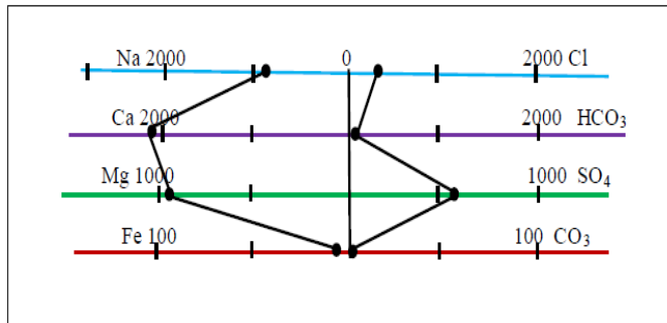


Fig. 10 Stiff diagram for GC5

9. Conclusion

Water analysis is very important for petroleum engineering industry from both upstream and downstream activities. The characteristics of Sarir and Nafora oil fields produced water revealed the following:

1. Oilfield produced waters contain elevated levels of salinity, TDS, cations and anions more than the standards characteristics of seawater.
2. Many characteristics of the formation water such as salinity, TDS, Na and Cl exhibit variable values, this may due to nature of reservoir formations.
3. According to the Sulin's system of water classification; the obtained results revealed that the investigated formation

water of both Sarir and Nafora oil fields were belonging to the CaCl_2 type and deep marine environmental deposition.

4. The comparing between the constructed Stiff diagrams for the studied formation water of gathering centers with Stiff diagrams patterns exhibit similarity, but some of them show slightly deviation, this may be attributed to the type of formation and the source of brine.

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