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Quantitative Analysis of Amorphous Silica and Its Influence on Reservoir Properties: A Case Study on the Shale Strata of the Lucaogou Formation in the Jimsar Depression, Junggar Basin, China

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Abstract: To establish a new quantitative analysis method for amorphous silica content and understand its effect on reservoir properties, the amorphous silica (SiO₂) in the shale strata of the Lucaogou Formation in the Jimsar Depression was studied by scanning electron microscopy (SEM) observation, X-ray diffraction (XRD), and X-ray fluorescence spectrometry (XRF). Amorphous silica shows no specific morphology, sometimes exhibits the spherical or ellipsoid shapes, and usually disorderly mounds among other mineral grains. A new quantitative analysis method for observing amorphous SiO₂ was established by combining XRD and XRF. On this basis, while the higher content of amorphous SiO₂ lowers the porosity of the reservoir, the permeability shows no obvious changes. The higher the content of amorphous SiO₂, the lower the compressive strength and Young's modulus and the lower the oil saturation. Thus, amorphous SiO₂ can reduce the physical properties of reservoir rocks and increase the reservoir plasticity, which is not only conducive to the enrichment of shale oil but also increases the difficulty of fracturing in later reservoir development.

Keywords: amorphous SiO₂; X-ray diffraction; X-ray fluorescence spectrometry; scanning electron microscope; quantitative analysis; reservoir properties

1. Introduction

The success of shale gas exploration and development in North America has promoted the development of the shale gas industry around the world. At present, successful exploration and development of shale gas in China is mainly concentrated in the Sichuan Basin and surrounding areas, such as the Weiyuan, Zhaotong, Zhengan, and Jiaoshiba areas [1–3]. The shale sections containing commercial scale gas in these areas are located at the top of the Wufeng and the bottom of the Longmaxi Formations, corresponding to the 2-3 graphitic biozones of the Wufeng Formation and the 1-4 graphitic biozones of the Longmaxi Formation [4,5]. These high-quality shale sections contain high content of silica: as much as 60% [6–11]. Although there are different opinions about the evidence of biogenesis, most researchers consider that the silica in these high-quality shale sections has biogenic sources [12–17]. Shale oil sources are mainly concentrated in basins in China, where lacustrine shale is widely developed, such as the Ordos, Songliao, and Bohai Bay Basins. Shale oil exploration has been particularly successful in the second member of the Kongdian Formation in

Cangdong Depression of Bohai Bay Basin, where commercial-scale oil has been obtained in several wells [18]. This quartz-feldspathic shale exhibits good quality, high TOC content, and high hydrocarbon potential [18]. The tuffaceous shale sections of the Lucaogou Formation shale strata in the oil reservoir of the Malang Depression contain high total organic carbon and exhibit high hydrocarbon generation potential. These tuffaceous shales are also mainly composed of quartz and feldspar [19]. In both marine shale gas and or lacustrine shale oil reservoirs, silica is an important component, having a significant impact on shale reservoir properties, organic matter enrichment, shale oil and gas accumulation, and fracturing potential [14,15,20–25]. Hence, silica is a hot spot in shale reservoir research at present.

Studies on silica diagenesis have shown that the end members are amorphous SiO₂ and crystalline quartz. The quartz can be further divided into authigenic quartz formed during diagenesis and detrital quartz from deposition. During diagenesis, amorphous SiO₂ will gradually change from the amorphous state (opal-A) to the cryptocrystalline state and finally to the fully crystalline state (α quartz), which is authigenic quartz. Amorphous SiO2 can be from biological organisms or an abioticearly diagenesis stage. Some studies suggest that amorphous SiO₂ has already transformed into crystalline quartz during early diagenesis (Ro is 0.35%~0.5%) [23,24]. Others suggest that the conversion of amorphous SiO₂ to crystalline quartz in shale reservoirs may be much later, because amorphous SiO₂ has been seen in the middle diagenetic stage A (Ro is 0.5%~1.3%) [21]. During clay mineral conversion, a large amount of silica is generated, and its content is closely related to mineral composition, crystallinity, and thermal conditions; it also affects the physical properties and brittleness of the reservoir [26–28]. The influence of amorphous SiO₂ on reservoir properties can make a large difference in different evolution stages. From the beginning of diagenesis to the cryptocrystalline state, formation porosity has been shown to be reduced from about 45% to less than 25%, and the permeability declines to be difficult to be measured [29]. In the authigenic quartz stage, reservoir physical properties and brittleness increases instead, which improves reservoir fracturability [29]. Thus, it can be seen that amorphous SiO₂ also plays a great impact on reservoir properties. If the influence of amorphous SiO₂ on reservoirs can be clarified, it will be of great significance for evaluating shale oil reservoirs and fracturing potential, especially for immature lacustrine shale oil reservoirs.

Accurate calculation of amorphous SiO₂ content is the key problem to understand the influence of amorphous SiO₂ on reservoir properties. There are currently four methods for the quantitative analysis of amorphous SiO₂ in heterogeneous systems. The first is chemical dissolution, which means removing minerals other than amorphous SiO₂. However, chemical dissolution incudes crystalline, which affects the accuracy of quantitative analysis. The second is quantitative analysis using XRD as proposed by Lin (1997) [30]. Although the method is correct in theory, human error enters into in the quantification [31]. Thirdly, Chu (1998) proposed a new quantitative XRD method based on the increment method proposed by Popović et al. (1983) [32,33], but this method required preparation of a standard sample having a known mineral composition and proportions; the error was relatively large in the actual experiment. Fourth, Huang et al. (2015) established a calculation method for amorphous SiO₂ in the Yanchang Formation shale of the Ordos Basin by using XRD combined with QEMSCAN analysis [34]. However, this method has two disadvantages. Firstly, it is too expensive to conduct large-scale tests. Secondly, the mineral composition obtained by QEMSCAN analysis can be understood as a volume percentage. Hence, it needs to be converted into a mass percentage, but the density of minerals was not determined in Huang et al. (2015) [34].

In view of the shortcomings of previous methods for calculating the content of amorphous silica [30–34], a new quantitative analysis method for amorphous silica content was established in this research, based on XRD and XRF analysis of core samples from the Lucaogou Formation in the Jimsar Depression. Through the analysis of the relationship between physical parameters, rock mechanical parameters, oil saturation, and amorphous silica content in shale strata, the effect of amorphous SiO₂ on reservoir properties and its geological significance was determined.

2. Geological Settings

The Junggar Basin is located in the northwestern part of China with an area of about 1.30×10^5 km² (Figure 1A); it is geotectonically located at the intersection of Kazakhstan, Siberian, and Tarim plates. The Jimusar Depression is in the southeast of Junggar Basin, covering an area of 1.278×10^3 km²; it is surrounded by the Shaqi Uplift to the north, the Guxi Uplift to the east, the Fukang faults zone to the south, and the Santai Uplift to the west (Figure 1B). The periphery of the Jimsar Depression is bounded by six faults (Figure 1B). The Permian Lusaogou Formation has a thickness of 200~350 m and is in conformable contact with the lower Jingjingzigou Formation and in unconformable contact with the upper Wutonggou Formation (Figure 1C). The Lucaogou Formation is mainly composed of deep and semideep lake facies formed of fine-grained, mixed sedimentary rocks [35,36]. It was formed in an intracontinental rifted saline lake basin environment, accompanied by volcanic eruptions and hydrothermal activity [37,38]. Since September 2011, J25, J23, J28, J30, and other exploration and evaluation wells have been successively drilled in the Jimusar Depression, oil testing shows industrial potential, and shale oil was discovered in the Lucaogou Formation. After nine years of development, the calculated reserves of shale reservoir have reached 11.12 × 10⁸ t [39].



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3. Materials and Methods

3.1. Materials

The samples of the Lucaogou Formation in this study are from four cored wells (S1–S4) in the Jimsar Depression (Figure 1B). We selected 42 samples that met experimental needs. Their lithology

includes tuffaceous shale (also called siliceous shale), shale, dolomite, and dolomitic mudstone. Generally, the lithology can be divided into three lithofacies: tuffaceous shale lithofacies, transitional lithofacies (also called mixed lithofacies), and carbonate lithofacies [41,42].

3.2. Experimental Method

XRD analysis was completed at Sichuan Keyuan Engineering Technology Testing Center. XRF analysis, rock mechanics experiments, and reservoir physical properties analysis were completed at the Experimental Research Center of East China Oil and Gas Branch of Sinopec.

3.2.1. XRD and XRF Analysis

The mineral composition of samples was obtained by XRD, which was determined on the premise of deducting background values through the Jade 5.0 software package. The principle of XRD analysis is that different minerals show different XRD diffraction effects. Data calculated by the XRD accurately represents the relative content of each mineral. However, XRD cannot measure the content of amorphous silica because it shows no diffraction peaks.

The secondary X-rays were emitted when the X-ray irradiated on the material. Different elements show their specific secondary X-ray with certain features or wavelength characteristics. XRF analysis uses secondary X-rays to convert the data into specific elements and their abundance. Elemental Si occurs in quartz, plagioclase, k-feldspar, clay minerals, and amorphous silica.

3.2.2. Rock Mechanics Experiment

Samples were tested using a TAW-2000 computer-controlled electrohydraulic servo testing machine under constant confining pressure conditions. The size of test samples is 25 mm (diameter) × 50 mm (length). In the process of testing, strain rate was controlled by the DUOLI microcomputer control system, mostly 0.01–0.03, which was convenient to obtain smooth stress–strain curves. The compressive strength, Young's modulus, and Poisson's ratio can be calculated by the stress–strain curves.

3.2.3. Reservoir Physical Properties

The total porosity was obtained by calculating the difference between the bulk density and the skeleton density. Permeability was obtained by calculating the expansion of He with increasing pressure (5 MPa–30 Mpa) at a constant temperature. Oil saturation was measured by nuclear magnetic resonance (NMR).

3.3. A New Method for Calculating the Content of Amorphous SiO2

In this study, a new method for quantitative analysis of amorphous SiO₂ in the Lucaogou Formation of the Jimusar Depression was established by using a combination of XRD and XRF. Through XRD analysis, the shale strata mainly consist of quartz, plagioclase, potash feldspar, dolomite, calcite, pyrite, and clay minerals (Figure 2A). Elemental Si is in quartz, plagioclase, potash feldspar, and clay minerals.

The combination of XRD and XRF can calculate amorphous silica as follows. Suppose the sample mass is M, where the mass of amorphous SiO₂, quartz, plagioclase, K-feldspar, and clay minerals are respectively represented by m_{SiO_2} , m_{quartz} , $m_{plagioclase}$, $m_{K-feldspar}$, and m_{clay} .



Figure 2. Mineral composition of different lithofacies samples in the Lucaogou Formation. (**A**) The mineral content of the different lithofacies; (**B**) clay mineral composition in the different lithofacies.

According to XRD analysis:

$$\frac{m_{quartz}}{M - m_{SiO_2}} = W_{quartz} .$$
⁽¹⁾

$$\frac{m_{plagioclase}}{M - m_{SiO_2}} = W_{plagioclase} \tag{2}$$

$$\frac{m_{K-feldspar}}{M - m_{SiO_2}} = W_{K-feldspar} \tag{3}$$

$$\frac{m_{clay}}{M - m_{SiO_2}} = W_{clay} \tag{4}$$

The W_{quartz} , $W_{plagioclase}$, $W_{K-feldspar}$, and W_{clay} represent the percentage of quartz, plagioclase, *k*-feldspar, and clay minerals measured by XRD analysis.

According to XRF analysis:

$$\frac{m_{SiO_2} \times P_{Si-SiO_2} + m_{quartz} \times P_{Si-quartz} + m_{plagioclase} \times P_{Si-plagioclase}}{M} + \frac{M}{m_{K-feldspar} \times P_{Si-K-feldspar} + m_{Clay} \times P_{Si-clay}}{M} = W_{Si}}$$
(5)

The mass percentages of Si in amorphous SiO₂, quartz, plagioclase, k-feldspar, clay minerals, and the sample are represented by P_{Si-SiO_2} , $P_{Si-quartz}$, $P_{Si-plagioclase}$, $P_{Si-K-feldspar}$, $P_{Si-clay}$, and W_{Si} , respectively.

Placing Formulas (1)-(4) into Formula (5), thus creating Formula (6)

$$\frac{m_{SiO_2} \times P_{Si-SiO_2} + W_{quartz} \times (M - m_{SiO_2}) \times P_{Si-quartz} + W_{plagioclase} \times (M - m_{SiO_2}) \times P_{Si-plagioclase}}{M} +$$
(6)

$$\frac{W_{K-feldspar} \times (M - m_{SiO_2}) \times P_{Si-K-feldspar} + W_{clay} \times (M - m_{SiO_2}) \times P_{Si-clay}}{M} = W_{Si}$$

Formula (6) can be changed to Formula (7):

$$\begin{aligned}
 W_{SiO_2} &= \frac{m_{SiO_2}}{M} = \\
 \frac{W_{Si} - W_{quartz} \times P_{Si-quartz} - W_{plagioclase} \times P_{Si-plagioclase} - W_{K-feldspar} \times P_{Si-K-feldspar} - W_{clay} \times P_{Si-feldspar} - W_{clay} \times P_{Si$$

In Formula (7), only the mass percentage of element Si in clay minerals is difficult to determine, because the molecular formulas of other minerals are known. The molecular formulas of clay minerals are variable. Therefore, the ideal molecular formulas of different types of clay minerals are applied in this research. For the mass percentage of Si in mixed clay minerals, it is calculated according to the mixed layer ratio based on XRD measurements. Molecular formulas used for kaolinite, montmorillonite, chlorite, and illite are respectively $Al_4(Si_4O_{10})(OH)_8$, $Al_4Si_8O_2(OH)_2$, $Al_6Si_4O_{10}(OH)_8$, and $Al_4(Si_8O_{20})(OH)_4$. The mass percentages of element Si in these are 21.7%, 56.3%, 19.6%, and 31.1%, respectively. The P_{clay} of the tuffaceous shale lithofacies, transitional lithofacies, and carbonate lithofacies samples can be calculated. Then, the contents of amorphous SiO₂ in these samples can be calculated by Formula (7).

4. Results

4.1. Occurrence and Characteristics of Amorphous SiO₂

The shale strata of the Lucaogou Formation in the Jimusar Depression can be divided into tuffaceous shale lithofacies, transitional lithofacies, and carbonate lithofacies [41,42]. The tuffaceous shale lithofacies is mainly composed of feldspathic minerals including quartz and feldspar. The carbonate lithofacies mainly consists of dolomite and includes dolomite and argillaceous dolomite. The mineral composition and lithology of the transitional lithofacies is primarily a hybrid of the other two lithofacies. It can be seen by SEM that in addition to the development of authigenic quartz in the shale strata (Figure 3A,B), amorphous SiO₂ is also present (Figure 3C–H). Amorphous SiO₂ shows no fixed form and usually fills randomly between mineral grains (Figure 3C–E). Some of the amorphous SiO₂ was wrapped in tuffaceous components (Figure 3F), and other forms were spherical or ellipsoid shapes having varying sizes (Figure 3G,H).





Figure 3. SEM images of quartz and amorphous silica in the Lucaogou Formation, Jimsar Depression. (**A**) Transitional lithofacies, S1 well, 3147.64 m; (**B**) tuffaceous shale lithofacies, S2 well, 3348.08 m; (**C**) transitional lithofacies, S1 well, 3147.64 m; (**D**) tuffaceous shale lithofacies, S2 well, 3343.00 m; (**E**) tuffaceous shale lithofacies, S2 well, 3348.08 m; (**F**) transitional lithofacies, S2 well, 3359.95 m; (**G**) tuffaceous shale lithofacies, S3 well, 2815.21 m; (**H**) tuffaceous shale lithofacies, S4 well, 2601.81 m; (**I**) energy spectrum analysis of point "+" in image H.

4.2. Composition Characteristics of Crystalline Minerals

Analysis of the XRD test results (Table 1) shows that the tuffaceous shale lithofacies samples exhibit the highest content of quartz-feldspathic minerals. The average content of quartz is as much as 40.26%; the average content of plagioclase and k-feldspar are as much as 16.68% and 5.26% respectively (Figure 2A). The carbonate lithofacies samples show the highest content of dolomite, reaching 63% on average. The transitional lithofacies samples present the highest content of clay minerals, which is as much as 26.14% (Figure 2A). In clay minerals, the content of the illite/smectite mixed layer is the highest, followed by illite. The average contents of the illite/smectite mixed layer in tuffaceous shale lithofacies, transitional lithofacies, and carbonate lithofacies are 41.37%, 59.86%, and 72.78%, respectively (Figure 2B). The tuffaceous lithofacies show the highest content of illite (average 37.89%), followed by transitional lithofacies (average 24.29%). The content of kaolinite, chlorite, and chlorite/smectite mixed layer is relatively low (Figure 2B).

4.3. Content of Amorphous SiO₂

Analysis of the XRF test results (Table 2) shows that the tuffaceous shale lithofacies samples have the highest content of Si, reaching 34.21% on average. As expected, the carbonate lithofacies samples exhibit the lowest content of Si, only 11.51% on average (Figure 4A). Moreover, the tuffaceous shale lithofacies samples also exhibit the highest values of Si in crystalline minerals calculated by the above method, reaching 33.18% on average (Figure 4A). According to the calculations, the shale strata of the Lucaogou Formation thereby contains a small amount of amorphous SiO₂. The tuffaceous shale lithofacies samples show the highest content of amorphous SiO₂, reaching an average of 7.07%, and the carbonate lithofacies samples show the lowest, only 1.52% (Figure 4A). Amorphous SiO₂ has a certain negative correlation with crystalline quartz (Figure 4B). During burial diagenesis, amorphous silica will gradually convert to crystalline quartz. The silica in the Lucaogou Formation is mainly derived from tuffaceous materials alteration in previous studies [17,21]. Therefore, the content of amorphous SiO₂ in the tuffaceous shale lithofacies sample is the highest among the three lithofacies. The content of silica in a sample is generally definite. Hence, the higher the content of crystalline quartz, the lower the content of amorphous SiO₂.



Figure 4. Diagrams showing (**A**) Si content tested by XRF, amorphous silica content calculated through the new method, and Si content in crystalline minerals. (**B**) cross plot of the amorphous silica content with crystalline quartz content in samples of the different lithofacies in the Lucaogou Formation.

Well	Depth (m)	Lithofacies	Clay Mineral/%	Quartz/%	K-Feldspar /%	Plagioclase/%	Calcite/%	Dolomite/%	Pyrite/%	Kaolinite/%	Chlorite/%	Illite/%	Illite/Smectite Mixed Layers/%	Chlorite/Smectite Mixed Layers/%	Illite/Smectite mixed Layer RATIO/%	Chlorite/Smectite Mixed Layers Ratio/%
S1	3063.80	Carbonate lithofacies	5	19	4	3	8	61	0	5	6	15	74	0	55	0
S1	3086.50	Carbonate lithofacies	2	10	4	5	0	79	0	0	0	50	50	0	56	0
S1	3136.85	Transitional lithofacies	28	22	0	13	3	34	0	8	9	16	67	0	70	0
S1	3147.64	Transitional lithofacies	31	21	4	8	0	33	3	25	29	14	32	0	44	0
S1	3149.00	Carbonate lithofacies	2	5	6	3	0	81	3	3	3	14	80	0	57	0
S2	3343.00	Tuffaceous shale lithofacies	22	23	5	22	0	25	3	0	0	64	36	0	40	0
S2	3315.00	Carbonate lithofacies	3	12	0	18	13	53	1	5	4	10	81	0	50	0
S2	3323.00	Transitional lithofacies	23	28	0	17	3	27	2	0	6	10	35	49	67	60
S2	3332.00	Carbonate lithofacies	2	13	0	18	5	60	2	9	6	9	76	0	49	0
S2	3344.00	Transitional lithofacies	17	17	9	12	7	35	3	0	11	24	65	0	56	0
S2	3347.00	Transitional lithofacies	39	26	0	10	10	12	3	0	7	15	78	0	55	0

Table 1. Test data table of mineral composition of different lithofacies samples in the Luchaogou Formation.

Energies	2020,	13,	6168
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		Tuffaceous														
S2	3348.08	shale	22	31	8	12	0	23	4	0	0	85	15	0	40	0
		lithofacies														
S2	3351.66	Carbonate lithofacies	2	19	0	10	24	45	0	4	4	32	60	0	52	0
S2	3353.14	Carbonate lithofacies	2	10	1	14	2	71	0	8	9	28	55	0	56	0
S2	3355.13	Transitional lithofacies	32	21	0	11	0	33	3	0	0	43	57	0	54	0
S2	3359.95	Transitional lithofacies	22	14	8	14	6	36	0	2	2	81	15	0	67	0
S2	3379.55	Transitional lithofacies	23	20	7	16	4	25	5	0	8	9	83	0	61	0
		Tuffaceous														
S2	3380.01	shale	16	48	11	20	0	0	5	12	15	0	73	0	90	0
		lithofacies														
S2	3463.00	Transitional lithofacies	27	15	6	14	0	36	2	0	0	38	62	0	51	0
S2	3477.00	Transitional lithofacies	26	20	9	14	0	28	3	0	7	13	80	0	65	0
S2	3600.34	Carbonate lithofacies	4	10	9	8	6	60	3	3	3	15	79	0	49	0
		Tuffaceous														
S3	2794.71	shale	18	40	4	21	12	4	1	13	15	9	63	0	87	0
		lithofacies														
		Tuffaceous														
S3	2795.80	shale	14	59	3	4	8	12	0	0	0	80	20	0	40	0
		lithofacies														
62	2805 21	Transitional	10	15	0	27	0	12	2	Б	6	0	80	0	70	0
33	2003.21	lithofacies	12	13	U	21	U	40	3	5	U	U	07	U	70	U
S3	2808.52	Carbonate lithofacies	7	6	10	10	10	57	0	0	0	0	100	0	52	0

Energies	2020,	13,	6168
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		Tuffaceous														
S3	2815.21	shale	27	30	9	18	6	5	5	0	0	94	6	0	40	0
		lithofacies														
		Tuffaceous														
S3	2817.81	shale	18	54	0	17	4	4	3	11	12	8	69	0	89	0
		lithofacies														
		Tuffaceous														
S3	2832.89	shale	22	38	7	10	19	1	3	0	0	98	2	0	40	0
		lithofacies														
		Tuffaceous														
S3	2857.65	shale	22	40	13	15	3	2	5	0	0	95	5	0	40	0
		lithofacies														
		Tuffaceous														
S3	2869.14	shale	23	38	8	5	0	26	0	0	0	86	14	0	40	0
		lithofacies														
		Tuffaceous														
S3	2870.01	shale	19	37	10	15	5	11	3	23	18	0	59	0	85	0
		lithofacies														
		Tuffaceous														
S3	2871.01	shale	18	35	13	23	3	4	4	14	17	0	69	0	88	0
		lithofacies														
		Tuffaceous														
S3	2873.01	shale	18	36	0	26	6	11	3	15	14	6	65	0	83	0
		lithofacies														
		Tuffaceous														
S3	2874.01	shale	23	38	0	26	4	5	4	14	16	0	70	0	86	0
		lithofacies														
		Tuffaceous														
S3	2876.81	shale	22	41	0	19	6	9	3	8	7	4	0	81	0	50
		lithofacies														
S4	2580 81	Transitional	25	20	0	16	0	37	2	12	11	49	28	0	40	0
51	_000.01	lithofacies	20	-0	U	10	Ũ	0,	-		••	17	-0	Ũ	10	Ũ

S4	2585.01	Transitional lithofacies	21	15	8	14	0	37	5	4	6	28	62	0	59	0
		Tuffaceous														
S4	2591.91	shale	21	51	0	13	8	5	2	13	16	4	67	0	86	0
		Tuffaceous														
S4	2592.21	shale	26	36	0	26	0	9	3	16	18	6	60	0	86	0
		lithofacies														
		Tuffaceous														
S4	2601.21	shale	17	50	3	15	8	3	4	12	14	7	67	0	86	0
		lithofacies														
		Tuffaceous														
S4	2601.81	shale	16	40	6	10	21	3	4	0	0	74	26	0	40	0
		lithofacies														
S 4	2607 60	Transitional	40	36	3	3	0	12	6	15	0	0	85	0	60	0
34	2007.00	lithofacies	-10	50	5	5	0	14	0	15	0	0	00	0	00	0

Well	Depth (m)	Lithofacies	Oil Saturation/%	Young`s Modulus/N*mm ⁻²	Poisson`s Ratio	Compressive Strength/Kg*cm ⁻²	Porosity/%	Permeability/mD	Si Content Test by XRF /%	Amorphous Silica Content Calculated Through the New	Calculated Si Content in Crystalline Minerals /%	Calculated Si Content in Clay Minerals /%
S1	3063.80	Carbonate lithofacies	17.6						13.535	1.017	13.184	40.196
S1	3086.50	Carbonate lithofacies	10.9				0.7041	0.0013	9.197	2.410	8.246	38.156
S1	3136.85	Transitional lithofacies							26.937	4.652	25.924	41.131
S1	3147.64	Transitional lithofacies	9.70				0.2367	0.0000923	24.348	6.921	22.612	28.963
S1	3149.00	Carbonate lithofacies					0.6041	0.00215	7.013	2.540	5.939	41.964
S2	3343.00	Tuffaceous shale lithofacies	5.80				0.5851	0.000043	29.234	11.885	26.743	34.728
S2	3315.00	Carbonate lithofacies	14.4	6.396	0.211	306.527			12.545	0.542	12.353	40.376
S2	3323.00	Transitional lithofacies	15.40	36.212	0.246	196.722			25.334	2.878	24.671	26.948
S2	3332.00	Carbonate lithofacies	1	16.381	0.353	368.663	1.1986	0.01	12.978	1.643	12.398	38.948
S2	3344.00	Transitional lithofacies	12.00						21.979	3.412	21.070	39.007
S2	3347.00	Transitional lithofacies	26.10	10.707	0.308	94.94			31.536	0.570	31.443	41.105

Table 2. Statistical table of calculated silica content, porosity, permeability, mechanical properties, and oil saturation in different lithofacies samples of the Lucaogou Formation.

		Tuffacoous										
52	2248 08	shalo		10 386	0 356	224 954			20.056	10 /8/	27 990	27 617
52	5540.00	lithofacies		40.500	0.550	524.754			50.050	10.404	27.550	52.012
		Carbonate										
S2	3351.66	lithofacies		12.181	0.225	234.695	0.9727	0.00236	13.028	0.472	12.836	38.126
		Carbonate										
S2	3353.14	lithofacies	22.7						10.224	0.522	10.027	37.074
60	22EE 12	Transitional	0.00	25 720	0 222	172 (12				2 251		20.05/
52	3333.13	lithofacies	9.90	35.739	0.333	173.012			26.236	2.251	25.762	38.836
52	3350 05	Transitional							22 9/3	8 594	20.615	33 21/
52	5557.75	lithofacies							22.745	0.074	20.015	55.214
52	3379 55	Transitional	8.60	14 625	0 202	184 049			26 952	2 838	26 346	42 938
02	0077.00	lithofacies	0.00	14.020	0.202	101.017			20.902	2.000	20.040	42.700
		Tuffaceous										
S2	3380.01	shale							39.916	6.143	39.407	44.803
		lithotacies										
S2	3463.00	Transitional					0.4650	0.0109	25.431	7.099	23.729	39.068
		lithotacies										
S2	3477.00	l'institional		18.328	0.269	114.190	1.1957	0.135	28.659	4.529	27.756	43.399
		litnoracies										
S2	3600.34	lithofacios	2.10	26.013	0.317	373.744			12.147	1.773	11.505	40.227
		Tuffaceous										
53	2794 71	shale	40 90	35 675	0 244	178 849	1 2807	0.012	34 525	2 676	34 163	41 965
00	2/ / 1./ 1	lithofacies	40.70	00.070	0.211	170.047	1.2007	0.012	04.020	2.070	04.100	41.705
		Tuffaceous										
S3	2795.80	shale	32.10	26.702	0.241	139.743	1.1727	0.0162	35.295	3.167	34.889	33.116
		lithofacies										
62	0005 01	Transitional					0 (000	0.0044	22 250	- 024		1= (00
\$3	2805.21	lithofacies					0.6000	0.0244	22.358	5.936	20.759	45.639
62	2000 E2	Carbonate		12 000	0.271	210.007	0.0104	0.00105	12.057	2 710	11.09/	44 204
53	2000.02	lithofacies		12.089	0.271	319.996	0.9194	0.00105	12.937	2./18	11.980	44.204

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		т ((
63	0015 01	Tuffaceous	0.00				0.0104	0.00044		11 505	21.007	01 504
\$3	2815.21	shale	8.90				0.9194	0.00244	32.932	11.535	31.006	31.704
		lithotacies										
~ ~		Tuffaceous										
S3	2817.81	shale	70.30	13.176	0.278	67.705	1.8524	0.0301	38.87632	0.589	38.824	44.161
		lithofacies										
		Tuffaceous										
S3	2832.89	shale							31.874	9.893	30.136	31.301
		lithofacies										
		Tuffaceous										
S3	2857.65	shale	36.60						35.511	7.841	34.474	31.604
		lithofacies										
		Tuffaceous										
S3	2869.14	shale	13.20	39.502	0.293	181.272	1.4511	0.076	31.016	8.205	29.524	32.511
		lithofacies										
		Tuffaceous										
S3	2870.01	shale	31.80						33.429	4.922	32.690	39.505
		lithofacies										
		Tuffaceous										
S3	2871.01	shale		18.927	0.327	128.162			36.128	6.621	35.307	43.130
		lithofacies										
		Tuffaceous										
S3	2873.01	shale							33.013	3.377	32.499	41.675
		lithofacies										
		Tuffaceous										
S3	2874.01	shale		35.684	0.355	132.212			36.425	4.705	35.868	43.114
		lithofacies										
		Tuffaceous										
S3	2876.81	shale	31.50	33.164	0.192	276.39			30.243	9.486	28.413	14.260
		lithofacies										
C/	2590.91	Transitional	1 70	20 504	0.287	125 174	0.5100	0 000080	<u></u>	6 616	22 22	21 520
34	2000.01	lithofacies	1.70	37.374	0.207	455.174	0.3100	0.000009	23.923	0.010	22.230	51.529

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S4	2585.01	Transitional					0.5600	0.043	22.973	3.681	22.027	39.252
		Tuffaceous										
S4	2591.91	shale	29.00						37.636	4.359	37.177	42.558
		Tuffaceous										
S4	2592.21	shale							35.947	3.382	35.535	40.529
		lithofacies										
		Tuffaceous										
S4	2601.21	shale	56.20	31.654	0.396	148.228			36.932	3.331	36.561	42.882
		lithofacies										
		Tuffaceous										
S4	2601.81	shale							31.012	9.318	29.297	33.721
		lithofacies										
C 4	2(07(0	Transitional	0.10				1 1 2 (1	0.05	26 124	1 1 (1	25.009	40 E 40
54	2607.60	lithofacies	9.10				1.1364	0.05	30.134	1.164	35.998	42.542

5. Discussion

5.1. Advantages and Disadvantages of the New Method

Compared with the previous quantitative analysis methods for amorphous SiO₂, the new method does not require chemical dissolution. The most important is that the cost of this method is much lower. The equipment required has already been widely used for a large-scale sample testing. This method also has some shortcomings: the ideal formula of clay mineral is used to calculate the mass percentage of elemental Si in clay minerals. Using illite as an example, its ideal structural molecular formula is Al₄(Si₈O₂₀)(OH)₄, and the mass percentage of Si is 31.1%. However, due to the fact that the illite in the actual sample contains impurities, its molecular formula is diverse, which introduces small errors into the calculated value.

5.2. The Influence of Amorphous SiO2 on Reservoir Properties

The silica content is mainly derived from the alteration of tuffaceous material in the shale strata. It was found through the cross plot between the calculated amorphous SiO₂ content and the reservoir physical property data that amorphous SiO₂ content was negatively correlated with reservoir porosity and permeability (Figure 5). The content of amorphous SiO₂ is negatively correlated with the content of crystalline quartz (Figure 4B). Hence, it indicates that the higher the content of crystalline quartz, the higher the porosity and permeability of the reservoir. Alteration is an important cause of pore formation in the Lucaogou Formation because it is a process of volume reduction for the total material [43,44]. From the perspective of density, it is easy to understand this process of volume reduction. The density of volcanic ash is only 2.3 g/cm³, while the mineral density after its alteration is much higher than 2.3 g/cm³, such as quartz 2.6–2.7 g/cm³. According to the law of conservation of mass, the overall volume must decrease. In other words, a large amount of silica was released during the alteration of tuffaceous components. Some silica crystallized to authigenic quartz, which increases the physical properties of the reservoir, while some silica did not crystallize and occurs between the grains in the form of amorphous SiO₂ cement, which reduces the storage space of the reservoir.



Figure 5. Cross plot of amorphous silica content with (**A**) porosity, (**B**) permeability of different lithofacies in Lucaogou Formation.

The rock mechanical parameters of the Lucaogou Formation were measured by triaxial stress experiment under given confining pressure (Table 2). The calculated content of amorphous SiO₂ was positively correlated with Young's modulus and compressive strength (Figure 6A,B). It indicates that the higher the content of amorphous SiO₂ was, the harder the samples were to be deformed and fractured. Amorphous SiO₂ cements various grains together, making the reservoir more compacted. Amorphous SiO₂ is negatively correlated with oil saturation (Figure 6D). It indicates that the existence of amorphous SiO₂ is unfavorable for hydrocarbon enrichment. Previous studies suggested that

volcanic ash would lead to algal blooms, and the alteration of volcanic ash would also generate a large number of pore spaces, which provided storage space for hydrocarbon enrichment. During volcanic eruptions, a large amount of volcanic ash was deposited with particulate organic matter and well preserved in a strong reduction environment. At last, they further condensed into kerogen and became source rocks with high organic matter. The organic matter type of Lucaogou Formation shale is mainly I~II1 type, which suggests an origin of bacteria, algae, and other aquatic organisms [19]. However, the presence of amorphous SiO₂ makes the tuffaceous shale lithofacies lack sufficient storage space. Furthermore, part of hydrocarbon migrated to the adjacent carbonate lithofacies. On the whole, amorphous SiO₂ in Lucaogou Formation in Jimsar Depression is not high in content (Figure 4A and Table 2), which is merely the same to that of K-feldspar. Therefore, the changes in reservoir properties are likely to be caused by other factors, such as the development of laminae, the direction of stress in triaxial stress experiments, and so on. In the early diagenetic stage (Ro is 0.35%~0.5%), amorphous SiO₂ has already started to crystallize to quartz in large quantities [23,24]. It can be inferred that the amorphous SiO₂ should have a greater physical influence on shale samples in the earlier diagenetic stage.



Figure 6. Cross plot of amorphous silica content with (**A**) Young's modulus, (**B**) Poisson's ratio, (**C**) compressive strength and (**D**) oil saturation of different lithofacies in Lucaogou Formation.

5.3. Factors Controlling the Conversion of Amorphous SiO₂ into Quartz

The conversion of amorphous SiO₂ into quartz in diagenesis was affected by many factors, including temperature, properties of fluid medium, burial, and formation pressure, etc. [45–48]. It was proposed that the hydrocarbon injection and formation overpressure can inhibit the formation of authigenic quartz [46–48]. However, in the same one sample, both authigenic quartz and

amorphous SiO₂ occur (Figures 3 and 7), the contents of amorphous silica in the four samples (Figure 7A–D) are 6.921%, 10.484%, 11.535%, and 9.318% (Table 2). It means temperature, fluid properties, and formation pressure was not the key factor. It was found that authigenic quartz tended to develop in pores, holes, or fractures through a large number of scanning electron microscope observations (Figure 7). It was a reasonable presumption that the authigenic quartz can only grow when there was space. Without growth space, it can only be amorphous SiO₂ without crystal morphological characteristics. The silica in shale strata of Lucaogou Formation mainly came from the tuffaceous material alteration. A large amount of silica was released. When these pores were filled with a large amount of amorphous SiO₂ merely existed in the amorphous state. Only when the silica-rich fluid entered one of those large pores, holes, or cracks was there enough space for silica to grow to authigenic quartz.



Figure 7. Scanning electron microscope of authigenic quartz in the pores, cavities, and cracks of Lucaogou Formation. (**A**) Transitional lithofacies, S1 well, 3147.64 m; (**B**) tuffaceous shale lithofacies, S2 well, 3348.08 m; (**C**) tuffaceous shale lithofacies, S3 well, 2815.21 m; (**D**) tuffaceous shale lithofacies, S4 well, 2601.81 m.

6. Conclusions

The amorphous SiO₂ in the shale strata of the Lucaogou Formation of the Jimusar Depression had no specific form and was usually mounded among mineral grains. XRD analysis measured the percentage of crystalline minerals, while XRF measured the percentage of elemental Si. Therefore, a new quantitative analysis method for calculating the percentage of amorphous SiO₂ was established by combining the two methods. The content of amorphous SiO₂ in the tuffaceous shale lithofacies of the Lucaogou Formation was the highest, with an average of 7.07%.

The calculation confirmed that the higher the content of amorphous SiO₂, the lower the porosity of the reservoir. Moreover, amorphous SiO₂ was found to be inversely proportional to the compressive strength, Young's modulus, and oil saturation of the reservoir. It indicates that amorphous SiO₂ reduces the physical properties of the reservoir, increases the plasticity, and increases the difficulty of fracturing during development for hydrocarbon extraction. The lack of growing space is the key factor affecting the conversion of amorphous SiO₂ into crystalline quartz. Thus, the existence of amorphous SiO₂ is harmful to shale reservoirs in many ways and has economic impact deleterious to oil and gas exploration and development.

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