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Molecular structures of 1-hydroxyethane-1,1diphosphonic acid for removing calcium sulfate scale under different pH conditions†

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As an important descaling agent, 1-hydroxyethane-1,1-diphosphonic acid (HEDP) is capable of dissolving calcium sulfate scale when the pH of HEDP solution is adjusted from acidic to weakly alkaline. The molecular structures of Ca-HEDP complexes were determined to understand the impact of pH on the dissolution of calcium sulfate scale. The structures of the complexes revealed that the two phosphonic acid groups and the hydroxyl group of HEDP each provide one O atom to coordinate with the Ca ion to form a stable three-coordinate configuration under alkaline conditions. The electronic structures were investigated by interaction region indicator analysis, atoms in molecules analysis, electron localization function and natural population analysis. The deprotonation of the phosphonic acid group enhances the binding of coordinated O atoms and Ca ions as the pH increases, and weak alkalinity is the optimal process condition as strong alkaline conditions result in the precipitation of hydroxide and Ca ions.

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1. Introduction

At present, a prevalent issue faced in the seawater desalination, reverse osmosis, boiler, oil production, and salt production is the scale formation of calcium sulfate in pipes and heatexchanger.¹⁻⁶ The precipitation of CaSO₄ is complicated, as it has three crystal forms known as gypsum (CaSO₄ \cdot 2H₂O), hemihydrate (CaSO₄ $\cdot \frac{1}{2}$ H₂O), and anhydrite (CaSO₄). Because the solubility of all three variants decreases with increasing temperature, CaSO₄ scale tends to precipitate from solution on heat-transfer surfaces.7 Unlike calcium carbonate scale, calcium sulfate scale has a hard texture and compact structure and cannot be directly dissolved by conventional inorganic acid, such as hydrochloric acid.8 Calcium sulfate scale is considered one of the hardest forms of scale and is difficult to remove. It can damage production equipment, such as pumps, pipes, valves, and heat-exchangers.9 The deposit can also cause the flow reduction, corrosion, and fouling of the equipment,

leading to decreased efficiency, increased maintenance costs, and even equipment failure. If the scale accumulates in the equipment, it can exacerbate blockages, increasing the risk of pressure buildup, which is undesirable for process automation and dangerous to the operator.

The common removal technique of calcium sulfate scale is that calcium sulfate is converted into calcium carbonate. The first step is to create Na₂CO₃ solution with mass fraction of 3% to 5% and to circulate under heating condition at 90 °C. The chemical reaction $(CaSO_4 + Na_2CO_3 = CaCO_3 \downarrow + Na_2SO_4)$ occurs at this time. Then, the carbonates generated in the reaction are removed by acid solution. The disadvantage of this technique is that the circulating time is long and the energy consumption is high. The alkaline solution is also used to remove calcium sulfate scale. This technique is to soak the calcium sulfate scale in NaOH solution with mass fraction of 20%, and the chemical reaction (CaSO₄ + 2NaOH = Ca(OH)₂ \downarrow + Na₂SO₄) occurs at this time. This method takes 5-9 days to remove the scale. The disadvantage of this method is to use a strong alkali and the cleaning solution is highly alkaline, which is dangerous to the operator. It is also necessary to add acid to neutralize, resulting in wastewater with high salt content. Another mechanical descaling technique is to use a pipeline pig using a mechanical knife accompanied by water to scrap the scale.^{10,11} However, calcium sulfate scale is too difficult to remove using a pipeline pig.12 Chelating agents, such as EDTA, have been successfully applied in scale removal.13,14 As a chelating agent, 1hydroxyethane-1,1-diphosphonic acid (HEDP) solution, adjusted to the weak alkaline condition, can effectively remove calcium sulfate scale, which is rarely reported.

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Paper

HEDP, a synthetic chelating agent, is utilized in a variety of applications, such as corrosion inhibitors, anti-scaling agents, dispersants in cooling water circulations, and builder components in industrial cleaners and household detergents.¹⁵ HEDP is a bisphosphonate consisting of C–PO(OH)₂ groups. The electron-rich groups of the hydroxyl and phosphonic acid groups of HEDP have strong interactions with metal ions.^{16,17} In addition, HEDP has low toxicity to aquatic organisms but causes eutrophication of water.¹⁸ So it should be avoided to be discharged into natural water bodies. HEDP solution can dissolve the calcium sulfate scale by adjusting the pH to the weak alkalinity, but cannot dissolve the scale in acidic conditions. This reaction occurs at room temperature and exhibits the benefits of high efficiency and thorough cleaning.

As a common water treatment agent, HEDP is readily available at a moderate cost and has been successfully employed in cleaning engineering. For instance, the volume of the equipment filled with cleaning solution is 100 m^3 , and the HEDP concentration is 1 wt%, which requires approximately 1 ton of HEDP. In addition, the Ca-HEDP solution after use is also easy to treat. The amount of the resulting cleaning waste is small, making it suitable for direct discharge into high pH ashflushing wastewater for treatment to avoid discharge into the natural water body.

To the best of our knowledge, there is no explanation for why HEDP has a different ability for dissolving calcium sulfate scale under different pH conditions. To uncover this phenomenon, the descaling rate of HEDP dissolved $CaSO_4$ under different pH conditions was tested, and the molecular structures of complexes formed by HEDP and Ca ions under different pH were calculated using wavefunction analysis at the atomic level. This study can provide new insights into how pH affects the stability of complexes.

2. Experimental and computational details

2.1. Experimental method

The calcium sulfate scale was collected from the brine pipeline of Pingdingshan Halon Salt Co., Ltd. The thickness of the scale layer was about 4 to 6 mm. After chemical analysis, the purity of CaSO₄ was 97.3 wt% and the rest was 0.4 wt% Fe₂O₃ and 2.3 wt% SiO₂. The samples were stored in a dry and cool place before use, and they were placed in a beaker and dried in an oven at 40 °C for 0.5 h.

HEDP was obtained from Shandong Taihe Technologies Co., Ltd with a purity of 97%. NaOH was obtained from Sinopharm Chemical Reagent Co., Ltd with a purity of 99%. To speed up the scale dissolution reaction, 5 wt% HEDP solution was used. At room temperature, 200 ml of 5 wt% HEDP solution was added to the beaker. The pH of the solutions was adjusted from 3 to 11 by adding 40 wt% sodium hydroxide solution. Then 3.00 g calcium sulfate scale samples were soaked in solutions with different pH for 6 h. After soaking, the samples were removed from the solutions and dried again in an oven at 40 °C for 0.5 h. The weight of the samples before and after soaking (*X* and *Y*, respectively) was recorded for each solution. The test was done three times in parallel, and the entire process was conducted at room temperature. The descaling rate was calculated as follows:

$$R = \frac{X - Y}{X} \times 100\%$$

where *R* is the descaling rate (unit in %), *X* and *Y* are the scale quality before and after adding solutions with different pH, (unit in g), respectively.⁸

2.2. Computational details

The structures of the Ca-HEDP complexes of all the molecules were determined by GaussView6, and the calculation was carried out by Gaussian16 software¹⁹ based on density functional theory. The B3LYP method^{20,21} with the def2-TZVP basis set²² was used for all atoms. The structure optimization and frequency calculation of the complexes were carried out in water using the SMD (solvation model based on density) implicit solvent model. The convergence threshold was set to the default value of the program. The vibration analysis of the obtained structure showed no imaginary frequency, indicating a stable structure on the potential energy surface. The optimized structures were analyzed by the Multiwfn program²³ for wave function analysis. The wave function analysis included the geometric configuration, interaction region indicator (IRI) analysis,24 extended charge decomposition (ECDA) analysis,25,26 and atoms in molecules (AIM) analysis,27 providing insights into the electronic structures of different forms of Ca-HEDP complexes. The geometric configuration, IRI, and AIM were graphically displayed using the VMD software.28

3. Results and discussion

3.1. Dissolution of calcium sulfate by adjusting the pH of HEDP solution

As shown in Fig. 1, HEDP can dissolve calcium sulfate scale in weak alkaline solution, which is more effective compared to acidic solution. The descaling efficiency is relatively high when the pH of the system is 7.0–9.0. However, the descaling rate of HEDP decreases under strongly alkaline conditions. The ability of HEDP to dissolve calcium sulfate scale is influenced by pH, suggesting that the chelating reaction between Ca ions and HEDP is significantly influenced by pH. In cleaning



Fig. 1 The effect of pH on the descaling rate.

engineering, the conditions for removing calcium sulfate scale by HEDP are consistent with those employed in this experiment. Therefore, controlling pH is crucial in the descaling process.

As it is well known, HEDP (often represented as H_4L) can ionize four protons H^+ in water. HEDP is a quaternary organic acid with multiple deprotonation states under different pH conditions. The pKa values, as reported in ref. 29, are 1.43, 2.70, 7.02, and 11.2, respectively. HEDP can form a 1:1 chelate complex with Ca ions, as reported in ref. 30. HEDP has five forms under different pH conditions. As shown in Fig. 2, HEDP is mainly in the form of H_4L in strongly acidic solution with pH < 1.43; in the solution of pH 1.43–2.70, it is mainly in the form of H_3L^- ; in the solution of pH 7.02–11.2, it is mainly in the form of HL³⁻; and in the solution of pH > 11.2, it is mainly in the form of L⁴⁻. Different forms of HEDP have different binding affinities with Ca ions.

 $Ca^{2+} + SO_4^{2-} = CaSO_4 \downarrow \log K = 5.04$ (1)

 $Ca^{2+} + 2OH^{-} = Ca(OH)_2 \downarrow \log K = 5.30$ (2)

 $Ca^{2+} + H_2L^{2-} = CaH_2L\log K = 7.63$, (taken from ref. 31) (3)

$$Ca^{2+} + HL^{3-} = CaHL^{-}\log K = 8.16$$
, (taken from ref. 31) (4)

$$Ca^{2+} + L^{4-} = CaL^{2-}\log K = 6.04$$
, (taken from ref. 31) (5)

The stability constants of $CaH_{3}L$ and $CaH_{4}L$ are not mentioned in ref. 31. The reaction trend can be estimated by comparing the size of stability constant log *K*. Higher log *K* values indicate a stronger reaction trend. HEDP has the strongest binding ability to Ca ions in weak alkaline conditions. The log *K* values of HEDP and Ca ion are significantly larger than that of eqn (1) or (2), and the log *K* values under acidic and strong alkaline conditions are smaller than those under weak alkaline conditions, which is consistent with the reaction phenomenon. The reason for this phenomenon is explored from a molecular perspective in the next section.



Fig. 2 Species distribution diagram of HEDP.

3.2. Electronic structures of different forms of complexes

The atomic type and label of the key atoms have been marked in a(1), a(3), and a(4) of Fig. 4. The molecular structures have been calculated and shown in Fig. 4(1). Under varying pH conditions, the complexes formed by HEDP and Ca ions also vary. The forms of CaH₄L, CaH₃L, and CaH₂L are two-coordinated complexes, while the forms of CaHL and CaL are threecoordinated complexes. The distances between Ca ion and hydroxyl O7 atom of HEDP in CaH₄L, CaH₃L, and CaH₂L are 0.3850, 0.3485, and 0.3652 nm, respectively, as shown in Table 1. These distances exceed the sum of the atomic radii of the Ca and O atoms, indicating that hydroxyl does not interact with Ca ions in these three configurations. The structural formation of CaHL and CaL complexes arises from the coordination of two phosphonic acid groups and the hydroxyl group of HEDP, with each group contributing one oxygen atom to interact with the calcium ion. This interaction leads to the formation of stable three-coordinated complexes. The distances of Ca-O7 in CaHL, and CaL are 0.2638 and 0.2614 nm, respectively. In general, the Ca-O2, Ca-O3, and Ca-O7 bond lengths shorten from CaH₄L to CaL, indicating that the binding ability of HEDP to Ca ions becomes stronger with increasing pH. HEDP forms a stable complex with Ca ions, which is the essential reason that HEDP can dissolve calcium sulfate scale.

The stability of the five complexes can be determined by the binding energies calculated using the formula $E_{\text{binding}} = E_{\text{complex}} - E_{\text{ligand}} - E_{\text{Ca}}$. The binding energies increase from CaH₄L to CaL, indicating that the binding ability between HEDP and Ca ions becomes stronger with increasing pH.

The interaction region indicator (IRI) method²⁴ can be used to graphically investigate chemical bonds and weak interactions. The position, intensity, and the type of weak interaction can be intuitively understood through the colour of the IRI isosurfaces. The corresponding chemical interpretations are shown in Fig. 3. The green region represents the van der Waals (vdW) interaction. The reddish region indicates the steric

Table 1 Partial calculated bond lengths (nm) and binding energies (kJ mol^{-1}) of the complexes

	-					
	Ca–O2	Ca-O3	Ca	a-07	Ebinding	
CaH ₄ L	0.2387	0.2368	0.	0.3850		
CaH ₃ L	0.2396	0.2350	0.3485		-68.628	
CaH_2L	0.2322	0.2332	0.	3652	-90.351	
CaHL	0.2296	0.2361	0.2638		-116.703	
CaL	0.2291	0.2308	0.2614		-140.984	
ρ λ	b > 0 sign($c_2 < 0$	$\lambda_2)\rho$ decreases	$\rho \approx 0$	$sign(\lambda_2)\rho$ increases	$\rho > 0$ $\lambda_2 > 0$	
-	0.04				+0.02	
$\rho \gg$	0					
chemical	bond notable H-bond, h	e attraction: alogen-bond	vdW interaction	notable repulsion: Steric effect in ring and cage		

Fig. 3 The color bar of the interaction region indicator.

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hindrance effect. The bright red region signifies a strong steric hindrance effect. The bluish region indicates the presence of significant attraction, such as hydrogen bonds. A completely blue region indicates the presence of either a relatively strong weak interaction or a chemical bond.

As can be seen from the IRI isosurfaces in Fig. 4(2), there is a distinct blue region between Ca–O2 and Ca–O3 bonds from CaH₄L to CaL. The difference between Ca–O2 or Ca–O3 of different configurations cannot be distinguished from the IRI isosurface. However, the electron density of the Ca–O bond, as shown in Table 2, increases from CaH₄L to CaL, suggesting an increasing force between them. There is no direct interaction between O7 and Ca ions from CaH_4L to CaH_2L , and there is no bond path formed between them in the topological analysis below, which indicates that the O atom of hydroxyl cannot effectively interact with Ca ion under acidic conditions. However, Ca–O7 forms a dark green region in CaHL and CaL, which is significantly different from the colour of the Ca–O2 and Ca–O3 bond and indicates that the interaction between them is essentially different. According to Table 2, the electron density of the Ca–O2 and Ca–O3 bonds (0.0423 and 0.0355e),



Fig. 4 (1) Optimized structures of five forms of complexes. (2) IRI isosurfaces. (3) Molecular graphs. (4) ELF colour-filled maps. (a) CaH_4L , (b) CaH_3L , (c) CaH_2L , (d) CaH_4L , and (e) CaL.

Table 2 The topological properties (a.u.) of bond critical points of complexes

	BCP	$\rho(r)$	$ abla^2 ho(r)$	V(r)	G(r)	H(r)	$H(r)/\rho(r)$	ELF
CaH_4L	Ca–O2	0.0310	0.1708	-0.0321	0.0374	0.0053	0.1710	0.0525
	Ca–O3	0.0324	0.1796	-0.0341	0.0395	0.0054	0.1667	0.0542
CaH_3L	Ca–O2	0.0308	0.1669	-0.0316	0.0367	0.0051	0.1656	0.0531
	Ca-O3	0.0352	0.1882	-0.0376	0.0423	0.0047	0.1335	0.0617
CaH_2L	Ca–O2	0.0380	0.2033	-0.0417	0.0463	0.0046	0.1211	0.0665
	Ca-O3	0.0369	0.1976	-0.0402	0.0448	0.0046	0.1247	0.0647
CaHL	Ca–O2	0.0423	0.2172	-0.0475	0.0509	0.0032	0.0757	0.0776
	Ca-O3	0.0355	0.1838	-0.0377	0.0418	0.0041	0.1155	0.0648
	Ca–O7	0.0183	0.0938	-0.0152	0.0193	0.0041	0.2240	0.0341
CaL	Ca-O2	0.0430	0.2199	-0.0485	0.0517	0.0033	0.0767	0.0788
	Ca-O3	0.0416	0.2107	-0.0462	0.0494	0.0032	0.0769	0.0774
	Ca–O7	0.0198	0.0995	-0.0169	0.0209	0.0040	0.2020	0.0385

suggesting a strong vdW interaction between Ca–O7. As the pH increases, the color of the intra-ring force within the complexes changes from green to dark red, indicating that the intra-ring force changes from vdW interaction to weak steric hindrance.

Atoms in molecules (AIM) theory²⁷ proposed by Bader et al. can quantitatively be used to describe ionic bonds, covalent bonds, and vdW interactions. A molecular graph is defined as the network of bond paths linking neighboring nuclei.³² The molecular graphs of the five complexes are displayed in Fig. 4(3). The purple point corresponds to the nucleus critical point, which is essentially the same as the nucleus position; the orange point is known as the bond critical point (BCP), located between two atoms with a force; the yellow point is the ring critical point (RCP), and the green point is the cage critical point (CCP). It can be seen from Fig. 4(3) that Ca-O7 pair gradually approaches with increasing pH and that the number of RCPs linking Ca to HEDP rises from 2 to 3 on going from CaH₂L to CaHL, concomitant with the appearance of a CCP, indicating that the Ca-HEDP complex changes from two-coordination to three-coordination. The increase in pH is favorable for the coordination of the hydroxyl group of HEDP with Ca ions.

According to Bader's theory, the electron density $\rho(r)$ and Laplacian electron density $\nabla^2 \rho(r)$ at the BCP are closely related to the chemical bond properties. A high value of $\rho(r) > 0.2$ a.u. and a negative value of $\nabla^2 \rho(r)$ indicate a covalent character, while a small value of $\rho(r)$ and a positive value of $\nabla^2 \rho(r)$ represent a non-covalent character.^{32,33} The values of $\rho(r)$ for the Ca– O2 and Ca-O3 bonds in Table 2 range from 0.0308 to 0.0430 a.u. The low value of $\rho(r)$ and positive value of $\nabla^2 \rho(r)$ indicate a noncovalent interaction and reflect the ionic nature of the Ca-O bonds. The $\rho(r)$ and $\nabla^2 \rho(r)$ at the BCP are closely related to the strength of the chemical bond. For homogeneous chemical bonds, the larger $\rho(r)$ and $\nabla^2 \rho(r)$ generally indicate a greater chemical bond strength. It can be seen that the values of $\rho(r)$ and $\nabla^2 \rho(r)$ for the Ca–O2 and Ca–O3 bonds gradually increase from CaH₄L to CaL, indicating that the strength of the Ca-O bond becomes stronger.

The energy density H(r) reflects the electron energy at a certain point, which is the sum of the kinetic energy density G(r) and the potential energy density V(r). According to ref. 34, H(r) > 0 indicates a non-covalent interaction, while H(r) < 0

0 indicates covalent interaction. The values of H(r) for the five complexes in Table 2 are all positive, suggesting that the Ca–O bond is an ionic interaction. In ref. 35, it is proposed that the physical meaning of $H(r)/\rho(r)$ is the energy density of the unit electron at the BCP position. For covalent interactions, the more negative $H(r)/\rho(r)$, the stronger the interaction. For non-covalent interactions, the greater the value of $H(r)/\rho(r)$, the weaker the interaction. Table 2 shows that the values of $H(r)/\rho(r)$ for the Ca–O bond are all positive and decrease gradually from CaH₄L to CaL, implying that non-covalent interactions of the Ca–O bond are becoming stronger.

The electron localization function (ELF) is the probability of finding an associated electron pair in space at a given point.³⁶ The ELF ranges from 0 to 1, where low values correspond to low electron localization and high values correspond to high electron localization. The colour coding in the ELF visualization represents electron localization levels, with high values shown in red, medium values shown in yellow to green, and low values shown in blue shades. The ELF colour-filled maps were created by plotting the region between Ca and O atoms, with Ca, O2, and O7 serving as the plane. It can be seen from Fig. 4(4) that the regions between Ca-O are shaded blue, indicating that electrons in these regions are delocalized. According to Table 2, the ELF values of Ca-O are relatively low, ranging from 0.0341 to 0.0788. This suggests that the interaction between Ca and O atoms is ionic in nature, contrasting the typical covalent interaction that is represented by the red region. The proximity of Ca and O7 caused the lone pair of electrons of O7 to shift towards the Ca ion, suggesting that the interaction between the two atoms strengthens with the increase in pH. The analysis confirms that the interaction between Ca and O is ionic, in contrast to typical covalent bonds. This finding is consistent with the results of $\rho(r)$, $\nabla^2 \rho(r)$, and $H(r)/\rho(r)$ analysis.

The natural population analysis (NPA) charges summarized in Table 3 demonstrate that the deprotonation of the phosphonic groups in the complex significantly changes its charge distribution from CaH_4L to CaL. With the deprotonation of phosphonic groups, the electrons initially attracted by H atoms shift to other O and P atoms, resulting in the more negative values of O2, O3, and O7. The donor ability of coordinated O atoms to Ca ion was continuously enhanced. The charge values

Table 3NPA charges and ECDA values of the partial atoms of fiveforms of complexes

	Ca	O2	O3	07	P4	P5	C6	ECDA
CaH_4L	1.969	-1.238	-1.238	-0.875	2.451	2.451	-0.477	0.1706
CaH_3L	1.951	-1.239	-1.299	-0.889	2.436	2.420	-0.521	0.2071
CaH_2L	1.932	-1.31	-1.304	-0.899	2.408	2.398	-0.530	0.2405
CaHL	1.895	-1.357	-1.343	-0.910	2.407	2.395	-0.559	0.3181
CaL	1.480	-1.376	-1.358	-0.954	2.359	2.352	-0.573	0.3618

of Ca ions decrease from 1.969 to 1.480, indicating that the electrons of the ligand flowed to the Ca ions with the increase in pH. With the enhanced electron donating ability of O2, O3, and O7 to Ca ions, Ca–O bond lengths become shorter.

Extended charge decomposition analysis (ECDA)^{25,26} can quantitatively describe the amount of charge transfer from ligands to metal ions. From the ECDA data in Table 3, it can be seen that the number of H atoms decreases with the increase of pH, and HEDP transfers more electrons to Ca ions, which implies that the interaction of O atoms of HEDP and Ca ions increases with the increase in pH.

The above analysis suggests that a stronger alkaline condition would be expected to have a more effective scale-dissolving effect. However, the actual observed effect does not align with this expectation. The formation of $Ca(OH)_2$ precipitation in strong alkalinity interferes with the reaction between HEDP and $CaSO_4$, leading to a better scale-dissolving effect under weak alkaline conditions. Ref. 9 also suggested that high pH can result in the occupation of coordination sites by hydroxide, thereby reducing the chelation ability of HEDP.

The effect of pH on calcium sulfate scale dissolved by HEDP reveals the influence of pH on the electronic structure of complexes, and lays a theoretical basis for the molecular design of cleaning agent for calcium sulfate scale. We consider the effects of stronger organic acid cleaners from the perspectives of pH, electron-donating groups, and molecular flexibility.

4. Conclusion

HEDP can dissolve calcium sulfate scale by adjusting the pH to weak alkalinity at room temperature. The structures of the complexes formed by HEDP and Ca ions under different pH were optimized. The hydroxyl O atom of HEDP is not effectively able to coordinate with Ca ions under acidic conditions, but is able to coordinate under alkaline conditions. Two phosphonic acid groups and the hydroxyl group of HEDP contribute an O atom that coordinates with Ca ions, forming a stable threecoordinated configuration under weak alkaline conditions. With the increase in pH, the phosphonic acid groups are gradually deprotonated, transferring electrons previously attracted by the hydrogen atoms to the P and O atoms. This enhances the electron-donating ability of the O atoms to the Ca ions, transitioning from CaH4L to CaL. AIM analysis showed that the Ca-O2 and Ca-O3 bonds reflect the nature of the ionic bond. IRI analysis showed that there is a strong vdW interaction between Ca-O7 bonds. Under strong alkaline conditions, the

reaction between hydroxide and Ca ions results in the precipitation of $Ca(OH)_2$, competing with the reaction between HEDP and $CaSO_4$. Therefore, HEDP has the best effect on dissolving calcium sulfate scale under weak alkaline conditions. This study provides a new perspective on the stability of complexes under different pH conditions.

Author contributions

Xiaoyang Zhao: investigation, methodology and writing. Xingzhe Zhu: test. Xin Xiao: validation. Junmin Nan: conceptualization and funding. Meng Xu: editing and writing. Chen Wu: supervision and writing.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- 1 M. Y. Ashfaq, M. A. Al-Ghouti, D. A. Da'na, H. Qiblawey and N. Zouari, *Sci. Total Environ.*, 2020, **703**, 134726.
- 2 D. J. Park, O. D. Supekar, A. R. Greenberg, J. T. Gopinath and V. M. Bright, *Desalination*, 2021, **497**, 114736.
- 3 M. Amiri, J. Moghadasi, M. Jamialahmadi and M. P. Shahri, *Energy Sources, Part A*, 2013, **35**, 648–658.
- 4 M. Jamialahmadi and H. Mueller-Steinhagen, *Heat Transfer Eng.*, 1991, **12**, 19–26.
- 5 Y. Liu, Y. Zhou, Q. Yao, J. Huang, G. Liu, H. Wang, K. Cao, Y. Chen, Y. Bu and W. Wu, *J. Appl. Polym. Sci.*, 2014, 131.
- 6 F. Rahman, Desalination, 2013, 319, 79-84.
- 7 D. Hasson and J. Zahavi, *Ind. Eng. Chem. Fundam.*, 1970, **9**, 1–10.
- 8 M. Murtaza, S. A. Alarifi, M. Y. Rasm, M. S. Kamal, M. Mahmoud and M. Al-Ajmi, *Sci. Rep.*, 2022, **12**, 10085.
- 9 M. S. Kamal, I. Hussein, M. Mahmoud, A. S. Sultan and M. A. Saad, J. Pet. Sci. Eng., 2018, 171, 127–139.
- 10 H. He and Z. Liang, Math. Probl. Eng., 2019, 2019, 1-10.
- 11 T. T. Nguyen, H. R. Yoo, Y. W. Rho and S. B. Kim, *ISIE 2001.* 2001 IEEE International Symposium on Industrial Electronics Proceedings, 2001, vol. 2, pp. 863–868.
- 12 M. Bader, J. Pet. Sci. Eng., 2007, 55, 93-110.
- 13 M. H. Al-Khaldi, A. M. Al-Juhani, S. H. Al-Mutairi and M. N. Gurmen, SPE European Formation Damage Conference, 2011.
- 14 S. Elmorsey, SPE Middle East Oil and Gas Show and Conference, 2013.

- 15 K. Fischer, Water Res., 1993, 27, 485-493.
- 16 S. Pu, M. Chen, Y. Chen, W. Zhang, H. Soliman, A. Qu, Q. Liu, X. Tang, N. Huang and G. Wan, *Corros. Sci.*, 2018, 144, 277–287.
- 17 M. Salasi, T. Shahrabi, E. Roayaei and M. Aliofkhazraei, Mater. Chem. Phys., 2007, **104**, 183–190.
- 18 E. Rott, H. Steinmetz and J. W. Metzger, *Sci. Total Environ.*, 2018, **615**, 1176–1191.
- 19 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, G. A. Petersson, H. Nakatsuji, X. Li, M. Caricato, A. V. Marenich, J. Bloino, B. G. Janesko, R. Gomperts, B. Mennucci, H. P. Hratchian, J. V. Ortiz, A. F. Izmaylov, J. L. Sonnenberg, D. Williams-Young, F. Ding, F. Lipparini, F. Egidi, J. Goings, B. Peng, A. Petrone, T. Henderson, D. Ranasinghe, V. G. Zakrzewski, J. Gao, N. Rega, G. Zheng, W. Liang, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, О. Kitao, H. Nakai, T. Vreven, K. Throssell, I. A. Montgomery Jr, J. E. Peralta, F. Ogliaro, M. J. Bearpark, J. J. Heyd, E. N. Brothers, K. N. Kudin, V. N. Staroverov, T. A. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A. P. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, J. M. Millam, M. Klene, C. Adamo, R. Cammi, J. W. Ochterski, R. L. Martin, K. Morokuma, O. Farkas, J. B. Foresman and D. J. Fox, Gaussian 16 Rev. B.01, Gaussian, Inc., Wallingford, CT, 2016.

20 A. D. Becke, Phys. Rev. A: At., Mol., Opt. Phys., 1988, 38, 3098.

- 21 C. Lee, W. Yang and R. G. Parr, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1988, 37, 785.
- 22 F. Weigend and R. Ahlrichs, *Phys. Chem. Chem. Phys.*, 2005, 7, 3297–3305.
- 23 T. Lu and F. Chen, J. Comput. Chem., 2012, 33, 580-592.
- 24 T. Lu and Q. Chen, Chem.: Methods, 2021, 1, 231-239.
- 25 S. Dapprich and G. Frenking, *J. Phys. Chem.*, 1995, **99**, 9352–9362.
- 26 M. Xiao and T. Lu, J. Adv. Phys. Chem., 2015, 4, 111-124.
- 27 R. F. Bader and T. Nguyen-Dang, in *Advances in Quantum Chemistry*, Elsevier, 1981, vol. 14, pp. 63–124.
- 28 W. Humphrey, A. Dalke and K. Schulten, *J. Mol. Graphics*, 1996, **14**, 33–38.
- 29 L. Dong, S. Zhu, M. Xia, Y. Chu, F. Wang and W. Lei, *J. Hazard. Mater.*, 2020, **383**, 121206.
- 30 I. Cukrowski, L. Popović, W. Barnard, S. O. Paul, P. H. Van Rooyen and D. C. Liles, *Bone*, 2007, 41, 668–678.
- 31 V. Deluchat, J.-C. Bollinger, B. Serpaud and C. Caullet, *Talanta*, 1997, **44**, 897–907.
- 32 R. F. Bader, J. Phys. Chem. A, 1998, 102, 7314-7323.
- 33 R. F. Bader, Chem. Rev., 1991, 91, 893-928.
- 34 D. Cremer and E. Kraka, Angew. Chem., Int. Ed. Engl., 1984, 23, 627–628.
- 35 E. Espinosa, I. Alkorta, J. Elguero and E. Molins, *J. Chem. Phys.*, 2002, **117**, 5529–5542.
- 36 A. D. Becke and K. E. Edgecombe, *J. Chem. Phys.*, 1990, **92**, 5397–5403.