Evaluation of Solid Minerals Inhibitory Potentials on Mildsteel Corrosion

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Abstract: The use of some solid minerals in the inhibition of mild steel corrosion in acidic media was investigated using the conventional weight loss and hydrogen gas evolution method. Those considered are Talc, Gypsum, Alumina, Iron chromite and Haematite. Scanning electron microscope (SEM) and Electron Dispersive X-ray spectroscopy (EDX) studies revealed the protective effectiveness of the adsorbed film of the mineral extracts on the steel surface. Results obtained indicated the minerals inhibited corrosion of the steel in the order Talc > Gypsum > Haematite > Alumina > Chromite > Feldspar as the order of Inhibition efficiency. The resultant effect of solid minerals with Sida acuta plant extract is inhibitorily antagonistic due to induced corrosion rather than improve on the inhibition efficiency. Elements in the minerals got adsorbed and reacted with the iron in steel to form a compound of effective protection of the metal against corrosion in its environment.

Keywords: Solid Minerals, Adsorption, Sida acuta, Metal Ore, Alumina, Gypsum, Haematite, Ironchromite and Talc

1. Introduction

Solid minerals are naturally occurring substances in the form of complexes and mixtures deposited in the ground with occasional outcrops. They are known to be abundant in nature, less expensive than their synthetic alternatives and most are non-toxic except few ones such as lead and chromates. They are categorized into metallic, non-metallic or carbonaceous. They are sources of raw materials for manufacturing industries and monumental [1].

Much energy is being expended in the extraction of metals from their ore especially the most electropositive ones due to strong bonds of elements involved in ore formation. In view of this, the pure form of the metals should have strong affinity to bond with highly electronegative elements that form ore constituents of a particular metal. The use of compound or ions of the elements contained in an ore as inhibitor for the pure form or already extracted metal is reactively expected to form a coat or complex by adsorption to the surface of the metal due to affinity specificity in reaction. Extracted metals are unstable in their pure state they tend to return to their ore form on exposure to the environment through oxidation reactions. This tendency of reversing the extraction process is corrosion. Metallic iron is extracted from ore of Magnetite Fe₃CO₃, Hematite Fe₂O₃. This pure iron have affinity for oxygen and moisture, it therefore corrodes. The oxide film of the reacting species formed on the metal surface act as a barrier for further redox reactions (adhesion). Some of them do contain inhibitive compounds or ions that may be synergized with plant extract to function as green inhibitor. On the other hand they may induce rather than inhibit corrosion in some medium. Some inhibitors retard corrosion by adsorption to form a thin, invisible film only a few molecules thick. Others form bulky
precipitates that coat the metal and protect it from attack [2]. A lot of methods have been employed in preventing the corrosion of metals but the use of inhibitor is the most efficient method in most chemical processing industries. [3, 4]. As an environmentally required condition to have inhibitors that are safe, non-toxic and cheap, attention is now been focused on alternatives to the known form of inhibitors such as chromates and dichromate which are adjudged to be toxic and not environmentally friendly.

Gypsum CaSO₄.2H₂O is a whitish chemical precipitate that occurred as thick stratified sediments associated as a deposit may occasionally be tinted with either of grey, yellow pink or buff colour depending on the nature of attached minerals. It is an intermediate material used in the manufacture of plasterboard and related products for building industries [5]. It can also be used in glass bath to produce SO₂ gas an essential fining agent. Oxygen gas also produced along for effective oxidation and subsequent reaction path way for sodium sulphate formation, which is an intermediary compound for further oxidation reactions and also acts as anti-scumming agent in glass melt. Certain metals are largely resistant to gypsum, too, because the surfaces are coated with an oxide film that is stable to sulphate [6].

Haematite Fe₂O₃ which contains about 70% of Fe is produced from Pig Iron through blast furnace by the reduction of iron ore by coke. They are used in mixture with silicate for the manufacture of various forms of pigments for paints and linoleum fillers.

Feldspar could be potash microcline K₂O.Al₂O₃.6SiO₂; soda albite Na₂O.Al₂O₃.6SiO₂; lime anorthite CaO.Al₂O₃.2SiO₂ or mixture of these three in various proportions. It is a general fluxing compound used in whiteware industry for the manufacture of ceramics and related products such as pottery. It improves scratch, thermal and breaking resistance of glass when included in the glass batch [7]. It increases the chemical durability and decrease the tendency of devitrification in glass products. It also increases an enamel’s viscosity at any given temperature.

Ironchromite (MgFe)O(CrAlFe)₂O₃ is used for ferroalloy and chromium electroplating, chromium chemical productions, chrome bearing refractories, colourant in production of bottles, pigment in manufacture of gray face brick, ingredient for foundry washes and major constituent of some refractory mortars.

Talc 3 MgO.4SiO₂.H₂O soft white or light greenish white material is a cheap source of MgO which act as flux and provides the required strength in ceramics and tiles. It imparts high thermal and electrical resistance when used as insulators and in refractories that requires low thermal expansion [8]. Its resistance to acid attacks made it a material suitable for casing of appliances and equipment that needs to be structured through acute environments.

Alumina Al₂O₃, a material of several crystalline forms can be produced and also found as mineral corundum. It occurs in combined state as silicates in clay, kyanite, bauxites and laterites. It is used as abrasive for jewels of rugby and sapphire. Also used in ceramic products and applied into glass batch through feldspar [9].

Sida acuta plant is from the family Malvaceae. This is a dicot genus of about 200 species originated from Central America and occurring all over the warm tropics. A Pan tropical plant distributed and considered a common weed in most regions of the world. Shrubby perennial semi woody herbs, commonly found along road sides, waste places and rough grazing land [10]. Alkaloids such as Crytolepine5-methylindolo (2-3b)-quinolic and Quidoline from Sida acuta isolated, characterized and investigated have been reported to be posses antibacterial activities that will inhibit microbiologically induced corrosion [11] Apart from phytoconstituents present in Sida acuta which made it a potential green inhibitor, it also contain the following minerals in mg/100mg: sodium 110; Calcium 85; phosphorus 65; Iron 4.87 and magnesium 24.5 [12].

This work aimed to study iron in their combined state with other elements to form an ore. The effect of some of the elements that freely reacted with iron to form an ore was been studied to confirm their corrosive or inhibitive role at ore formation and after.

2. Materials and Method

A steel pipe of 3.0 mm thickness was sliced into 18 mm x 9 mm, using saw on lathe machine. The coupons were degreased in absolute ethanol and washed in acetone and then dried in a desiccator. [13] Elemental composition of the steel was analysed using Metal Spectrometer Analyser of Model ARL Quanko Desk OTD 226 (Optical emission) as shown in Table 1.

<table>
<thead>
<tr>
<th>Elements</th>
<th>C</th>
<th>Co</th>
<th>Mo</th>
<th>V</th>
<th>Fe</th>
<th>Si</th>
<th>Mn</th>
<th>Nb</th>
<th>Cr</th>
<th>Ni</th>
<th>Cu</th>
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<td>%</td>
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<td>0.0106</td>
<td>0.0169</td>
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<td>0.1499</td>
<td>0.0792</td>
<td>0.0445</td>
<td>0.0280</td>
</tr>
</tbody>
</table>

2.1. Digestion of Solid Minerals

Small quantity of about 5g of Gypsum, Hematite, Feldspar, Ironchromite and Alumina samples are finely ground separately from the representative sample. 1 g was taken out of this sample and weighed into a Platinum Dish moistened with distilled water. 4 mL of ratio [1:1] distilled water with concentrated H₂SO₄ and 14 mL of 40% HF were added. The solution was heated on sand bath for 3 h. The sample was cooled in water, open the lid and rinse into the dish with the sides of the dish rinsed with water, then 7 mL of 40% HF is added. It was heated on a sand bath again for about 2 h until white fumes persist for 5-10 minutes. The lid was opened to allow cool. The content was transferred into a 250 mL Conical Flask, 20 mL of ratio [1:1] Distilled water with concentrated HCl and 30 mL of water was added (the water
added was used to wash the sides of the flask). The solution was heated on the water bath without covering it for 20 minutes. [14, 15, 16]

2.2. Extraction from Plant Leaves

Fresh leaves of *Sida acuta* were dried and ground into powder. 10 g was weighed into 250 mL conical flask containing 100 mL ethanol using analytical balance Ohaus PA4102. The flask was properly corked and left to stand for 48 h with occasional shakings. The resultant mixture was filtered and residue thoroughly washed with ethanol. It was filtered and both filtrates added together. The extract was concentrated by use of rotary evaporator and the ethanol recovered. The concentrated extract was evaporated to oily solid on water by gently heating at 75°C to finally expel the leftover ethanol.

1 g of the extract was dissolved in 1 L of 1.0 M HCl to make 1 gdm⁻³ of the extract solution. All reagents were of Analar grade from (BDH Chemicals, England) and distilled water was used.

2.3. Determination of Inhibition Rate of the Minerals by Weight Loss Method

Weighed steel coupons of 18 mm by 8.5 mm by 3 mm thickness were immersed separately in 50 mL beaker containing 40 mL of digested solutions of 3 g / 100 mL of Gypsum, Hematite, Feldspar, Ironchromite, Alumina and a control of 1.0 M HCl at 30°C. They were removed at 2 h and dropped in water to discontinue corrosion reaction. It was scrubbed with brittle brush under running water to remove the corrosion products. They were rinsed in ethanol and in acetone before final air drying and weighed. Each reading was recorded up to fourth decimal place using Ohaus PA4102 electronic analytical balance. The differences in weight of the coupons were taken as the weight loss.

2.4. Determination of Inhibition Rate of the Minerals by Gasometric Method

50 mL of 1.0 M HCl solution (control) was introduced into Mylius Cell as shown in Figure 1. The coupon was dropped into the corrodent and corked airtight with vaseline. The hydrogen gas evolved by download displacement of water was recorded on the calibrated tube (inverted burrette). Readings were taken at every hour for 8 h at 30°C. The experiment was repeated for digestion solutions containing 1.0 gdm⁻³ of Talc and Gypsum. At the end of each experiment of Hydrogen gas Evolution (HE), the coupon was withdrawn from the tested solution, washed thoroughly with water followed by acetone and dried with air. [17] Both talc and gypsum were further synergized with plant *Sida acuta* plant extract at ratio 1:4 (extract: mineral).

Readings were taken in duplicates and recorded to 0.1 cm³ to compensate for errors that might have been encountered in the course of the experiment. Schematic diagram of the experiment as described elsewhere. [18].

3. Results and Discussion

With a solid minerals concentration of 3 g per 100 mL in acid solution, mildsteel in acidic talc solution deposited about the same quantity of material loss in control, other minerals deposited varying more quantities. The weight loss by the mildsteel in various solid minerals took the order of talc < gypsum < haematite < alumina < ironchromite < feldspar.

3.1. Adsorption Characteristics

In Figure 2, Talc without plant extract inhibited more than when plant extract was added as the hydrogen gas released by the three different solutions were in the order talc < talc with extract < acid solution. Thesame order was recorded for gypsum in Figure 3.

Inhibition efficiency of both talc and gypsum falls within the 1st hour but improved by the next hour. For the period of 8 hours the inhibition efficiency of Talc was averagely maintained at 70% and gypsum was on an average of 50% as indicated in Figure 6. Corrosion rate of steel in various media are in the order Control > SA+Gypsum > SA+Talc > Gypsum > Talc as shown in Figure 6. As corrosion of SA+Gypsum, SA+Talc and Gypsum increases with time that of the talc decreases with time. Inhibition efficiency of Gypsum mixed with *Sida acuta* extract initially was higher than that of Talc mixed with *sida acuta* extract but at 4th hour there was an interchange of reaction where inhibition efficiency of Talc mixed with *sida acuta* extract increases from 10% to 41% in 8 hours. While there was gypsum mixed with same plant extract depreciated continuously from 59% to 16% as indicated in Figure 4. It was observed that among the five minerals gypsum, hematite, feldspar, ironchromite and alumina that were tested for weight loss. The order of their Inhibition efficiency is talc > gypsum > haematite > alumina > chromite > feldspar. Talc deposited least material and then followed by gypsum. This implies that talc inhibited the metal most and followed by gypsum. It is expected that if the solid minerals were to be in neutral medium and not in acid or highly alkaline medium, they could got adsorbed more effectively. Some of the minerals are ore of metals and compounds of non-metals expected to have affinity to bind with Iron. Therefore elements contained in these minerals are required to be attracted to steel and adhered to its surface by chemical reactions thereby preventing direct attack by the medium.

Figure 5 indicated the higher inhibition efficiency potential of talc to be higher than that of gypsum as both were compared together.

In Figure 6, Corrosion rate of acid increased above the control at the addition of mineral extract. The adsorption of these solid minerals on the metals with plant extract reduces the rate of corrosion. It implies iron has affinity for elements that made up its ore rather than organic constituents of plant extract. The addition of extracts hindered the affinity of the iron for the elemental components of the mineral thereby reduced the adsorption of the mineral components on the metal. Plants extracts have been reported to inhibit corrosion.
of metals [19, 20, 21, 22]. The *sida acuta* extracts on its own is an effective green inhibitor on metals. The mechanism of hindrances between both potential adsorbates is yet to be well understood, but could be trace from the assumptions that: Gypsum (CaSO$_4$.2H$_2$O) is a very soluble compound, an introduction of its extract containing high concentration of sodium as mineral recorded in *sida-acuta* plant extracts [12]. Due to interaction between the two soluble compounds there could be formation of sodium sulphate a soluble compound which may stimulate corrosion rather than favour inhibition. Also the properties responsible for effective inhibition such as the number of adsorption active centres in the molecule, their charge density, the molecular size and the mode of adsorption may have been restructured.

The fact that sodium compound are not prone to corrosion inhibition of metal, hence the soluble calcium level decreases which reduces the amount of it available to be adsorped unto the surface of the metal and the ability to reduce corrosion decrease as shown in Figure 6. Plant extract containing high sodium content within temperature of range of 25°C to 60°C reduces the potency of solid mineral and hence reduces the adsorption or inhibiting properties of mineral solids by precipitating out of solution the soluble mineral that are adsorbable on metal surface. This makes the resultant solution to induce rather than inhibit corrosion. This confirms that pure metals are ready to return to their ore or oxide state in the presence of required substances and in the absence of organic extracts from plant with significant sodium content will retard adsorption of inhibitors on metals.

Figure 1. Graph of weight loss from mildsteel within 1hr in 1.0M HCl acidic solution of Talc, Gypsum, Haematite, Chromite and Feldspar.

Figure 2. Graph of Hydrogen gas evolution cm$^3$ against Time (hr) for synergized and unsynergized talc in acidic media at 30°C.

Figure 3. Graph of hydrogen gas evolution cm$^3$ against Time (hr) synergized gypsum and unsynergised gypsum in 1.0 M HCl acidic media at 30°C.

Figure 4. Graph of Inhibition efficiency % against Time(hr) of Talc, Gypsum, SA synergized with Talc and SA synergized with Gypsum solutions on steel in acidic 1.0 M HCl media.

Figure 5. Graph of % Inhibition efficiency against Time (hr) of Talc and Gypsum on steel in 1.0 M HCl acidic media.

Figure 6. Graph of Corrosion rate (cm$^3$ per hour) of steel against Time (hr) in Talc, Gypsum, SA with Talc, SA with Gypsum 1.0M HCl acidic solution.
3.2. Surface Morphological Analysis

Scanning electron microscope (SEM) image in Figure 7 shows attacks of acids on several spots on the metal. The surface of steel metal was characterized by patches of corrosion products. It has made much of α-Fe phase to dissolve into the acid solution.

Figure 8 Energy dispersive X-ray spectroscopy (EDX) spectrum. Prominent peak of iron with peaks of other impurities such as zirconium with oxygen and carbon. Low value of 61.49% Fe and 33.38% shows high activities of corrosion on steel by the acid without extract.

SEM image in Figure 9 the high degree of protection provided by synergized extract with mild depletion of material surface covered with oxide films. This inertness down the inner part of the alloy in the acidic environment was provided for by the extract thereby reduced the depreciation rate of the material.

Figure 10 EDX spectrum showed low carbon with high oxygen peaks 84% iron low value of 4% carbon and 11% oxygen which implies high corrosion rate.

Figure 7. SEM image of steel pipe in 1.0 M HCl (control).

Figure 8. EDX spectrum of mild steel in 1.0M HCl (control).
4. Conclusion

The inhibitory potential of talc on the corrosion of steel was the most effective and closely followed by gypsum on steel in acidic media. The inhibition efficiency of the minerals are in the order talc > gypsum > haematite > alumina > chromite > feldspar.

Plant extract and solid mineral were individually effective as inhibitors on metal but of different adsorption mechanism. The inhibiting power of solid minerals such as haematite, alumina, ironchromite and feldspar got depreciated when synergized with plant extract. This implies the corrosion inhibition effectiveness of resultant reaction product of plant extract with solid minerals expected to be complimentary in potency was confirmed lower compared with that of reactants. Elements under same chemical condition combined complimentary to form compound. Elements in the minerals got adsorbed unto the iron in steel for effective protection against corrosion or environmental aggression. Iron in the presence of other required elements under same chemical conditions combined to form ore.
References


