## **Original Research Article**

### D<sup>C</sup> Calculations of Mesembryanthemum nodiflorum compounds as Corrosion Inhibitors of Aluminum

#### 5 Abstract

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DFT-B $\mathbb{O}^{P}$  calculations have been performed on Mesembrine, Mesembrenone, 6 and Tortuosamine, using  $G^{(n)}$  program with complete optimization of Mesembrenol 7 geometries. Quantum parameters and thermodynamic Gibbs function have been used to 8 9 investigate the efficiency of the corrosion inhibition of each compound. Mesembrenone has 10 been found to have very good corrosion inhibition efficiency as compared to the other compounds. Quantum parameters and frontier orbitals together with calculated thermodynamic 11 12 function  $\Delta G$  for adsorption show spontaneous physical adsorption of the Mesembrenone on aluminum. 13

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15 Keywords: DFT, Aluminum, Inhibitors, Mesembryanthemum nodiflorum, Corrosion

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#### 18 **1- Introduction**

Extracts of several natural products of plant origin, containing organic compounds 19 20 with multiple bonds and having the hetero atoms (O, N, S, and P), are useful and widely used as effective corrosion inhibitors [1-6]. Aluminum is one of the most important metals 21 22 and has been used in a wide range of alloys [7]. It is the second to iron in terms of production and consumption. This is attributed to the following distinguished Aluminum 23 24 characteristics: low atomic mass, being inexpensive, environmentally friendly, pleasing appearance, and its industrial application [8,9]. This area of research is of much importance 25 due to naturally friendly, plant results are readily available, renewable sources of materials and 26 inexpensive. Plant products are organic in nature and some of the constituents including 27 28 tannins (organic and amino acids) alkaloids, and pigments are known to exhibit inhibiting action. Therefore, plant extract has become important because it is a rich source 29 of natural products which can be extracted by simple methods at low cost [10]. Recently, leaf 30 31 extract of Mesembryanthemum nodiflorum has been reported as natural and green inhibitors for aluminum corrosion [11]. 32

The leaf extract of Mesembryanthemum nodiflorum is organic and contains four major chemical constituents, which are classified as indole alkaloids (Figure 1), namely, Mesembrine, Mesembrenone, Mesembrenol and Tortuosamine.



- 37MesembrineMesembrenoneMesembrenolTortuosamine38Figure 1: Structure of major chemical constituent's indole alkaloids in the Mesembryanthemum<br/>nodiflorumnodiflorum
- In this study, the major compounds have not been isolated, instead the whole plant extract has been dealt with. Moreover, the quantum chemical parameters and molecular dynamics simulations have been performed to investigate the effect of Mesembryanthemum nodiflorum indole alkaloids as corrosion inhibitors of aluminum using density functional theory DFT to find their inhibitors activity. Furthermore, these theoretical calculations have been used to investigate the efficiency of each compound as a corrosion inhibitor of Al.
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#### 48 **2-** Computational

The molecules were optimized using density functional theory DFT/B3LYP (Gaussian 03, 49 Revision B.03) [12], the corresponding geometries of all molecules under investigation were 50 optimized without any geometric constraints for full geometric optimizations [13]. No 51 52 imaginary frequency was found, indicating minimal energy structures. Self- consistent reaction field (SCRF) was used to perform calculations in the presence of a solvent by open up an 53 inhibitor cavity within the solvent reaction field [14a]. To accelerate the calculation process, 54 PM3 semi-empirical method was used in optimizing the molecular structures of the indole 55 compounds. After that, re-optimizations for the structures were repeated by DFT/B3LYP 56 57 method using 6-31G (d) main set. This method is a Becke's three-parameter functional (B3)

and including mixture of Hartree – Fock (HF) wave function and DFT energy calculation that
adds exchange term correlation in DFT terms together with the functional of Lee, Yang,
and Parr (LYP) [15-17].

E<sub>HOMO</sub> and E<sub>LUMO</sub> (the energy of the frontier molecular orbitals) [18], the energy gap ( $\Delta$ E), the hardness ( $\eta$ ), the softness ( $\sigma$ ), the fraction of the electron transferred ( $\Delta$ N), the electrophilicity index ( $\omega$ ), and the  $\Delta$ G of adsorption of inhibitors on aluminum have been calculated for these compounds. The absolute electronegativity (X), the absolute hardness ( $\eta$ ) of the inhibitor, the softness ( $\sigma$ ), and the electrophilicity index ( $\omega$ ) are given as follow [19]:

66 X = (I+A)/2  $\eta = (I-A)/2$   $\sigma = 1/\eta$   $\omega = \mu^2/2\eta$ 

67 where the ionization potential (I) and the electron affinity (A) are calculated by the 68 following relations according to molecular orbital theory [18]:

69  $I = - E_{HOMO}$  and  $A = - E_{LUMO}$ 

and  $\mu$  represents the chemical potential and is assumed to be equal to the negative of the X [18a].  $\omega$  was proposed by Parr [19] as a measure of the electrophilic power of a molecule.

Herein, electrons flow from lower X (inhibitor) to higher X (metal) until the chemical potentials become equal.  $\Delta N$  has been calculated from the obtained values of X and  $\eta$ , from the inhibitor to metallic surface as follow [20, 21]:

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$$\Delta N = (X_{\text{metal}} - X_{\text{inh}}) / 2 (\eta_{\text{metal}} + \eta_{\text{inh}})$$

where X metal and X inh denote the absolute electronegativity and  $\eta$  metal and  $\eta$  inh denote the absolute hardness of metal and the inhibitor, respectively. The difference in electronegativity drives the electron transfer, and the sum of the hardness parameters acts as resistance [19].

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#### 80 **3- Results and Discussion**

The compounds under investigation are Mesembrine, Mesembrenone, Mesembrenol and Tortuosamine [11]. The calculated results of the energies of frontier molecular orbitals for the inhibitors are shown in Table 1.

Compound	E <sub>HOMO</sub> (eV)	E <sub>LUMO</sub> (eV)
Al	-5.98 <sup>a</sup>	0.43 <sup>a</sup>
Mesembrine	-5.54	-0.40
Mesembrenone	-5.41	-1.60

84 Table 1. Calculated HOMO – LUMO energies of the inhibitors by the DFT method

Mesembrenol	-5.16	-0.22
Tortusamine	-5.31	-0.59
a from Ref. [24]		

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The chemical reactivity is a function of the interaction between the HOMO and LUMO 87 levels of the reacting species [22] based on the frontier molecular orbital (FMO) theory. The 88 energy of the highest occupied molecular orbital E<sub>HOMO</sub> shows the electron donating ability of 89 the molecule. High value of E<sub>HOMO</sub> indicates a tendency of the molecule to donate electrons 90 to the appropriate acceptor molecule of low empty molecular orbital energy [23]. On the 91 other hand, ELUMO represents the energy of the lowest unoccupied molecular orbital and 92 indicates the ability of the molecule to accept electron [24]. Consequently, the lower the 93 94 value of E<sub>LUMO</sub>, the more the molecule accepts electrons. Therefore, when increasing HOMO and decreasing LUMO the binding ability of the inhibitor to the metal surface increases. 95 The values of the energies of HOMO and LUMO for metal- Al [25] have been compared to 96 97 the calculated values calculated for Mesembryanthemum nodiflorum major compounds to 98 determine the type of the interaction. LUMO - HOMO gaps for the interaction aluminuminhibitors (given in Table 2), show that aluminum will act as a Lewis base while the inhibitor 99 100 Mesembrenone act as a Lewis acid. So, aluminum utilizes the HOMO orbital to initiate the reaction with LUMO orbital of Mesembrenone. The interaction has a certain amount of ionic 101 102 character because the values of LUMO inh – HOMO Al gap approximately fall between 4 to 5 eV. Strong covalent bond can be expected only if the LUMO inh - HOMO Al gap is approximately 103 104 zero [26].

105 The inhibitors Mesembrine, Mesembrenol and Tortuosamine act as a Lewis base and 106 aluminum acts as Lewis acid (Table 2). In this case Mesembrenone act as cathodic 107 inhibitors while the other inhibitors act as anodic inhibitors.

108 Table 2. HOMO – LUMO gap interaction of Al- inhibitor by the DFT method

Inhibitors	$LUMO_{inh} - HOMO_{Al} (eV)$	LUMO $_{Al}$ – HOMO $_{inh}$ (eV)
Mesembrine	5.58	5.11
Mesembrenone	4.38	4.98
Mesembrenol	5.75	4.73
Tortusamine	5.39	4.88

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110 In Table 3, the energy separation,  $\Delta E_{gap} = (E_{LUMO} - E_{HOMO})$ , is an important parameter 111 and it is a function of the reactivity of the inhibitor molecule towards the adsorption on

112 metallic surface. As  $\Delta E_{gap}$  decreases, the reactivity of the molecule increases leading to an increase of the inhibitor efficiency [27]. The effectiveness of Mesembryanthemum 113 114 nodiflorum compounds under investigation as inhibitors has been further addressed by evaluating the global reactivity parameters. The electronegativity, X, the global chemical 115 116 hardness, n, the global softness,  $\sigma$ , the fraction of electrons transferred,  $\Delta N$ , and the electrophililcity,  $\omega$ , are shown in Table 3. 117

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120	Quantum	Mesembri	Mesembren	Mesembren	Tortusami
121	E <sub>HOMO (eV)</sub>	-5.54	-5.41	-5.16	-5.31
123	E <sub>LUMO (eV)</sub>	-0.40	-1.60	-0.22	-0.59
124	ΔEgap	5.14	3.81	4.94	4.72
125	I (eV)	5.54	5.41	5.16	5.31
126	A (eV)	0.40	1.60	0.22	0.59
127	X(eV)	2.97	3.50	2.69	2.95
129	η (eV)	2.57	1.90	2.47	2.36
130	σ	0.38	0.52	0.40	0.42
131	ΔΝ	0.02	0.03	0.04	0.02
132	ω	1.72	3.22	1.47	1.84
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 Table 3. Calculated quantum chemical parameters for the inhibitors

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 $X_{\rm Al} = 3.20$  ,  $\eta_{\rm Al} = 2.77$ 

136 Hard-Soft-Acid-Base (HSAB) terms have been used to discuss the bonding tendencies 137 of the inhibitors towards the metal atom and the frontier-controlled interaction concepts [28, 29]. The principle of HSAB says that hard acids prefer to co-ordinate to hard bases and soft 138 acids prefer to co- ordinate to soft bases. Metal atoms are known as soft acids [30]. Therefore, 139 140 hard molecules have a high HOMO-LUMO gap and soft molecules have a small HOMO-LUMO gap [31], and thus soft bases inhibitors are the most effective ones for metals [27]. 141 So, Mesembrenone which has the lowest energy gap and the highest softness, is expected 142 to have the largest inhibition efficiency as compared to Mesembrine, Mesembrenol and 143 Tortuosamine. This could also be confirmed by calculating another quantum chemical 144 parameter,  $\sigma$ , which measures the softness of the molecule and so its reactivity. In Table 145 3, it is shown that Mesembrenone has the larger  $\sigma$  values than the others. Table 3 also 146 presents the hardness values,  $\eta$ , obtained for the inhibitors. We note that Mesembrine, 147 Mesembrenol, and Tortuosamine have the larger hardness values than Mesembrenone. This 148 noticed tendency is the reverse of what has been obt with d for softness. Therefore, the 149

150 inhibitor with the smallest value of global hardness (hence the highest value of global 151 softness) is the best<sub>x</sub> is because a soft molecule is more reactive than a hard molecule [32].

152 The fraction of transferred electrons ( $\Delta N$ ) is also calculated and tabulated in Table 3. values of Mesembrenone and Mesembrenol are greater than the  $\Delta N$  values of ΔN 153 Mesembrine and Mesembrenone. The electrophilicity index,  $\omega$ , shows the ability of the 154 inhibitor molecules to accept electrons from aluminum (Table 3). Mesembrenone exhibits 155 the highest electrophilicity value as compared to the electrophilicity values of Tortuosamine, 156 Mesembrine and Mesembrenol; consequently, this observation confirms its high capacity to 157 accept electrons. The observed electrophililcity value for Mesembrenone is attributed to the low 158  $E_{LUMO}$  of Mesembrenone ( $E_{LUMO}$ = -1.60 eV) compared to other compounds. That is, 159 aluminum acts as Lewis base while Mesembrenone act as Lewis acids (cathodic inhibitor). 160 Also, the Al atoms can accept electrons from inhibitor molecule to form a coordinated bond 161 (anodic inhibitor). The inhibitor molecule can accept electrons from aluminum atom to form 162 back-donating bonds depending on the orientation of optimized structure of the inhibitor on the 163 spatial (Figure 2). These donation and back-donation processes strengthen the adsorption 164 165 of Mesembrenone onto the aluminum surface and increase the inhibition efficiency.



176 calculated and given in Table 4.

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Inhibitors	Cathodic Inhibitors	Anodic Inhibitors	Dipole Moment
			(Debye)
Mesembrine		22.4	2.7
Mesembrenone	-28.8		2.2
Mesembrenol		49.5	1.2
Tortuosamine		24.4	2.9

177 Table 4. Calculated  $\Delta$  G (kJ mol<sup>-1</sup>) values of the investigated inhibitors

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All compounds show positive  $\Delta G$  values except Mesembrenone. The difference between 179 physical adsorption and chemical adsorption depends on the magnitude of Etbs free energy 180 changes [33-35]. For physical adsorption,  $\Delta G$  value is in the range of 0 to 40 KJ.mol<sup>-1</sup>, whereas 181 for chemical adsorption,  $\Delta G$  value is in the range of -80 to 400 KJ.mol<sup>-1</sup> -80 to 400 KJ.mol<sup>-1</sup>. T 182 suggested mechanism for Mesembrenone is physical adsorption because  $\Delta G$  value is -28.8 183 KJ.mol<sup>-1</sup>, whereas the other compounds show non-spontaneous process. The measured  $\Delta G$  value 184 for nodiflorum extract at temperature 298K is  $-11 \bigcirc$  KJ.mol<sup>-1</sup>, which suggests a physical 185 adsorption [11]. It is difficult to make comparison between the nodiflorum extract [11] and the 186 present theoretical results because we perform individual calculations for each compounds. 187

#### 188 **4-** Conclusion

It can be concluded that theoretical calculations gave a good picture about the 189 190 Mesembryanthemum nodiflorum leaf extract by suggesting that Mesembrenone shows the 191 most inhibition efficiency as compared to the other compounds, because it has low E<sub>LUMO</sub> that can form a strong interaction with aluminum to act as cathodic inhibitor. Also, the highest 192 electrophilicity of Mesembrenone as compared to the other compounds which represents a 193 measure of the electrophilicity power of the molecule. Gibbs free energy while other 194 compounds show non-spontaneous Gibbs free energy [11,36]. The leaf extract shows 195 spontaneous energy which may suggests that is corrosion inhibition is mainly from 196 Mesembrenone. 197

#### 198 **References**

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