Preparation and Characterization of Mg/Fe-LDH Nano Hybrids with (O, O-Diethyl O-3, 5, 6-Trichloro-2-Pyridyl Phosphorothioate (Chlorpyrifos) by Direct and Indirect Ion Exchange Method

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Abstract: The study present the synthesis nano compound th rough a process intercalation of insecticide (chlorpyrifos) in Mg/Fe-LDHs by indirect ion exchange. Characterization of this compound (chlorpyrifos, Mg/Fe-LDHs and nano compound) done by DXR technique  and FT-IR spectrophotometer before and after intercalation formation of nano compound, then cooperation to sure the form nano hybrid organic-inorganic compound and the two and three dimensional image of the above compound in Atomic Force Microscope (AFM) for the layer and nano compound. The aim of this study, is the synthesis hybridnano compounds can be controlled release insecticide (Chlorpyrifos) by ion exchange process for limited of environment pollution. The results appeared the release of hybrids to specific liquids in different (Na₂CO₃, Na₂PO₄) concentration (0.5 m) by direct and indirect  ion exchange method. It was found to be contorted by parabolic diffusion at the opening of the process then again the whole process is governed by pseudo-second order rate appearance according to Lagregrans equation addition the studied percentage of rate release for both anionic (Chlorpyrifos) from Mg/Fe-LDHs The results high percentage of release in different liquid follows this arrangement: Na₂CO₃>Na₂PO₄.

Key words: Image, insecticide, compound, cooperation, FT-IR

INTRODUCTION

Insecticides can be biological activity molecules that have extrusive use in agriculture, leading to direct contact by uses it is also possible that these particles seep into soil and groundwater one of the problems experienced by this sector it also has effects on human health which affects the human nervous system as well as the growth and immunity of children adding to the risk of causing lung cancer in order to reduce the phenomenon of pollution by these materials it is possible to use a method to control release of these molecules in the form of ions and applying the technique of nanoparticles, represented by the (LDHs) are a group of compounds carry positively charged layers with anions and water molecules intercalated in the interlayer and often called anionic clay comparing with the more conservative cationic clay is the most commonly investigated anionic clay and is rarely found in nature (Cavani et al., 1991). The chemical composition of LDHs is represented by general formula (Vaccari, 1998; Khan and O’Hare, 2002; Rives, 2001; Evansa and Duan, 2006) [M^{2+}_{1-x}M^{3+}_x(OH)_x]^{(3-x)/2} [An_{n-x}YH_2O]^{x-} divalent cation such as M^{2+} = Mg^{2+}, Zn^{2+}, Co^{2+}, Ni^{2+}, Mn^{2+} is a trivalent cation such as M^{³+} = Al^{3+}, Fe^{3+}, Cr^{3+}, Ga^{3+}, V^{3+}. The M^{0}/M^{Ⅲ} molar ratio usually lies between (2 and 5). The layers as a result, take a positive charge (Ookubo et al., 1993). This value is attributed to the charge density of the hydroxide basal layer, namely, Anion Exchange Capacity (AEC). LDHs layer have a positive charge diffusion on the surface of them due to the M^{³+} substituted for the M^{⁺²} and the interlayer space is neutralized by the intercalation of anions with water molecules. The intercalation of various anions into LDHs has been classify by the following methods: calcination-rehydration (reconstruction), ion exchange, co-precipitation, thermal reaction and hydrothermal reactions (Aisawa et al., 2001), (Aⁿ⁻) is an ion exchangeable anion such as OH⁻, Cl⁻, NO₃⁻, CO₃⁻, SO₄²⁻ and various organic anions (Arizaga et al., 2009; Raki et al., 1995; Whilton et al., 1997) but also organic anions, complex anions, herbicides, insecticides (Franklin et al., 1995; Carlino, 1977), pharmaceuticals (Ookubo et al., 1993).

These layers (Mg/Fe-LDHs) have the capability to capture active molecular such as phosphorus organic insecticide (chlorpyrfooe) in the form anion bonded between the layer to form nano crystalline hybrid organic inorganic where the layer represent the host part but the packed anion represent the guest as shown in Fig. 1.

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MATERIALS AND METHODS

Experimental part: *Synthesis of Mg/Fe-LDHs by co-precipitation method. The natural NO₃-LDHs were prepared by a standard Co-precipitation method. A mixed dissolved weight of (2.5 g) of (Mg(NO₃)₂.6H₂O) and (2.01 g) of (Fe(NO₃)₃.9H₂O) when R = 2 in 100 mL ethanol, solution at 44°C with stirring and the postponement for 18 h. The solution pH was adjusted at (Raki et al., 1995) for the Mg/Fe-LDHs by drop wise addition of (2M) NaOH solution during the instillation of the mixed solution separate the precipitate the washing by DW and drying at 40°.

*The nano compound prepare by two methods

First method (indirect ion exchange): Dissolved (2.5 g) of Mg(NO₃)₂, 6H₂O (2.01 gm) Fe(NO₃)₃.9H₂O in 100 mL ethanol when molar Ratio (R = 2), adjusted the pH at (10) by (2M) sodium hydroxide, gradually addition 500 ppm of (chlorpyrifos) dissolved in 100 mL ethanol incubate the solution in water bath with shaking at 44°C for 18 h, separate the precipitate the washing by DW and drying at 40°.

Second method (direct ion exchange): Dissolve (0.4 g) of Mg/Fe-LDHs which prepare by Co-precipitation method. In 100 mL ethanol, gradually addition 1000 ppm of (chlorpyrifos) dissolved in 100 mL ethanol, adjusted the pH at (5.5) by (2M) sodium hydroxide. Incubate the solution in water bath with shaking at 44°C for 18 h, separate the precipitate the washing by DW and drying at 40°.

*Kinetic controlled release of (ascorbic acid): Using direct ion exchange to release (chlorpyrifos) from host (Mg/Fe-LDHs) to dilute solution (0.5 M) from Na₂SO₄, Na₂PO₄ then measure the release concentration of anion ion by U.V ray at the λₘₐₓ = 260 nm.

*Study of percentage to release (chlorpyrifos) from nano hybrid compound: Using UV ray to study percentage to release anion by taking solution (0.001 g) nano compound dissolve in 3.5 mL from HCl (2M) then measure the absorbance at λₘₐₓ to determine the total concentration of anion in layer C_T as the following equation.

RESULTS AND DISCUSSION

Figure 2 explains of X-ray diffraction through the different in the value of thin layer before and after intercalation the anion ion in the layer by using the Bragg low (Bragg, 1993) (2dsinθ = nλ) while (λ = 1.540562Å), n = 1 is the different in value of the thin layer causes by the affect of volume and the chemical structure for the organic molecular in the layer double hydroxide and figure show the crystal level is (hkl) from (003), (006) and (009), the before the intercalation d = 8.8Å of (003) is angle (5°), d = 4.4Å of (006) is angle (10°) and d = 2.9Å but after the intercalation of (chloropyrophos) d = 29.5Å at the same of the crystal level the angle is deflection (1.5°), d = 7.3 of (006) is angle (6°) and d = 4.9 Å of (009) is angle (9°) this result show the succeed intercalation of (chloropyrophos) between Mg/Fe LDHs. Figure 3-5, explains was characterized by using Atomic Force Microscope (AFM) for the two and three-dimensional image to layer (Mg/Fe NO₃-LDH) and hybrid nano compounds recorded in direct indirect ion exchange method (Mg/Fe-chlorpyrifose-LDH).
Fig. 2: Powder DXR patterns for (A) Mg/Fe NO₃-LDH and (B) Mg/Fe-chlorpyrifos-LDH

Fig. 3(a, b): AFM image of (Mg/Fe-LDH), (a) Two-dimensional image and (b) Three-dimensional image

Figure 3a the image shows two-dimensional layer before intercalation process where molecular clusters appear while Fig. 3b were show the three-dimensional image high molecular assemblies which has limits...
Table 1: The diameters, sizes and aggregation of the molecular in (Mg/Fe NO3-LDH)

| Diameter (nm) | Volume (%) | Cumulation (%) | Diameter (nm) | Volume (%) | Cumulation (%) | Diameter (nm) | Volume (%) | Cumulation (%) |
|--------------|------------|----------------|--------------|------------|----------------|--------------|------------|----------------|
| 65.00        | 4.46       | 4.460          | 85.00        | 9.82       | 41.96          | 105.00       | 6.25       | 84.82          |
| 70.00        | 10.71      | 15.180         | 90.00        | 14.29      | 56.25          | 110.00       | 8.04       | 92.86          |
| 75.00        | 8.04       | 23.210         | 95.00        | 13.39      | 69.64          | 115.00       | 6.25       | 99.11          |
| 80.00        | 8.93       | 32.140         | 100.00       | 8.93       | 78.57          | 120.00       | 0.89       | 100.00         |

Fig. 4(a, b): AFM image of (Mg/Fe-(chloropyrophose)-LDH) prepared by indirect ion exchange method, (a) Two-dimensional image and (b) Three-dimensional image

(0.39-0.61 nm), shows the diameters sizes and aggregation of the molecular in layer (Mg/Fe NO3-LDH) the means of particle size (87.5 nm) Table 1 before the intercalation process anion.

Figure 4a the image shows two-dimensional of nano compound prepared by indirect ion exchange method showing molecular clusters of spherical shapes while Fig. 3b were show the three-dimensional image for section of the surface of the nano compound Mg/Fe-chlorpyrifose-LDH showing the high of the molecular clusters that are within limits (0.28-1.28), suggesting (Shaheed et al., 2014) the manufacture of nano
compound from insecticide (chloropyrifos) and layer (Mg/Fe-DH) as shown in Table 2 means of particle size of nano compound (108.08 nm).

Figure 5a the image shows two-dimensional of nano compound prepared by direct ion exchange method showing molecular clusters of spherical shapes while Fig. 3b were show the three-dimensional image for section of the surface of the nano compound Mg/Fe-chloropyrifos-LDH showing the high of the molecular clusters that are within limits (1.35-5.18), suggesting (Shaheed et al., 2014) the manufacture of nano compound from insecticide (chloropyrifos) and layer (Mg/Fe-DH) as shown in Table 3 means of particle size of nano compound (112.05 nm).

Figure 6-8 explains FT-IR spectrum of insecticide (chloropyrophos) layer (Mg/Fe-LDH and (Ni/Al-chloropyrifos-LDH).

The chloropyrophos shows the assimilation peaks of Aliphatic $\nu$C-H stretch in the (2966 cm$^{-1}$), the absorption peaks of Aromatic $\nu$C-H stretch in the 3059 cm$^{-1}$ at 844 cm$^{-1}$ of absorption peaks of $\delta$C:H aromatic, absorption peaks of acidic $\nu$C-O stretch at 1163 cm$^{-1}$, absorption
Table 3: The diameters, sizes and aggregation of the molecular in (Mg/Fe(cloropyrifose-LDH) prepared by direct ion exchange method

| Diameter (nm) | Volume (%) | Cumulation (%) | Diameter (nm) | Volume (%) | Cumulation (%) | Diameter (nm) | Volume (%) | Cumulation (%) |
|---------------|------------|----------------|---------------|------------|----------------|---------------|------------|----------------|
| 60.00         | 1.67       | 1.67           | 110.00        | 10.00      | 45.00          | 160.00        | 6.67       | 95.00          |
| 70.00         | 6.67       | 8.33           | 120.00        | 15.00      | 60.00          | 170.00        | 3.33       | 98.33          |
| 80.00         | 10.00      | 18.33          | 130.00        | 8.33       | 68.33          | 180.00        | 1.67       | 100.00         |
| 90.00         | 11.67      | 30.00          | 140.00        | 11.67      | 80.00          |               |            |                |
| 100.00        | 5.00       | 35.00          | 150.00        | 8.33       | 88.33          |               |            |                |

Fig. 6: FT-IR spectrum of chlorpyrifose

Fig. 7: FT-IR spectrum of (Mg/Fe-NO$_3$-LDH)

Fig. 8: FT-IR spectrum of nano hybrid compound

peaks of (υC = C) aromatic in the (1477-1589 cm$^{-1}$) region, absorption two peaks of υ (P-O-alkyl) and (P-O-aryl) at (1024 cm$^{-1}$) and (1240 cm$^{-1}$). (18) at assimilation N = C apper (1606 cm$^{-1}$) but peak at the (966 cm$^{-1}$) it is Cl-aryl Table 4 and Fig. 6.

The layers, (Mg/Fe-NO$_3$-LDH) show the absorption peaks to υO-H group stretches at (3454 cm$^{-1}$) (Cardoso et al., 2006), the NO$_3$ absorption peak at (1390 cm$^{-1}$) in the LDHs (Hussein et al., 2010), absorption peaks to υ(Mg-O) at (437 cm$^{-1}$) in layers and absorption peaks to υ (Fe-O) at (578 cm$^{-1}$) in layers (Cornejo et al., 2000) Table 5 and Fig. 7.

The nano compound (Mg/Fe-chloropyrifose-LDH) appears many a new peaks this means successful for
intercalation an ion (chloropyrophospose) between the layers, absorption peaks of uO-H stretch in the (3483 cm⁻¹). The peaks of aliphatic uC-H stretch in the (2362 cm⁻¹), peaks of (uC = C) aromatic appprase at (1516 and 1464 cm⁻¹), absorption peaks of uC = N stretch ataromatic 1690 cm⁻¹. The NO₃ absorption peak at 1381 cm⁻¹ in the LDHs absorption two peaks of u(P-O-alkyl) and (P-O-aryl) at (1030 and 1250 cm⁻¹) but peak at the (690 cm⁻¹) it is Cl-aryl, the peak δC-H aromatic apper at (669 cm⁻¹), absorption peaks to u(Mg-O) at (434 cm⁻¹) in layers and absorption peaks to u(Fe*O) at (586 cm⁻¹) in layers (Cornejo et al., 2000) Table 5 and Fig. 8.

The kinetic of ion exchange: To study the kinetic for controll release (Ho and Mckay, 2000) used Lagergran equation to zero order, first order and second order as the following:

\[ C_t = K_0 t \]  \hspace{1cm} \text{(1)}

\[-\log \left( 1 - \frac{C_t}{C^*} \right) = \frac{K_1 t}{2.303} \]  \hspace{1cm} \text{(2)}

\[ \frac{t}{C_t} = \frac{1}{K_2 C^*} + \frac{t}{C^*} \]  \hspace{1cm} \text{(3)}

Where:

- \( C_t \): Concentration at each time (mg L⁻¹)
- \( C^* \): Concentration at equilibrium (mg L⁻¹)
- \( K_0 \): Constant rate ion exchange zero model (mg L⁻¹ min⁻¹)
- \( K_1 \): Constant rate ion exchange first model (min⁻¹)
- \( K_2 \): Constant rate ion exchange second model (mg⁻¹ L min⁻¹)
- \( t \): Reaction time (min)

Through the note of Table 6 and the kinetic figures for (chloropyrophose) release from the nano compound prepare by (direct ion exchange and in direct ion exchange) (Fig. 9a-f) in sodium carbonate medial and (Fig. 10a-f) for (chloropyrophose) release from the nano compound prepare by (direct ion exchange and in direct ion exchange) in sodium phosphate medial and the value (r²) we note that the kinetic of ion exchange is model pseudo-second order.

When we note the value for the constant rate second (K₂) at (Fig. 9 and 10) result the rate kinetic release in different medial is as the following Na₂CO₃>Na₂PO₄.

The causes is that carbonate ion more electro file compare with sulfate ion and the figure of carbonate is triangular, so is easily intercalation with LDH than phosphate ion which is tetragonal.
Study percentage release for (chloropyrifos): The percentage release of (chloropyrifos) from the hybrid nano compound prepare in ethanol by direct and in direct ion exchange at (0.5M) from sodium carbonate and sodium phosphate medial used the linear equation as shown in Fig. 11a-d. Figure 11 and Table 7 from the results, we note the high value for percentage is the following Na$_2$CO$_3$>Na$_2$ PO$_4$. The percentage for in ion is high value in direct method than in direct method.
Table 7: Value for (percentage for release anion (VC)) by ion exchange method from the nano compound layer prepare in ethanol by direct and in direct ion exchange in (0.5M) sodium carbonate and sodium phosphate

| K   | Time (min) | Chloropyrifos (%) | Concentration Mol. L⁻¹ | Samples | R     |
|-----|------------|------------------|------------------------|---------|-------|
| 2.505 | 40       | 99               |                        | Na₂CO₃  |       |
| 0.761 | 110      | 85               |                        | Na₂PO₄  |       |
| 2.5   | 55        | 90               |                        | Na₂CO₃  |       |
| 0.411 | 140      | 77               |                        | Na₂PO₄  |       |

Fig. 11(a-d): Percentage of release (chloropyrifos) between the nano hybrid compound in aqueous media from sodium carbonate (0.5M) (a) prepare nano compound by direct ion exchange method (b) prepare nano compound by indirect ion exchange method; Percentage of release (chloropyrifos) between the nano hybrid compound in aqueous media from sodium phosphate (0.5M) (c) Prepare nano compound by direct ion exchange method and (d) Prepare nano compound by indirect ion exchange method.

**CONCLUSION**

The result appeared can be prepare nano compound hybrid by intercalation of insecticide (chloropyrifos) as anion between Mg/Fe double layer hydroxide by two method direct ion exchange and indirect ion exchange and study control release by direct ion exchange process to limited the problem of environment pollution and the healthly human. Where the kinetic control release is second order at (0.5M) from sodium carbonate and sodium phosphate. The high value kinetic release and percentage release at sodium carbonate. The study suggested ability use Mg/Fe double layer hydroxide as gents chemical and agriculture material and control release inter the body of the insectico limited the pollution, this method is inexpensive in economic terms where this method is to give the insecticide in small quantities, so as not to harm health on the other does not leaked to the soil or ground water and thus causing pollution.

**REFERENCES**

Aisawa, S., S. Takahashi, W. Ogasawara, Y. Umetsu and E. Narita, 2001. Direct intercalation of amino acids into layered double hydroxides by coprecipitation. J. Solid State Chem., 162: 52-62.

Arizaga, G.G.C., J.E.F.D.C. Gardolinski, W.H. Schreiner and F. Wypych, 2009. Intercalation of an oxalatoxoniobate complex into layered double hydroxide and layered zinc hydroxide nitrate. J. Colloid Interface Sci., 330: 352-358.

Bragg, W.L., 1993. The diffraction of short electromagnetic waves by a crystal. Proc. Cambridge Philos. Soc., 17: 43-57.

Cardoso, L.P., R. Celis, J. Cornejo and J.B. Valim, 2006. Layered double hydroxides as supports for the slow release of acid herbicides. J. Agric. Food Chem., 54: 5968-5975.

Carlino, S., 1997. The intercalation of carboxylic acids into layered double hydroxides: A critical evaluation and review of the different methods. Solid State Ionics, 98: 73-84.

Cavani, F., F. Trifiro and A. Vaccari, 1991. Hydrotalcite-type anionic clays: Preparation, properties and applications. Catal. Today, 11: 173-301.

Cornejo, J., R. Celis, I. Pavlovic, M.A. Ulibarri and M.C. Hermosin, 2000. Structural changes in phenol-intercalated hydrotalcite caused by heating. Clay Miner., 35: 771-779.

Evansa, D.G. and X. Duan, 2006. Preparation of layered double hydroxides and their applications as additives in polymers, as precursors to magnetic materials and in biology and medicine. Chem. Commun., 1: 485-496.

Franklin, K.R., E. Lee and C.C. Nunn, 1995. Preparation and characterisation of layered double hydroxides containing monovalent and divalent ions derived from 5-benzoyl- 4-hydroxy-2- methoxybenzenesul fonic acid. J. Mater. Chem., 5: 565-569.
Ho, Y.S. and G. McKay, 2000. The kinetics of sorption of divalent metal ions onto sphagnum moss peat. Water Res., 34: 735-742.

Hussein, M.Z., F.A. Bahar and A.H. Yahya, 2010. Synthesis and characterization of hippurate-layered double hydroxide nanohybrid and investigation of its release property. J. Iran. Chem. Soc., 7: S42-S51.

Khan, A.I. and D. O’Hare, 2002. Intercalation chemistry of layered double hydroxides: Recent developments and applications. J. Mater. Chem., 12: 3191-3198.

Ookubo, A., K. Ooi and H. Hayashi, 1993. Preparation and phosphate ion-exchange properties of a hydroxalbite-like compound. Langmuir, 9: 1418-1422.

Raki, L., D.G. Rancourt and C. Detellier, 1995. Preparation, characterization and Moessbauer spectroscopy of organic anion intercalated pyroaurite-like layered double hydroxides. Chem. Mater., 7: 221-224.

Rives, V., 2001. Layered Double Hydroxides: Present and Future. Nova Science Publishers, Hauppauge, New York, USA., ISBN-13: 9781590330609, Pages: 439.

Shaheed, S.H., A.A.K. Al-Ghanimi and A.M. Bashi, 2014. Preparation of nanohybrid compound from the food preservative octyl gallate and studying some of its biological activities. Karbala J. Pharm. Sci., 5: 277-289.

Vaccari, A., 1998. Preparation and catalytic properties of cationic and anionic clays. Catal. Today, 41: 53-71.

Whilton, N.T., P.J. Vickers and S. Mann, 1997. Bioinorganic clays: Synthesis and characterization of amino-and polyamino acid intercalated layered double hydroxides. J. Mater. Chem., 7: 1623-1629.