Synthesis of Al-Based Metal-Organic Framework in Water With Caffeic Acid Ligand and NaOH as Linker Sources With Highly Efficient Anticancer Treatment

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In this study, novel nanostructures of aluminum base metal-organic framework (Al-MOF) samples were synthesized using a sustainable, non-toxic, and cost-effective green synthesis route. Satureja hortensis extract was used as an effective source of linker for the development of the Al-MOF structures. The Fourier-transformed infrared (FTIR) spectrum confirmed the presence of characterization bonds related to the Al-MOF nanostructures synthesized by the green synthesis route. The scanning electron microscopy (SEM) and transmission electron microscopy (TEM) analyses revealed that the sample synthesized by Na2-CA was composed of multilayers, although it was agglomerated, but it had dispersed and occurred in spherical particles, indicating active organic matter. N2 adsorption/desorption isotherms demonstrated the significant porosity of the Al-MOF samples that facilitate the high potential of these nanostructures in medical applications. The anticancer treatment of Al-MOF samples was performed with different concentrations using the MTT standard method with untreated cancer cells for 24 and 48 h periods. The results exhibited the significant anticancer properties of Al-MOF samples developed in this study when compared with other MOF samples. Thus, the development of a novel Al-MOF and its application as a natural linker can influence the anticancer treatment of the samples. According to the results, the products developed in this study can be used in more applications such as biosensors, catalysts, and novel adsorbents.

Keywords: novel Al-MOF, green synthesis, S. hortensis extract, caffeic acid ligand, anticancer properties

INTRODUCTION

One of the latest and important classes of porous materials are metal-organic framework (MOF) nanostructures, with industrial importance in diverse fields. These classes of organic and inorganic frameworks composed of metallic or metal oxide structural units are bound together by organic molecules and form interconnected porous structures (Langseth et al., 2019).

It is important to exploit and commercialize MOF production methods that are sustainable, mass-produced, and cost-effective (Julien et al., 2017; Reinsch and Stock, 2017; Zhu et al., 2022). Therefore,
green synthesis is possible to produce MOFs using non-toxic and efficient materials (Thornton et al., 2016; Taddei, 2017). One of the most important advantages of green synthesis is the non-use of toxic solvents such as dimethylformamide (DMF) and synthesis at ambient temperature and pressure, as well as lower activation temperatures to eliminate the byproducts and guest molecules in MOFs (Reinsch and Stock, 2017; Zhang et al., 2020a; Yang, 2021). Therefore, we need to use water for the green synthesis of MOFs because water is more accessible and vibrant than organic solvents. However, the important point here is that most of the water-soluble organic bonds are weak and cause the instability of MOFs (Gelfand and Shimizu, 2016; Shih et al., 2017). This method may hence not be suitable for mass production, but the time of MOFs synthesis involving metal ions bonding, and the natural organic framework is shorter and less significant than that in other methods. To date, the aqueous synthesis of MOFs using chemistry offers major benefits. In these methods, the water-solubility issues have been mitigated mainly by using sodium salts of the organic linkers and by increasing the length of the tubing systems to increase the overall reaction time (Bayliss et al., 2014; Sánchez-Sánchez et al., 2015; Avcı-Camur et al., 2018).

Therefore, we use Satureja hortensis extract of organic compounds, such as chlorogenic acid, caffeic acid (CA), gallic acid, ferulic acid, and γ-hydroxymethylcinnamic acid as the organic framework (Movahhedkhah et al., 2019) and used sodium salts for a faster and better reaction. Most of the ingredients in the extract of savory are CA, which converts to CA despite the presence of a large amount of organic matter in the extract (Milos, 2001). As mentioned, one of the polyphenols produced through the secondary metabolism of vegetables is CA (Espíndola et al., 2019). CA participates in the defense mechanism of plants against predators, pests, and infections, as it has an inhibitory effect on the growth of insects, fungi, and bacteria (Tosovic, 2017) as well as promotes the protection of plant leaves against ultraviolet radiation B (UV-B) (Gould et al., 2000; Tosovic, 2017). In vitro and in vivo experiments have been performed and demonstrated innumerable physiological effects of CA and its derivatives, such as antibacterial (Verma and Hansch, 2004; Genaro-Mattos et al., 2015), anti-hepatocarcinoma (HCC) (Zhang et al., 2017), antioxidant (Lin and Yan, 2012; Huang et al., 2013; Genaro-Mattos et al., 2015), anti-inflammatory (Lin and Yan, 2012; Huang et al., 2013), anticancer (Lin and Yan, 2012; Huang et al., 2013), and anti-hepatocellular carcinoma activity (Gu et al., 2016). Among these properties, the anti-HCC activity is highlighted because HCC is one of the main causes of cancer mortality in the world (McGlynn et al., 2015). Therefore, further studies on the chemical and pharmacological aspects of CA are necessary to contribute to the future development of a new drug and, consequently, to the expansion of therapeutic possibilities (Zhang et al., 2017).

Aluminum (Al) is one of the most abundant metals with non-toxic salts that are easy to use and also economical. In addition, Al nanoparticles (Al-nano) may be a good choice for anticancer immunotherapy, which should possess good safety and efficacy. Al-nano such as aluminum hydroxide and phosphate or hydroxy phosphate aluminum have excellent safety properties for systemic vaccination. In fact, it has been used as an adjunct for over half a century and is now one of the most widely used adjuncts in animal and human vaccines (Lindblad, 2004; Mesa and Fernández, 2004). Muehmann and Sun et al., for example, demonstrated that aluminum oxide nanoparticles as drug carriers are promising for photodynamic cancer therapy as well as for the activation of the immune system in cancer diseases (Lin and Yan, 2012). Wang et al. revealed the use of aluminum anode nanotubes as a carrier of anticancer drugs is shown on the MDA cancer cell line. Protein from the family of tumor necrosis factors that induce apoptosis was used as a model drug. Anodized alumina nanotubes that have been structurally engineered can be used as nano-carriers for the delivery of anticancer therapeutics (Hou et al., 2021). So, the oxidation of aluminum and copper has anticancer and antimicrobial properties due to its highly porous structure. Therefore, we have chosen one type of Al-MOFs as a carrier for the anti-proliferative herbal extract (Zeraati et al., 2021a). The Al-MOFs seem to be more effective against the proliferation of breast cancer cells when compared with herbal extraction. Therefore, Al-MOFs have received much attention (Senkovska et al., 2009; Moreno et al., 2018). A challenge in the investigation of Al-MOFs is the rich solution chemistry of Al3+, which gives rise to a variety of inorganic building units ranging from isolated Al3+ ions that are exclusively connected by the carboxylate groups to trimeric building units, rings of edge- and corner-sharing AlO6 polyhedra, chains of trans- or cis-corner sharing AlO6 polyhedra, chains of edge-sharing AlO6 polyhedra, or flexible inorganic building units of corner-sharing Al13-oxo clusters. Therefore, the study of Al-MOF can provide a new step toward cancer treatments.

As a result, the use of green sources as a link precursor can be regarded as a revolution in MOF preparation. Ionic liquids, natural oils, and plant extracts are examples of such sources, all of which have a high level of acceptability in terms of green synthesis.
technology (Liu et al., 2013; Zeraati et al., 2021a). Green MOF synthesis also reduces the need for organic solvents, operates at low temperatures and pressures, and requires less time to react. These benefits have increased the efficiency of the green technique (Zeraati et al., 2021b).

In this study, the green synthesis of Al-MOF was developed by using S. hortensis extract as an organic framework and NaOH as a salt linker. The products were characterized by Fourier-transformed infrared (FTIR) spectroscopy, field-emission scanning electron microscopy (SEM), transmission electron microscopy (TEM), and N₂ adsorption/desorption isotherm. The final Al-MOF products were investigated for their anticancer properties in detail. Figure 1 illustrates the flowchart of the study method.

MATERIALS AND METHODS

The reagents used for the chemical synthesis of Al-MOF are aluminum nitrate [Al(NO₃)₃·9H₂O], deionized (DI) water (H₂O), and S. hortensis extract as the green organic linker. The S. hortensis extract was prepared by taking 10 g of dry Satureja cut (harvested from Ghobk in Kerman province of Iran) and immersing it in 100 ml of distilled boiled water for 5 min, followed by cooling and filtering through centrifugal spinning at 4,000 rpm thrice. The Markham method was used to extract flavonoids (Couderchet et al., 1997). Primarily, 250 ml of 2% AlCl₃·6H₂O solution was mixed with 40 ml S. hortensis aqueous extract. After storage for 15 min at room temperature, the absorbance response was measured at 430 nm. This data was expressed as milligram per gram of dry matter. The synthesis process was performed in an ambient atmosphere. First, 3.3 g of Al precursor [Al(NO₃)₃·9H₂O] was dissolved in 12.5 ml of DI water as the solvent at 80°C. Then (S1) 12.5 ml of the Satureja extract was added to the Al solution, and (S2) 12.5 ml of S. hortensis extract and 1 ml of 50% NaOH solution were added to the Al solution with aggressive stirring at 300 rpm. In this step, the transparent colloidal of the initial solution changed in color to light yellow. The synthesis process was prolonged to 60 min until the formation of white precipitate from the yellow solution at the same temperature. The precipitate was then collected and dried. Finally, considering the results of the thermal analyses, the prepared sample was maintained at 120°C in a vacuum oven for 5 min to active its structure. Morphological and structural characteristics of the prepared samples were performed using the Nanosem 450. The field-emission SEM (FE-SEM) images were acquired with the Inspect F SEM (FEI) operating at 3 kV and equipped with an energy dispersive X-ray (EDX) spectroscopy and TEM. The FTIR spectra of the samples were measured on the Varian LS Chemical Imaging Microscope at 4,000–400 cm⁻¹ and the Brunauer–Emmett–Teller (BET) model from the nitrogen adsorption/desorption isotherms of the samples were obtained using 196°C (77 K) after separation at 200°C and 10–5 mm Hg for at least 4 h. The surface-specific regions (SBETs) were calculated from the adsorption isotherm data using the BET method. The mesoporous size distribution and total mesoporous volume were determined using the modified Barrett–Joyner–Halenda (BJH) method of isotherm adsorption data. The human breast cancer (MDA-MB-468) cell line was provided from the Iranian Biological Resource Center (IBRC; Tehran, Iran). Dulbecco’s modified Eagle’s medium (DMEM), fetal bovine serum (FBS), phosphate-buffered saline (PBS), trypsin/EDTA solution, 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT), and dimethyl sulfoxide (DMSO) were purchased from Gibco BRL and Sigma, respectively. They were cultured in DMEM (Gibco, United Kingdom) supplemented with 10% FBS (Gibco) and 1% penicillin–streptomycin (Gibco) and incubated in a 5% CO₂ atmosphere at 37°C. For treatment, 5 × 10³ cells/well were seeded in 96-well flat-bottomed plates overnight, then the cells were exposed to various concentrations of the herbal extract (0–100 μM) and Ag-MOF (0–100 μM) for 24 and 48 h. Subsequently, the medium was removed, and 200 μl of MTT solution (5 mg/ml in PBS) was added to each well and incubated for 4 h at 37°C. After discarding the solution, 100 μl of DMSO was added and the plates were shaken for 15 min. The absorbance of each sample was read at 570 nm using an ELISA microplate reader. The outcomes were affirmed as a percent of cell viability with respect to the untreated control cells.

RESULTS AND DISCUSSIONS

Characterization of Al-MOF

The green synthesis of two samples of Al-MOFs at room temperature was obtained by mixing CA and 1 ml of 50% NaOH solution as Na₂-CA, and their structures were activated at 120°C for 5 min. The physical and chemical properties of the samples S1 and S2 were investigated.

Figure 2 shows the FTIR spectrum of Al-MOFs synthesized samples. S1 shows the spectrum of the corresponding sample composed of CA, and S2 shows the spectrum of the sample composed of Na₂-CA. As expected, both the samples were quite similar. The peaks near 3,700–3,800 cm⁻¹ are related to O-H stretching vibration of water (Mihaylov et al., 2015; Tong et al., 2019). The frequency at 3,420 cm⁻¹ is ascribed to the stretching
vibration of aromatic C-H (Seoane et al., 2013; Mihaylov et al., 2015; Hu et al., 2018). The absorption bands around 2,423 and 1,640 cm$^{-1}$ are assigned to the C=C group and methylene bonds, respectively (Jing et al., 2021; Xue et al., 2021). The peaks near 1,378 cm$^{-1}$ as well as the 645 cm$^{-1}$ and 829 cm$^{-1}$ are attributed to the aliphatic CH bond (Biswas et al., 2019). The spectrum corresponding to S2 showed less hydrogen bonding due to the addition of NaOH to CA, which caused H$_2$O to be eliminated by the heating process (Sánchez-Sánchez et al., 2015). The FTIR characterization related to the formation of Al-MOF nanostructures agrees to that reported previously (Reinsch et al., 2012; Li et al., 2019). It is therefore important evidence for the successful synthesis of nanostructural compounds.

Investigating and predicting the stability of Al-MOFs produced by the green synthesis in water is particularly important because moisture is the main challenging factor of unstable Al-MOFs. Moreover, the mechanism of this system in which the bonds containing benzene, carboxylate, and water-soluble aromatic groups are largely unknown in the field of Al-MOFs (Sánchez-Sánchez et al., 2015). Figure 3 schematically affording a sodium salt from the corresponding available carboxylic acid is relatively easy and follows the reaction for the generation of the depicted organic linker anion CA3-depicted.

Because of the availability of sodium salts and natural organic frameworks, this method is of particular interest and application. The structure and morphology of the specimens were examined as illustrated in Figure 4. Figures 4(A,B) correspond to the samples S1 and S2, respectively. According to Figure 4(A), the sample synthesized with CA showed an agglomerated structure and multilayers that also gave rise to polycrystalline agglomerates. The Al-MOF sample synthesized by Na$_2$-CA occurred in multilayers, although it was agglomerated, but the sample had dispersed in spherical particles, indicating active organic matter and the absence of any guest element.

Figures 5A,B represents the TEM image of Al-MOF samples with a spherical shape distribution. The high accuracy of the image in TEM depicts the nature of the porosity of the structure. As is evident, it seems that the porosity can be affected by the surface area of the products. Considering the surface area as a criterion, the fraction of pores (dark area) to the total surface in the TEM image was estimated by using an image analyzer software. As shown in Figures 5C,D, the background is blue and the pore phase is indicated in red and green, with phase percent of samples S1 and S2 being equal to 70 and 83%, respectively. This result, according to the mechanism and results of sample S2, showed better purity and efficiency (Yan et al., 2015; Moreno et al., 2018). The homogenous morphology of the sample S2 was significantly improved when compared to that of the previous Al-MOF sample (Lin and Yan, 2012; Aleksandrzak et al., 2019). According to the results, there is no evidence of agglomeration procedure in the structures. This stability in the structures can be related to the development of a novel natural linker as well as the application of the green synthesis route used in this study.

Figure 6 and Figure 7 depict the N$_2$ desorption/desorption isotherms of S1 and S2. The difference between the two isotherms indicates the distinct nature of the two samples in terms of textural properties. There is also a different hysteresis cycle depicted in Figure 6, ending in sample S1 at p/p$_0$ = 0.45 and
sample S2 at \( p/p_0 = 0.3 \). The larger hysteresis cycle of sample S1 is related to the presence of guest molecules in the pores (Zhang et al., 2020b; Li et al., 2020). As Figure 7 shows the size distribution of its pores, we obtained the narrow particle size of samples with an average of 20 nm. While the sample S2 pore size distribution was <20 nm, the average value was 12 nm. To estimate the surface area of the prepared S1 and S2 samples used BET analysis (Figure 6). As shown, the adsorption isotherms of both samples are similar to the isotherm III type (Wang et al., 2019), while the average pore size of S1 and S2 samples was 1,245.2 m\(^2\)/g and 2,452.8 m\(^2\)/g, respectively. The high surface area for samples can be attributed to the select novel synthesis route as well as the optimization process. Accordingly, the prepared Al-MOF was accepted as mesoporous martial as reported by Moreno et al. (Zhang et al., 2020c). The decrease in surface area compared to previous reports is probably due to the presence of organic extract molecules that are not completely removed by activation. In addition, the molecules of solvent trapped in the MOF pores reduce the specific surface area. On the other hand, the Al-MOF powders have been agglomerated and the gas penetration does not occur completely (Rahmani and Rahmani, 2018; Nowroozi-Nejad et al., 2019).
Due to the volume of N₂ in Figure 7(A), they are present layer to layer between the layers of Al-MOFs powder as S1 samples. However, the sample S2 had no layered structure (Figure 6B), and the adsorption was not multilayered due to the low volume of N₂ gas, and the adsorption was of the surface type, indicating more numbers of active holes.

Anticancer Properties
To evaluate the biocompatibility and cytotoxicity of samples S1 and S2, we used different concentrations and performed the MTT standard method with untreated cancer cells for 24 and 48 h. As shown in Figure 8, the survival rate of cancer cells treated by the dose-dependent method decreased after 24 and 48 h when compared to the untreated cancer cells. The inhibitory effect of sample S1 on cell proliferation was significantly higher than on sample S2. Parallel treatment of normal cells with these components demonstrated a higher inhibitory effect on the survival of normal human cells. The primary property of proliferating cancer cells was uncontrolled. Therefore, tumor growth control is considered a valid treatment approach in cancer treatment. Numerous studies have shown that MOFs confer beneficial aspects such as host-guest interaction, low toxicity, hydrophobic/hydrophobic equilibrium, body distribution, and biodegradation tissue aggregation and irritability. They can, therefore, be used in biological applications such as cancer treatment (Gao et al., 2018; GAO, 2019; Uthappa et al., 2019; Jiang et al., 2021). Several studies have shown that the metalorganic frameworks have beneficial aspects, including low cytotoxicity, host-guest interactions, hydrophobic/hydrophilic balance, biodegradability, body distribution, tissue accumulation, and excitability, allowing them to be used in biological applications for cancer treatment (Tuncer et al., 2013; Ebrahimi et al., 2017). Therefore, we have chosen one type of Al-MOFs as a carrier for the anti-proliferative herbal extract. The Al-MOFs seem to be more effective against the proliferation of breast cancer cells when compared with herbal extraction (Nowroozi-Nejad et al., 2019).

Therefore, we selected a type of S2 as a carrier of the anti-proliferative plant extract. It appears that Al-MOF is more effective than breast extraction against the proliferation of breast cancer cells. In addition, the small concentration of these biological structures (7 mg/ml) significantly reduced the growth of MDA-MB-468 cells after 48 h. Therefore, Al-MOF as an anticancer anti-cancer drug against human breast cancer is a
potent component that is being further investigated for its cytotoxicity properties. On the other hand, CA, which is an active ingredient of S. hortensis extract, offers promising therapeutic potential against a wide range of biological conditions, especially cancer (Butt et al., 2019; Skubij and Dzida, 2019). Fazary et al. reported that the Na₂-CA monoprotonated in basic media is diprotonated in acidic media and that they possess better anticancer properties (Fazary et al., 2019). Therefore, it can be concluded that sample S2 in a shorter time and with smaller pores possess better anticancer properties than the prototype of sample S1.

The anticancer properties of samples were significant compared to previous samples. It can be related to selecting the novel nanostructures and also optimization the procedure (Ji et al., 2020; Sun et al., 2021; Zhang et al., 2021).

**CONCLUSION**

The novel Al-MOF was prepared by using S. hortensis extract as an organic framework with NaOH as a salt linker. The final products were characterized by relevant analyses. The main group was confirmed by FTIR spectroscopy. TEM exhibited the spherical morphology of the structure with porosity nature arranging the compound. The N2 adsorption/desorption isotherm of Al-MOF nanostructure was similar to the IV type-classical isotherms, which confirmed the significant porosity of Al-MOF nanostructures. The final Al-MOF sample was examined for anticancer treatment, and the results confirmed the remarkable activity of this novel compound. The present study can open a new window to the introduction of new anticancer materials. The nature of Al-MOF, the natural S. hortensis extract linker, and the purification procedure were explored in this work. The physico-chemical properties of the samples obtained in this study can be attributed to select novel nanostructure as well as an optimization process. The anticancer features obtained in this study can be opened a new window for developing new material.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are included in the article/Supplementary Materials, further inquiries can be directed to the corresponding authors.

**AUTHOR CONTRIBUTIONS**

Conceptualization, MZ, AR, DM, and GS; methodology, MZ, AR, DM, and GS; validation, MZ; formal analysis, MZ, AR, DM, and GS; investigation, MZ, AR, DM, and GS; data curation, MZ, AR, DM, and GS; writing—original draft preparation, MZ, AR, DM, and GS; writing—review and editing, MZ, AR, DM, and GS; supervision, MZ, AR, DM, and GS; project administration, MZ, AR, DM, and GS; funding acquisition, MZ, AR, DM, and GS. All authors have read and agreed to the published version of the manuscript.

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