Some Nano Bio-Mofs Evaluated For Storage Purpose of Drugs

Tabinda Sattar and Muhammad Athar*

Assistant professor, Institute of Chemical Sciences, Pakistan

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*Corresponding author: Muhammad Athar, assistant professor, institute of chemical sciences, Bahaudin zakariya university Multan, Pakistan, Tel: 03234548374; Email:Atharq@hotmail.com

Abstract

Nano bio-MOF compounds 3-5 (copper serinate, copper prolinate and copper threoninate) have been evaluated for successful in vitro drugs adsorption of three drugs, named terazosine, telmisartan and glimpiride. Pore sizes and BET surface areas were recorded on N2 adsorption isotherms. In order to elaborate the phenomenon of in vitro drugs adsorption in all these compounds we obtained TGA and PXRD spectra of them. The high performance liquid chromatography (HPLC) was used to monitor the amounts of adsorbed drugs and its slow release from all these compounds.

Keywords: Nano bio-MOFs; Invitro; Drugs adsorption; TGA; PXRD; HPLC

Abbreviations: MOFs: Metal Organic Frameworks; SEM: Scanning Electron Microscopic; HPLC: High Performance Liquid Chromatography; TGA: Thermo Gravimetric Analysis

Introduction

Metal Organic Frameworks (MOFs) are the materials which are significant due to different applications especially in the field of drugs storage [1,2]. Still there is need of designing more compatible drugs carriers for successful delivery applications both for in vitro and in vivo purposes. In these respect bio-metal organic frameworks (Bio-MOFs) is a useful extension of MOFs [3]. Actually bio molecule linkers and bio compatible metal cations can be utilized for the construction of bio-MOFs. These materials have great potential utility as targeted drugs carrier materials inside human body containing no harmful effects at all for health [4]. The MOF materials which were previously used for drugs carrying purposes faced some compatibility problems inside human body due to their large sizes [5]. So with the passage of time use of nano sized MOF materials became more feasible for drugs storage and release. Kitağawa et al. have reported the use of some large size MOF materials used successfully for drugs storage purposes [6].

Latter on Rieter et al. have synthesized some nano scale MOFs and used them for drugs carrying, so having much better results as compared to previously reported drugs carrier MOFs. Wöttki et al. [7] have reported some MOFs materials which were designed and tuned accordingly for usage as more potential nano carrier materials as compared to previously used drugs carrier materials [8,9]. Roder et al. [10] have also reported the successful use of nano sized MOF materials for drugs delivery. The smaller size and high surface areas these nano sized MOF materials make them the targeted drugs delivery vehicles [10]. The bio MOFs can be scaled down to the nano-regime as well in order to form nanoscale bio metal-organic frameworks (nano bio-MOFs). Nano bio MOFs are the nano sized MOF materials containing bio-molecules and biocompatible metal cations. This emerging class of nano-bio-MOFs can be considered as promising candidates for the drug adsorption and its controlled release due to their large surface areas, high porosity, and presence of bio compatible linker molecules [11]. Several research reports reveal the use of nano sized MOFs in the field of drugs delivery [12-14].

Three hydrothermally synthesized nano bio-MOFs compounds 3-5 (copper serinate, copper prolinate and copper threoninate) have been reported by Sattar et al.[15]. These compounds have been hydrothermally synthesized and characterized by Scanning Electron Microscopic (SEM) studies. Previously these materials were used as drug carrying for rosuvastatin drug. These materials were found feasible for drug storage with its slow release from the materials after intervals of time as monitored through High Performance Liquid Chromatography (HPLC). The present work is designed to elaborate the pore sizes and BET surface area calculations of these materials and also to evaluate their drug storage potential for three more drugs like terazosine hydrochloride, telmisartan and glimpiride. In vitro drugs adsorption studies of these drugs and their release may lead to the in vivo usage of these drugs delivery inside human body as these drugs have applications for hypertension and
control blood sugar levels. Thermogravimetric analysis (TGA) and powder X-rays Diffraction Analysis (Powder XRD) patterns of all these new compounds in pure form and after drugs adsorption have been recorded. The amounts of adsorbed drugs and its slow release after intervals have been monitored through the High Performance Liquid Chromatography (HPLC).

Figure 1: a) N2 adsorption isotherm of compound 3, b) A plot of BET surface area of compound 3, c) N2 adsorption isotherm of compound 4, d) A plot of BET surface area of compound 4, e) N2 adsorption isotherm of compound 5, f) A plot of BET surface area of compound 5.

BET specific surface areas of compounds 3, 4 and 5 have been determined as 2800 m$^2$g$^{-1}$, 2000 m$^2$g$^{-1}$ and 2900 m$^2$g$^{-1}$ respectively with a pore size of 10Å by using BET-method based calculations on N$_2$ adsorption isotherm data. The Figure 1 represents the N$_2$ adsorption isotherm of compound 3, 4 and 5. Four types of drugs (terazosine hydrochloride, telmisartan, glimpiride and rosuvastatin) have been absorbed into compounds 3, 4 and 5. It was also elaborated that some pores were blocked after the drugs adsorptions thus decreasing the specific surface areas of this material as shown in Figure 1. TGA plots of the as synthesized and drugs adsorbed compounds 3-5 were carried out on a SDT Q600 by heating the compounds from 0oC to 600oC at a heating rate of 10oC per minute. TGA plot (Figure 2a) before drugs adsorption shows first weight loss of 10% at 120oC which is due to the removal of water molecule. Up to 340oC the framework shows stability with no weight loss. Second weight loss is observed at 340oC which continues up to 400oC with a 40% weight loss due to the framework decomposition along with ligands. Then framework continues to decompose gradually until the whole framework decomposes till 590oC with the formation of metal oxides (Figure 2).

Figure 2: TGA plot of compound 3 a) before drugs loading, b) after terazosine hydrochloride loading, c) after telmisartan loading, d) after glimpiride loading.
After the terazosine adsorption the TGA plot of compound 3 (Figure 2b) shows a weight loss of 9% at 40°C, which may be attributed towards the loss of drug molecules. After that the framework remains intact for some time and then at 140°C the framework shows stability up to 180°C. Second weight loss of 4% at 180°C is observed due to the loss of water molecule. After that at 280°C the framework starts to decompose up to 380°C with a weight loss of 58%. No further weight loss observed for this material. TGA plot of telmisartan adsorbed compound 3 (Figure 2c) shows a weight loss of 4% at 40°C which is due to the loss of telmisartan contents. Another weight loss of 4% at 150°C shows the loss of water molecule. Then the framework starts to decompose at 250°C which continues till 550°C with a maximum weight loss of 59%. Similarly TGA plot of glimpiride adsorbed compound 3 (Figure 3) shows first weight loss of 4% at 50°C which is due to the removal of glimpiride contents. At 170°C the framework shows a weight loss of 4% up to 190°C which is due to removal of water molecule. At 280°C the decomposition of the framework and ligands starts which continues gradually up till 560°C with maximum weight loss of 58%.

Figure 3: TGA plot of compound 4 a) Pure, b) after terazosine loading, c) after telmisartan loading, d) after glimpiride loading.

Figure 3a shows the TGA plot of compound 4, before drugs adsorption shows first weight loss of 4% at 180°C which is due to the removal of water molecules. Second weight loss is observed at 280°C which continues up to 380°C due to the start of framework decomposition along with ligands. Then framework continues to decompose gradually until the whole framework decomposes till 580°C with a maximum weight loss of 49% along with the formation of metal oxides. The TGA plot of compound 4 after the terazosine adsorption shows (Figure 3b) a weight loss of 5% at 60°C, which is due to the loss of drug molecules. Then the framework shows stability up to 180°C. Second weight loss of 9% at 180°C is due to the loss of coordinated water molecule. The framework remains intact till 280°C, after that it starts to decompose and this decomposition continues up to 380°C with a weight loss of 58%. Similarly TGA plot of telmisartan adsorbed compound 4 (Figure 3c) shows a weight loss of 4% at 40°C which is due to the loss of telmisartan contents. Another weight loss of 8% at 150°C shows the loss of water molecules. At 250°C the gradual decomposition of the framework starts which continues till 300°C with a maximum weight loss of 59%.

Figure 4: TGA plot of compound 5 a) Before drugs adsorption, b) after terazosine loading, c) after telmisartan loading, d) after glimpiride loading.
A TGA plot of glimpiride adsorbed compound 4 (Figure 3d) shows first weight loss of 4% at 50°C which is due to the removal of glimpiride contents. At 170°C framework shows a weight loss of 8% up to 190°C which is due to removal of water molecule. At 280°C the decomposition of the framework and ligands starts which continues till 330°C with weight loss of 58% (Figure 3). A plot of rosuvastatin adsorbed compound 4 (Figure 3e) shows first weight loss of 3% at 45°C which is due to the removal of drug contents. At 171°C the framework shows a weight loss of 8% which is due to the removal of water molecule. At 285°C the decomposition of the framework and ligands starts which continues till 560°C with ultimate weight loss of 58% (Figure 3).

Figure 5: PXRD pattern of compound 3, a) in pure form, b) after terazosine loading, c) after telmisartan loading, d) after glimpiride loading.

PXRD pattern of compound 4, e) in pure form, f) after terazosine loading, g) after telmisartan loading, h) after glimpiride loading.

PXRD pattern of compound 5, i) in pure form, j) after terazosine loading, k) after telmisartan loading, l) after glimpiride loading.

TGA plot of terazosine adsorbed compound 5 (Figure 4b) shows a weight loss of 6% at 40°C, due to the loss of drug molecules. Second loss of mass at 180°C is observed and this 4% weight loss is due to the loss of water molecules. At 280°C the framework starts to decompose and end up to 580°C with a final weight loss of 58%. TGA plot of telmisartan adsorbed compound 5 (Figure 4c) shows a weight loss of 4% at 40°C which is due to the loss of telmisartan contents. Another weight loss of 4% at 150°C shows the loss of water molecule. Then the framework starts to decompose at 250°C which continues till 550°C with a maximum weight loss of 59%. TGA plot of glimpiride adsorbed compound 5 (Figure 4d) shows first weight loss of 4% at 25°C which is due to the removal of glimpiride contents. Then up to 120°C the framework remains intact and at this temperature framework shows a weight loss of 4% up to 190°C which is due to removal of water molecule. At 280°C the decomposition of the framework and ligands starts which continues till 560°C with ultimate weight loss of 58% (Figure 4).

PXRD patterns of compounds 3-5 were recorded as described in Figure 5. Powder XRD pattern of all these newly synthesized materials after drugs adsorption has revealed the permanent crystalline integrity of these compounds as these have retained its crystallinity even after soaking in water for several days (Figure 5). For the sake of estimation of the drugs contents in the synthesized materials, high Performance Liquid Chromatography was performed on a Waters 2695 separation module. All the details of HPLC parameters used for HPLC studies have been given in Table 1 (supporting information). 0.095g/g, 0.083g/g and 0.041g/g material of terazosine hydrochloride, telmisartan, and glimpiride has been estimated in compound 3 with a maximum time of release of 5, 3 and 3 days respectively. 0.200g/g, 0.160g/g and 0.138g/g material of terazosine, telmisartan and glimpiride were detected in compound 4. These adsorbed amounts of drugs were slowly released from the compound after different time intervals. The maximum time of release of these drugs was estimated at 7, 5 and 3 days respectively.

The adsorbed amounts of drugs terazosine, telmisartan and glimpiride have been estimated as 0.316g/g, 0.086g/g, 0.138g/g, 0.094g/g material of compound 5. Slow release of these drugs from compound 8 was observed through HPLC. The maximum time of release of these drugs from compound was estimated at 9, 3 and 3 days respectively. Table 3 gives summative information.
about the drugs adsorption capacities of compounds 3-5. (Figures 6-17) (supporting information) indicate the HPLC peaks for the estimation of adsorbed drugs in the channels of compound 3-5 and its slow release after different time periods (Table 1).

**Figure 6:** HPLC Peak of standard (Terazosine), b) HPLC peak of samples #1 (aliquot of water containing released drug collected after one day of soaking the drug adsorbed material in distilled water, c) sample #2 collected after three days, d) Sample #3: collected after five days, e) Collected after seven days.

**Figure 7:** HPLC Peak of standard (Telmisartan), b) HPLC peak of samples #1 (aliquot of water containing released drug collected after one day of soaking the drug adsorbed material in distilled water, c) sample #2 collected after three days, d) Sample #3: collected after five days, e) Collected after five days.
Figure 8: HPLC Peak of standard (glimpiride), b) HPLC peak of samples #1 (aliquot of water containing released drug collected after one day of soaking the drug adsorbed material in distilled water, c) sample #2 collected after three days, d) Sample #3: collected after five days, e) Collected after five days.

Figure 9: HPLC Peak of a) HPLC Peak of standard (rosuvastatin), b) HPLC peak of samples #1 (aliquot of water containing released drug collected after one day of soaking the drug adsorbed material in distilled water, c) sample #2 collected after three days, d) Sample #3: collected after five days, e) Collected after five days.
Figure 10: HPLC Peak of a) HPLC Peak of standard (Terazosine), b) HPLC peak of samples #1 (aliquot of water containing released drug collected after one day of soaking the drug adsorbed material in distilled water, c) sample #2 collected after three days, d) Sample #3: collected after five days, e) Collected after seven days, f) Samples #5: collected after nine days.

Figure 11: HPLC Peak of a) HPLC Peak of standard (Telmisartan), b) HPLC peak of samples #1 (aliquot of water containing released drug collected after one day of soaking the drug adsorbed material in distilled water, c) sample #2 collected after three days, d) Sample #3: collected after five days, e) Collected after seven days, f) Samples #4: collected after seven days.
Figure 12: HPLC Peak of a) HPLC Peak of standard (glimpiride), b) HPLC peak of samples #1 (aliquot of water containing released drug collected after one day of soaking the drug adsorbed material in distilled water), c) sample #2 collected after three days, d) Sample #3: collected after five days.

Figure 13: HPLC Peak of a) HPLC Peak of standard (rosuvastatin), b) HPLC peak of samples #1 (aliquot of water containing released drug collected after one day of soaking the drug adsorbed material in distilled water), c) sample #2 collected after three days, d) Sample #3: collected after five days, Sample #4: Collected after seven days.
Figure 14: HPLC Peak of a) HPLC Peak of standard (Terazosine), b) HPLC peak of samples #1 ( aliquot of water containing released drug collected after one day of soaking the drug adsorbed material in distilled water), c) sample #2 collected after three days, d) Sample #3: collected after five days, Sample #4: Collected after seven days, f) Sample #5: collected after nine days, g) Sample #6: collected after eleven days.

Figure 15: HPLC Peak of a) HPLC Peak of standard (Telmisartan), b) HPLC peak of samples #1 ( aliquot of water containing released drug collected after one day of soaking the drug adsorbed material in distilled water), c) sample #2 collected after three days, d) Sample #3: collected after five days.
Figure 16: HPLC Peak of a) HPLC Peak of standard (glimpiride), b) HPLC peak of samples #1 (aliquot of water containing released drug collected after one day of soaking the drug adsorbed material in distilled water), c) sample #2 collected after three days, d) Sample #3: collected after five days.

Figure 17: HPLC Peak of a) HPLC Peak of standard (rosuvastatin), b) HPLC peak of samples #1 (aliquot of water containing released drug collected after one day of soaking the drug adsorbed material in distilled water), c) sample #2 collected after three days, d) Sample #3: collected after five days, e) Sample #4: collected after seven days.
Table 1: A comparison of drugs storage capacities of compounds 3-5.

| Synthesized porous material | Specific surface areas (m²g⁻¹) | Pore size (Å) | Name of drug absorbed in materials | Drug loading capacity (%) | Time of released (days) |
|-----------------------------|---------------------------------|---------------|----------------------------------|--------------------------|-------------------------|
| Compound 3                  | 2800                            | 10            | Terazosine hydrochloride         | 0.095                    | 5                       |
|                             |                                 |               | Telmisartan                      | 0.083                    | 3                       |
|                             |                                 |               | Glimpiride                       | 0.041                    | 3                       |
| Compound 4                  | 3000                            | 10            | Terazosine hydrochloride         | 0.200                    | 7                       |
|                             |                                 |               | Telmisartan                      | 0.160                    | 5                       |
|                             |                                 |               | Glimpiride                       | 0.138                    | 3                       |
| Compound 5                  | 2900                            | 10            | Terazosine hydrochloride         | 0.316                    | 9                       |
|                             |                                 |               | Telmisartan                      | 0.086                    | 3                       |
|                             |                                 |               | Glimpiride                       | 0.138                    | 3                       |

Conclusion

In conclusion the present work directs towards the use of some nanosized bio-MOFs which were aimed to know their potential for in vitro drugs adsorption. This work elaborates the more potential use of nanoscale bio-MOFs for drugs storage and its release in comparison to previously reported carrier materials. In medical grounds the successful in vitro drug adsorption experiments on these nanosized materials may lead to their usage for in vivo drugs adsorption as well.

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