The Impact of Synthesis Method on the Properties and CO₂ Sorption Capacity of UiO-66(Ce)

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Abstract: A series of cerium-based UiO-66 was obtained via hydrothermal and sonochemical methods, using the same quantities of reagents (cerium ammonium nitrate (CAN), terephthalic acid (H₂BDC)) and solvents) in each synthesis. The impact of synthesis method and metal to linker ratio on the structural and textural properties of obtained UiO-66(Ce), as well as their composition in terms of Ce⁴⁺/Ce³⁺ ratio, structure defects resulting from missing linker, and CO₂ adsorption capacity was discussed. By using typical characterization techniques and methods, such as XRD, N₂ and CO₂ sorption, TGA, XPS, and SEM, it was shown that the agitation of reacting mixture during synthesis (caused by stirring or ultrasounds) allows to obtain structures that have more developed surfaces and fewer linker defects than when MOF was obtained in static conditions. The specific surface area was found to be of minor importance in the context of CO₂ adsorption than the contribution of Ce³⁺ ions that were associated with the concentration of linker defects.

Keywords: UiO-66; cerium; CO₂ sorption; sonochemical synthesis

1. Introduction

Gas adsorption and separation [1], gas storage [2], chemical sensing [3], drug delivery [4] and heterogeneous catalysis [5] are the most popular fields of metal-organic framework (MOFs) utilization. Among all classified MOF structures (over 60,000 reported in Cambridge Crystallographic Data Centre, but many of them are repeated), there are some that attract special attention owing to very high—if not outstanding—surface area, chemical properties or thermal and chemical resistance. The structure instability upon heating or in the presence of acids/bases is the main reason for hindering MOFs application in many catalytic processes. One well-recognized, stable metal-organic structure is UiO-66 that is built from hexanuclear zirconium clusters connected with 1,4-benzenedicarboxylate (H₂BDC). The UiO-66 was first reported by Cavka et al. [6]. It is characterized by good thermal and chemical stability, thanks to strong Zr-O bonds. Moreover, the UiO-66 structure exhibits excellent stability in water [7] and promising features for CO₂/CH₄ gas separation [8]. By changing metal in the node or
introducing/substituting the functional group in an organic linker, one can change the properties of the parent MOF. Modifications in UiO-66 usually affect linker, which often improves its adsorptive properties but also reduces its thermal resistance [9]. The exchange of Zr to other metals, like Hf [10] and Ti [11], has been reported, while introducing Ce$^{4+}$ to an MOF structure is a promising way to obtain redox-active materials [12–15].

Nouar et al. [16] reported that partial substitution of Zr by Ce in UiO-66 enhanced binding of methanol and led to its catalytic decomposition to CO$_2$ which may be due to the combination of structural defects and redox activity. The authors found that cerium occurred in two oxidation states: Ce (IV) (predominantly) and Ce (III). Also, Dalapati et al. [14] detected by XPS both Ce (IV) and Ce (III) ions in UiO-66 type Ce-based MOF synthesized from cerium (IV) nitrate and 3,4-dimethylthieno[2–b] thiophene-2,5-dicarboxylic acid (H$_2$DMTDC). They used cerium-based MOF in the oxidation of thiophenol to 1,2-diphenyldisulfide and found that it revealed excellent oxidase-like catalytic properties, thanks to the existence of redox-active cerium atoms in the framework. In contrary, Lammert et al. [12] characterized obtained in their laboratory UiO-66(Ce) by XANES and did not detect any traces of Ce$^{3+}$. That material was tested as co-catalyst in aerobic oxidation of benzyl alcohol, and provided significantly higher alcohol conversion than zirconium based UiO-66 or CeO$_2$. The Ce$^{4+}$ cations in the hexanuclear cluster in MOF can catalyze redox reactions through reduction to Ce$^{3+}$[15]. Cerium (IV) ammonium nitrate (CAN), which is often used as a Ce precursor for UiO-66 based metal-organic frameworks, can act as a homogeneous catalyst in oxidation of alcohols, thioethers, epoxides, alkylbenzenes, and active methylene compounds [17].

Owing to its unique properties (high oxygen mobility, easy and reversible Ce$^{4+}$ $\rightarrow$ Ce$^{3+}$ transition), cerium (IV) oxide has attracted significant interest in a wide range of reactions [18–20], including CO$_2$ conversion [21,22]. Oxygen vacancies in CeO$_2$ act as sites for activation of adsorbing CO$_2$ molecules. In general, the presence of Ce$^{3+}$ (Lewis base) enhances CO$_2$ adsorption and conversion, promotes CO$_2$ disproportionation, and stabilizes CO-containing intermediates [21,23]. The global importance of CO$_2$ in terms of greenhouse gas emissions from fossil fuel-based power generators drives the need for developing new materials that can be used for its adsorption, storage, and/or conversion. Carbon dioxide is a thermodynamically stable compound with very low reactivity. Utilization of CO$_2$ is considered challenging because its activation requires overcoming a thermodynamic barrier. One of many possibilities of CO$_2$ utilization is its conversion to methanol, which is a popular solvent and key substrate in the chemical industry, e.g., for the synthesis of liquid fuels [24]. Hydrogenation of CO$_2$ to methanol is favored at low temperatures and high pressure. The most popular catalyst for this reaction is Cu/ZnO (also supported and/or promoted) [25]. It was reported that CeO$_2$-containing catalysts exhibit much-higher methanol yield in comparison with those without CeO$_2$ [26,27].

The issue of UiO-66 doping with cerium to obtain mono- or bimetallic MOFs that may reveal better catalytic properties than parent, Zr-based MOFs, is still to be addressed, both in the context of the improvement of particular properties of these materials (such as textural, thermal, presence of structure defects, structure homogeneity) and their redox activity. Excellent performance of ceria in CO$_2$ activation justifies the choice of cerium-based MOFs as potential catalysts for CO$_2$ conversion. It is also essential to study the surface of MOF, since structures with open metal sites characterize better CO$_2$ adsorption capacity [28].

Among various possibilities of MOFs preparation, i.e., solvothermal (hydrothermal when carried out in aqueous solutions), sonochemical, mechanochemical, electrochemical or microwave assisted [29], the most popular is the first one. The synthesis method, as well as its parameters, such as temperature, duration time, the type of used metal salt or solvent, the pH of the solution, may influence the properties of the final product.

In this work, we study the influence of the synthesis method on the properties of cerium-based UiO-66. The material was obtained in three different ways by using hydrothermal and sonochemical syntheses. The UiO-66(Zr) is typically obtained via solvothermal route, but a very good outcome of the sonochemical synthesis was reported by Kim et al. [30]. The use of ultrasounds during MOF
preparation allows accelerated nucleation; hence, at low reaction temperature and ambient pressure, crystallization time is short. Moreover, obtained particles are more uniform and smaller than in the case of solvothermal synthesis [31]. To our best knowledge, UiO-66(Ce) has not been obtained yet via the sonochemical route. According to recently published [12–14,32], cerium-based UiO-66 structures were obtained only by solvothermal synthesis.

2. Results and Discussion

2.1. Crystallographic structure of UiO-66(Ce) materials

The XRD patterns of UiO-66(Ce) are presented in Figure 1. It has been proven that all of the synthesized materials are crystalline and are similar to the XRD patterns of UiO-66(Zr) [33] and UiO-66(Ce), as reported in the literature [13,14]. The XRD patterns exhibit sharp reflexes around $\theta$ of 7.2°, 8.3°, 11.7°, 13.8°, 14.4° corresponding to reflections for the (111), (200), (220), (311) and (222) planes, respectively. All of these reflexes are somewhat shifted towards lower $\theta$ values compared to the UiO-66(Zr) [6], which is due to higher radii of Ce$^{4+}$ (0.97 Å) than Zr$^{4+}$ (0.84 Å) [34]. All of the presented diffractograms are typical for metal-organic structures. However, the diffractogram of UiO-66(Ce) obtained under continuous mixing and excess of the metal precursor (sample M (2:1)), displays intensive reflexions at 16.5°, 28.9° and 33.5°, that can be ascribed to cerium formate (Ce(HCOO)$_3$), which is formed as a result of deprotonation of carboxylic acid. Significant amounts of cerium formate in the M (2:1) sample affects the textural properties and CO$_2$ sorption, as discussed in the next section. Besides, the difference between M (2:1) and other samples was visible during the filtering-off step. The M (2:1) sample contained two fractions—white microcrystalline powder, which was cerium formate, and yellow powder which was UiO-66(Ce). It seems that unlike other methods, the hydrothermal one with mixing is sensitive to metal: linker ratio, which results in a homogeneous product only at equimolar M:L. Moreover, the sonochemical syntheses yielded the most homogeneous product, which could be due to lower temperatures applied (50 °C) and good mass transport provided by the ultrasounds. Crystallization of cerium formate is facilitated at higher temperatures and according to [35] occurs when synthesis is carried out for longer than two hours. Formation of Ce(HCOO)$_3$ is a consequence of DMF hydrolysis to formic acid and dimethyl amine. That reaction is impossible to avoid in the presence of water, which was used as one of the solvents in this work. As was calculated [36], hydrogen bonds between water and amides are stronger than water molecules. The rate of DMF hydrolysis increases with temperature; therefore, the risk of cerium formate crystallization was higher during hydrothermal syntheses of UiO-66(Ce), than during sonochemical procedure. Superior formation of cerium formate during synthesis of M (2:1) was probably caused by the simultaneous occurrence of three factors: the presence of H$_2$O, the excess of Ce precursor and the fleeting overheating of solution, which is often a more or less serious concern when a magnetic stirrer is used. Overheating was not an issue in the case of synthesis of M (1:1), which was Ce(HCOO)$_3$-free, simply because cerium precursor was not in excess (as in the case of M(2:1)).

The crystallite sizes calculated from Scherrer equation for the [111] plane were as follows: 128 nm for UiO-66, 83.5 and 83.7 nm for H(1:1) and H(2:1); 65.8 and 59.7 nm for M(1:1) and M(2:1) and 43.4 and 46.9 nm for S(1:1) and S(2:1), respectively. Therefore, the synthesis method had an impact on the size of UiO-66(Ce) crystallites, yielding the biggest when carried out hydrothermally in static conditions (H). Smaller crystals were obtained when mixing was applied during hydrothermal synthesis (M), whereas the employment of ultrasounds (S) led to the smallest crystals.

In solution, crystals grow from molecules that must be transported from the bulk of the solution to the interface between the crystal and the solution. Therefore, mass transport is a key factor in crystal growth. Depending on the crystallization technique, the effectiveness of mass transfer is different because in each method, supersaturation is attained in a unique way. Hence, the applied procedure may affect the final product. For example, crystallization from motionless solution should lead to bigger crystals since the molecules from solution have more chances to fit already-existing crystals,
making them bigger, whereas agitation may cause molecules from the solution to hit the crystals, smashing off some molecules, so the crystals get smaller. Thus, the type of agitation affects the motion in solution; and therefore, the effectiveness of mass transfer. The rate of agitation, besides other factors such as temperature, or trace impurities, influences the rate of nucleation and the rate of crystal growth [37,38]. It can be observed in the presented study that motion of cerium precursor and linker molecules in the solution during syntheses M and S led to UiO-66(Ce) crystallites of decreased size compared to stationary conditions (H). In our case, bigger crystals were obtained when agitation was caused by stirring (M) than by ultrasounds (S). When the liquid is subjected to acoustic cavitation caused by ultrasounds, the microscopic bubbles are set into radial oscillation. They collapse very quickly, within several nanoseconds and the phenomenon repeats for a high number of bubbles in each acoustic period. Since there is possibly no mechanical equilibrium under these conditions at the microscopic scale in the liquid, local supersaturation may occur and result in more numerous, small crystals that nucleate more rapidly. The time of nucleation is the time required for the clusters to reach critical size. It can be reduced by any microscopic effect that enhances the formation of the smallest clusters. In addition, nucleation can be enhanced by segregation that is driven by cavitation bubbles [39]. It was reported by Enomoto et al. [40] that crystallization under ultrasounds from less concentrated solutions leads to smaller and more uniform particles than when the process is carried out under stirring, whereas there is no significant difference in particle size distribution between those two methods for solutions of high concentrations.

Figure 1. XRD patterns of UiO-66 and that of UiO-66(Ce) obtained in this work. The “*” is Ce(HCOO)₃.

The crystallographic data of UiO-66(Ce) samples determined from XRD patterns are set in Table 1. The lattice parameters for all samples are similar to those reported in [13,14] and are slightly higher than calculated for UiO-66(Zr), which arises from higher ionic radii of Ce⁴⁺ than Zr⁴⁺. It can also be observed that unit cell parameters of UiO-66(Ce) samples differ from each other, which can be due to structure defects, such as vacancies, interstitial atoms, substitutional atoms, local structure transformations, etc. The defects may cause strains in the crystal lattice of UiO-66(Ce) resulting from planar stress—either compressive (corresponding to the increase of unit cell and a peak shift to lower 2θ angles) or tensile (decrease of the unit cell, peak shift to higher 2θ angles). The strain in crystal lattice of UiO-66(Ce) samples can be caused by structure defects, such as linker or metal cluster vacancies. Whereas the occurrence and concentration of those vacancies can be affected by the method of synthesis. Moreover, the occurrence of tensile stress, and thus the decrease of unit cell parameters can be caused by the presence of Ce³⁺ ions that have bigger radius than Ce⁴⁺ (1.143 Å against 0.97 Å [34]). In the case of studied in this work UiO-66(Ce) obtained at M:L=1:1, the following

| Sample | a [Å]   | Volume [Å³] | D₁₁₁ [nm] |
|--------|---------|-------------|-----------|
| H(1:1) | 21.0469 | 9323.2      | 83.5      |
| M(1:1) | 21.2110 | 9543.0      | 65.8      |
| S(1:1) | 21.2797 | 9636.0      | 43.4      |
| H(2:1) | 21.4095 | 9813.4      | 83.7      |
| M(2:1) | 21.4544 | 9875.4      | 59.7      |
| S(2:1) | 21.3468 | 9727.4      | 46.9      |
| UiO-66 | 20.7004 | 8870.3      | -         |
| UiO-66(Ce) | 21.4727 | 9900.6 | -         |

Formula: [Ce₆O₄(OH)₄(BDC)₆] Space group: Fm-3m
increase of unit cell size is observed: H < M < S, and it corresponds to the decrease in Ce\(^{3+}\) contribution (determined by XPS), being 47.9, 46.1 and 45.1%, respectively. For the samples obtained at the excess of metal precursor (M:L=2:1), the impact of Ce\(^{3+}\) contribution was less significant than the impact of such structural defects as linker vacancies that caused compressive stress in the lattice, thus increasing the size of the unit cell. The issues of the presence of Ce\(^{3+}\) cations and linker vacancies in the UiO-66(Ce) samples are discussed further in this paper.

| Sample | a [Å] | Volume [Å\(^3\)] | \(D_{111}\) [nm] |
|--------|-------|-------------------|------------------|
| H(1:1) | 21.0469 | 9323.2 | 83.5 |
| M(1:1) | 21.2110 | 9543.0 | 65.8 |
| S(1:1) | 21.2797 | 9636.0 | 43.4 |
| H(2:1) | 21.4095 | 9813.4 | 83.7 |
| M(2:1) | 21.4544 | 9875.4 | 59.7 |
| S(2:1) | 21.3468 | 9727.4 | 46.9 |

For UiO-66 [6] and UiO-66(Ce) [12], the materials obtained in this work possess a face-centered cubic (fcc) framework structure and crystallize in the Fm-3m space group. According to detailed studies performed by others [13,14], these materials consist of hexanuclear \([\text{Ce}_6\text{O}_4\text{(OH)}_4\text{(BDC)}_6]^{12+}\) building blocks that are connected by the carboxylate groups of twelve BDC ligands. The framework contains two types of polyhedral cages: tetrahedral (smaller) and octahedral (larger). Each central octahedral cage is connected with eight tetrahedral cages through narrow triangular windows [14]. The topologies and structures corresponding to UiO-66(Ce) materials obtained in this work are presented in Figure 2.

The XRD analyses suggest that, as reported in [6,10,12], the UiO-66(Ce) materials obtained in this work possess a face-centered cubic (fcc) framework structure and crystallize in the Fm-3m space group. According to detailed studies performed by others [13,14], these materials consist of hexanuclear \([\text{Ce}_6\text{O}_4\text{(OH)}_4\text{(BDC)}_6]^{12+}\) building blocks that are connected by the carboxylate groups of twelve BDC ligands. The framework contains two types of polyhedral cages: tetrahedral (smaller) and octahedral (larger). Each central octahedral cage is connected with eight tetrahedral cages through narrow triangular windows [14]. The topologies and structures corresponding to UiO-66(Ce) materials obtained in this work are presented in Figure 2.
The SEM pictures of M (1:1) and M (2:1) samples are shown in Figure 3. It can be seen that both contain small crystals of UiO-66(Ce), but the sample obtained at the equimolar metal to linker ratio is homogeneous (Figure 3a), while the one obtained with the excess of CAN also contains big rods of cerium formate (Figure 3b), whose presence was also confirmed by XRD.

![SEM pictures of M(1:1) and M(2:1) samples](image)

**Figure 3.** SEM pictures of M(1:1) (a) and M(2:1) (b) samples.

### 2.2. Textural properties of UiO-66(Ce)

The experiments of adsorption-desorption of N\(_2\) (at 77K), CO\(_2\) and benzene (both at 298 K) were conducted in order to determine the specific surface areas of obtained materials and to evaluate their porosity. Results are presented in Table 2, Figures 4 and 5.

![N\(_2\) adsorption-desorption isotherms at 77K for UiO-66(Ce) materials](image)

**Figure 4.** N\(_2\) adsorption-desorption isotherms at 77K for UiO-66(Ce) materials (adsorption—filled symbols; desorption—empty symbols).

![Distribution of the volume of sub-micropores (<0.4–2 nm), micropores (0.4–2 nm) and mesopores (2–50 nm) in UiO-66(Ce) obtained at M:L=1:1 (a) and M:L=2:1 (b).](image)

**Figure 5.** Distribution of the volume of sub-micropores (<0.4–2 nm), micropores (0.4–2 nm) and mesopores (2–50 nm) in UiO-66(Ce) obtained at M:L=1:1 (a) and M:L=2:1 (b).

As is shown in Table 2, the highest specific surface area (SSA) of 606 m\(^2\)/g and 560 m\(^2\)/g was noticed for UiO-66(Ce) obtained by hydrothermal synthesis on mixing M(1:1) and in sonochemical synthesis S(1:1), respectively. The material obtained via static hydrothermal synthesis H(1:1) was...
characterized by significantly lower SSA. Hence, the motion of molecules in the solution, caused by stirring or ultrasounds, facilitated crystallization of structures having more developed surface than in the case of traditional, static hydrothermal synthesis. As was mentioned in the previous section, the agitation of solution during crystallization affects the nucleation and crystal growth; therefore, crystal size and size distribution must be determined. Stirring and sonication influence nucleation by shifting size distribution towards small particles and modifying the morphology of the obtained crystals, whereas the impact of ultrasounds is more important. In general, the UiO-66(Ce) materials obtained at M:L=2:1 had lower surface areas than those synthesized at M:L=1:1. However, the impact of metal: linker ratio was more pronounced for syntheses carried out under continuous stirring and in ultrasounds than in static hydrothermal synthesis. The lowest SSA (111 m²/g) was noticed for the M(2:1) sample that contained unwanted cerium formate, as was proven by the XRD (Figure 1).

The shapes of N₂ adsorption-desorption isotherms (Figure 4) of obtained materials are similar and reveal both adsorption at low relative pressures assigned to micropore filling, and adsorption at high relative pressures, which corresponds to N₂ adsorption in the interparticle voids.

The parent UiO-66(Zr) contains tetrahedral and octahedral cages of 0.8 nm and 1.1 nm width, respectively [41], and is microporous. The experiments of CO₂ sorption at 298 K showed that UiO-66(Ce) materials obtained in this work contain both micro- and mesopores (Table 2). In most of the samples, the volume fraction of micropores was predominant. It can be seen in Table 2 and Figure 5a that synthesis method affected pore size distribution for the samples obtained at M:L=1:1, i.e., the UiO-66(Ce) obtained in static hydrothermal synthesis (H) had the highest fraction of micropores, whereas increasing contribution of mesopores was observed in the samples obtained under stirring (44%) and sonication (67%). Combining the results of XRD (Table 1) with CO₂ sorption (Table 2), it can be noticed that the content of mesopores in the sample increased with decreasing crystallite size. For example, UiO-66(Ce) obtained in sonochemical synthesis was characterized by the smallest crystallite size and highest fraction of mesopores. It is very likely that the crystallites of UiO-66(Ce) generate mesoporous interparticle voids that are more numerous when smaller crystals are randomly packed. Moreover, the decrease in crystallite size should lead to a decrease of the internal surface area and increase of the external surface area, accompanied by a decrease in micropore volume [42].

Table 2. Pore volume, mean mesopores dimension, and fraction of micro- and mesopores in UiO-66(Ce) materials determined from CO₂ and benzene sorption (slit-pore model).

|                  | S_{BET} [m²/g] | \(V_{mic}\) [cm³/g] | \(V_{mes}\) [cm³/g] | \(d_{mes}\) [nm] | \(\phi_{mic}\) | \(\phi_{mes}\) |
|------------------|----------------|---------------------|---------------------|-----------------|----------------|----------------|
| H(1:1)           | 391            | 0.1810              | 4.37                | 0.84            | 0.16            |
| M(1:1)           | 606            | 0.3079              | 7.94                | 0.56            | 0.44            |
| S(1:1)           | 560            | 0.3120              | 10.42               | 0.33            | 0.67            |
| H(2:1)           | 360            | 0.1970              | 5.98                | 0.70            | 0.30            |
| M(2:1)           | 111            | 0.0679              | 5.42                | 0.78            | 0.22            |
| S(2:1)           | 381            | 0.1860              | 7.99                | 0.66            | 0.34            |

UiO-66 1187 [6]
1086 [9]
1290 [41]
1080 [43]

UiO-66(Ce) 1282 [12]
Samples obtained at M:L=2:1, especially H(2:1) and S(2:1), are characterized with similar contribution of micro- and mesopores. Those two samples also had similar SSA, but they significantly differed in terms of crystallite size (Table 1). Syntheses carried out with the deficiency of organic linker are more likely to yield products with a higher concentration of structure defects, among which the most probable to occur are linker vacancy, i.e., microporous intraparticle void. Thus, for UiO-66(Ce) samples obtained at M:L=2:1, the fraction of micropores predominated over mesopore interparticle voids (thanks to the higher concentration of linker defects), regardless of the crystallite size.

UiO-66(Ce) materials reported in this work have significantly lower surface areas than other Zr and Ce based UiO-66 structures reported in the literature. This might be due to a considerably larger scale of synthesis than compared to [6,9,12,41], which also influenced the porosity of UiO-66(Ce) reported in this paper.

2.3. XPS of UiO-66(Ce)

Determination of cerium content and contribution of its oxidized and reduced state can be performed by using XPS, which is a surface sensitive technique that probes the top 10–12 nm of a film. It allows quantitative analysis of particular elements in the sample, including their oxidation state. The analysis of Ce3d photoemission spectra in Ce$^{3+}$ and Ce$^{4+}$ containing samples is not straightforward because it is composed of 10 peaks corresponding to both oxidation states. In addition, sample sputtering with Ar$^+$ can cause reduction of some Ce$^{4+}$ owing to loss of the lighter component, such as oxygen [44]. Thus, the artificial increase of the Ce$^{3+}$ concentration can be observed. Nevertheless, it was found that the shorter the time of Ar$^+$ sputtering, the lower is the degree of Ce reduction [45,46]. UiO-66(Ce) samples studied in this work were not subjected to Ar sputtering; hence any Ce$^{3+}$ presence is not from oxygen scavenging. Figure 6 shows the Ce3d spectra for all synthesized UiO-66(Ce) samples, excluding M(2:1). In the ideal UiO-66 structure, the concentration of cerium is 5.55 at%, but in presented samples, it ranged from 6.07 to 7.19 at.%. Higher Ce concentration arises from the presence of linker vacancies. All spectra reveal 10 peaks that can be assigned to Ce$^{3+}$ and Ce$^{4+}$ cations, proving the presence of redox-active cerium atoms in the framework. The concentrations of particular cerium cations were similar, but Ce$^{3+}$ predominated. In the ideal UiO-66 structure, cerium will exclusively occur in Ce$^{4+}$. The presence of Ce$^{3+}$ cations may be due to the occurrence of such structural defects as linker vacancies. As was presented in Figure 6, the synthesis method affected the contribution of Ce$^{3+}$ cations, both in samples obtained at M:L=1:1 and M:L=2:1. In each group, the highest contribution of Ce$^{3+}$ was noticed for samples obtained in static hydrothermal synthesis (H), and it slightly decreased when the synthesis was carried out under stirring (M) and ultrasounds (S). The presence of both oxidation states of cerium is important for the catalytic and adsorptive properties of UiO-66(Ce). Nonetheless, the detailed features of such a structure containing both Ce$^{4+}$ and Ce$^{3+}$ cations in the nodes have not been determined yet. However, it is known that under ambient conditions, both Ce(III) and Ce(IV) are stable valence states. Recently, Smolders et al. [15] published the outcome of their study on the redox behavior of cerium-based UiO-66 during oxidation of benzyl alcohol in the presence of TEMPO (2,2,6,6-Tetramethylpiperidin-1-yl)oxyl). In the initial UiO-66(Ce), cerium was in its oxidized (Ce$^{4+}$) state. Based on the Extended X-Ray Absorption Fine Structure (EXAFS), the authors reported that Ce$^{4+}$ can be reduced to Ce$^{3+}$ upon reaction conditions, perpetuating the structural integrity of the material. It was also found that larger ionic radii of Ce$^{3+}$ (1.02 vs. 0.92 for Ce$^{4+}$) led to a distortion in Ce cluster; thus, two different Ce-Ce distances occurred.
linker deficiency in the mixture (M:L=2:1) also facilitates DMF attachment to cerium cluster. Coordination of DMF to cerium cations was more important for materials obtained in syntheses, carried out under continuous stirring (M) and ultrasounds (S). XPS analysis showed that those samples contained from 1.2 at.% to 6.5 at.% of nitrogen (Table 3), proving the same DMF presence. The TG and DTG curves for Uio-66(Ce) materials obtained at M:L=1:1 and 2:1 using three different synthesis methods are presented in Figure 7. According to TGA, all obtained structures are thermally stable up to ca. 260–280 °C (Table 3). The stability of Uio-66(Ce) up to 240 °C was proved by temperature-dependent PXRD—as reported by Lammert et al. [12]. Observed in Figure 7, weight loss at ca. 100 °C is due to moisture evaporation from the sample. The DMF evaporates at 156 °C; hence, the lack of weight loss at this temperature proves that solvent exchange after synthesis (DMF to chloroform) in the pores of Uio-66(Ce) was completed. However, it is quite possible that during synthesis, some DMF molecules formed an adduct with Ce precursor and were not exchanged with the BDC linker. Such a strongly coordinated to cerium cation DMF may be removed from the sample at a higher temperature. In the case of all Uio-66(Ce) samples (except for M(2:1)), some weight loss, that can be attributed to degradation of strongly coordinated DMF, was observed at ca. 240 °C. The XPS analysis showed that those samples contained from 1.2 at.% to 6.5 at.% of nitrogen (Table 3), proving the same DMF presence. Coordination of DMF to cerium cations was more important for materials obtained in syntheses, carried out under continuous stirring (M) and ultrasounds (S). Therefore, agitation of solution during synthesis helped DMF coordination to Ce clusters. In addition, linker deficiency in the mixture (M:L=2:1) also facilitates DMF attachment to cerium cluster.

### Table 3. The average amount of BDC and DMF molecules coordinated to one Ce cluster in Uio-66(Ce).

| Sample  | Molecules per Ce Cluster | N [At. %] (from XPS) |
|---------|--------------------------|----------------------|
|         | BDC                     | DMF                  | BDC+DMF   |
| H(1:1)  | 11.1                     | 0.0                  | 11.1      | 1.2       |
| M(1:1)  | 10.2                     | 1.2                  | 11.4      | 2.1       |
| S(1:1)  | 10.8                     | 0.8                  | 11.7      | 4.1       |
| H(2:1)  | 9.8                      | 0.4                  | 10.2      | 1.4       |
| M(2:1)  | 10.3                     | 0.0                  | 10.3      | -         |
| S(2:1)  | 9.1                      | 2.4                  | 11.5      | 6.5       |
Figure 7. TG (a,b) and DTG (c,d) profiles for UiO-66(Ce) synthesized at M:L=1:1 (a,c) and M:L=2:1 (b,d).

Thermal decomposition of BDC linker in the obtained UiO-66(Ce) materials starts at ca. 260 °C and is followed by complete decomposition of the framework. Oxidation of BDC is completed at ca. 470 °C for the samples obtained in hydrothermal syntheses (H and M), and above 520 °C for samples obtained via the sonochemical method (S). Thermal decomposition of all samples leads to CeO₂. The DTG plots of samples obtained in hydrothermal syntheses (H and M) show that there are two weight loss regions above 260 °C: (i) sharp, negative peaks at ca. 320 °C, corresponding to 10–16% weight loss (depending on the sample), that can be attributed to degradation of terminal (surface) linker molecules, which have one unoccupied carboxylic group; therefore, are less stable thermally, and (ii) broad peaks at higher temperatures, corresponding to 15–25% weight loss, that can be assigned to degradation of more thermally resistant BDC that is connected to two Ce clusters. Hence, we may expect that surfaces of samples obtained in hydrothermal syntheses (H and M) are terminated with cerium clusters and BDC. The DTG profiles for samples obtained in sonochemical syntheses do not show peak at ca. 320 °C; therefore, those materials are probably terminated on their surfaces with cerium clusters.

The weight loss from 260 °C corresponds to the percentage of BDC in particular UiO-66(Ce) sample. In a defect-free UiO-66(Ce) structure, each Ce₆O₄(OH)₄ cluster coordinates 12 ligand molecules, which corresponds to 46.25% weight loss on TGA. For UiO-66(Ce) samples obtained in this work, the weight loss in the 260–600 °C range was lower, indicating the presence of structural defects due to missing linker molecules. It was reported [33,47] that removing some BDC linkers from UiO-66 yields a stable framework. According to our calculations (done based on Katz et al. [48]), one inorganic Ce₆O₄(OH)₄ cluster coordinated average from 9.8 to 11.1 BDC ligands (Table 3). This number of BDC ligands was affected by the synthesis method and applied M:L ratio. As was discussed above, in some of the samples, DMF coordinated with Ce clusters, occupying the place of BDC linker. More linker vacancies were noticed for samples synthesized with the excess of Ce precursor (the M(2:1) is not taken into consideration owing to the presence of cerium formate).

As was proven by XRD and TGA, linker vacancies are present in all UiO-66(Ce) samples. On one hand, missing linker means lower thermal stability of the structure, but on the other hand, it can positively influence the catalytic properties of MOF because defected (Figure 8b) or open metal sites (Figure 8c) are the adsorption sites for reagents. The removal of one BDC molecule from the original UiO-66 structure affects four Zr clusters and results in a reduction of its coordination from 12 to 11. The overall charge of such a defected structure is +2. In order to ensure charge neutrality, the dangling
bonds of the positively charged metal cluster may be saturated, e.g., with OH⁻ groups, anions of the acidic modulator (if used during synthesis) [49,50], or neutral molecules such as H₂O or DMF. Basically, linker removal is promoted by acidic modulators [51]; however, even if the synthesis is carried out without those modulators (like in this paper), the UiO-66 can have linker defects [50]. Linker vacancies in UiO-66 introduce coordinatively unsaturated Zr atoms, which results in open Lewis acid sites that is important for the adsorptive and catalytic properties of that material [52–54]. In the case of cerium-based UiO-66, the formation of linker vacancy may result either in lower valent cerium, i.e., Ce(III), or that vacancy may be compensated by -OH, H₂O or DMF, preserving the Ce(IV) and introducing Bronsted acid sites. The concentration of structural defects as well as the chemistry of defective metal cluster, either hydrated as shown in Figure 8b or dehydrated as displayed in Figure 8c, has an impact on the catalytic activity of MOF. For example, Caratelli et al. [55] reported that the activity of UiO-66 in esterification of levulinic acid with ethanol was higher for MOF having more linker defects and when the defective cluster was hydrated.

![Figure 8. Ce cluster in UiO-66: defect-free 12-coordinated (a) and defective (after removal of 1 linker molecule) 11-coordinated, hydrated (b) and dehydrated (c).](image)

According to TGA (Figure 7), the less-defected Ce sites were present in S(1:1) and S(2:1), where an average of 11.5 and 11.7 BDC+DMF molecules coordinated to one cerium cluster, respectively. In addition, as was shown in Figure 9, increasing concentration of linker vacancies (i.e., a decrease of the average number of BDC and BDC+DMF coordinated to one Ce cluster) went together with the increasing content of Ce³⁺ cations.

![Figure 9. Coordination of BDC linker and DMF to one Ce cluster in UiO-66(Ce) vs. concentration of Ce³⁺. On the right, the exemplary UiO-66(Ce) structure with linker vacancies corresponding to the average coordination of 10 linker molecules to one Ce cluster.](image)
2.5. CO₂ adsorption on UiO-66(Ce)

The CO₂ adsorption on UiO-66(Ce) samples obtained by different methods was performed at 273 K and 298 K (Table 4). Higher CO₂ adsorptions were noticed for materials synthesized at M:L=1:1. In general, the amounts of CO₂ adsorbed in both temperatures varied significantly and were differently depended on factors such as specific surface area of the sample, content of Ce³⁺ and Ce⁴⁺ ions or concentration of structural defects caused by the missing linker (Figure 10.). It was found that CO₂ adsorption on UiO-66(Ce) samples at 273 K was not influenced by their SSA (Figure 10a), whereas it increased with Ce³⁺ content (Figure 10b). In contrary, CO₂ adsorption at 298 K increased with SSA of the sample, but did not depend on the Ce³⁺ contribution. It is known that carbon dioxide can adsorb both Ce³⁺ and Ce⁴⁺ ions, but it bounds more strongly to Ce³⁺. According to our research, CO₂ adsorption on reduced cerium cations in UiO-66(Ce) is favored only at low temperature (273 K).

Table 4. CO₂ adsorption and contribution of reversible and irreversible CO₂ adsorption on UiO-66(Ce) samples.

| Sample   | CO₂ Adsorption at 273 K [mmol/g] | CO₂ Adsorption at 298 K [mmol/g] | Reversible [%] | Irreversible [%] |
|----------|---------------------------------|---------------------------------|----------------|-----------------|
| H(1:1)   | 1.90                            | 0.7640                          | 83.6           | 16.4            |
| H(2:1)   | 1.34                            | 0.6125                          | 84.6           | 15.4            |
| M(1:1)   | 1.68                            | 0.9011                          | 92.9           | 7.1             |
| M(2:1)   | -                               | 0.4328                          | 58.5           | 41.5            |
| S(1:1)   | 0.83                            | 0.7545                          | 74.9           | 25.1            |
| S(2:1)   | 0.47                            | 0.7498                          | 75.1           | 24.9            |
| CeO₂     | -                               | 0.6528                          | 47.8           | 52.2            |

Figure 10. CO₂ adsorption at 273 K (▲) and 298 K (●) as a function of specific surface area (a), Ce⁴⁺ /Ce³⁺ ratio (b) and missing linker defect concentration (c) in the obtained UiO-66(Ce) materials.

The CO₂ adsorption-desorption tests carried out at 298 K revealed that some of CO₂ adsorbed on MOFs irreversibly (i.e., did not desorb entirely under vacuum at 298 K). To our best knowledge, the irreversible CO₂ adsorption on UiO-66(Zr) was reported only by Ethiraj et al. [56]. The contribution of reversible and irreversible CO₂ adsorption on UiO-66(Ce) samples is presented in Table 4. The M(2:1) sample was excluded from the discussion owing to the presence of cerium formate that probably transformed to cerium oxide upon heating under vacuum prior to CO₂ sorption. In that sample, the contribution of reversible and irreversible CO₂ adsorption was similar to nanocrystalline, commercial CeO₂ (Rhodia).

As was presented in Table 4, the synthesis method had a bigger impact on the amount of reversibly and irreversibly adsorbed CO₂ than the M:L ratio. The percentage of CO₂ that adsorbed irreversibly
Catalysts 2019, 9, 309 on UiO-66(Ce) samples obtained by different methods decreased in the following order: S (∼25%) > H (∼16%) > M (∼7%). Slightly higher contribution of irreversible CO₂ adsorption can be observed for samples synthesized at equimolar cerium to BDC ratio.

The CO₂ sorption capacities at 298 K are similar for H and S samples (Table 4) and somewhat higher for the M sample, which has the highest SSA. The concentration of cerium in all UiO-66(Ce) samples (calculated from XPS spectra) is also similar, but the concentration of Ce³⁺ ions, believed to be responsible for irreversible CO₂ adsorption, is the lowest for the materials obtained in sonochemical synthesis (S(1:1) and S(2:1)) (Figure 6). These samples, therefore, reveal the highest contribution of irreversible CO₂ adsorption probably due to the termination of S(1:1) and S(2:1) surfaces with cerium clusters (as was indicated by TGA). The concentration of Ce³⁺ at the surface of those samples was higher than in samples obtained in methods H and M, allowing more CO₂ to strongly (irreversibly) bond with cerium. D’Amato et al. [57] speculate that accumulation of Ce³⁺ near the external surface of UiO-66(Ce) can be due to water oxidation occurring at the solid/liquid interface during synthesis. Good argument for that can be the fact that cerium ammonium nitrate (CAN) is a stoichiometric oxidant in iridium catalyzed H₂O oxidation (4Ce⁴⁺ + 2H₂O → 4Ce³⁺ + 4H⁺ + O₂) [38]. Moreover, in acidic aqueous media, Ce⁴⁺ is an oxidizing reagent in the reaction of H₂O oxidation over ruthenium complexes [59].

Another factor that was found to have an impact on CO₂ adsorption was the concentration of missing linker molecules in the UiO-66(Ce) structure (Figure 10c). The concentration of this kind of structural defects is presented as the number of molecules of BDC linker (and DMF molecules, as discussed in Section 2.4) coordinated to one Ce cluster. The presence of missing linker defects (as well as missing cluster defects) in UiO-66 is normal and does not influence the framework, but may lead to different mechanical, adsorptive, and catalytic properties of this material [33]. Wu et al. [60] reported that linker vacancies enhanced the porosity of UiO-66 and had profound effects on the CO₂ adsorption. Whereas Thornton et al. [61], who computationally studied the role of defects, their variation, and heterogeneity, found that CO₂ uptake at low pressures (up to 1 bar) decreased with defects concentration and at high pressures (up to 35 bar) increased with defect concentration. Moreover, large cavities and pore volumes formed in UiO-66 owing to missing linker molecules led to higher CO₂ uptake. In the case of UiO-66(Ce) samples discussed here (Figure 10c), CO₂ adsorption at 298 K was the highest for the structure containing approximately one missing linker per Ce cluster, i.e., H(1:1). A lower concentration of linker vacancies (higher average coordination of Ce cluster) resulted in a gradual decrease in CO₂ sorption, which is in line with [61] and can be explained by the lower concentration of Ce³⁺ ions in less-defective samples, as was presented in Figure 9.

3. Materials and Methods

3.1. Synthesis of UiO-66(Ce)

A series of cerium-based UiO-66 were synthesized via three methods: hydrothermal (denoted H), hydrothermal with mixing (denoted M) and sonochemical (denoted S) (Figure 11). The Ce precursor was cerium (IV) ammonium nitrate (NH₄)₂Ce(NO₃)₆ (CAN, Fluka, ≥99.9%) while benzene-1,4-dicarboxylic acid (H₂BDC, Sigma Aldrich, 98%) was used as an organic linker. For each method, two metal: linker molar ratios (M:L), i.e., 1:1 and 2:1, were applied.

The UiO-66(Ce) materials were obtained according to the following procedure: The (NH₄)₂Ce(NO₃)₆ (5 mmol) was dissolved in deionized water (10 mL), whereas H₂BDC was dissolved in dimethylformamide (DMF) (30 mL for M:L=1:1 or 15 mL for M:L=2:1). Next, both solutions were mixed and subjected to three different procedures according to three different synthesis methods, i.e.:

1. Hydrothermal synthesis (H): The mixture of Ce precursor and linker was placed in a Teflon lined steel autoclave and heated up to 110 °C in a dryer. After 30 min, the autoclave was slowly cooled down to room temperature.
2. **Hydrothermal synthesis with continuous mixing (M)**: The reagents mixture was placed in a Teflon lined steel autoclave and heated up to 110 °C in an oil bath that was placed on a magnetic stirrer. After 30 min, the autoclave was slowly cooled down to room temperature. The mixture was stirred throughout the whole synthesis, including heating and cooling steps.

3. **Sonochemical synthesis (S)**: The mixture of Ce precursor and linker was placed in a glass vessel and heated up to 50 °C in an ultrasound bath (35 kHz). After 30 min, the mixture was cooled down to room temperature.

Obtained products were filtered off under vacuum and washed with H₂O and DMF in order to remove unreacted inorganic and organic substrates. In the next step, the DMF molecules were removed from UiO-66(Ce) pores and exchanged to chloroform (CHCl₃, Sigma Aldrich, ≥99.9%). The UiO-66(Ce) products were dispersed in 40 mL of CHCl₃, left for 24 h, and filtered off. This procedure was repeated three times. Finally, obtained powders were dried at 80°C for 12 h. Synthesized UiO-66(Ce) materials were denoted X(M:L), where X is the method of synthesis (X= H, M or S) and M:L is the metal: linker molar ratio, i.e., 1:1 or 2:1. The yields of particular products were as follows: H(1:1)-50.9%, H(2:1)-46.0%, M(1:1)-49.3%, M(2:1)-45.2%, S(1:1)-43.5% and S(2:1)-40.0%.

**3.2. Characterization of UiO-66(Ce)**

In order to determine the textural properties, structure, morphology and thermal stability, the UiO-66(Ce) materials were characterized using various techniques and methods, including N₂, CO₂, and benzene adsorption/desorption, X-ray diffraction (XRD). Scanning electron microscopy (SEM), X-ray photoelectron spectroscopy (XPS) and thermogravimetric analysis (TGA).

The N₂ adsorption experiments were carried out on Autosorb 1C apparatus (Quantachrome Instruments) at 77 K. Prior to the measurements, the samples were outgassed in vacuum at 100 °C for 12 h. Specific surface areas were determined by the multipoint BET method. Specific total pore volume was evaluated from N₂ uptake at a relative pressure of P/P₀ = 0.99.

The porous structure of obtained UiO-66(Ce) was determined by CO₂ and benzene adsorption/desorption at 298 K. The CO₂ adsorption/desorption isotherms were obtained at a pressure ranging from 0 mmHg to 700 mmHg. Application of the Dubinin-Radushkevich (DR) equation allowed to determine the volume of micro- and mesopores accessible for CO₂. Moreover, the amount of reversibly and irreversibly adsorbed CO₂ was determined. Benzene adsorption/desorption isotherms (BET)
were obtained at a relative pressure \( P/P_0 \) ranging from 0 to 1. The desorption curve was used for determining the distribution of the volume and surface of slit type mesopores as a function of their dimension. Preceding \( \text{CO}_2 \) and benzene adsorption, the sample was degassed in vacuum at 100 °C. The \( \text{CO}_2 \) sorption capacity was also determined at 273 K, at relative pressure ranging from 0 to 1 bar. Prior to the measurement, the sample was degassed at 100 °C for 12 h.

The powder XRD analyses were conducted using X'Pert Pro apparatus (PANalitycal) using Ni-filtered Cu Kα radiation (\( \lambda = 1.54056 \) Å) equipped with specific optics to perform analysis by Small-angle X-Ray scattering (SAXS) in transmission geometry) with X'Pert HighScore Plus program used for display and analysis of the diffraction patterns. The scanning was performed from 5° to 80° with a rate of 0.03°/step.

The X-ray photoelectron spectroscopy (XPS) of the catalyst was performed using a ThermoScientific K-ALPHA spectrometer equipped in Al-K radiation (1486.6 eV) monochromatized by a twin crystal monochromator, yielding a focused X-ray spot with a diameter of 400 µm, at 3 mA × 12 kV when charge compensation was achieved with the system flood gun that provides low energy electrons and low energy argon ions from a single source. The hemispherical analyzer was operated in the constant energy mode with survey scan pass energies of 200 eV to measure the whole energy band, and 50 eV in a narrow scan to selectively measure particular elements. An estimation of the intensities was done after a calculation of each peak integral. S-shaped background subtraction and fitting of the experimental curve to a combination of a Lorentzian (30%) and Gaussian (70%) lines was done. Binding energies, referenced to the C1s line at 284.6 eV, had an accuracy of ± 0.1 eV.

The thermogravimetric analysis (TGA) of the catalyst was performed using Metter-Toledo apparatus. The change of sample mass was registered both in flowing air at a temperature increasing from 25 °C to 900 °C with a 10 °C/min heating rate.

Scanning electron microscopy (SEM) analyses were carried out on a Jeol JSM-6610LVnx integrated with energy dispersive detector (EDS; Oxford Aztec Energy) unit with accelerating voltage ranging from 300 V to 300 kV.

4. Conclusions

A series of UiO-66(Ce) metal-organic frameworks were obtained by using three methods: static hydrothermal, hydrothermal under continuous stirring and (for the first time) sonochemical. The obtained samples were characterized for their composition, structure, morphology, textural properties, thermal stability, presence of structural defects, and \( \text{CO}_2 \) adsorption capacity.

It was found that the synthesis method affected the physicochemical properties of UiO-66(Ce). The agitation of the reaction mixture during synthesis (stirring or sonication) facilitated the formation of more developed surfaces and led to the formation of smaller crystallites of UiO-66(Ce) than when the synthesis was carried out in static conditions. All samples contained structural defects (coming from missing H₂BDC linker) that resulted in the presence of a lower valent cerium. It was found that the contribution of Ce³⁺ was linked to the concentration of linker vacancies (more defects = more Ce³⁺). However, the charge on the defective Ce cluster was partly compensated, either by H₂O or DMF. As was determined by TGA, the coordination of DMF to Ce clusters was enhanced when synthesis was carried out under stirring or ultrasounds. The \( \text{CO}_2 \) sorption capacity at 273 K was the lowest for samples obtained by the sonochemical synthesis, and it increased with Ce³⁺ concentration. The \( \text{CO}_2 \) sorption on UiO-66(Ce) was found to take place in reversible as well as irreversible ways. The synthesis method significantly influenced the contribution of a particular type of adsorption; however, the important factor was not the concentration of Ce³⁺ ions (sites for strong \( \text{CO}_2 \) adsorption) but the termination of the surface sample with Ce clusters. High contribution of irreversible \( \text{CO}_2 \) adsorption was noticed for UiO-66(Ce) obtained using the sonochemical method. This method led to MOF having the smallest particles, which, in addition, were terminated with Ce clusters; therefore, the amount of Ce³⁺ accessible for strong \( \text{CO}_2 \) adsorption was higher than in the UiO-66(Ce) samples obtained by other methods and terminated, both with metal clusters and linker molecules.
In this work, we have shown that all of the used methods led to UiO-66(Ce) frameworks containing redox-active cerium atoms in the nodes (i.e., Ce⁴⁺ and Ce³⁺ cations), providing both Bronsted and Lewis acid centers, whose presence is important for MOF's performance in a given reaction. By using different synthesis methods, we can cause an impact on the character of active sites, textural properties, or even surface chemistry of MOF. Considering unique properties of cerium and excellent performance in many reactions, its introduction to metal nodes in Zr-based MOFs is an interesting area of research. The syntheses presented in this work were conducted on a scale ten times higher than those already reported [6,9,12,41] and will be improved in order to enhance UiO-66(Ce) properties, principally the size of the surface area. Moreover, the obtained UiO-66(Ce) will be tested in CO₂ hydrogenation to methanol.

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