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Synthesis, Crystallography, and Physical Properties**

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Temperature-Dependent Supramolecular Isomerism of Lutetium-Aminoterephthalate Metal-Organic Frameworks: Synthesis, Crystallography and Physical Properties.

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KEYWORDS: Lutetium, metal-organic framework, temperature-dependent supramolecular isomerism, gas adsorption, luminescence

ABSTRACT

Three supramolecular isomers of lutetium metal-organic framework, $\{\text{Lu}_2(\text{H}_2\text{O})_4(\text{ATA})_3 \cdot 4\text{H}_2\text{O}\}_n$ (**Lu-ATA@RT**), $\{\text{Lu}_2(\text{H}_2\text{O})_2(\text{C}_3\text{H}_7\text{NO})_2(\text{ATA})_3\}_n$ (**Lu-ATA@100**) and $\{\text{Lu}_2(\text{C}_3\text{H}_7\text{NO})(\text{ATA})_3\}_n$ (**Lu-ATA@150**), have been obtained from the reaction of $\text{Lu}(\text{NO}_3)_3 \cdot 6\text{H}_2\text{O}$ with 2-aminoterephthalic acid (ATA) at different temperatures. The resulting structures of Lu-ATA MOFs depend on the temperature applied during the synthesis, revealing a temperature-susceptible supramolecular isomerism. Single-crystal X-ray diffraction analyses suggest that new compounds with formula $\{\text{Lu}_2(\text{S})_x(\text{ATA})_3\}_n$ (S = solvent: H_2O , DMF) display different three-dimensional architectures which consist on dinuclear lutetium building units. The supramolecular isomer **Lu-ATA@RT**, formed at room temperature, has a pcu-net topology while its double interpenetrated analogue, **Lu-ATA@100**, assembles at 100 °C under hydrothermal conditions. Hydrothermal synthesis at 150 °C affords formation of the dense **Lu-ATA@150** cage-like framework displaying a new hexagonal-packed net topology. All Lu-ATA isomeric phases are porous and display different gas-uptake behaviour towards carbon dioxide as a function of polymeric network arrangement. The luminescent properties of Lu-ATA frameworks in solid state as well as in suspension in the presence of different solvents reveal a solvent dependent emission.

Introduction

During the past decades, the research field focused on porous materials underwent a revolution with the appearance of metal-organic frameworks (MOFs).¹⁻³ These porous solids constitute a subclass of three-dimensional (3D) coordination polymers formed by combination of metal ions or metal-oxygen clusters with organic ligands. These highly crystalline microporous materials have received enormous attention due to their manifold properties, with application in many different fields,⁴ from gas adsorption and separation to catalysis,^{5,6} sensing and medicine.^{7,8}

In general, practical applications of MOFs are directly related to their structural features.⁹⁻¹² Therefore, the rational design of MOFs' structures at molecular as well as nano- or macro-levels is still one of the main activities in the topic.^{13,14} Polymorphism or supramolecular isomerism is becoming an increasingly important subject in the field of crystal engineering, as it may facilitate the design of porous polymeric materials with targeted physical properties.¹⁵⁻¹⁷ In this context, the unique properties of MOF polymorphs, displaying both different crystal structures and identical molecular composition, allow for the derivation of structure-property relationships solely based on structural parameters or topology of the polymeric network.¹⁸ There are various key factors governing the formation of MOFs' polymorphic forms, including reaction time, temperature, concentration of precursors, pH, solvent, modulators, and so on. For example, Shi *et al.* report two polymorphic frameworks consisted of Eu^{III}, 2,5-furandicarboxylic and oxalic acids, which formation is determined by pH of reaction solution.¹⁹ In addition, the nature of the solvent is also an important factor that influences on the topology of resulted polymeric structure.²⁰ Therefore, several polymorphic forms of Pb^{II} and Cu^{II}-based MOFs were successfully obtained using different solvents or their mixtures.^{21,22} Similarly, the polymorphism of 3D MOFs was found to be instigated by modulators or structure directing agents.^{23,24} As

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example, Zhu *et al.* isolated two polymorphic forms of augmented “Cu₂-paddlewheel”-based MOFs with identical compositions but different **tbo** and **pto** topologies which are strongly correlated with the nature of the structure directing modulator used for the synthesis.²⁵ Recently, Friščić *et al.* have highlighted the importance of *in-situ* solid-state routes for the monitoring of MOF’s polymorphs formation.²⁶

Although numerous individual factors²⁷ or their conjunction²⁸ affect the occurrence of polymorphic forms, temperature is the one of the most important variables.²⁹ In particular, the use of temperature enables to take advantages of entropic differences between alternative products.³⁰ As illustrated by several studies,³¹⁻³⁴ different polymorphic phases of MOFs can be obtained from a single reaction mixture held at different temperatures. Among them, the experimental and theoretical studies carried out on series of polymorphic La-based MOFs reveal that the kinetic metastable and thermodynamic phases could be isolated as pure products over certain temperature ranges.³⁵ Additionally, the temperature-dependent ground states in the energy landscape for Zn^{II}-imidazole polymorphs³⁶ were also discussed recently.³⁷ The study of polymorphism is not only important in producing novel materials with target properties but may also be helpful in developing a fundamental understanding of the factors influencing crystal growth, such as reaction temperature.

Despite the numerous MOFs reported in the last decades, the number of lutetium based MOFs is very scarce.^{38, 39}

In this work, we report the syntheses, crystal structures, adsorption and fluorescence properties of three novel lutetium(III) 2-aminoterephthalate MOFs, namely **Lu-ATA@RT**, **Lu-ATA@100** and **Lu-ATA@150**. These three compounds are supramolecular isomeric forms of 3D lutetium(III) 2-aminoterephthalate framework, {Lu₂(S)_x(ATA)₃}_n (S = solvent: H₂O, DMF; ATA

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3 = 2-aminoterephthalic acid), which were obtained from identical starting materials in equal metal-
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5 to-ligand ratios under different temperatures.
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8 9 **Experimental Section**

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12 **Materials and characterization methods.** All starting materials were of analytical grade,
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14 obtained from Sigma-Aldrich and used without further purification.
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17 **Syntheses.** The synthetic condition for **Lu-ATA@RT** (room temperature supramolecular
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19 isomer) was previously optimized (see Supporting Information). 0.1 g (0.26 mmol) of
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21 $\text{Lu}(\text{NO}_3)_3 \cdot 6\text{H}_2\text{O}$ was dissolved in 10 mL H_2O and mixed with an aqueous solution (5 mL)
22
23 containing 0.14 g (0.77 mmol) of 2-aminoterephthalic acid (ATA) and 0.04 g (1.54 mmol) of
24
25 LiOH under magnetic stirring. The resulting solution was allowed to complete the reaction for 1
26
27 hour at room temperature upon continuous stirring. The formed pale-yellow precipitate was
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29 separated by vacuum filtration, washed several times with distilled water and air dried. Yield,
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31 0.14 g (99% based on Lu).
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37 For **Lu-ATA@100** (supramolecular isomer formed at 100 °C) synthesis, a mixture containing
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39 0.1 g (0.26 mmol) of $\text{Lu}(\text{NO}_3)_3 \cdot 6\text{H}_2\text{O}$, 0.05 g (0.3 mmol) of 2-aminoterephthalic acid (ATA), 2
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41 mL of DMF and 8 mL of H_2O was placed after dissolution in a Teflon-lined autoclave and
42
43 heated for 5 h at 100 °C in an oven under static conditions. The yellow precipitate was separated
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45 by vacuum filtration, washed several times with distilled water and air dried. Yield, 0.12 g
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47 (93% based on Lu).
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52 The synthesis of **Lu-ATA@150** polymorph was realized using the same reagent amounts and
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54 procedures as for **Lu-ATA@100**, but the reaction was performed under hydrothermal conditions
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56 at 150 °C for 5 h yielding 0.10 g yellow-brown powdered product (90 % yield based on Lu). The
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3 solid was separated by vacuum filtration, washed several times with distilled water and air
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5 dried.

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8 **Single crystal growth.** Single crystals of **Lu-ATA@RT** were obtained by crystallization in a
9
10 tetremethoxysilane (TMOS) gel containing (0.77 mmol) of 2-aminoterephthalic acid (ATA) and
11
12 0.04 g (1.54 mmol) of LiOH. When the gel is formed, the aqueous solution of $\text{Lu}(\text{NO}_3)_3 \cdot 6\text{H}_2\text{O}$
13
14 was placed as upper layer. After one week, prismatic yellow crystals of **Lu-ATA@RT** were
15
16 formed in gel phase (Figure S1a, Supporting Information).
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21 **Scanning Electron Microscopy (SEM)** was used to study the crystal size and morphology of the
22
23 Lu-MOF polymorphs. SEM micrographs were acquired in a Philips XL20 (15–30 kV)
24
25 microscope after sputtering the sample with a conductive Au layer.
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28 **Thermogravimetric analysis.** Mettler-Toledo TGA/SDTA851e was used for the thermal
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30 analyses in air dynamic atmosphere at heating rate of 10 °C/min in the range of temperatures
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32 from 30 °C to 1000 °C (blank runs were performed).
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35 **CO₂ adsorption.** Low-pressure adsorption isotherms of CO₂ (purity of 99.995%) were measured
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37 at 273 K in a device built by Bruker based on the volumetric technique. High-pressure adsorption
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39 isotherms of CO₂ (purity of 99.995%) were determined using the volumetric technique with an
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41 apparatus from BEL Japan (Belsorp HP). Around 0.2 g of the samples was placed in the sample
42
43 container. Before every measurement, the adsorbent was pre-treated by increasing the
44
45 temperature to 473 K at a rate of 10 K/min under vacuum and maintaining the temperature for 2
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47 hours. The measurements for carbon dioxide adsorption at high pressure were carried out at 273
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49 K for the complete series of Lu-MOF supramolecular isomers.
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52 **Diffuse reflectance** (BaSO_4 as a white standard) and **absorption UV/Vis** spectra were collected
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54 using a Perkin–Elmer Lambda 900 spectrophotometer equipped with an integrating sphere
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3 (“Labsphere”). Steady-state emission spectra were acquired using the photospectrometer Quanta
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5 Master QM1 of Photon Technology International (PTI) provided with a 100 W Xenon lamp, a
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7 double excitation monochromator (model PTI 121 A), a sample box and an emission
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9 monochromator (model PTI 101 A) and a photon counting detector PMT (model 710).
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13 **X-ray structure determination.** The single crystal X-ray diffraction data for **Lu-ATA@RT**,
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15 **Lu-ATA@100** and **Lu-ATA@150** were collected at 293 K on a an Oxford-Gemini X-ray
16
17 diffractometer equipped with graphite-monochromatic Cu-K α ($\lambda = 1.5418 \text{ \AA}$) and Mo-K α ($\lambda =$
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19 0.7107 \AA) radiation. The CrysAlisPro program was used for cell refinement and data reduction.
20
21 Images were collected at a 55 mm fixed crystal-detector distance, using the oscillation method,
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23 with 1° oscillation and variable exposure time per image. All the structures were solved by direct
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25 methods using the program SIR92⁴⁰ and refined by full-matrix least-squares techniques against
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27 F^2 using the SHELXTL-97⁴¹ crystallographic software package. All non-hydrogen atoms were
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29 found from the difference Fourier maps and refined anisotropically, whereas the hydrogen atoms
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31 of the organic molecules were placed by geometrical considerations and were added to the
32
33 structure factor calculations. In case of **Lu-ATA@150** structure, nitrogen atoms of amino-groups
34
35 are refined isotropically. Moreover, terminal amino-groups of 2-aminoterephthalic ligand are
36
37 disordered with occupancy ratio near to 0.5. For compounds **Lu-ATA@RT**, **Lu-ATA@100** and
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39 **Lu-ATA@150**, the high residual peaks and holes are observed in the vicinity of Lu³⁺, which are
40
41 normal for compounds containing heavy atoms. Absorption corrections were applied using
42
43 XABS2 program.⁴² The crystallographic data and structure refinement parameters for **Lu-**
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45 **ATA@RT**, **Lu-ATA@100** and **Lu-ATA@150** are summarized in the Table 1.
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55 **Table 1.** Crystallographic parameters and refinement data for **Lu-ATA@RT**, **Lu-ATA@100**
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57 and **Lu-ATA@150** supramolecular isomers.
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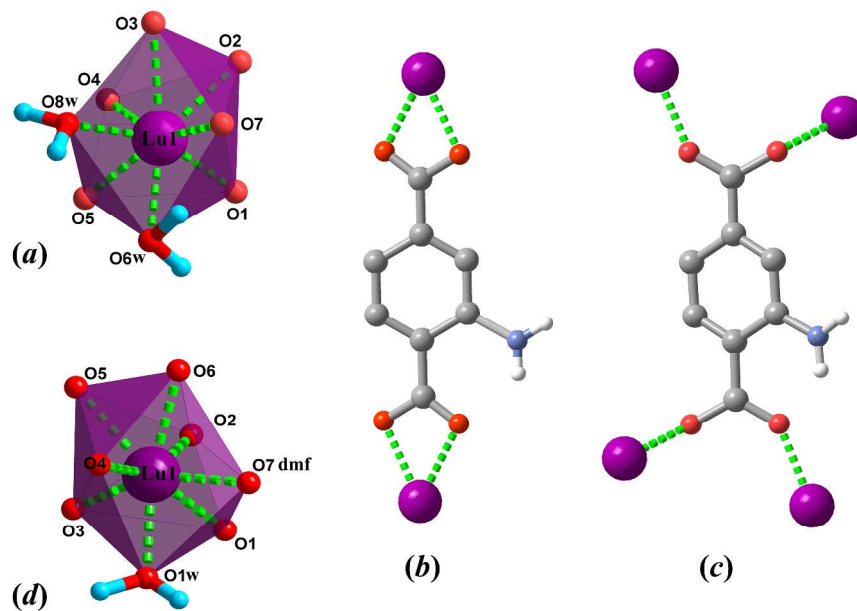
Compound	Lu-ATA@RT	Lu-ATA@100	Lu-ATA@150
Formula	C ₂₄ H ₃₄ Lu ₂ N ₆ O ₂₀	C ₃₀ H ₃₆ Lu ₂ N ₈ O ₁₆	C ₂₇ H ₂₅ Lu ₂ N ₄ O ₁₃
FW /g·mol ⁻¹	1076.5	1114.6	963.44
Temperature /K	296	296	300
Wavelength	CuKα (1.54180 Å)	MoKα (0.71073 Å)	MoKα (0.71073 Å)
Crystal system	triclinic	triclinic	monoclinic
Space group	<i>P</i> -1	<i>P</i> -1	<i>P</i> 2 ₁ / <i>a</i>
Unit cell dimensions			
<i>a</i> (Å)	10.0669(12)	8.1181(3)	20.5758(8)
<i>b</i> (Å)	10.3562(17)	10.2300(5)	16.6308(4)
<i>c</i> (Å)	10.538(2)	11.7103(5)	20.5719(8)
α (°)	103.158(17)	114.980(4)	90
β (°)	115.524(17)	99.906(3)	119.801(5)
γ (°)	92.546(12)	98.258(3)	90
Cell volume/ Å ³	952.6(3)	842.54(7)	6108.6(5)
<i>Z</i>	2	2	8
Calc. Density/g·cm ⁻³	1.87	2.19	2.08
μ /mm ⁻¹	10.49	5.90	6.50
F(000)	522	542	3624
Crystal size (mm ³)	0.06 × 0.04 × 0.02	0.39 × 0.24 × 0.20	0.13 × 0.06 × 0.06
2θ range data collection/°	4.9 ÷ 61.2	3.4 ÷ 31.5	3.4 ÷ 28.4
Index ranges	-12 ≤ <i>h</i> ≤ 10; -12 ≤ <i>k</i> ≤ 12; 0 ≤ <i>l</i> ≤ 12	-11 ≤ <i>h</i> ≤ 11; -15 ≤ <i>k</i> ≤ 13; 0 ≤ <i>l</i> ≤ 16	-29 ≤ <i>h</i> ≤ 28; -19 ≤ <i>k</i> ≤ 24; -28 ≤ <i>l</i> ≤ 27
Reflection collected	3497	5155	17095
Independent reflections	3497 [R _{int} = 0.11]	5155 [R _{int} = 0.04]	17095 [R _{int} = 0.08]
T _{max} , T _{min}	0.813, 0.614	0.247, 0.169	0.985, 0.554
Data/restraints/parameters	3497/36/180	5155/34/260	17095/13/810
Goodness-of-fit on <i>F</i> ²	1.01	1.08	1.02
R ₁ , wR ₂	0.090, 0.257	0.032, 0.082	0.058, 0.103
Δρ _{max} , Δρ _{min} / e Å ⁻³	2.29, -1.67	1.58, -2.40	1.59, -1.66
CCDC #	1449678	1449680	1449688

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3 X-ray powder diffraction patterns were collected at room temperature on a Bruker Advance D8
4 diffractometer using Co-K α ($\lambda = 1.78897 \text{ \AA}$), equipped with a LynxEye detector. The powder
5 diffractograms for the synthesized compound were recorded in the $5 - 95^\circ$ range of 2θ with a
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10 step of 0.019° and a counting time of 0.4 s per step.

11 12 13 14 15 16 **Results and Discussion**

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19 Three supramolecular isomeric forms of Lu-ATA (ATA = 2-aminoterephthalic acid)
20 coordination frameworks were obtained through the reaction of lutetium(III) nitrate with ATA
21 linker at different temperatures using similar intervals of precursors concentration. Although
22 they share the same compositional characteristics, $\{\text{Lu}_2(S)_x(\text{C}_8\text{H}_6\text{NO}_4)_3\}_n$ ($S = \text{solvent: H}_2\text{O}$ or
23 dimethylformamide, DMF), the crystal structures of these new metal-organic networks reveal
24 temperature-dependent supramolecular isomerism. Using optimized temperature ranges,
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supramolecular isomers **Lu-ATA@100** and **Lu-ATA@150** were obtained hydrothermally as
pure, highly crystalline phases (Figure S2a-b) composed of polyhedra-shaped crystals with sizes
ranging from 20 to 500 μm (Figure S4). Consequently, the quality of the crystals isolated
hydrothermally allows direct structure solution using single-crystal X-ray diffraction (SCXRD).
In contrast to above mentioned compounds, the supramolecular isomeric phase **Lu-ATA@RT**,
prepared at room temperature (Figure S3 and S4a,d), is composed of smaller crystals (1–3 μm).
The crystal growth in gel media is very useful and widely applied technique for obtaining high
quality MOFs' single crystals.⁴³⁻⁴⁸ In order to carry out detailed structural analysis for **Lu-**
ATA@RT, the crystal growth in TMOS gel media has been applied. After one week in gel-
phase, the yellow dendritically-organized crystals were formed and used for further SCXRD
characterization (Figure S1a).

Single crystal X-ray diffraction analysis of **Lu-ATA@RT** reveals that this coordination polymer of formula $\{\text{Lu}_2(\text{H}_2\text{O})_4(\text{C}_8\text{H}_5\text{NO}_4)_3 \cdot 4\text{H}_2\text{O}\}_n$ crystallizes in a triclinic space group $P-1$ with unit cell parameters summarized in Table 1. The asymmetric unit of **Lu-ATA@RT** contains one lutetium atom, one and half aminoterephthalic acid (ATA), two coordinated and two crystallization water molecules (Figure S6).



Scheme 1. Coordination environment of Lu(III) ion in (a) **Lu-ATA@RT** and (d) **Lu-ATA@100** polymorphs, and coordination modes of the 2-aminoterephthalic ligand (b) $\mu-(\eta_2:\eta_2)$ and (c) $\mu_4-(\eta_1:\eta_1:\eta_1:\eta_1)$ in both frameworks. Hydrogen atoms of benzene ring were omitted for clarity. Lutetium atoms are depicted as violet spheres.

As shown in Scheme 1a, the lutetium center adopts a distorted bicapped trigonal prismatic geometry and is surrounded by eight oxygen atoms where six of them belong to four carboxylic groups of ATA ligands and two to coordinated water molecules. In the structure of **Lu-ATA@RT**, the 2-aminoterephthalic acid (ATA) exhibits two different tetradentate coordination modes (Scheme 1b-c): the mode (b) $\mu-(\eta_2:\eta_2)$, where two oxygen atoms of one carboxylic group chelate one lutetium ion; the mode (c) $\mu_4-(\eta_1:\eta_1:\eta_1:\eta_1)$, where the aminoterephthalate dianion coordinates four lutetium centers by each available oxygen atoms proceeding from two

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3 carboxylate groups. Consequently, ATA ligands in μ -($\eta_2 : \eta_2$) and μ_4 -($\eta_1 : \eta_1 : \eta_1 : \eta_1$)
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5 coordination modes link lutetium ions in alternately manner to form one-dimensional (1D)
6
7 chains of Lu polyhedra extended along the *b*-crystallographic direction and arranging into the
8
9 two-dimensional (2D) substructure of the **Lu-ATA@RT** framework (Figure S7d). Additionally,
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11 the ATA ligands behaving only in μ -($\eta_2 : \eta_2$) mode link lutetium ions in parallel mode and define
12
13 a second 2D substructure of **Lu-ATA@RT** (Figure S7e). Such 2D-substructures are further
14
15 associated together through the angular coordination to Lu polyhedra chain resulting in a 3D
16
17 framework with large rhombohedral channels running along *a*- and *b*-crystallographic axis
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21 (Figure 1a-b).

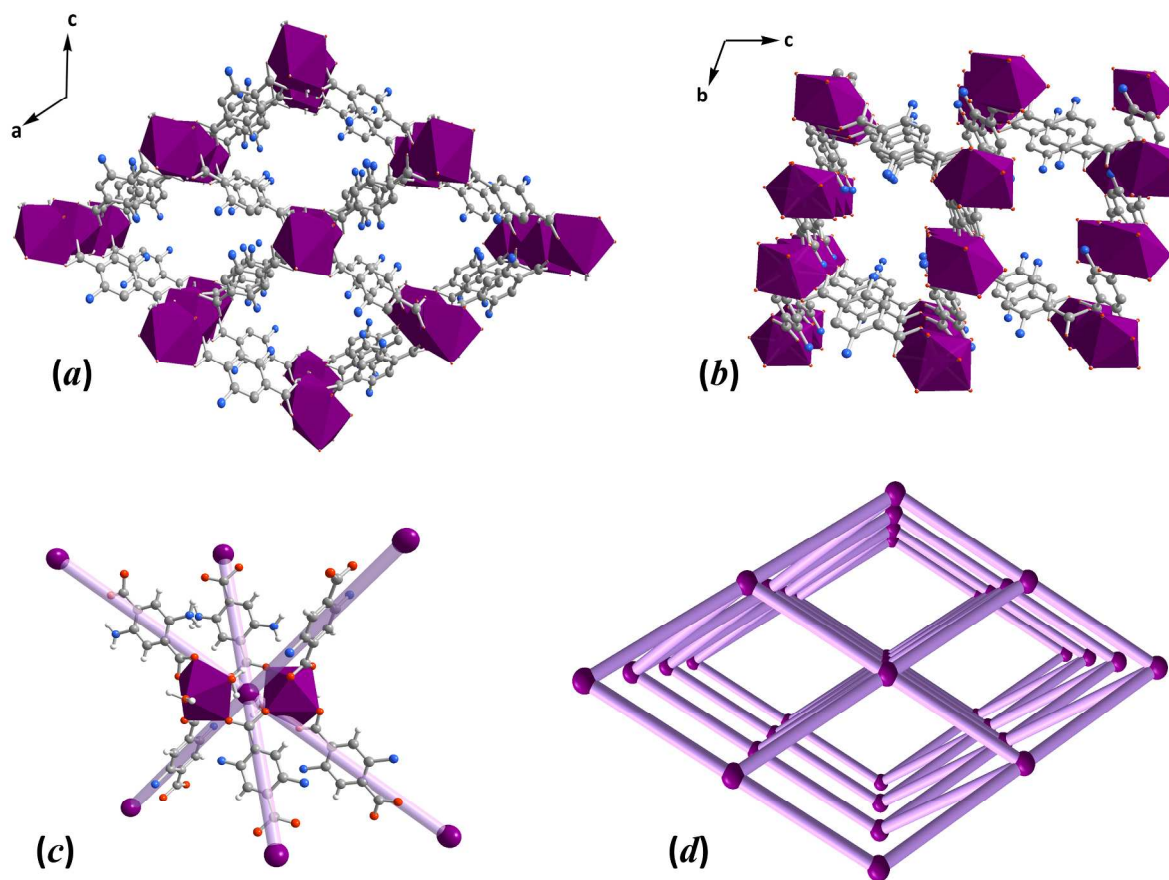


Figure 1. Representation of 3D polymeric structure of **Lu-ATA@RT** that possesses large rhombohedral channels: views (a) along *b*-axis and (b) along *a*-axis. (c) The dinuclear lutetium

1
2 motif acted as SBU with linear ATA connectors, and (d) the topological representation of **pcu**
3 simple net of **Lu-ATA@RT**.
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5
6 According to the void analysis performed with PLATON program,⁴⁹ using a probe molecule
7 radius of 1.4 Å and a grid interval of 0.2 Å, the extended 3D structure of **Lu-ATA@RT**
8 framework possess 150 Å³ (15.8 %) of free void space. It is worth noting that the structure of
9
10 **Lu-ATA@RT** supramolecular isomer is formed by a dinuclear lutetium motif acting as
11 elementary structural building unit (SBU) and can be defined as a simple node during the
12 topological analysis of the framework (Figure 1c). Thus, according to the analysis performed by
13 TOPOSPro,⁵⁰ the resulting 3D arrangement corresponds to a 6-connected uninodal net of **pcu**
14 alpha-Po primitive cubic type with (4¹².6³) point (Schläfli) symbol (Figure 1d), where the
15 dinuclear lutetium motif act as a nodes and ATA ligands as a bidentate rods.
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29 Compound **Lu-ATA@100**, with formula {Lu₂(H₂O)₂(C₃H₇NO)₂(C₈H₅NO₄)₃}_n, crystallizes in a
30 triclinic space group *P*-1 with unit cell parameters summarized in Table 1. The asymmetric unit
31 of **Lu-ATA@100** is composed of one lutetium atom, one and a half aminoterephthalic acid
32 (ATA), one coordinated water and one coordinated DMF molecules (Figure S8). The
33 coordination environment of lutetium ion in **Lu-ATA@100** is formed by eight oxygen atoms,
34 where six of them (O1–O6) belong to three carboxylic groups of ATA ligands, one (O1w) to
35 coordinated water and one (O7dmf) to coordinated DMF molecules, and adopts a distorted
36 bicapped trigonal prismatic geometry (Scheme 1d). Notably, in **Lu-ATA@100**, the 2-
37 aminoterephthalic acid (ATA) structure exhibits same (b) μ -($\eta_2 : \eta_2$) and (c) μ_4 -($\eta_1 : \eta_1 : \eta_1 : \eta_1$)
38 coordination modes (Scheme 1b-c) as in **Lu-ATA@RT** structure. However, the ATA linker
39 behaving in μ_4 -mode bridges two nearest metal centers, forming a dimeric lutetium unit as
40 shown in Figure 2a.
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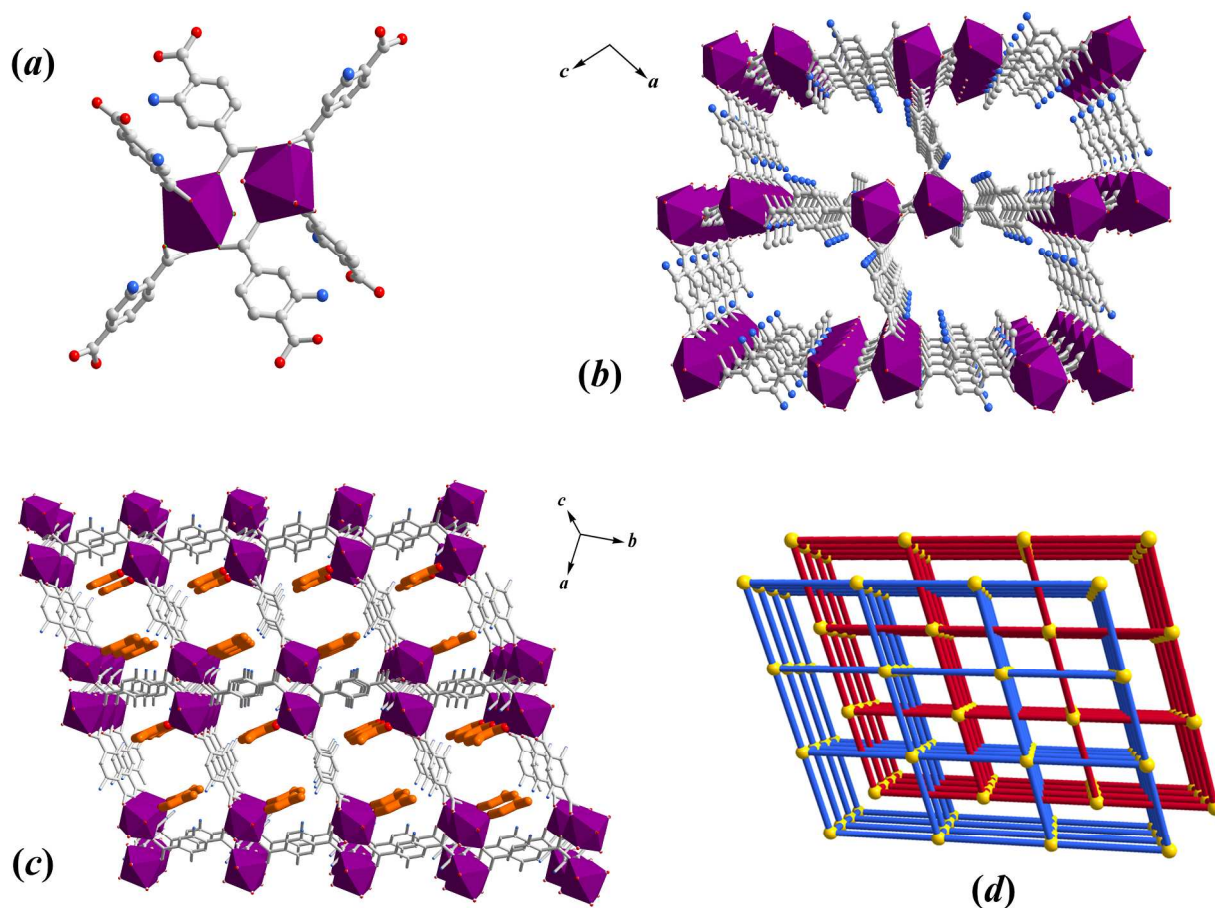
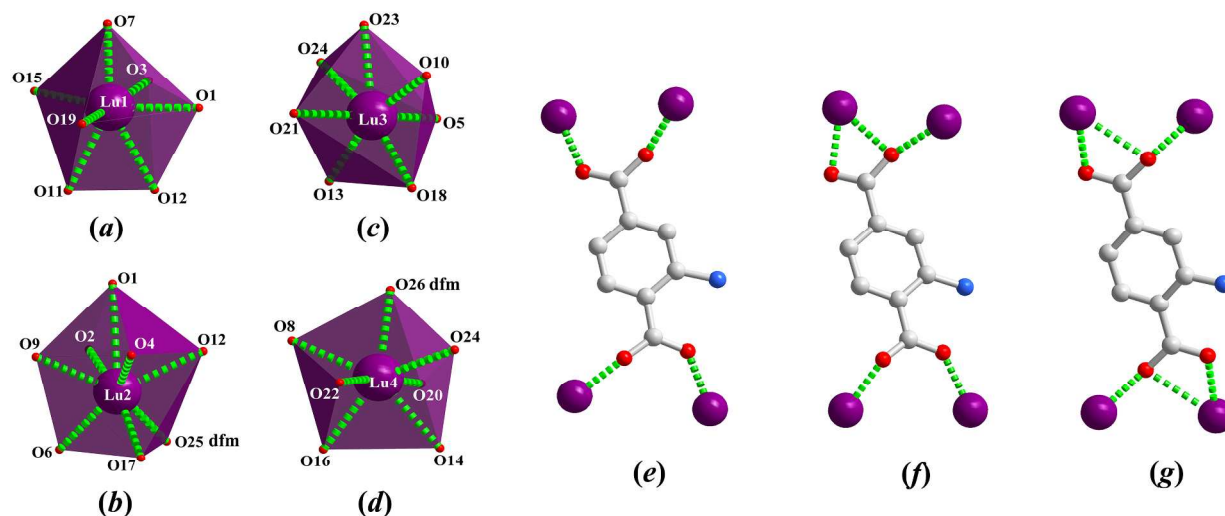


Figure 2. (a) Dimeric lutetium structural unit that forms the 3D extended framework of **Lu-ATA@100** with (b) large rhombic channels running along *b*-axis. (c) View along [111] direction of **Lu-ATA@100** showing the coordinated DMF molecules (orange units) pointed into the channels' of the frame. (d) The simplified representation of **Lu-ATA@100** network of *pcu* topological type with interpenetration.

Such secluded lutetium dimers are linked through μ_4 -bridging ATA linkers to give a 1D substructure (Figure S9). Furthermore, adjacent chains are further associated together *via* the linear ATA exhibiting μ -chelation, resulting in a 3D framework. As shown in Figure 2b-c, the polymeric network of **Lu-ATA@100** possesses open rhombohedral channels running along *b*-axis. Notably, the coordinated DMF molecules of dimeric lutetium units pointing into channels restrict the aperture of the framework in the [111] direction (Figure 2c). The void analysis performed by PLATON⁴⁹ using a probe molecule radius of 1.4 Å and a grid interval of 0.2 Å reveals that **Lu-ATA@100** framework not possess externally accessible surface volume due to

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3 interpenetration as has been proved below. According to the topological analysis, **Lu-ATA@100**
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5 framework is a uninodal six-connected net of **pcu** alpha-Po primitive cubic type expressed by
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7 $(4^{12}.6^3)$ point (Schläfli) symbol. Moreover, the framework structure consists of two independent
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9 simple nets of **pcu** topology with the interpenetration belonging to the class Ia,⁵¹ where each net
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11 are symmetry-equivalent and related both by translation vector [100] and inversion (Figure 1d).
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13 Although an increase of the densities of the obtained Lu-ATA supramolecular isomers with
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15 temperature was observed, **Lu-ATA@100** showed an out-of-trend result due to network
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17 interpenetration.
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21 The hydrothermal synthesis of **Lu-ATA@150** supramolecular isomer resulted in a dense
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23 hexagonally-packed framework structure. As revealed by single crystal X-ray diffraction, the
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25 **Lu-ATA@150** of formula $\{\text{Lu}_2(\text{C}_3\text{H}_7\text{NO})(\text{C}_8\text{H}_5\text{NO}_4)_3\}_n$ crystalizes in monoclinic space group
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27 *P21/a* with unit cell parameters given in Table 1. The asymmetric unit of **Lu-ATA@150**
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29 contains four crystallographically independent lutetium atoms, six aminoterephthalic acid (ATA)
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31 and two coordinated DMF molecules (Figure S10). Three lutetium ions (Lu1, Lu3 and Lu4)
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33 adopt a distorted monocapped trigonal prismatic coordination geometry (Scheme 2a-c,d) by
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35 participation of seven oxygen atoms: (a), (c) all of them from carboxylate groups of ATA ligands
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37 (Lu1 and Lu3 centers); (d) six from carboxylate groups of ATA ligands and one (O26dmf) from
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39 coordinated DMF molecule (Lu4 center).
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Scheme 2. Coordination environments of four Lu(III) ions in **Lu-ATA@150** with corresponding atom labeling scheme: (a) and (c)–(d) monocapped and (b) bicapped trigonal truncated prisms. The coordination modes of the 2-aminoterephthalic ligands: (e) $\mu_4-(\eta_1 : \eta_1 : \eta_1 : \eta_1)$, (f) $\mu_4-(\eta_1 : \eta_1 : \eta_1 : \eta_2)$, (g) $\mu_4-(\eta_1 : \eta_2 : \eta_1 : \eta_2)$ in **Lu-ATA@150**. Hydrogens were omitted for clarity and lutetium atoms are depicted as violet spheres.

The coordination environment of central lutetium atom Lu2 consists of a distorted bicapped trigonal prismatic geometry comprising eight oxygen atoms: seven of them are originated from carboxylate groups of ATA ligands and one (O25dmf) from a coordinated DMF molecule (Scheme 2b). In the structure of the **Lu-ATA@150** supramolecular isomer, the 2-aminoterephthalic acid (ATA) exhibits three different tetradentate coordination modes (Scheme 2e–g): the mode (e) $\mu_4-(\eta_1 : \eta_1 : \eta_1 : \eta_1)$, where the aminoterephthalate linker coordinates four lutetium centers by each available oxygen atoms from two carboxyl groups; the mode (f) $\mu_4-(\eta_1 : \eta_1 : \eta_1 : \eta_2)$, where three oxygen atoms of two carboxyl groups coordinate to three lutetium centers and chelate fourth one; the mode (g) $\mu_4-(\eta_1 : \eta_2 : \eta_1 : \eta_2)$, where each of carboxyl groups coordinates to one lutetium center and chelates the second one.

As seen in Figure 3a, the neighboring lutetium atoms are connected together through the carboxylate groups of ATA ligands (Scheme 2) in (e)–(g) coordination fashions to form an infinite chain of lutetium polyhedra running along crystallographic *b*-axis. Notably, the

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3 polymeric 1D chain consists of two types of dinuclear motifs (Figure 3b-c): (b) edge-sharing and
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5 (c) corner-sharing units. Furthermore, the 1D skeleton of lutetium centers are extended by ATA
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7 linkers resulting in hexagonally packed 3D supramolecular architecture. As shown in Figure 3d,
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9 the polymeric network of **Lu-ATA@150** is a cage-like structure with triangular windows which
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11 are partially obstructed by coordinated DMF molecules of 1D lutetium skeleton. The solvent-
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13 accessible volume of **Lu-ATA@150** calculated by PLATON⁴⁹ is 374.6 Å³ (a probe molecule
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15 radius of 1.4 Å and a grid interval of 0.2 Å were used) and corresponds to 6.1 % of the unit cell
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17 volume. Moreover, the polymeric network of **Lu-ATA@150** is four-nodal 3,4,4,9-connected net
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19 of new topological type with $(4.5^2)_2(4^4.5.6)(4^4.5^2)(4^8.5^{12}.6^{16})_2$ point (Schläfli) symbol (Figure
20
21 3e). In the simplification procedure, the dinuclear lutetium motifs act as nine-connected nodes
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23 while the ATA ligands as bidentate rods, three-connected and four connected nodes (Figure
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25 S11). Interestingly, due to the cage-like structure of **Lu-ATA@150**, the empty space of the
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27 network can be represented by tiling with [4 9 11 6] tile transitivity (Figure 3f).
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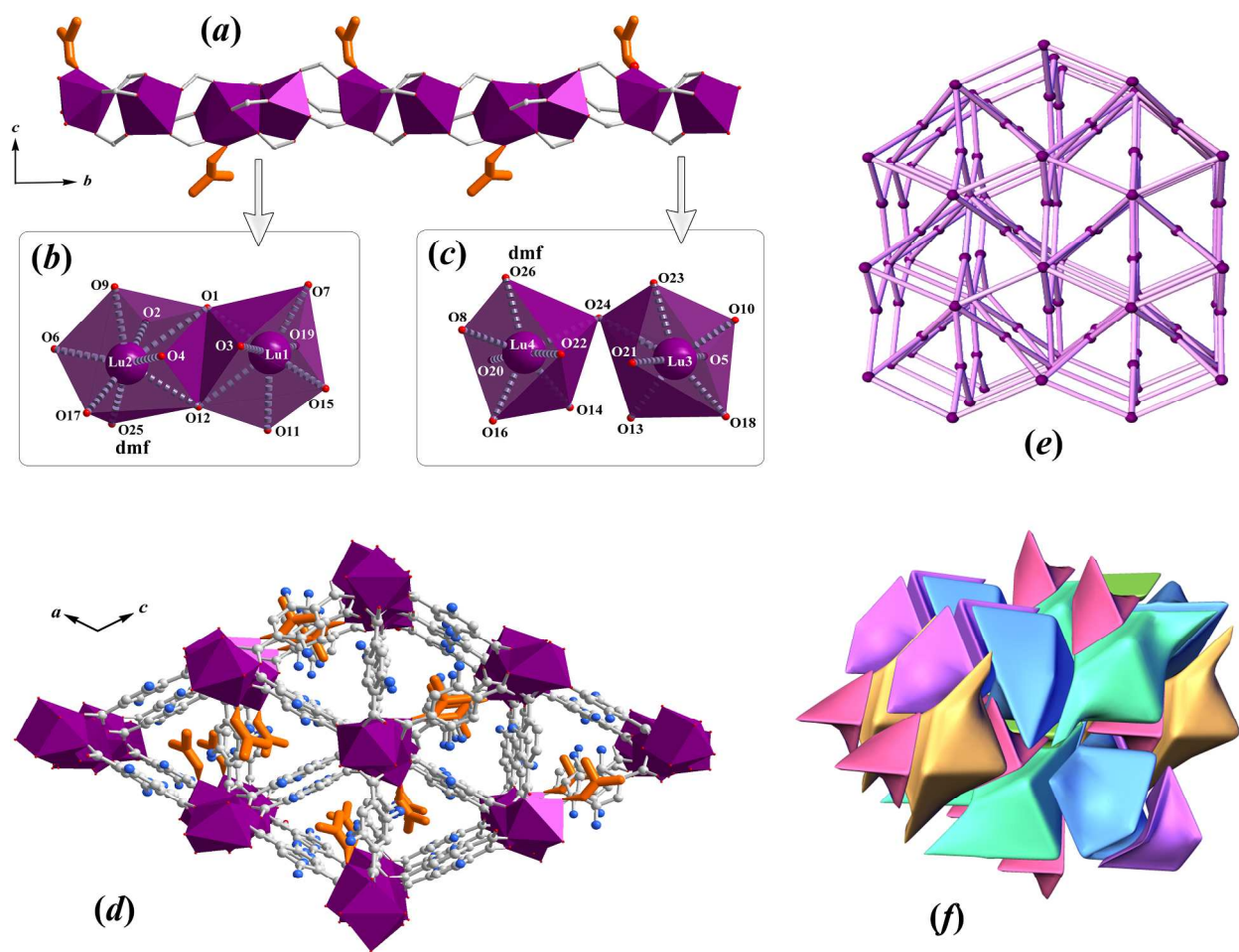


Figure 3. (a) One dimensional chain of lutetium polyhedra in **Lu-ATA@150** composed of two types of dinuclear: (b) edge-sharing and (c) corner-sharing units. Coordinated DMF molecules are drawn in orange. (d) View along *b*-axis to the 3D cage-like framework of **Lu-ATA@150** with triangular windows. (e) The simplified net of **Lu-ATA@150** of new hexagonal-packed topological type and (f) its tilled structure of [4 9 11 6] transitivity.

Thermal stability in air atmosphere of the three Lu-ATA supramolecular isomers was investigated and the thermal decomposition profiles are shown in Figure S12. The degradation of **Lu-ATA@RT** proceeds in two stages, as depicted in Figure S12a. The first mass loss between 30 and 120 °C with a total mass loss of 15.5 % (calculated 14.8 %), corresponds to the simultaneous release of four water molecules trapped in the pores and four coordinated to lutetium atoms, which is accompanied by an endothermic peak in the SDTA curve. Up to 300 °C, the resulted anhydrous compound remains stable until the second stage, which takes place in

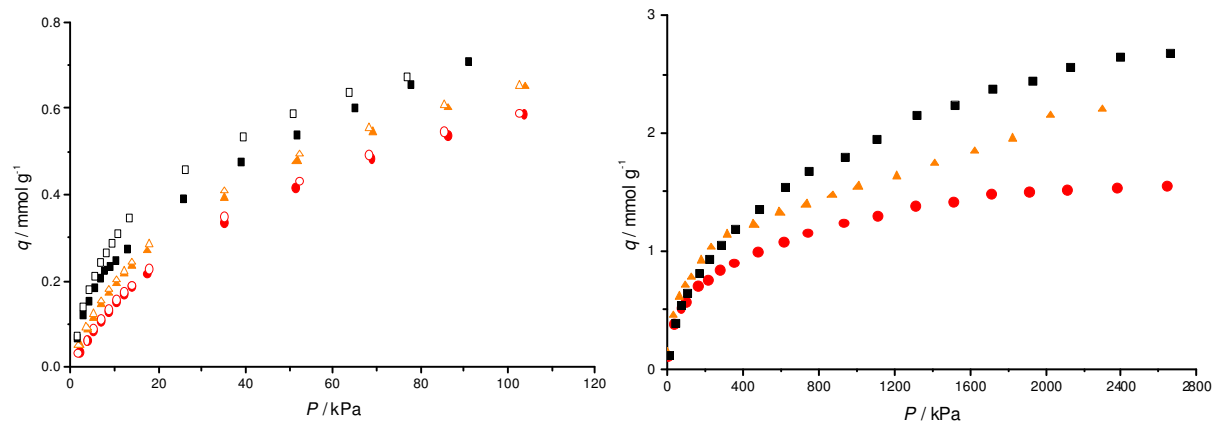
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3 the range 320–540 °C (continuous process) with the total mass loss of *ca.* 44.4 %, and
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5 corresponds to the progressive decomposition of the organic part of the framework, associated
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7 with exothermic peaks in SDTA graph. According to the mass of the remaining solid, the
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9 residual product of decomposition is Lu₂O₃
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13 In turn, the thermal decomposition profile of **Lu-ATA@100** shows a different behavior and is
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15 composed of two well-separated steps. As shown in Figure S12b, the first mass loss step occurs
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17 at higher temperature in the range 170–230 °C with the mass loss of 16.3 % and consists of two
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19 contributions which correspond to evacuation of two coordinated water and two DMF molecules
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21 (calculated 16.5 %). These processes are conformed by an endothermic peak in SDTA plot. The
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23 anhydrous framework of **Lu-ATA@100** remains stable up to 340 °C. The second decomposition
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25 stage takes place in the range 350–550 °C with the mass loss of *ca.* 61.2% and attributed to
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27 oxidative degradation of the organic part of the polymeric frame, associated with exothermic
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29 peaks in SDTA curve.
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37 Similarly, the degradation process for **Lu-ATA@150** supramolecular isomer proceeds in two
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39 stages (Figure S12c). The first mass loss between 140 and 230 °C with a mass loss of 7.6 %
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41 (calculated 7.3 %), corresponds to the evacuation of two DMF molecules coordinated to lutetium
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43 atoms, which is accompanied by an endothermic peak in SDTA curve. In air atmosphere, the
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45 anhydrous **Lu-ATA@150** is stable in the 240–320 °C temperature plateau, followed by a second
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47 degradation step, which takes place in the range 330–700 °C. The total mass loss of *ca.* 57.3 %
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49 corresponds to the decomposition of organic ligands and collapse of the structure (exothermic
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51 process).
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In order to explore any structural transformation that can take place upon activation of the Lu-ATA isomers and be related to the solvent removal, additional powder X-ray diffraction studies have been performed. As shown in Figures S13 and S14, PXRD patterns of **Lu-ATA@RT** and **Lu-ATA@100** treated in vacuo at 200 °C show a loss of crystallinity without any notable structural transformations. In contrast, **Lu-ATA@150** exhibits irreversible structural change upon activation at 200 °C in vacuo, as illustrated in Figure S15. Powder X-ray diffraction patterns of **Lu-ATA@150** desolvated form has been indexed in the triclinic crystal system with the following unit cell dimensions: $a = 10.56(2) \text{ \AA}$, $b = 10.33(2) \text{ \AA}$, $c = 7.48(1) \text{ \AA}$, $\alpha = 95.6(1)^\circ$, $\beta = 109.2(1)^\circ$, $\gamma = 113.3(2)^\circ$, $V = 683(2) \text{ \AA}^3$. The results reveal that the unit cell contracts after desolvation process lowering the crystal symmetry to $P-1$.

The porosity of the three supramolecular isomers could not be studied with nitrogen physisorption due to the very narrow porosity that these materials exhibit. That fact, in combination with the big crystal size led to diffusional problems and made carbon dioxide adsorption a suitable choice for their textural characterization as it has been proposed several times in literature.⁵² The adsorption of carbon dioxide on the series of materials was analyzed at 273 K at low and high pressure.



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3 **Figure 4.** Carbon dioxide adsorption at 273 K at low (left) and high pressure (right) in **Lu-**
4 **ATA@RT** (black squares), **Lu-ATA@100** (orange triangles) and **Lu-ATA@150** (red circles).
5 Full and empty symbols represent adsorption and desorption, respectively. The amount of CO₂
6 adsorbed is designated as q .
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10 At carbon dioxide pressure up to 100 kPa **Lu-ATA@RT**, **Lu-ATA@100** and **Lu-ATA@150**
11 adsorb a similar amount, around 0.7, 0.65 and 0.59 mmol/g respectively. It is important to notice
12 that the removal of coordinated DMF molecules, which restricts the accessibility to vacant pore
13 space in **Lu-ATA@100** and **Lu-ATA@150** frameworks, is essential to obtain adsorption of
14 carbon dioxide. When the pretreatment was carried out at 323 K the materials did not show any
15 uptake, in contrast the porosity was accessible if temperatures higher than 373 K were applied to
16 pretreat the samples. At higher pressures the same trend is confirmed, being **Lu-ATA@150** the
17 material with lower adsorption capacity with a maximum value of 1.56 mmol/g at 2636 kPa. **Lu-**
18 **ATA@100** adsorbs a higher amount reaching 2.2 mmol/g at 2299 kPa. The material with the
19 highest adsorption capacity for carbon dioxide of the supramolecular isomers series is **Lu-**
20 **ATA@150**, adsorbing a maximum of 2.7 mmol/g at 2656 kPa. The adsorption trend of the three
21 supramolecular isomers is in agreement with the obtained pore volumes from the
22 crystallographic information, exhibiting the three of them a very narrow porosity.
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41 The optical properties of the Lu-ATA MOFs series were analyzed by light absorption and
42 fluorescence spectroscopy. The three isomeric structures show fluorescence emission in solid
43 state as well as in suspension of acetonitrile and toluene (Table 2). Lu(III) has full f orbitals and
44 thus the typical lanthanide metal luminescence cannot occur. The band at longest wavelength in
45 the free linker, assigned to a highest occupied molecular orbital to lowest unoccupied molecular
46 orbital (HOMO-LUMO) transition with a $n \rightarrow \pi^*$ nature,^{53, 54} appears in the MOFs' spectra
47 without important shifts. This seems to indicate that the contribution of the metal orbitals to the
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frontier orbitals might be negligible and that this transition remains linker based in the MOFs.⁵⁴
⁵⁵ Nevertheless, computational studies are needed to gain insights on the location of HOMO and LUMO in each Lu-ATA system and thus to confirm if the bands around 370-380 nm arise from linker based or metal-to-linker charge-transfer (MLCT) transitions.⁵⁶⁻⁵⁹

Table 2. Absorption (λ_{abs}) and emission (λ_{em}) maxima along with derived Stokes shifts for studied supramolecular isomers **Lu-ATA@RT**, **Lu-ATA@100** and **Lu-ATA@150**.

	Compound	λ_{abs} (nm)	λ_{em} (nm)	Stokes shift (nm)
Activated powder	Lu-ATA@RT	377 (br)	456	79
	Lu-ATA@100	386 (br)	460	79
	Lu-ATA@150	384 (br)	456	70
Toluene	Lu-ATA@RT	378	448	70
	Lu-ATA@100	380	450	70
	Lu-ATA@150	385	451	66
Acetonitrile	Lu-ATA@RT	366	430	64
	Lu-ATA@100	379	440	61
	Lu-ATA@150	387	450	63
	Linker ester	367	428	61

(br) = broad signal

Of particular interest is the trend observed in both the absorption and the emission maxima of the different Lu-ATA supramolecular isomers in acetonitrile (Figure 5), which is not observed in the powder or in suspension of an unipolar solvent like toluene (Supporting Information S16-S21). The spectra exhibit a clear hypsochromic shift of 10 nm step following the order **Lu-ATA@RT** < **Lu-ATA@100** < **Lu-ATA@150**.

It is known that increasing the polarity of the solvent leads to blue shifts in $n\pi^*$ transitions.⁶⁰ In the ground state, the dipoles of the solvent are oriented with their positive endings towards the n

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3 orbital bearing the lone pair of electrons. Upon excitation the n orbital losses electron density
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5 (which is injected into the π^*) but the solvent dipoles do not switch in this timescale and thus a
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7 repulsive effect between the solvent molecules and the electron-poor nitrogen destabilizes the
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9 excited state.

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11 Thus a possible explanation for the different blue shift of each Lu-ATA MOF suspended in
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13 acetonitrile compared to the corresponding spectra in powder and in toluene could be that the
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15 polar solvent molecules interact with a different strength with each supramolecular isomer, thus
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17 leading to different degrees of destabilization of their excited state. That is, solvatochromism^{55, 61}
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19 might be different for each Lu-ATA structure. Nevertheless, MOFs can undergo several
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21 mechanisms that lead to optical changes⁶² and further studies would be required to unequivocally
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23 explain this trend.
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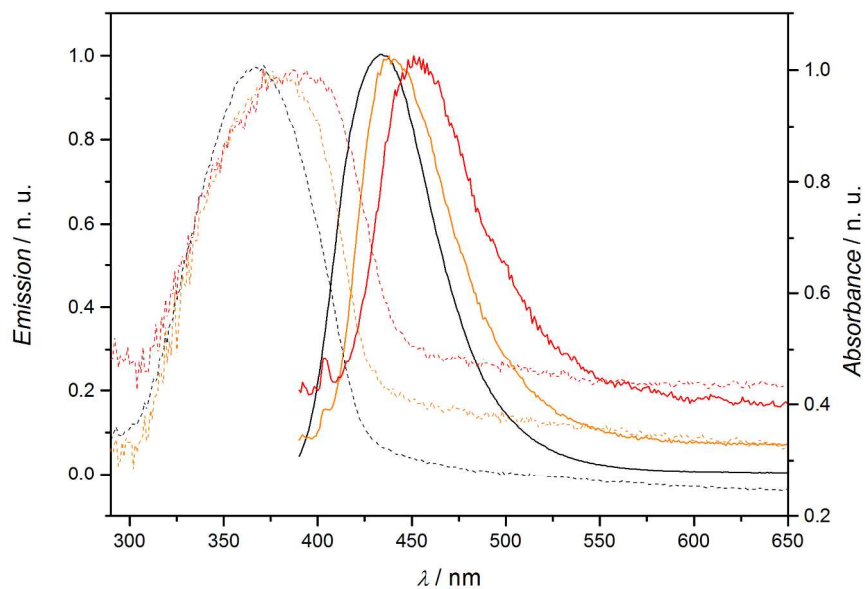


Figure 5. Absorption (dashed lines) and emission (solid line) spectra of **Lu-ATA@RT** (black), **Lu-ATA@100** (orange), and **Lu-ATA@150** in acetonitrile suspension ($5 \cdot 10^{-2}$ g/L). Bands are normalized for better visualization. The non-zero backgrounds results from light scattering.

Conclusions

Three supramolecular isomeric forms of lutetium-based MOF, **Lu-ATA@RT**, **Lu-ATA@100** and **Lu-ATA@150**, built by 2-aminoterephthalic linker were synthesized starting from the same precursors. The structure of the final isomeric form of Lu-ATA framework as well as its crystal packing is influenced by the temperature applied during the synthesis. Thus, the room temperature approach leads to formation of **Lu-ATA@RT** framework having **pcu** topology while the synthesis at 100 °C results in **Lu-ATA@100** forming an interpenetrated (class Ia) analogue of the **pcu**-net. A dense cage-like structure with a new topology is formed at 150 °C under hydrothermal conditions (**Lu-ATA@150**). Notably, the structural differences between three supramolecular isomers of Lu-ATA MOF expressed by general formula of $\{\text{Lu}_2(\text{S})_x(\text{ATA})_3\}_n$ are reflected on their absorption behaviour towards CO₂. High-pressure CO₂ absorption results reveal that the three Lu-ATA supramolecular isomers possess narrow porosity, the maximum capacity follows the trend **Lu-ATA@RT** > **Lu-ATA@100** > **Lu-ATA@150**, in agreement with the calculated pore volumes extracted from the crystallographic information. Lu-ATA series suspended in acetonitrile exhibited a blue shift of different magnitude in their absorption and emission bands compared to the spectra in toluene or in powder, which might be an indication of different solvation efficiency in the diverse structures.

ASSOCIATED CONTENT

Supporting Information. Optical and electron microscopy images for Lu-ATA polymorphs, additional structure images, PXRD of synthesized materials, TG/SDTA data plots, additional

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3 absorption and emission spectra. This material is available free of charge via the Internet at
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5 <http://pubs.acs.org>.
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9 The supplementary crystallographic data for this paper were deposited to Cambridge
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11 Crystallographic Data Centre with CCDC numbers of 1449678, 1449680 and 1449688. These
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13 data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing
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15 data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre,
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17 12, Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.
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32 **Author Contributions**

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34 ‡These authors contributed equally. The manuscript was written through contributions of all
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36 authors. All authors have given approval to the final version of the manuscript.
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ABBREVIATIONS

MOF, metal-organic framework; **ATA**, 2-aminoterephthalic acid; **DMF**, N,N'-dimethylformamide; **RT**, room temperature; **SEM**, scanning electron microscopy; **SCXRD**, single-crystal X-ray diffraction; **SBU**, structural building unit; **SDTA**, simultaneous difference thermal analysis; **HOMO**, highest occupied molecular orbital; **LUMO**, lowest unoccupied molecular orbital; **MLCT**, metal-to-linker charge-transfer; **CCDC**, Cambridge crystallographic data center.

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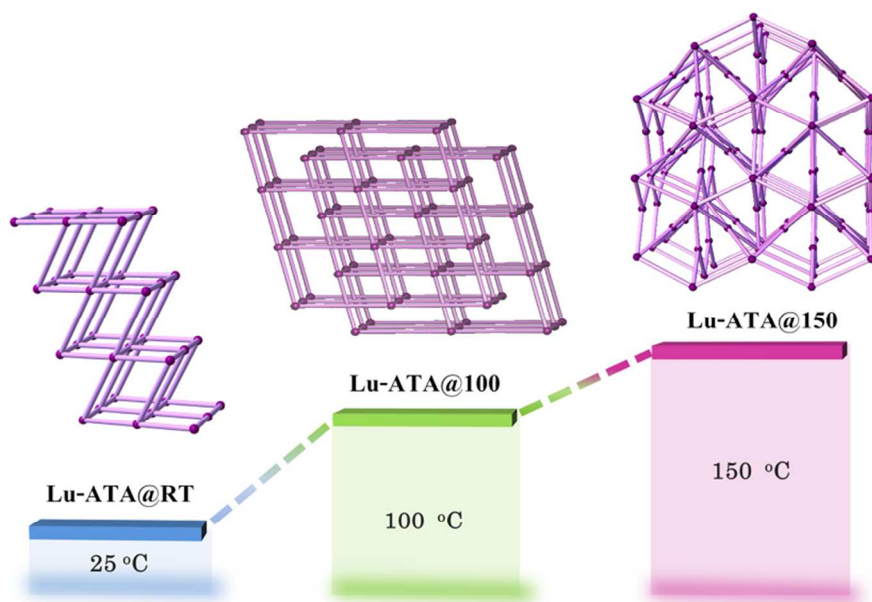
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37
38 Three new porous metal-organic framework (**Lu-ATA@RT**, **Lu-ATA@100** and **Lu-**
39 **ATA@150**) with general formula of $\{\text{Lu}_2(S)_x(\text{ATA})_3\}_n$ (where $S = \text{H}_2\text{O}$, DMF; ATA = 2-
40 aminoterephthalic acid) were obtained. The self-assembled systems composed of
41 aminoterephthalate linker and lutetium ions demonstrate a case of temperature-dependent
42 supramolecular isomerism of their crystal structures. Moreover, the difference between these
43 crystalline isomeric forms is reflected on their gas adsorption behavior.
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