

Cite this: *Chem. Sci.*, 2022, 13, 9321

All publication charges for this article have been paid for by the Royal Society of Chemistry

Amino group induced structural diversity and near-infrared emission of yttrium-tetracarboxylate frameworks†

Hai-Lun Xia,^a Kang Zhou,^a Jiandong Guo,^a Jian Zhang,^{ab} Xiaoxi Huang,^{id a} Dawei Luo,^b Xiao-Yuan Liu^{id *a} and Jing Li^{id *ac}

Near-infrared (NIR)-emitting materials have been extensively studied due to their important applications in biosensing and bioimaging. Luminescent metal–organic frameworks (LMOFs) are a new class of highly emissive materials with strong potential for utilization in biomedical related fields because of their nearly unlimited structural and compositional tunability. However, very little work has been reported on organic linker-based NIR-MOFs and their emission properties. In the present work, a series of yttrium-tetracarboxylate-based LMOFs (HIAM-390X) are prepared *via* judicious linker design to achieve NIR emission with diverse structures. The introduction of an amino group not only offers the remarkable emission bathochromic shift from 521 nm, 665 nm to 689 nm for the resultant MOFs, but also influences the linker conformations, leading to the topology evolution from (4,12)-c ftw, (4,8)-c scu, which is rarely reported in rare earth element-based MOFs, to an unprecedented topology hlx for HIAM-3901 (without an amino group), HIAM-3905 (with one amino group) and HIAM-3906 (with two amino groups). Among these MOFs, HIAM-3907 shows an emission maximum at ~790 nm, with the emission tail close to 1000 nm. The NIR emission may be attributed to the combination of the strongly electron-donating amino group and the strongly electron-withdrawing acceptor naphtho[2,3-*c*][1,2,5]selenadiazole. This work sheds light on the rational design of organic linker-based LMOFs with controlled structures and NIR emission, and inspires future interest in biosensing and bioimaging related applications of NIR-MOFs.

Received 13th May 2022

Accepted 21st July 2022

DOI: 10.1039/d2sc02683j

rsc.li/chemical-science

Introduction

Luminescent metal–organic frameworks (LMOFs) built on emissive organic linkers exhibit strong promise in various applications, including, but not limited to, solid-state-light,^{1–6} sensing^{7–13} and bioimaging,^{14,15} owing to their easily tunable linker structures and highly accessible functional sites.¹⁶ However, most LMOFs reported to date emit light within the visible light spectrum except those using a rare-earth-element as the metal node, particularly Yb, Nd and Er, which usually need organic chromophores as sensitizers and exhibit limited emission tunability due to the specific electronic structures of rare earth elements.^{17–19} There is a lack of study on near-infrared emissive MOFs (NIR-MOFs) based on organic linkers that emit

light in the NIR region.²⁰ The major challenge is to design and synthesize organic linkers that are suitable for the synthesis of MOFs with NIR emission.

In recent years, different kinds of NIR emissive materials, such as carbon nanotubes,²¹ quantum dots,²² rare earth element-based nanoparticles^{23,24} and small molecules,^{25,26} have been gradually developed to explore their bio-related applications. To enhance the performances, it is important to extend the emission to a longer wavelength range, to ensure negligible tissue absorption and minimum auto-fluorescence and reduced scattering, particularly for materials that emit in the NIR-II region. Among them, molecular probes with the emission wavelengths within the 700–1700 nm range, such as benzo[1,2-*c*:4,5-*c'*]bis([1,2,5]thiadiazole) (BBTD) and its derivative-based compounds, have been extensively studied, particularly for bioimaging and biosensing related research,^{25,27–29} due to their high spatiotemporal resolution, good penetration depth and non-invasiveness. However, to achieve NIR emission, most of the organic compounds are designed with large conjugated backbones, which usually suffer from low quantum efficiency due to strong intermolecular interactions and the aggregation-caused quenching (ACQ) effect.^{29–31} Recent studies have shown that ACQ could be suppressed by confining or immobilizing organic linkers or emissive guests in the MOF matrix.^{32,33} In

^aHoffmann Institute of Advanced Materials, Shenzhen Polytechnic, 7098 Liuxian Blvd, Nanshan District, Shenzhen, 518055, P. R. China. E-mail: liuxiaoyuan1989@szpt.edu.cn

^bSchool of Materials and Environmental Engineering, Shenzhen Polytechnic, 7098 Liuxian Blvd, Nanshan District, Shenzhen, 518055, P. R. China

^cDepartment of Chemistry and Chemical Biology, Rutgers University, 123 Bevier Road, Piscataway, New Jersey 08854, USA. E-mail: jingli@rutgers.edu

† Electronic supplementary information (ESI) available. CCDC 2165296 and 2151211. For ESI and crystallographic data in CIF or other electronic format see <https://doi.org/10.1039/d2sc02683j>



addition, the unique properties of MOFs, such as high porosity and pore surface tunability, will also endow NIR-MOFs with more potential for practical applications, *e.g.* drug delivery and light-triggered treatment of diseases. Therefore, developing organic linker-based NIR-MOFs is an important and much desired research area due to their applications in bioimaging and sensing.

The key to success is to design suitable linker structures to achieve broad light absorption and emission, and to use them to prepare NIR-MOFs. It has been reported that introduction of electron-donating amino groups into BTD-based molecules can lead to a significant emission red-shift.^{34–36} We have also discovered a 229 nm red-shift when carboxylic acid groups were substituted by amino groups, from 4,4'-(benzo[*c*][1,2,5]thiadiazole-4,7-diyl)dibenzoic acid (H₂BTBA, 480 nm) to 4,4'-(benzo[*c*][1,2,5]thiadiazole-4,7-diyl)dianiline (BTDA, 709 nm) (Fig. 1a and b). Furthermore, in principle, NIR emissive organic linkers can be designed and synthesized *via* increasing the energy of the highest occupied molecular orbitals (HOMO) or decreasing the energy of the lowest unoccupied molecular orbitals (LUMO). Particularly, for donor–acceptor–donor (D–A–D) type compounds, three different strategies can be employed to achieve a bathochromic shift in emission: (i) decreasing the electron density of acceptors, (ii) increasing the electron density of donor groups and (iii) elongating the conjugated chain between the donor and acceptor.^{26,37,38} For example, when the acceptor group was changed from benzo[*c*][1,2,5]thiadiazole in H₂BTBA to naphtho[2,3-*c*][1,2,5]selenadiazole to form H₂NSB, a 154 nm bathochromic shift was observed (Fig. 1b).³⁹

One useful approach to introduce amino groups into organic linkers for synthesizing MOFs is to use tetracarboxylate-based compounds as the skeleton.⁴⁰ On the other hand, tetraprotic carboxylate linkers used to prepare MOFs usually lead to various

topologies, thereby enriching the structural diversity which leads to enhancement in the performance or extension of the applications.^{40–42} In the case of benzo[*c*][1,2,5]thiadiazole and its derivative-based tetraprotic carboxylate linkers, *i.e.* 5',5'''-(5,6-dimethylbenzo[*c*][1,2,5]thiadiazole-4,7-diyl)bis([1,1':3',1''-terphenyl]-4,4''-dicarboxylic acid) (H₄DTTC), 5',5'''-(benzo[*c*][1,2,5]thiadiazole-4,7-diyl)bis([1,1':3',1''-terphenyl]-4,4''-dicarboxylic acid) (H₄BTTC), 5',5'''-(benzo[*c*][1,2,5]selenadiazole-4,7-diyl)bis([1,1':3',1''-terphenyl]-4,4''-dicarboxylic acid) (H₄BSTC), 5',5'''-(naphtho[2,3-*c*][1,2,5]thiadiazole-4,9-diyl)bis([1,1':3',1''-terphenyl]-4,4''-dicarboxylic acid) (H₄NTTC), and 5',5'''-(naphtho[2,3-*c*][1,2,5]selenadiazole-4,9-diyl)bis([1,1':3',1''-terphenyl]-4,4''-dicarboxylic acid) (H₄NSTC),⁴³ three topologies (**scu**, **scp** and **csq**) have been reported with the Zr₆ cluster as the secondary building unit (Fig. 2).⁴⁴ However, apart from the fact that the **ftw** topology of Zr-based MOFs has strict requirement regarding the linker size (*i.e.*, aspect ratio), we find that it is very difficult to obtain the **ftw** structure even when using linkers that have a suitable size. Attempts to form Zr₆-based **ftw** structures have failed even after using numerous combinations of three common organic solvents and six acids under different reaction conditions. However, the Y₆ cluster shows high flexibility to the aspect ratio of linkers in forming the **ftw** topology.^{45,46} Therefore, yttrium was chosen as the metal in the present work to investigate the structural diversity and achieve NIR-MOFs. In addition, as it is well known that addition of substituents, such as amino, hydroxyl and methyl groups, on the molecular skeleton will introduce steric hindrance, which would dictate the resulting MOFs' structures,^{40,47,48} we thus speculate that the introduction of amino groups may not only induce the emission red-shift, but also lead to diverse structures of the resultant MOFs due to steric hindrance, which is rarely studied in Y-MOFs.

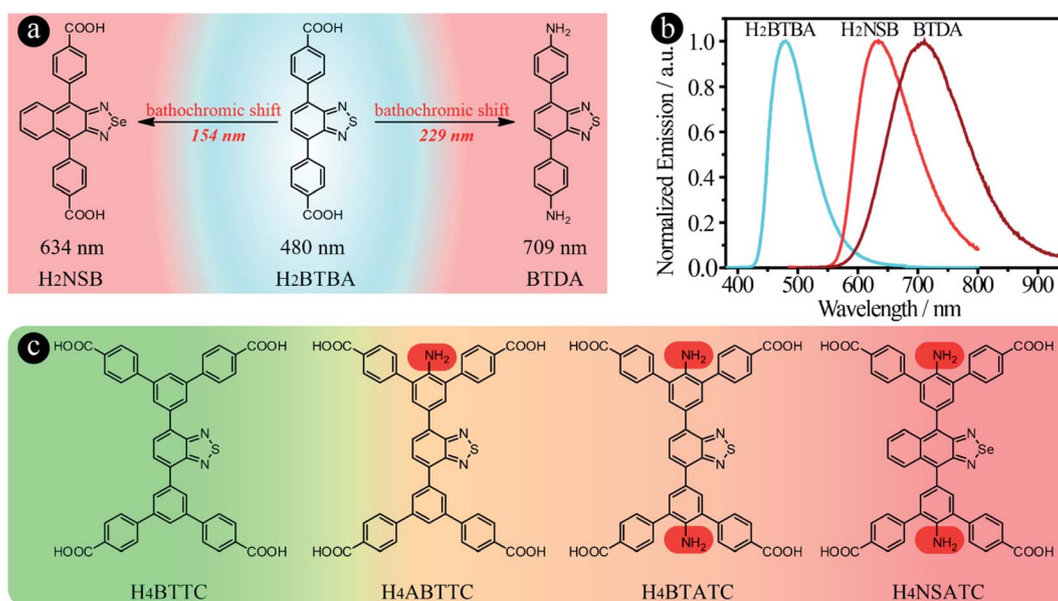


Fig. 1 The molecular structures of H₂NSB, H₂BTBA and BTDA with the corresponding emission wavelengths (a); the normalized emission spectra of H₂NSB, H₂BTBA and BTDA in DMF under 365 nm excitation (b); the molecular structures of H₄BTTC, H₄ABTTC, H₄BTATC and H₄NSATC used in this work to prepare LMOFs with controllable structures and emission toward the NIR range (c).



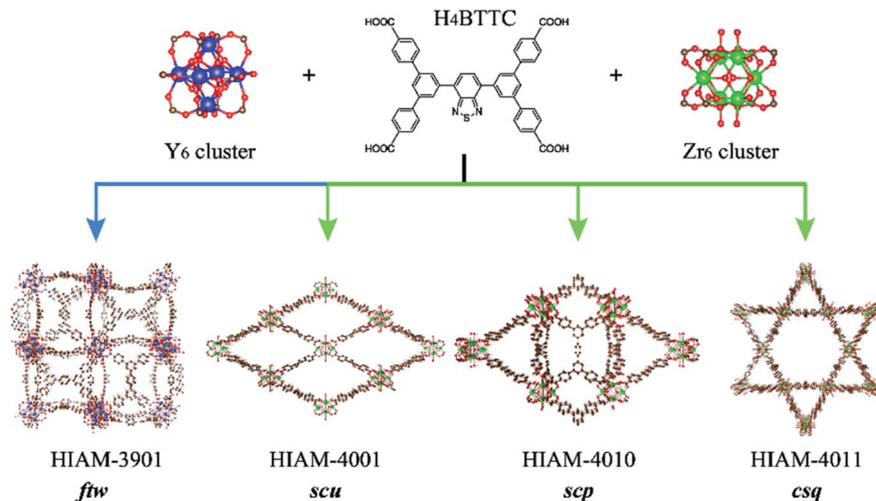


Fig. 2 The topology evolution of LMOFs prepared using Zr_6/Y_6 clusters and H_4BTTC as the organic linker (color scheme: brown, C; green, Zr; dark blue, Y; red, O; yellow, S; blue, N).

Results and discussion

ftw-type Y-MOFs with tunable emission toward the NIR region

Bearing the aforementioned discussions in mind, we first attempted to construct Y-MOFs with the **ftw** topology using H_4DTTC , H_4BTTC , H_4BSTC , H_4NTTC and H_4NSTC as the organic linkers (Fig. 3a). Single crystals of five Y-MOFs, HIAM-390X (HIAM = Hoffmann Institute of Advanced Materials; 39 = yttrium), were obtained according to the reported procedures with some modifications.⁴⁵ A typical synthesis is schematically shown in Fig. 3a: the reaction was carried out in a 5 mL vial containing 2.5 mL DMF, $Y(NO_3)_3 \cdot 6H_2O$ (13.0 mg, 0.034 mmol), organic linker and varying amounts of 2-fluorobenzoic acid in a preheated oven at 120 °C for 24 hours. The synthetic procedures for each MOF are described in detail in the Experimental section. Depending on the properties of the organic linker,

colorless (HIAM-3900), light yellow (HIAM-3901), yellow (HIAM-3902), light orange (HIAM-3903) and red (HIAM-3904) cube-shaped single crystals were obtained with blue, green, yellow, orange and red emission (Fig. 3b). Although we were able to obtain single crystals of appropriate sizes for all five MOFs, it was still very difficult to solve their crystal structures due to the weak diffraction. However, we found that the powder X-ray diffraction (PXRD) patterns of these MOFs match well with those of simulated Y-ftw-MOF-2 (Fig. 3c),⁴⁵ where a similar linker, 5',5'''-bis(4-carboxyphenyl)-[1,1':3',1'':4'',1''':3''',1''''-quinque-phenyl]-4,4''''-dicarboxylic acid, was used as the organic linker, indicating that they are isorecticular structures. Therefore, the structural characterization of the HIAM-390X (X = 0, 1, 2, 3, 4) series was based on Y-ftw-MOF-2, which crystallizes in the cubic system with the $Im\bar{3}$ space group. Each Y_6 cluster is

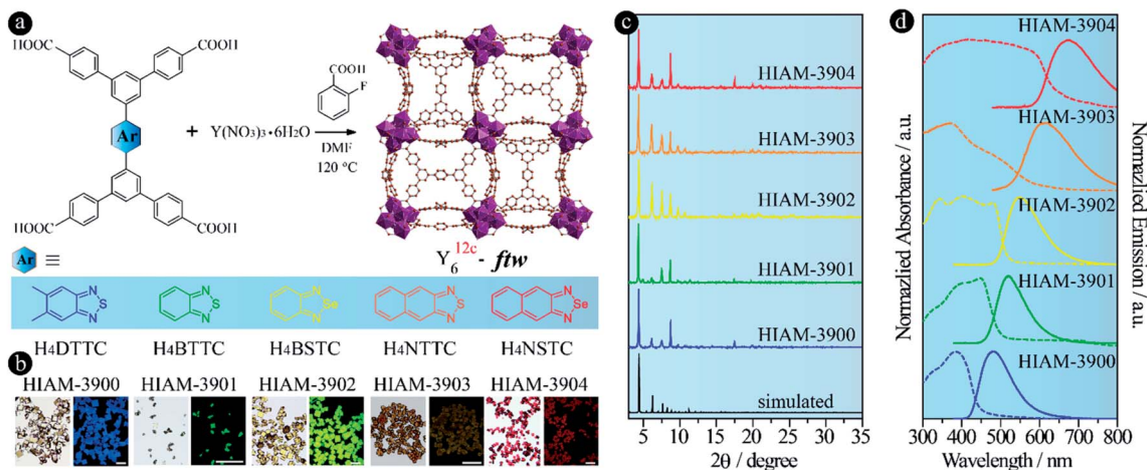


Fig. 3 The molecular structures of the linkers, synthetic conditions and single crystal structure of HIAM-390X (a) (color scheme: brown, C; dark blue, Y; red, O); crystal images under the daylight and irradiation (excitation under 365 nm for HIAM-3900, HIAM-3901 and HIAM-3902, and excitation under 450 nm for HIAM-3903 and HIAM-3904, scale bar: 200 μ m) (b); the PXRD patterns (c) and the normalized solid-state emission spectra under 365 nm excitation and UV-vis absorption spectra (d) of HIAM-3900, HIAM-3901, HIAM-3902, HIAM-3903 and HIAM-3904.



coordinated by twelve linkers and each linker is connected to four Y_6 clusters, leading to a (4,12)-c **ftw** topology.

The optical absorption and emission spectra of HIAM-390X were measured at room temperature. As shown in Fig. 3d, HIAM-3900, HIAM-3901, HIAM-3902, HIAM-3903 and HIAM-3904 emit at different energies with the peak maxima at 480 nm, 521 nm, 550 nm, 615 nm and 675 nm, respectively, consistent with the emission color of the single crystals (Fig. 3b). The corresponding solid-state photoluminescence quantum yields (PLQY) are 28.9%, 31.7%, 26.1%, 5.30% and 6.20% for HIAM-3900, HIAM-3901, HIAM-3902, HIAM-3903 and HIAM-3904, respectively, under 365 nm excitation, and are much higher than the 6.5%, 1.3%, 0.7%, 0.7% and 0.8% for the corresponding linkers H_4 DTTC, H_4 BTTC, H_4 BSTC, H_4 NTTC and H_4 NSTC, respectively. These data further confirm that immobilization and confinement of linkers in the MOF matrix is an efficient strategy to prevent them from aggregation, leading to increased PLQY.^{1,49} It should be noted that HIAM-3901, HIAM-3902, HIAM-3903 and HIAM-3904 can also be excited under blue light (450 nm) with PLQY of 27.0%, 20.0%, 4.10% and 4.70%, respectively. In addition, the emission tails of HIAM-3903 and HIAM-3904 extend into the energy region (>700 nm) of near-infrared light (NIR-I, 700–900 nm), which might be useful for bio-related applications if they have aqueous stability under specific conditions. A monotonic bathochromic shift is also observed in the absorption spectra from HIAM-3900 to HIAM-3904 (Fig. 3d). These results demonstrate that using acceptor groups with decreased electron density is an effective approach to achieve NIR-MOFs, and it has no effect on the underlying net of the resultant LMOFs. When acceptors with electron density much lower than that of naphtho[2,3-*c*][1,2,5]selenadiazole, such as benzo[1,2-*c*:4,5-*c'*]bis([1,2,5]thiadiazole), can be incorporated to the linker skeleton, we believe that LMOFs with NIR-I, and even second near-infrared region (NIR-II) emission can be achieved, which is an ongoing study in our group.

Structural analysis and description of amino group induced topology evolution of Y-MOFs

In addition to the use of acceptors with lower electron density to red-shift emission as discussed above, pre-synthesis linker modifications, such as the introduction of amino groups, have also been utilized to prepare MOFs with emission/absorption red-shift behaviour, *i.e.* MIL-125-NH₂ and MIL-125-(NH₂)₂.⁵⁰ To introduce amino groups into organic linkers and confirm their effect on the emission properties of the resultant LMOFs, three benzo[*c*][1,2,5]thiadiazole-based tetratopic carboxylate linkers, H_4 BTTC without an amino group, 2'-amino-5'-(7-(4,4'-dicarboxy-[1,1':3',1''-terphenyl]-5'-yl)benzo[*c*][1,2,5]thiadiazol-4-yl)-[1,1':3',1''-terphenyl]-4,4''-dicarboxylic acid (H_4 ABTTC) with one amino group and 5',5'''-(benzo[*c*][1,2,5]thiadiazole-4,7-diyl) bis(2'-amino-[1,1':3',1''-terphenyl]-4,4''-dicarboxylic acid) (H_4 BTATC) with two amino groups, were designed and synthesized as shown in Fig. 1c. To further extend the light absorption into the lower energy region to realize NIR emission, the acceptor was changed from benzo[*c*][1,2,5]thiadiazole to

naphtho[2,3-*c*][1,2,5]selenadiazole to produce 5',5'''-(naphtho[2,3-*c*][1,2,5]selenadiazole-4,9-diyl)bis(2'-amino-[1,1':3',1''-terphenyl]-4,4''-dicarboxylic acid) (H_4 NSATC). Since either adding amino groups or using electron-deficient acceptors can red-shift the emission wavelength of benzo[*c*][1,2,5]thiadiazole-based linkers,^{32,43} we envision that such a combination would lead to a much larger bathochromic shift. In addition, introduction of amino groups will offer more options for post-synthesis alteration *via* modification of the amino groups, which can provide multiple possibilities for tuning the emission and facilitate related applications, such as biosensing.^{51–54} As expected, a remarkable emission bathochromic shift is observed for H_4 ABTTC ($\lambda_{em} = 717$ nm) with one amino group and H_4 BTATC ($\lambda_{em} = 697$ nm) with two amino groups compared with the one without an amino group, H_4 BTTC ($\lambda_{em} = 498$ nm), in DMF solution (Fig. S1†). This reveals that introducing an amino group into the linker skeleton indeed is a powerful approach to realize emission red shift.

Subsequently, HIAM-3905 and HIAM-3906 were grown as orange and red single crystals following a similar method to that for HIAM-3901 except using H_4 ABTTC and H_4 BTATC as the organic linkers (Fig. 4a1–a3 and b1–b3). However, single crystal X-ray diffraction (sc-XRD) analysis revealed that HIAM-3905 and HIAM-3906 possess totally different structures compared with HIAM-3901. As shown in Fig. 4a4 and a5 and Table S1,† HIAM-3905 crystallized in the orthorhombic crystal system with the *Cmmm* space group. Each Y_6 cluster is coordinated by eight ABTTC linkers and eight terminal H₂O/OH[−] groups. Each ABTTC is connected to four Y_6 clusters, resulting in a (4,8)-c **scu** net for HIAM-3905, which is rarely reported in rare-earth element-based MOFs.^{55–57} Four types of one-dimensional open channels co-exist in HIAM-3905, *i.e.* rhombic channels and hexagonal channels along the *a*-axis, and two different rhombic channels along the *b*- and *c*-axis (Fig. S2†). However, in the structure of HIAM-3906 (Fig. 4b4, b5, S3 and Table S2†), each Y_6 cluster is coordinated with ten BTATC and four terminal H₂O/OH[−] groups. Each deprotonated BTATC is connected to four Y_6 clusters. Further analysis demonstrates that HIAM-3906 is a 3-nodal (4,4,10)-c net with the point symbol of $\{4^{24} \cdot 6^{20} \cdot 8\}_2\{4^4 \cdot 6^2\}_5$, which is an unprecedented topology in MOFs and denoted as **hlx** in the present work.

The amino group induced structural diversity can be considered as the most interesting structural feature of HIAM-3901, HIAM-3905 and HIAM-3906, where three types of linker conformations co-exist in HIAM-3906 (Fig. 4b6), while only one and two types of conformations are found in HIAM-3905 and HIAM-3901, respectively (Fig. 4a6 and S4†). In fact, the dihedral angles between the outer phenyl ring (with carboxylate groups) and the inner phenyl ring (with amino groups) in HIAM-3906 are 53.22°, 58.66° and 90.00° (Fig. 4b6), respectively. The three types of linkers all adopt the *C*_{2v} symmetry, in which the pairs of phenyl rings rotate toward each other. The dihedral angles are 15.19° and 37.45° for HIAM-3901 with both linker types adopting the *C*_{2v} symmetry (Fig. S4†), and it is 45.29° for HIAM-3905 with the *C*_{2h} symmetry (Fig. 4a6). Regarding the structural effect, substituting the hydrogen atoms in H_4 BTTC with a bulkier amino group would introduce steric hindrance, which



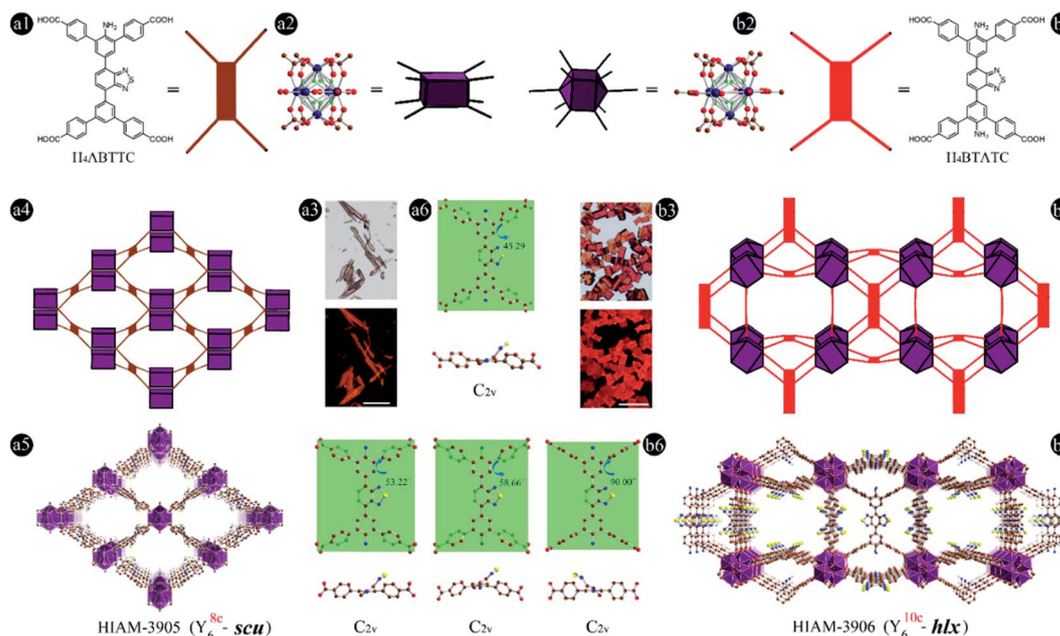


Fig. 4 The molecular structure of 4-connected H_4ABTTC in HIAM-3905 (a1) and H_4BTATC in HIAM-3906 (b1); the 8-connected Y_6 cluster in HIAM-3905 (a2) and the 10-connected Y_6 cluster in HIAM-3906 (b2); the bright-field (above) and photoluminescence (below) single-crystal images of HIAM-3905 (a3) and HIAM-3906 (b3) under 450 nm excitation; the *scu* net topology (a4) and single-crystal structure (a5) of HIAM-3905; the *hlx* net topology (b4) and single-crystal structure (b5) of HIAM-3906; conformations of ABTTC in HIAM-3905 (a6) and BTATC in HIAM-3906 (b6) with the dihedral angle between the inner phenyl ring and outer phenyl ring (color scheme: brown, C; green, F; dark blue, Y; red, O; yellow, S; blue, N).

will change the rotation of the phenyl rings during the MOF synthesis and dictate the structures of the resultant MOFs.^{40,47,48} The experimental PXRD patterns of the as-synthesized HIAM-3906 match well with the simulated ones, indicative of phase purity (Fig. 5a). However, for HIAM-3905, due to the structural flexibility of the (4,8)-c *scu* net, notable peak shifts were observed between the simulated and experimental patterns. It should also be noted that the secondary building units (SBUs) in HIAM-3905 and HIAM-3906 are fluoro-bridged clusters, $Y_6(F_8)(COO)_8$ and $Y_6(F_8)(COO)_{10}$ (Fig. 4a2 and b2), which is consistent with the recent work which reported that using μ_3 -F rather than μ_3 -O will offer a much better crystallographic fit when 2-fluorobenzoic acid is employed as the modulator.⁵⁸ The fluorine presence in HIAM-3905 and HIAM-3906 was confirmed by SEM-EDS analysis (Fig. S5 and S6[†]).

Characterization of amino group induced NIR emissive Y-MOFs

The single crystals of HIAM-3905 and HIAM-3906 show red and dark red emissions with peak maxima at 665 nm and 689 nm, which are bathochromically shifted by 144 nm and 168 nm compared with that of HIAM-3901 (Fig. 5b), confirming that the amino group has the capacity to red-shift the emission energy/wavelength. To further extend the emission wavelength to a lower energy region, we used the acceptor group naphtho[2,3-*c*][1,2,5]selenadiazole, with even lower electron density than benzo[*c*][1,2,5]thiadiazole in H_4BTATC to form H_4NSATC , which was then utilized as the organic linker to prepare HIAM-3907.

The PXRD pattern of HIAM-3907 shows excellent agreement with that of simulated HIAM-3906 (Fig. 5a), indicating its isotropic nature and phase purity. Not surprisingly, HIAM-3907 emits at 790 nm, with a red-shift of 115 nm and 101 nm compared with that of HIAM-3904 (the one without the amino group) and HIAM-3906, respectively, which brings HIAM-3907 to the NIR-I range. Together, a 269 nm bathochromic shift has been achieved by the combination of amino and naphtho[2,3-*c*][1,2,5]selenadiazole groups in the linker skeleton of H_4BTTC to form HIAM-3907.

HIAM-3907 is one of the few organic-linker-based LMOFs with the emission maximum beyond 750 nm, covering the entire NIR-I range with the emission tail towards the NIR-II window.²⁰ These results further demonstrate that the use of acceptors with lower electron density is an effective method to achieve low energy emissions. The corresponding PLQY are 9.0%, 5.0% and 2.0% for HIAM-3905, HIAM-3906 and HIAM-3907, respectively, under 450 nm excitation, which are also much higher than the 3.1%, 0.40% and 0.10% for the corresponding linkers H_4ABTTC , H_4BTATC and H_4NSATC , respectively. In addition, the absorption edges of HIAM-3906 and 3907 reach 650 and 750 nm (Fig. 5c). The broad light harvesting energy range makes these MOFs promising for application in photocatalysis. To further investigate the amino group effect on the emission behavior of the designed linkers, density functional theory (DFT) calculation was utilized to study the electronic structures of these linkers. As depicted in Fig. 5d, after addition of one or two amino groups on H_4BTTC to form H_4ABTTC and H_4BTATC , the LUMO energies gradually



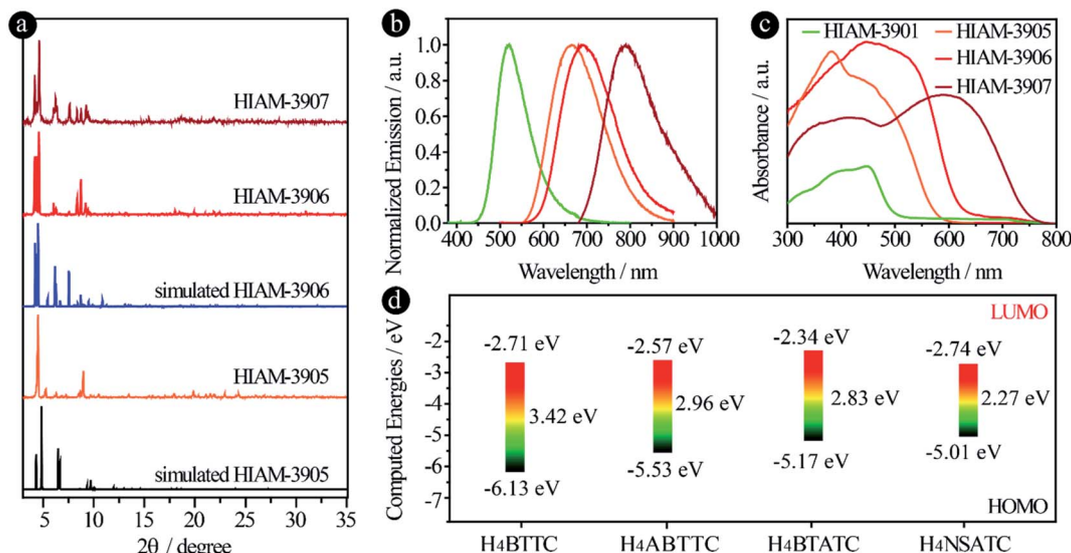


Fig. 5 The PXRD patterns of simulated and synthesized HIAM-3905, HIAM-3906 and HIAM-3907 (a); the normalized solid-state emission (365 nm excitation for HIAM-3901, and 450 nm excitation for HIAM-3905, HIAM-3906 and HIAM-3907) (b) and UV-vis absorption spectra (c) of HIAM-3901, HIAM-3905, HIAM-3906 and HIAM-3907; the calculated HOMO–LUMO energy levels of H₄BTTC, H₄ABTTC, H₄BTATC and H₄NSATC (d).

Table 1 Summary of the structures and emission properties of the MOFs reported in the present work

Linker	MOFs	Connectivity of the Y ₆ cluster	Topology	Emission wavelength (nm)	PLQY (%; excitation wavelength/nm)
H ₄ DTTC	HIAM-3900	12	ftw	480	28.9 (365 nm)
H ₄ BTTC	HIAM-3901	12	ftw	521	31.7 (365 nm)
H ₄ BSTC	HIAM-3902	12	ftw	550	27.0 (450 nm)
H ₄ NTTC	HIAM-3903	12	ftw	615	26.1 (365 nm)
H ₄ NSTC	HIAM-3904	12	ftw	675	20.0 (450 nm)
H ₄ ABTTC	HIAM-3905	8	scu	665	5.30 (365 nm)
H ₄ BTATC	HIAM-3906	10	hlx	689	4.10 (450 nm)
H ₄ NSATC	HIAM-3907	10	hlx	790	6.20 (365 nm)
					4.70 (450 nm)
					9.0 (450 nm)
					5.0 (450 nm)
					2.0 (450 nm)

increased from -2.71 eV to -2.57 eV and to -2.34 eV, while a much larger increase was observed for the HOMO energies, from -6.13 eV to -5.53 eV and to -5.17 eV. As a result, the energy gaps between the HOMO and LUMO are significantly decreased from 3.42 eV for H₄BTTC to 2.96 eV for H₄ABTTC, and 2.83 eV for H₄BTATC, which are consistent with the experimental results. These results demonstrate that the amino group exhibits a much more significant effect on the HOMO than on the LUMO. When an acceptor group possessing much lower electron density than benzo[*c*][1,2,5]thiadiazole was used, such as naphtho[2,3-*c'*][1,2,5]selenadiazole, to form H₄NSATC, a 0.40 eV lower LUMO energy and a 0.16 eV higher HOMO energy were achieved, which leads to a further decreased energy gap of 2.27 eV for H₄NSATC.

As summarized in Table 1, although a series of Y-MOFs were prepared with emission toward the near-infrared range and

good water stability, for example HIAM-3907 (Fig. S7†), it is still difficult to apply them in bioimaging and biosensing in the present work due to the inability to achieve nanocrystallization. Attempts to achieve nano-sized NIR-MOFs with excellent aqueous compatibility, such as stability and dispersibility, for bio-related applications are ongoing.

Conclusion

In conclusion, a series of luminescent Y-MOFs with the **ftw** topology, HIAM-3900 to HIAM-3904, have been prepared using 2,1,3-benzothiadiazole and its derivative-based tetrapotic carboxylate linkers. The emission tails of HIAM-3903 and HIAM-3904 fall in the NIR-I region. In addition, introducing an amino group and acceptor with lower electron density into the linker skeleton will not only redshift the emission of the resultant LMOFs to cover the entire NIR-I window (*i.e.* HIAM-



3907, $\lambda_{em} = 790$ nm), but also induce topology evolution, from (4,12)-c **ftw**, (4,8)-c **scu**, which is rarely reported in rare-earth element-based MOFs, to an unprecedented topology (4,10)-c **hlx**, as a result of steric hindrance caused by the amino group. The present work sheds light on the rational design of emissive organic linkers to obtain NIR-MOFs with controllable structures, which no doubt will facilitate their applications in biosensing and bioimaging.

Data availability

All the data have been included in the ESI.†

Author contributions

X.-Y. L. conceived the idea and designed the experiment; H.-L. Xia and J. Zhang worked on the synthesis and characterization of all materials. K. Zhou carried out and analyzed the single crystal structures. J. Guo did the density functional theory calculations. X. Huang and D. Luo provided detailed structural analysis and discussion. X.-Y. L. and J. Li wrote the paper with the help from all authors.

Conflicts of interest

The authors declare no competing financial interests.

Acknowledgements

X.-Y. Liu acknowledges the financial support from start-up funding for Shenzhen High-Caliber Personnel of Shenzhen Polytechnic (6022310053K), Guangdong Basic and Applied Basic Research Foundation (2020A1515110420), and Shenzhen Science and Technology Program (RCBS202007141114941230). D. Luo acknowledges the Special Funds for the Science and Technology Innovation Project of Guangdong Provincial Department of Education (2021KTSCX278). H.-L. Xia gratefully acknowledges the support from Shenzhen Polytechnic (6021330018K).

References

- Q. Gong, Z. Hu, B. J. Deibert, T. J. Emge, S. J. Teat, D. Banerjee, B. Mussman, N. D. Rudd and J. Li, *J. Am. Chem. Soc.*, 2014, **136**, 16724.
- J. Cornelio, T. Y. Zhou, A. Alkas and S. G. Telfer, *J. Am. Chem. Soc.*, 2018, **142**, 15470.
- W. P. Lustig and J. Li, *Coord. Chem. Rev.*, 2018, **373**, 116.
- W. J. Newsome, S. Ayad, J. Cordova, E. W. Reinheimer, A. D. Campiglia, J. K. Harper, K. Hanson and F. J. Uribe-Romo, *J. Am. Chem. Soc.*, 2019, 11298.
- W. P. Lustig, Z. Shen, S. J. Teat, N. Javed, E. Velasco, D. M. O'Carroll and J. Li, *Chem. Sci.*, 2020, **11**, 1814.
- W. J. Newsome, A. Chakraborty, R. T. Ly, G. S. Pour, D. C. Fairchild, A. J. Morris and F. Uribe-Romo, *J. Chem. Sci.*, 2020, **11**, 4391.
- Z. Hu, B. J. Deibert and J. Li, *Chem. Soc. Rev.*, 2014, **43**, 5815.
- Z. Hu, W. P. Lustig, J. Zhang, C. Zheng, H. Wang, S. J. Teat, Q. Gong, N. D. Rudd and J. Li, *J. Am. Chem. Soc.*, 2015, **137**, 16209.
- A. Mallick, A. M. El-Zohry, O. Shekhah, J. Yin, J. Jia, H. Aggarwal, A. H. Emwas, O. F. Mohammed and M. Eddaoudi, *J. Am. Chem. Soc.*, 2019, **141**, 7245.
- J. Li, S. Yuan, J.-S. Qin, J. Pang, P. Zhang, Y. Zhang, Y. Huang, H. F. Drake, W. R. Liu and H.-C. Zhou, *Angew. Chem., Int. Ed.*, 2020, **59**, 9319.
- C. R. Martin, P. Kittikhunnatham, G. A. Leith, A. A. Berseneva, K. C. Park, A. B. Greytak and N. B. Shustova, *Nano Res.*, 2020, **14**, 338.
- Y. Zhao, H. Zeng, X.-W. Zhu, W. Lu and D. Li, *Chem. Soc. Rev.*, 2021, **50**, 4484.
- W. P. Lustig, S. Mukherjee, N. D. Rudd, A. V. Desai, J. Li and S. K. Ghosh, *Chem. Soc. Rev.*, 2017, **46**, 3242.
- J. Park, Q. Jiang, D. Feng, L. Mao and H. C. Zhou, *J. Am. Chem. Soc.*, 2016, **138**, 3518.
- J. Park, M. Xu, F. Li and H. C. Zhou, *J. Am. Chem. Soc.*, 2018, **140**, 5493.
- S. Ali Akbar Razavi and A. Morsali, *Coord. Chem. Rev.*, 2019, **399**, 213023.
- D. Zou, J. Zhang, Y. Cui and G. Qian, *Dalton Trans.*, 2019, **48**, 6669.
- T.-Y. Luo, P. Das, D. L. White, C. Liu, A. Star and N. L. Rosi, *J. Am. Chem. Soc.*, 2020, **142**, 2897.
- P. F. Muldoon, G. Collet, S. V. Eliseeva, T.-Y. Luo, S. Petoud and N. L. Rosi, *J. Am. Chem. Soc.*, 2020, **142**, 8776.
- F. Wang, J. Wang, S. F. Maehrlein, Y. Ma, F. Liu and X. Y. Zhu, *J. Phys. Chem. Lett.*, 2020, **11**, 762.
- K. Welsher, Z. Liu, S. P. Sherlock, J. T. Robinson, Z. Chen, D. Daranciang and H. Dai, *Nat. Nanotechnol.*, 2009, **4**, 773.
- O. T. Bruns, T. S. Bischof, D. K. Harris, D. Franke, Y. Shi, L. Riedemann, A. Bartelt, F. B. Jaworski, J. A. Carr, C. J. Rowlands, M. W. B. Wilson, O. Chen, H. Wei, G. W. Hwang, D. M. Montana, I. Coropceanu, O. B. Achorn, J. Kloepper, J. Heeren, P. T. C. So, D. Fukumura, K. F. Jensen, R. K. Jain and M. G. Bawendi, *Nat. Biomed. Eng.*, 2017, **1**, 0056.
- D. J. Naczynski, M. C. Tan, M. Zevon, B. Wall, J. Kohl, A. Kulesa, S. Chen, C. M. Roth, R. E. Riman and P. V. Moghe, *Nat. Commun.*, 2013, **4**, 2199.
- P. Pei, Y. Chen, C. Sun, Y. Fan, Y. Yang, X. Liu, L. Lu, M. Zhao, H. Zhang, D. Zhao, X. Liu and F. Zhang, *Nat. Nanotechnol.*, 2021, **16**, 1011.
- A. L. Antaris, H. Chen, K. Cheng, Y. Sun, G. Hong, C. Qu, S. Diao, Z. Deng, X. Hu, B. Zhang, X. Zhang, O. K. Yaghi, Z. R. Alamparambil, X. Hong, Z. Cheng and H. Dai, *Nat. Mater.*, 2016, **15**, 235.
- Z. Lei and F. Zhang, *Angew. Chem., Int. Ed.*, 2021, **60**, 16294.
- J. Li and K. Pu, *Chem. Soc. Rev.*, 2019, **48**, 38.
- S. Zhu, R. Tian, A. L. Antaris, X. Chen and H. Dai, *Adv. Mater.*, 2019, **31**, e1900321.
- S. Liu, Y. Li, R. T. K. Kwok, J. W. Y. Lam and B. Z. Tang, *Chem. Sci.*, 2020, **12**, 3427.
- W. Xu, D. Wang and B. Z. Tang, *Angew. Chem., Int. Ed.*, 2021, **60**, 7476.



- 31 S. Xu, Y. Duan and B. Liu, *Adv. Mater.*, 2020, **32**, e1903530.
- 32 S. Wu, D. Ren, K. Zhou, H.-L. Xia, X.-Y. Liu, X. Wang and J. Li, *J. Am. Chem. Soc.*, 2021, **143**, 10547.
- 33 X.-Y. Liu, K. Xing, Y. Li, C.-K. Tsung and J. Li, *J. Am. Chem. Soc.*, 2019, **141**, 14807.
- 34 Y. Sun, M. Ding, X. Zeng, Y. Xiao, H. Wu, H. Zhou, B. Ding, C. Qu, W. Hou, A. G. A. Er-bu, Y. Zhang, Z. Cheng and X. Hong, *Chem. Sci.*, 2017, **8**, 3489.
- 35 L. e. Zhang, C. Liu, S. Zhou, R. Wang, Q. Fan, D. Liu, W. Wu and X. Jiang, *Adv. Healthcare Mater.*, 2020, **9**, 1901470.
- 36 Y. Fang, J. Shang, D. Liu, W. Shi, X. Li and H. Ma, *J. Am. Chem. Soc.*, 2020, **142**, 15271.
- 37 B. Li, M. Zhao and F. Zhang, *ACS Mater. Lett.*, 2020, **2**, 905.
- 38 H.-L. Xia, K. Zhou, S. Wu, D. Ren, K. Xing, J. Guo, X. Wang, X.-Y. Liu and J. Li, *Chem. Sci.*, 2022, **13**, 8036.
- 39 G. Han, S. Wu, K. Zhou, H.-L. Xia, X.-Y. Liu and J. Li, *Inorg. Chem.*, 2022, **61**, 3363.
- 40 J. Pang, S. Yuan, J. Qin, C. Liu, C. Lollar, M. Wu, D. Yuan, H.-C. Zhou and M. Hong, *J. Am. Chem. Soc.*, 2017, **139**, 16939.
- 41 Z. Chen, S. L. Hanna, L. R. Redfern, D. Alezi, T. Islamoglu and O. K. Farha, *Coord. Chem. Rev.*, 2019, **386**, 32.
- 42 Y. Chen, X. Zhang, M. R. Mian, F. A. Son, K. Zhang, R. Cao, Z. Chen, S. J. Lee, K. B. Idrees, T. A. Goetjen, J. Lyu, P. Li, Q. Xia, Z. Li, J. T. Hupp, T. Islamoglu, A. Napolitano, G. W. Peterson and O. K. Farha, *J. Am. Chem. Soc.*, 2020, **142**, 21428.
- 43 D. Ren, H.-L. Xia, K. Zhou, S. Wu, X.-Y. Liu, X. Wang and J. Li, *Angew. Chem., Int. Ed.*, 2021, **60**, 25048.
- 44 H.-L. Xia, K. Zhou, L. Yu, H. Wang, X.-Y. Liu, D. M. Proserpio and J. Li, *Inorg. Chem.*, 2022, **61**, 7980.
- 45 R. Luebke, Y. Belmabkhout, L. J. Weselinski, A. J. Cairns, M. Alkordi, G. Norton, L. Wojtas, K. Adil and M. Eddaoudi, *Chem. Sci.*, 2015, **6**, 4095.
- 46 H. Wang, X. Dong, V. Colombo, Q. Wang, Y. Liu, W. Liu, X.-L. Wang, X.-Y. Huang, D. M. Proserpio, A. Sironi, Y. Han and J. Li, *Adv. Mater.*, 2018, **30**, 1805088.
- 47 B. Wang, X. L. Lv, D. Feng, L. H. Xie, J. Zhang, M. Li, Y. Xie, J. R. Li and H. C. Zhou, *J. Am. Chem. Soc.*, 2016, **138**, 6204.
- 48 X. L. Lv, S. Yuan, L. H. Xie, H. F. Darke, Y. Chen, T. He, C. Dong, B. Wang, Y. Z. Zhang, J. R. Li and H. C. Zhou, *J. Am. Chem. Soc.*, 2019, **141**, 10283.
- 49 Z. Hu, G. Huang, W. P. Lustig, F. Wang, H. Wang, S. J. Teat, D. Banerjee, D. Zhang and J. Li, *Chem. Commun.*, 2015, **51**, 3045.
- 50 C. H. Hendon, D. Tiana, M. Fontecave, C. Sanchez, L. D'arras, C. Sassoie, L. Rozes, C. Mellot-Draznieks and A. Walsh, *J. Am. Chem. Soc.*, 2013, **135**, 10942.
- 51 Z. Wang and S. M. Cohen, *Angew. Chem., Int. Ed.*, 2008, **47**, 4699.
- 52 Y. Zhang, X. Feng, H. Li, Y. Chen, J. Zhao, S. Wang, L. Wang and B. Wang, *Angew. Chem., Int. Ed.*, 2015, **54**, 4259.
- 53 A. M. Fracaroli, P. Siman, D. A. Nagib, M. Suzuki, H. Furukawa, F. D. Toste and O. M. Yaghi, *J. Am. Chem. Soc.*, 2016, **138**, 8352.
- 54 Y. Wen, J. Zhang, Q. Xu, X.-T. Wu and Q.-L. Zhu, *Coord. Chem. Rev.*, 2018, **376**, 248.
- 55 X.-L. Lv, L. Feng, L.-H. Xie, T. He, W. Wu, K.-Y. Wang, G. Si, B. Wang, J.-R. Li and H.-C. Zhou, *J. Am. Chem. Soc.*, 2021, **143**, 2784.
- 56 L. Feng, J. Pang, P. She, J.-L. Li, J.-S. Qin, D.-Y. Du and H.-C. Zhou, *Adv. Mater.*, 2020, **32**, 2004414.
- 57 F. Saraci, V. Quezada-Novoa, P. R. Donnarumma and A. J. Howarth, *Chem. Soc. Rev.*, 2020, **49**, 7949.
- 58 J. P. Vizuet, M. L. Mortensen, A. L. Lewis, M. A. Wunch, H. R. Firouzi, G. T. McCandless and K. J. Balkus, *J. Am. Chem. Soc.*, 2021, **143**, 17995.

