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Cite this: *Org. Chem. Front.*, 2022, **9**, 3084

Triazolated calix[4]semitubes: assembling strategies towards long multicalixarene architectures†

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Cone and 1,3-alternate calix[4]arenes bearing pairs of 2-azidoethyl or propargyl groups, and 1,3-alternate calix[4]arenes having four 2-azidoethyl and four propargyl groups or pairs of 2-azidoethyl and silylated propargyl groups were explored as the components for the CuAAC-syntheses of triazolated calix[4]semitubes having three calixarene cores in the structures. Though condition tuning for each of the calixarene combinations was performed, the four-fold CuAACs using the tetrafunctional calixarenes returned the respective tris(calixarenes) in good yields only in the syntheses involving the calixarene tetrakis(alkyne) and hydroxylated bis(azides). The stepwise approach towards the triazolated calix[4]semitubes by using the 2-azidoethylated/propargylated calix[4]arene as the key synthon and de-silylation as the intermediate step was found to be a good alternative, which allowed the preparation of the semitubes with different substituents in the terminal calixarene units and can also be used for further extension of the semi-tubular assemblies.

Received 16th March 2022,
Accepted 14th April 2022

DOI: 10.1039/d2qo00432a
rsc.li/frontiers-organic

Introduction

The well-developed approaches for chemical transformations of calixarenes and related macrocycles allow us to treat them as molecular platforms within a chemical 'building kit' for constructing diverse receptor molecules in which the required number of certain functional groups can be introduced and mutually pre-organized in a predictable manner.¹ When several platforms are involved in chemical reactions, more diverse and complicated, and, thus, much more attractive architectures may be created by using such multicalixarene conjugates. To date, many examples have been published on the covalently joining 'classical' calixarene macrocycles,² and those of thiocalixarenes,³ calixpyrroles,⁴ calixresorcinarenes,⁵ and heteromacrocyclic conjugates from the calixarene family have also been known.⁶ Most often, these conjugates comprise only two calix-

ene macrocycles (or/and related ones), which are connected to each other by one or several linkers. In the latter case, the calixarenes and the linkers form an additional macrocyclic site or cavity, which may possess special or even outstanding receptor abilities. Tris- and even larger multicalixarene assemblies are also known, of which those having a tubular shape are the most attractive,⁷ though they are less available. These compounds inherit all the structural benefits from the above biscalixarene assemblies and allow allocation of more than one macrocyclic site/cavity within one molecule and manage their connectivity in the range from pure independent mode where simple binding/recognition of multiple guests is possible, up to tightly bound mode, which is prospective for constructing sophisticated supramolecules, *e.g.* multipositional molecular switchers.

We have recently shown⁸ that diverse biscalixarene semitubes may be constructed from the respective bisazide and bisalkyne calixarene precursors using copper(i)-catalyzed azide-alkyne cycloaddition (CuAAC).⁹ Each of these assemblies comprises a single binding site formed by two 1,4-disubstituted 1,2,3-triazole groups surrounded by the calixarene cores and targeted to transition cations. Herein we explored the applicability of this synthetic approach for the construction of larger semi-tubular assemblies having three calix[4]arene cores in their structures and possessing two bistriazole receptor sites separated from each other by the central 1,3-alternate calix[4]arene unit.

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† Electronic supplementary information (ESI) available: Experimental details and the NMR spectra of novel compounds. CCDC 2157422. For ESI and crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d2qo00432a>

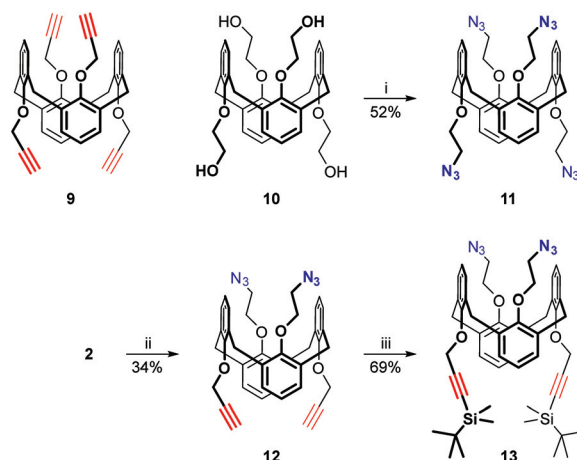
Results and discussion

Alkyne and azide components of calix[4]semitubes

The design of the long triazolated calix[4]semitubes requires azide/alkyne-containing calixarenes of at least two types to be involved in CuAAC. The calixarenes of the first type are the respective bis(azides) or bis(alkynes) which can be used as the terminal units at both sides of a semi-tube, while the calixarenes of the other type are those having two pairs of azide or/and alkyne groups separated from each other by the macrocyclic core. Calix[4]arenes **1**,¹⁰ and **2–4**⁸ bearing 2-azidoethyl groups (Fig. 1) were selected as the bisazide terminal units in the triazolated calix[4]semitubes, and the structurally related calixarene dipropargyl ethers **5**,¹¹ **6**,¹² **7**,¹³ and **8**⁸ were used as the bisalkyne terminal units. Within each of the series, the bifunctional molecules have free phenolic OH groups or those propylated to fix the calix[4]arene macrocycle in a cone or 1,3-alternate molecular shape. Notably, the bis(azides) or bis(alkynes) of each type have been shown to react with each other to form the respective biscalixarene semitubes,⁸ so they were expected to be suitable for building longer semitubes as well.

As for the central macrocycles of the triscalixarene semitubes, calixarene **9**¹⁴ having two pairs of propargyl groups arranged by the two ‘sides’ of the 1,3-alternate core was selected for the CuAAC reactions with the bis(azides) **1–4**. To study the ‘inverted’ azide/alkyne-combinations involving dipropargyl ethers **5–8**, the respective tetrakis(azide) **11** was prepared for the first time from the tetrakis(alcohol) **10**^{7d} and diphenylphosphoryl azide (DPPA) under the Mitsunobu reaction conditions (Scheme 1).

The attractive idea to prepare the hybrid central core having pairs of propargyl and 2-azidoethyl groups within the same molecule was successfully implemented by alkylating the bis(azide) **2** with propargyl bromide in the presence of Cs_2CO_3 . To avoid any self-reactions of the prepared calixarene **12** under the CuAAC conditions, its propargyl groups were de-protonated with lithium bis(trimethylsilyl)amide (LiHMDS) and then sily-



Scheme 1 Calixarenes used as the central units in the calix[4]semitubes: (i) DPPA, DIAD, Ph₃P, THF, rt; (ii) propargyl bromide, Cs₂CO₃, DMF, rt; (iii) TBDMSCl, LiHMDS, THF, rt.

lated with *tert*-butyldimethylsilyl chloride (TBDMSCl) to obtain the protected calixarene **13** in which only the azide groups can participate in the Cu-catalyzed cycloaddition.

One-step assembling of triscalixarene semitubes

It has been shown for the preparation of calix[4]semitubes from bis(azides) and bis(alkynes) under CuAAC conditions that the formation of polymeric/oligomeric byproducts during the reaction could not be avoided, so the selection of proper reaction conditions for every bis(azide)/bis(alkyne) combination was crucial for obtaining targeted semitubes in more or less satisfactory yields, which did not exceed 46%.⁸ Reasonably, successful preparation of the triscalixarene semitubes using tetrafunctional central cores **9** or **11** requires the same or even more strict tuning of the reaction conditions. Still, the conditions obtained previously for the preparation of a semitube from 1,3-alternate bis(alkyne) **8** and cone bis(azide) **1**⁸ were found suitable also for the reaction between the bis(azide) **1** and the tetrakis(alkyne) **9** taken in a 2 : 1 molar ratio (Scheme 2). Indeed, the reaction conducted in the presence of 0.3 equiv. of CuI (per bis(azide)) and triethylamine in toluene at room temperature returned the triscalixarene semitube **14** in 38% yield, which was impressively high considering the formation of four triazole heterocycles in one molecule. The remaining products of the reaction were polymeric/oligomeric compounds as determined from the ¹H NMR spectrum of the reaction mixture after removal of copper salts, in which no other sharp signals except for those from the semitube **14** were observed (Fig. 2a). Deceptively, it may seem that the reaction mixture consisted mainly of the semitube **14**, while the amount of the polymeric/oligomeric product was relatively small. But in fact, the amount of separated polymeric/oligomeric fraction which showed a very broadened ¹H NMR spectrum (Fig. 2c) was nearly equal to that of the pure calix[4]semitube **14** having a sharp NMR pattern (Fig. 2b).

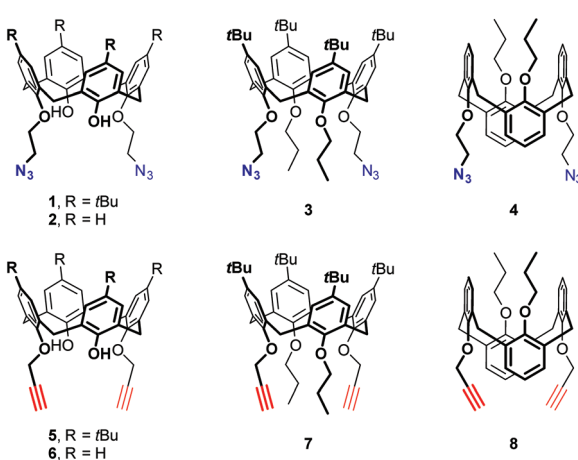
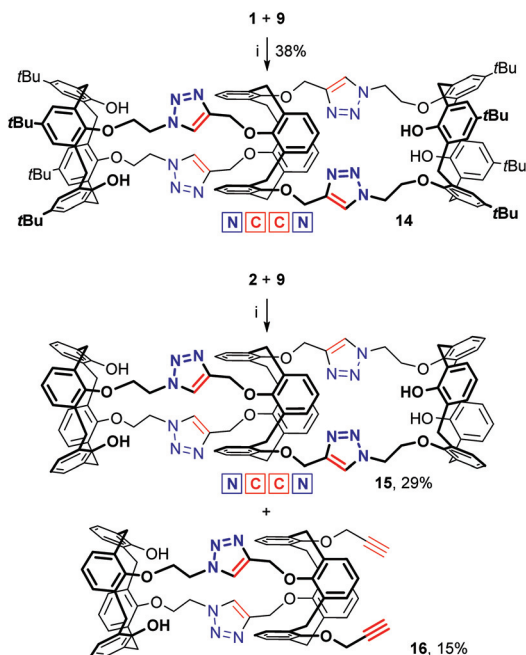


Fig. 1 Calixarenes tested as the terminal units in the calix[4]semitubes.



Scheme 2 Syntheses of calix[4]semitubes **14–16**: (i) CuI (0.3 equiv. per bis(azide)), Et₃N, toluene, rt, 48 h; here and below, for a better distinguishability of the quite similar structures, the combination of **C** and **N** symbols in the squares denotes the arrangement of the triazole groups around the central calixarene core, where **C** (carbon) and **N** (nitrogen) mean the acetylene and azide ‘parts’ of the triazole ring.

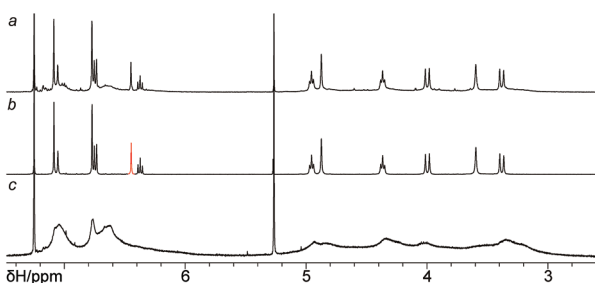


Fig. 2 Parts of the ¹H NMR spectra of (a) the mixture obtained from the reaction between calixarenes **1** and **9** after removal of copper salts, (b) pure triscalixarene semitube **14**, and (c) separated polymeric/oligomeric fraction (scaled up); 400 MHz, CDCl₃; the singlet from the triazole protons in the semitube is colored red.

At slow evaporation of a dichloromethane/methanol solution of calix[4]semitube **14** single crystals were collected and subjected to X-ray diffraction analysis.¹⁵ The collected data confirm the triscalixarene molecular structure of compound **14** with the 1,3-alternate central calix[4]arene core linked to two terminal cone calix[4]arene cores by two pairs of triazole groups (Fig. 3). Due to the absence of a second pair of substituents at the narrow rims of the terminal calixarenes, the whole system seems relatively flexible, so the overall shape of the molecule, at least in the crystalline state, is far from a straight tubular one.

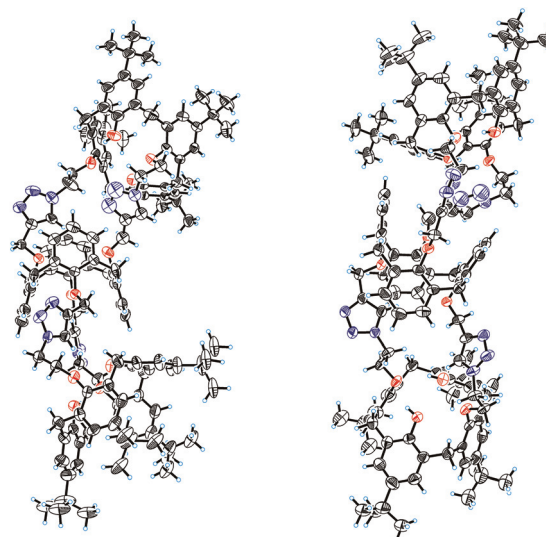
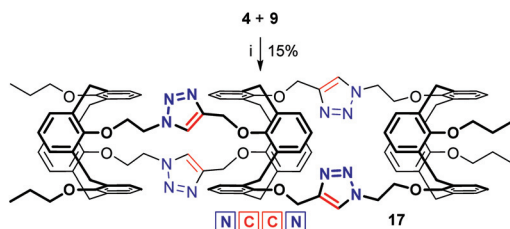


Fig. 3 Molecular structure of the calix[4]semitube **14** in two projections; thermal ellipsoids are drawn at a 50% probability level.

When de-*tert*-butylated bis(azide) **2** was involved in CuAAC with the tetrakis(alkyne) **9** under the conditions used for the preparation of calix[4]semitube **14**, the respective tris(calixarene) **15** was obtained in a good yield of 29% (Scheme 2). But in this case, a part of bis(azide) **2** returned unreacted and a non-polymeric by-product was formed. The latter one was separated and confirmed to be the bis(calixarene) **16** formed at the addition of only one molecule of bis(azide) **2** to the tetrakis(alkyne) **9**. The reasons for the incomplete conversion of the bis(azide) **2** during the reaction are not clear and may be related to its less solubility in toluene at room temperature. Still, the appearance of bis(calixarene) **16** allows one to suggest it to intermediate the stepwise formation of the triscalixarene semitube **15**.

Next, the pair of calixarenes **3** and **9** was studied for the ability to form a semitube. Following our previous observations, the CuAAC between the propylated cone bis(azide) **3** and bis(alkyne) **8** (having the same 1,3-alternate molecular shape as that of the calixarene **9** but only two of four propargyl groups) has given the respective bis(calixarene) separated from a complex mixture with other triazolated huge macrocycles composed of several 3/8 calixarene pairs.⁸ In the case of calixarenes **3** and **9**, the direct implementation of the previously found reaction conditions (CuI (0.3 equiv. per bis(azide)), Et₃N, toluene/1,2-dichloroethane, 60 °C, 9 h) failed, as the obtained reaction mixture consisted of both starting calixarenes, a polymeric/oligomeric material, and several triazolated multimacrocycles from which the desired semitube could not be separated using column chromatography with different eluents. Several alterations of the reaction conditions were performed (see Table S1 in the ESI†), which included the changing of solvents (toluene, toluene/1,2-dichloroethane, THF, and THF/H₂O) and the catalytic system (CuI·P(OEt)₃, CuI, and CuSO₄·5H₂O/sodium ascorbate), but none of the conditions



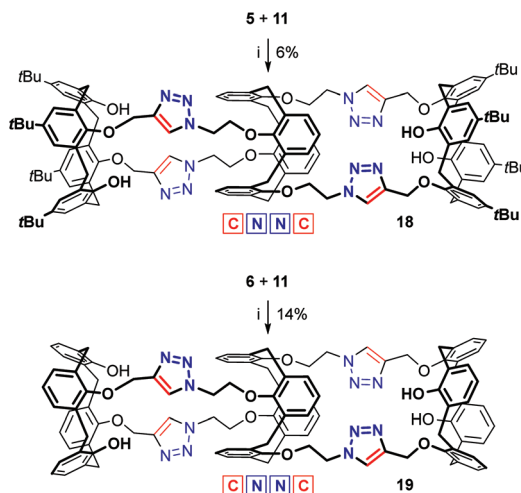
Scheme 3 Synthesis of the calix[4]semiture 17: (i) CuI (0.3 equiv. per bis(azide)), Et₃N, toluene, 60 °C, 24 h.

tested gave a satisfactory outcome. In the best run (CuI (0.3 equiv. per bis(azide)), Et₃N, toluene, 75 °C, 24 h) the complete conversion of the starting calixarenes was achieved and the formation of undesired multimacrocycles was suppressed, but the semiture was formed at nearly a trace level among non-separable polymeric products.

When the 1,3-alternate bis(azide) 4 was reacted with the tetrakis(alkyne) 9 (Scheme 3), additional tuning of the reaction conditions was required (see Table S2 in the ESI[†]). Again, the CuAAC conditions (CuSO₄·5H₂O/sodium ascorbate (1.0 equiv. per bis(azide)), THF/H₂O, 60 °C, 24 h), which were found perfect to assemble two 1,3-alternate calixarenes (4 and 8) into a biscalixarene semiture, turned non-suitable for the synthesis of the tris(calixarene) 17 from the calixarenes 4 and 9. The best result was obtained under CuI/Et₃N-catalysis with the catalyst loading decreased back to 0.3 equiv. per bis(azide). After purification by column chromatography the calix[4]semiture 17 was obtained in 15% yield.

The difference in the CuAAC conditions suitable for the formation of bis- and triscalixarene semitures can be rationalized in terms of the stepwise addition of the calixarenes to each other (see the example of the semiture 15) and by the formation of an efficient copper-binding site in the intermediate bis(calixarene). This site can entrap the cation and prevent it from migration to the next pair of reacting propargyl groups at the other side of calixarene 9, or/and prevent the reaction between the cation(s) from the bulk solution and the propargyl groups due to the Coulomb repulsion, or/and can spatially rearrange the propargyl groups in a way they turn more suitable for polymer formation rather than for the desired two-point connection to the second molecule of a bis(azide).

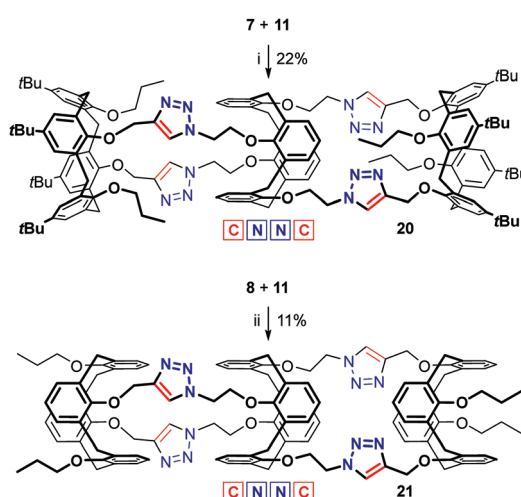
When used as the central core in the triscalixarene semitures, the tetrakis(azide) 11 enables a more distant arrangement of the pairs of the reacting functional groups than the tetrakis(alkyne) 9 does. Accordingly, this can not only diminish the above repulsive interactions between the two Cu-containing bisazide/bistriazole sites, but also make the azide groups within each site less pre-organized, which can hamper the semiture formation. For the CuAAC reaction between the tetrakis(azide) 11 and bis(alkyne) 5 (Scheme 4), condition tuning (see Table S3 in the ESI[†]) gave the best outcome in the case when CuI/Et₃N (0.3 equiv. of Cu per bis(azide)) was used under heating. Nevertheless, the yield of the tris(calixarene) 18 was as small as 6%. Analogous synthesis involving de-*tert*-butylated



Scheme 4 Syntheses of calix[4]semitures 18 and 19: (i) CuI (0.3 equiv. per bis(azide)), Et₃N, toluene, 60 °C, 12 h.

bis(azide) 6 furnished the respective calix[4]semiture 19 in 14% yield. Notably, as in the case of tris(calixarene) 17, in both the above cases the Cu-catalyst loading was decreased (from 1.0 to 0.3 equiv. per bis(azide)) with respect to the reference reaction between the bis(azide) 4 and bis(alkyne) 5.⁸

In contrast, the CuAAC between the propylated bis(alkyne) 7 and the tetrakis(azide) 11 (Scheme 5) was successfully conducted under the conditions found previously for the reference reaction between compounds 4 and 7 (1.0 equiv. of CuI and a huge excess of Et₃N at heating),⁸ and no improvement was achieved upon varying the catalyst loading (see Table S4[†]). The tris(calixarene) 20 was obtained in 22% yield which is the highest one among the syntheses involving the tetrakis(azide) 11. The full-1,3-alternate calix[4]semiture 21 was obtained in 11% yield from calixarenes 8 and 11 using 1.0 equiv. of the



Scheme 5 Syntheses of calix[4]semitures 20 and 21: (i) CuI (1.0 equiv. per bis(azide)), Et₃N, toluene, 70 °C, 9 h; and (ii) CuI (1.0 equiv. per bis(azide)), Et₃N, toluene/1,2-dichloroethane, rt, 48 h.

CuI/Et₃N catalyst at room temperature, which were found to be the best conditions (Table S5;† 1,2-dichloroethane was used as the co-solvent due to the lower solubility of the calixarene **8** in neat toluene).

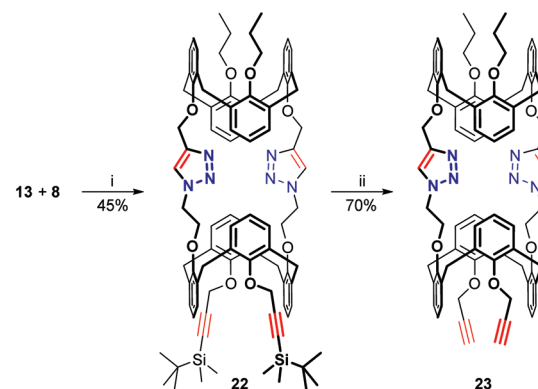
Lower yields of the hydroxylated calix[4]semitubes **18** and **19** in comparison with those of the isomeric semitubes **14** and **15** allow us to propose that two azide groups (which are weak copper-targeted ligands) in calixarenes **1** and **2** are relatively well pre-organized by the reacting Cu-cations during CuAAC to allow their simultaneous connection to the same bisalkyne site of the tetrakis(alkyne) **9**, whereas two propargyl groups in calixarenes **5** and **6** are not held together by the reacting cation(s) near the bisazide site of the calixarene **11**, thus favoring the formation of polymeric/oligomeric products. As for the propylated terminal units of the semitubes (of either a cone or 1,3-alternate shape), the difference in the reactivities of the respective bis(azides) and bis(alkynes) with the central cores **9** or **11** is not easy to rationalize, because too many factors must be taken into account, including shielding the copper cation(s) by the propyl groups (cone isomers) or inverted calixarene aromatics (1,3-alternate isomers), repulsion between the cations bound to bistriazole sites,^{9d} and their transfer between pairs of closely located azide^{9e} or acetylene groups.^{9a}

In general, though triazolated calix[4]semitubes can be obtained by CuAAC using the symmetrical central cores **9** and **11**, only the hydroxylated semitubes (especially the tris(calixarenes) **14** and **15** prepared in good yields) seem to have synthetic perspectives upon further chemical transformations of the OH groups. As for the semitubes constructed from the exhaustively alkylated calixarenes, further development in this direction can be seriously hampered by the need for condition tuning for each combination of the CuAAC counterparts, by the low yields in the syntheses, and by the difficulties in the separation of triazolated by-products from the targeted semi-tubular assemblies.

Stepwise assembling of triscalixarene semitubes

Unlike the tetrafunctional molecules **9** and **11**, the partially protected azide/alkyne-containing calixarene **13** allows a stepwise synthesis of long triazolated calix[4]semitubes when involved in CuAAC with bis(acetylenes) followed by de-protection and further reaction with bis(azides). The obvious advantage of this approach is that two but not four triazole heterocycles are formed during each of the CuAAC steps, which shrinks the range of triazolated by-products and thus improves the yield of the targeted semi-tubular bis(calixarenes). On the other hand, the stepwise syntheses using the calixarene **13** cannot replace the one-step preparation of the *C*_{2v}-symmetrical calix[4]semitubes **14**, **15**, and **17–21**, as only the *C*_s-symmetrical calix[4]semitubes can be obtained by the stepwise approach.

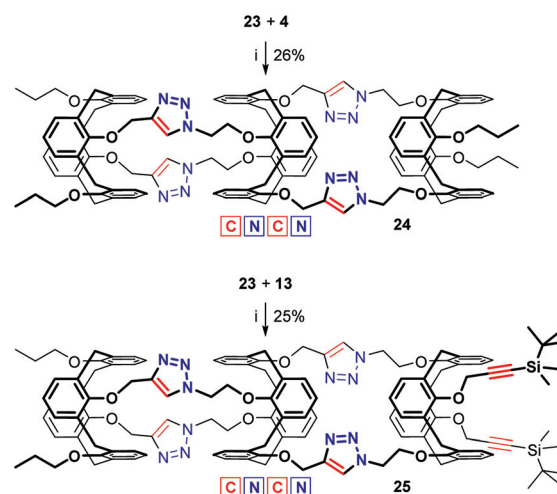
The CuAAC between calixarenes **13** and **8** (Scheme 6) was conducted using a copper(II) sulphate/sodium ascorbate catalytic system following the conditions previously found for assembling calixarenes **4** and **8** into a biscalixarene semitube,⁸ which were inefficient for the preparation of the tris(calixarenes) **17** and **21** (see above). The yield of the silylated bis(calix-



Scheme 6 Syntheses of calix[4]semitubes **22** and **23**: (i) CuSO₄·5H₂O (1.0 equiv. per bis(azide)/bis(alkyne)), sodium ascorbate, THF/H₂O (5 : 1), 60 °C, 24 h; (ii) nBu₄NF-3H₂O, THF/H₂O (100 : 1), 50 °C, 12 h.

arene) **22** reached 45%, so no further tuning of the reaction conditions was required in this case. *tert*-Butyldimethylsilyl protecting groups were removed by treating the compound **22** with tetrabutylammonium fluoride in aqueous THF, and the calix[4]semitube **23** bearing two terminal acetylene groups ready for further CuAAC was obtained.

With no additional tuning of the reaction conditions, the bis(calixarene) **23** was reacted with the bis(azide) **4** for obtaining the triscalixarene semitube **24** in 26% yield (Scheme 7). This compound is the isomer of both the calix[4]semitubes **17** and **21** but has lower symmetry as the two pairs of triazole linkers have different orientations with respect to the central calixarene core. The difference in the molecular symmetry immediately affects the patterns in the NMR spectra of the semitubes, thus allowing clear distinction of these three structures (Fig. 4). In particular, the ¹H NMR spectrum of the less symmetrical calix[4]semitube **24** contains two signals from the



Scheme 7 Syntheses of calix[4]semitubes **24** and **25**: (i) CuSO₄·5H₂O (1.0 equiv. per bis(azide)/bis(alkyne)), sodium ascorbate, THF/H₂O (5 : 1), 60 °C, 24 h.

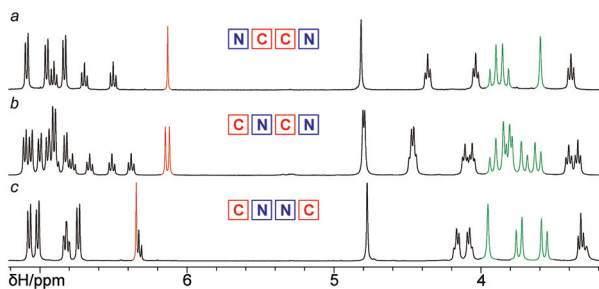


Fig. 4 Parts of the ^1H NMR spectra of (a) the calix[4]semiture 17, (b) calix[4]semiture 24, and (c) calix[4]semiture 21; 400 MHz, CDCl_3 ; the signals from the triazole protons are colored red and those from the methylene groups of the calixarene cores are colored green.

triazole protons and three sets of signals from the methylene bridges of three different calixarene cores (Fig. 4b), while for the semitubes 17 (Fig. 4a) and 21 (Fig. 4c) single resonance from the triazole protons and only two sets of signals from the methylene bridges appear in each case.

Similar to the synthesis of the calix[4]semiture 24, the propargylated bis(calixarene) 23 was reacted with the bis(azide) 13, which was the synthetic precursor of the bis(calixarene) 23 (Scheme 7). As a result, the triscalixarene semiture 25 having protected acetylene units in the structure was obtained in 25% yield. This allows us to expect that further growth of the tubular architectures will be possible by step-by-step attachment of the bis(azide) 13 to the propargylated semitubes using alternating de-silylation and CuAAC procedures. It seems that the stepwise approach enables more possibilities for the preparation of the long triazolated calix[4]semitures including those having additional functional groups in the terminal calixarene units, which can be achieved by replacing the propyl groups with the functional ones in the bis(alkyne) 8 or/and in the bis(azide) 4.

The prepared triscalixarene semitures of different types seem to be able to form complexes with transition metal cations, which follows from our previous studies on the receptor properties of triazolated calix[4]arenes^{9d} as well as from the preliminary data on the complexation ability of the related bis-calixarene semitures towards silver and zinc cations.⁸ Still, taking into account the homo/heteroditopic nature of the triscalixarene semitures and possible inter/intramolecular migrations of the bound cations between the triazole-containing binding sites, the evaluation of their receptor abilities requires the introduction of additional reporter (e.g. fluorescent) or other functional/receptor units to the molecules, which can be achieved using the synthetic approaches developed in this work.

Conclusions

We have shown that long triazolated calix[4]semitures having three calixarene units in the structures can be assembled by using bifunctional and tetrafunctional calixarene azides and

alkynes under CuAAC conditions. The four-fold CuACCs between the bifunctional terminal units of the cone or 1,3-alternate shape and the tetrafunctional 1,3-alternate central core have been shown to have limited applicability as condition tuning is required for each of the calixarene combinations, while the separated yields of the targeted assemblies have been modest in most cases. As an alternative, the two-step syntheses of the triscalixarene semitures, which involved the calix[4]arene bearing two azide groups and two silylated acetylene groups as the key synthone, have been shown to be successfully conducted under similar conditions at each of the two-fold CuAAC steps and have allowed the preparation of the semitures with different substituents in the terminal units or/and further growth of the semi-tubular assemblies.

It is believed that the obtained results will find application in the development of multitopic tubular hosts having different mutual orientations of the bistriazole cation-binding sites within the molecule, which in turn will allow the creation of stimuli-switchable and other sophisticated supramolecules operating by intramolecular oscillations of a bound cation between the binding sites.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The financial support from the Russian Science Foundation (Grant No. 20-73-00109) is gratefully acknowledged. X-ray diffraction studies were performed at the Centre of Shared Equipment of IGIC RAS.

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