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1 About the Author



Oliver Sutcliffe graduated from the University of Liverpool with a BSc (Hons) in Chemical Sciences (1996) and subsequently obtained a PhD in Synthetic Organic Chemistry working on *"New Extended Dipolar Systems"* under Drs Richard C. Storr and Thomas L. Gilchrist from the same institution. In 1999, he joined Professor Martin R. Bryce's group at the University of Durham as a postdoctoral fellow investigating the application of enantiopure ferrocene ligands in asymmetric synthesis, before moving to industry and working as a medicinal chemist on the design, development and optimisation of drug molecules for a broad-range of biological targets, including cancer. He moved to the University of Strathclyde in 2006, obtained an MSc in Pharmaceutical Analysis and after a period at the European Directorate for the Quality of Medicines and Healthcare (EDQM) in

Strasbourg was appointed as a lecturer in the Strathclyde Institute of Pharmacy and Biomedical Sciences (SIPBS). In 13 14 January 2012, Dr Sutcliffe joined the staff at Manchester Metropolitan University as both a Senior Lecturer in Pharmaceutical and Analytical Chemistry and Co-Director of the Manchester Centre for the Study of Legal Highs 15 (MCSLH). Dr Sutcliffe's research interests include the isolation, identification and analysis of "new psychoactive 16 17 substances" (formally known as "legal highs") such as mephedrone and its derivatives, and the development of simple, rapid and selective presumptive tests and devices which can be applied by "non-specialists" for the detection 18 19 of illicit substances in the field. He is the co-inventor of 10 international patents and co-author of 33 peer-reviewed 20 research articles.

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33 **Key term 1:** Cucurbit[*n*]uril

Cucurbit[*n*]urils are supramolecular structures made of glycoluril monomers linked by methylene bridges. The oxygen atoms are located along the peripheral edge of the macrocycle and are tilted inwards, forming a partly enclosed hydrophobic cavity that has been shown to bind to a range of organic molecules and inorganic ions. The name is derived from the resemblance of this molecule with a pumpkin of the family of *Cucurbitaceae*.

38

39 **Key term 2:** Hemicucurbit[*n*]uril

Hemicucurbit[n]urils are supramolecular structures made of N-substituted ethyleneureas monomers linked by methylene bridges. These macrocycles resemble a cucurbit[n]uril bisected along its equator where the ethyleneurea subunits adopt an alternating *"zigzag"* conformation. Unlike cucurbit[n]urils, hemicucurbit[n]urils have a limited ability to form inclusion complexes with organic compounds and do not to bind to inorganic ions.

44

45 **Key term 3:** Bambus[*n*]uril

Bambus[*n*]urils are supramolecular structures made of 2,4-disubstituted glycoluril monomers linked by methylene bridges. These macrocycles combines the features of both cucurbit[*n*]urils and hemicucurbit[*n*]urils to give a hyperbolic three-dimensional structure where the glycouril subunits adopt an alternating *"zigzag"* conformation and allow the formation of inclusion complexes. The name is derived from the resemblance of this molecule with the bamboo plant stem of the family of *Bambusoidea*.

51

52 Key term 4: Microwave Synthesis

53 Microwave synthesis is a rapidly expanding field in both organic and inorganic chemistries where microwave energy 54 is focussed directly (and efficiently) into the reactive components of a mixture, rather than being transferred from 55 the bulk solvent by convection/conduction (as occurs in conventionally heated reactions). This technique has been 56 applied to a wide-range of chemical processes, using both solution- and solid-phase reagents, and is able to greatly 57 enhance the rate and overall yield of many reactions leading to products, which in general, require minimal work-up 58 and purification.

61

Classical and microwave-assisted synthesis of cucurbit[n]urils, hemicucurbit[n]urils and bambus[n]urils

62

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64

65 Abstract

This chapter gives a concise overview of the synthetic and supramolecular chemistry of cucurbit[*n*]uril (CB[*n*]) family of macrocycles and expands it to hemicucurbit[*n*]urils (HemCB[*n*]) and bambus[*n*]urils (BU[*n*]). The application of microwave-assisted synthesis to access these macrocycles is reviewed. Several extensive papers describing the synthesis, physical and recognition properties and applications of CB[*n*]s have been published and the reader is directed to these key sources throughout this chapter.

71

72 1. Introduction

The toroidal cucurbit[n]uril (CB[n]; n = 5, 6, 7, 8 or 10) (1) family of macrocycles represent a unique class of 73 74 compounds that display a variety of host-guest interactions [1 - 7]. Their symmetrical, oxygen-lined portals give 75 access to a hydrophobic cavity that has been shown to bind organic or inorganic compounds, gases and alkali earth metal salts with binding constants of up to 10^{15} M⁻¹. Hemicucurbit[*n*]urils (**2**) and bambus[*n*]urils (**3**) can be viewed as 76 structurally-related to cucurbit[n]urils where the former are bisected along their equators and adopt a "zigzag" 77 78 conformation [1], whilst the latter combines the structural features of both cucurbit[n]urils and hemicucurbit[n]urils 79 to give a hyperbolic three-dimensional arrangement [8] (Figure 1). Cucurbit[n]urils have been extensively reviewed 80 in terms of their synthesis, properties and potential applications in both chemical and pharmaceutical sciences and 81 the author directs the reader to the recent series of articles published in the Israel Journal of Chemistry for more in-82 depth information regarding selected aspects of CB[n] chemistry [7]. This chapter seeks to provide a concise overview of these macrocycles, their synthesis and applications and how the field has expanded to the structurally-83 84 related hemicucurbit[n]uril and bambus[n]uril families.



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Figure 1. Structures of cucurbit[n]uril (1), hemicucurbit[n]uril (2) and bambus[n]uril (3)

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90 **2.** Synthesis of cucurbit[*n*]urils, hemicucurbit[*n*]urils and bambus[*n*]urils

The large-scale production of CB[n]s for commercial and industrial applications, particularly drug delivery, requires 91 reliable methods of synthesis that are effective in terms of both the ready availability of cheap starting materials and 92 energy consumption. Cucurbit[n]urils (6) are typically made through the condensation of glycoluril (5, R = H) and 93 formaldehyde (or paraformaldehyde) in hot concentrated mineral acids (Scheme 1, Method A) [1 – 7, 9]. By changing 94 95 the type of acid used, the molarity of the acid and/or the concentration of glycoluril, the distribution of the CB[n]96 products can be adjusted. For instance, synthesis in concentrated sulfuric acid typically produces samples containing 97 88% CB[6] with approximately 12% CB[5] and inconsequential amounts of CB[7] and CB[8] [9]. Alternatively, the use of 5M hydrochloric acid produces mixtures containing up to 35% CB[7] through the use of very high concentrations 98 of glycoluril (*circa*. 1700 mg mL⁻¹) and concentrated hydrochloric acid solutions produce mixtures containing up to 8% 99 100 CB[8] [9].



103

104	Scheme 1.	Synthesis of cucurbit[n]urils($n = 5, 6, 7, 8$ or 10) and functionalised cucurbit[n]urils ($n = 5, 6$ or 7).
105		Method A ($n = 5 - 10$ [R = H); $n = 5$ or 6 [R = Me or cyclohexyl-]); Method B ($n = 5 - 10$, R= R ₁ = H);
106		Method C ($n = 5 - 10$, R = H, see Scheme 6 for conditions).

107

Efforts to further extend the supramolecular chemistry and improve the solubility of CB[n] in common solvents led to 108 109 the development of efficient synthetic methods to prepare functionalized CB[n] derivatives. Principally three routes were envisaged: (i) condensation of glycoluril ($\mathbf{5}$, $\mathbf{R} = \mathbf{H}$) with substituted aldehydes to introduce substituents at the 110 111 methylene bridgehead (4, Scheme 1, Method B); (ii) acid-catalysed condensation of substituted glycolurils (5, R = alkyl, aryl etc.) with formaldehyde (Scheme 1, Method A) and finally (iii) direct functionalization of cucurbit[n]uril 112 itself (for example: $6 \rightarrow 7$) (Scheme 1). The first route was unsuccessful and the second has met with little success 113 with only Me₁₀CB[5] (**6**, n = 5, R = Me), Me₁₀CB[6] (**6**, n = 6, R = Me) and the cyclohexano-derivatives (**6**, n = 5 or 6, R = 114 cyclohexano-) reported via this route [1]. CB[n] (n = 6 or 7) analogues utilising bis(phthalylhydrazide) and substituted 115 116 glycoluril(bis-cyclic ether) as the building blocks have also been reported and show significant improvements in solubility in organic and aqueous media depending on the substituents present and have a similar binding capacity to 117 CB[6] [1]. 118

119

Partially substituted CB[*n*] derivative, Ph₂CB[6] (**9a**, R = Ph) was first prepared by Nakamura *et al.* by sulphuric acidcatalysed mixed condensation of formaldehyde with diphenylglycoluril (**5**, R = Ph) and glycoluril in a 1:5 ratio [1, 3] with Kim *et al.* patenting the synthesis, *via* di(*o*-nitro)phenylCB[6] (**9b**, R = *o*-NO₂Ph) and di(*p*-nitro)phenylCB[6] (**9c**, R = *p*-NO₂Ph-), of di(*o*-amino)phenylCB[6] (**10a**, R = *o*-NH₂Ph) and di(*p*-amino)phenylCB[6] (**10b**, R = *p*-NH₂Ph-) shortly thereafter [3] (Scheme 2). Other partially substituted derivatives, namely Me₄CB[6] and (Me₂CyP)_nCB[6] (cyclopentano, CyP), have been reported [1, 3].



an oxidant to provide the multi-functionalised derivative, $(HO)_{2n}CB[n](7, n = 5, 6 \text{ or } 7, R = H)$, in low to moderate yield 132 (5 - 45% yield) (Scheme 1). Subsequent functionalization with either an alkyl bromide or anhydride yielded the 133 corresponding alkoxy (7, R = R₁) or acyloxy- (7, R = COR₁) products which have exploited in numerous biological 134 applications [3, 10]. Scherman et al. refined the persulfate oxidation of CB[6] facilitating isolation of (HO)₁CB[6] (11) 135 in 12% yield (Scheme 3). Concomitant transformation into the propargyloxyCB[6] (12) facilitates Huisgen azide-136 acetylene click reaction to generate the self-complexing CB[6] derivative (13) [11]. Isaacs et al. who recently 137 reported the gram scale synthesis of a number of monofunctionalized CB[6] derivatives by the reaction with 138 substituted phthalaldehydes, have built on this work and published the preparation of phenol-substituted CB[6] (14), 139 its transformation into propargyloxy compound (15). Cycloaddition of (15) in the presence of tris(triazolyl)methanol-140 141 Cu(I) (Pericás' catalyst) [12] to give stimuli-responsive triazole (16) (Scheme 3) [13].



Functionalization of cucurbit[n]urils using Huisgen azide-acetylene click reaction



144

145

Scheme 3.

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147 Miyahara and Buschmann have independently reported that acid-catalysed condensation reactions using equimolar 148 amounts of ethyleneureas (18, R = H) with formaldehyde in 4N hydrochloric acid (at room temperature) provides access to hemicucurbit[n]urils (Scheme 4, Method A, n = 6) in 94% yield [1, 14, 15]. Hemicucurbit[n]urils are cyclic 149 and structurally-related to cucurbit[n]urils where the former are bisected along their equators as demonstrated by 150 their X-ray structures which feature an alternate (or "zigzag") arrangement of the six ethylene urea units and a 151 chloride ion (which is believed to act as a templating agent during the cyclocondensation process) present within the 152 cavity. The larger macrocycle, HmCB[12], can be obtained in excellent yield and purity by heating an equimolar 153 mixture of (18, R = H) and 37% formaldehyde in 1N HCl at 55 °C for three hours (Scheme 4, Method B, n = 12, 93% 154 yield) and subsequent purification by recrystallization with chloroform [14]. The X-ray crystal structure of 18 (R = H) 155 has been determined and indicates cyclic structural architecture exhibiting an analogous "zigzag" conformation as 156 the hexamer with several chloroform molecules in its crystal lattice. In contrast to CB[n]s, hemicucurbit[n]urils are 157 soluble in non-polar solvents and exhibit significantly different behaviour in terms of their ability to selectively 158 complex to small molecules (e.g. formamide, CHCl₃ and propargyl alcohol), anions (e.g. Cl⁻, SCN⁻, Br⁻ and I⁻) and 159 cations (Co^{2+} , Ni^{2+} and UO_2^{2+}) but not alkali or alkaline earth metals, Ag^+ or NH_4^+ ions [1, 3, 5, 14, 15]. 160



162

163

Scheme 4. Synthesis of hemicucurbit[*n*]urils and chiral hemicucurbit[*n*]urils

165

166 Recently Aav et al. have disclosed the chiral (R, R)-cyclohexylhemicucurbit[6]uril and (S, S)cyclohexylhemicucurbit[6]urils (17), prepared (as their corresponding hydrochloride and hydrobromide salts) from 167 equimolar amounts of the enantiopure perhydrobenzimidazolines (18, R = (S, S)- or (R, R)-cyclohexyl) and 168 paraformaldehyde in 4M hydrochloric (or hydrobromic) acid at 70 °C in 75 – 85% yield after flash chromatography 169 (Scheme 4, Method C). The enantiopure cyclohexylhemicucurbit[6]urils formed 1:1 complexes with Cl., Br., 170 carboxylic and thiolacetic acids and amines. Diastereomeric complexes with (R)- and (S)-O-methylmandelic acid in 171 172 organic media were also prepared [16].

173

Bambus[n]urils (BU[n]s), which combined the structural features of both cucurbit[n]urils and hemicucurbit[n]urils, 174 were first disclosed by Sindelar et al. in 2010 and subsequently patented in 2011 [8, 17]. The cyclic hexamer, 175 Me₁₂BU[6] (20, R = Me, as the HCl salt) was prepared as a white powder, in 30% yield, by heating 2,4-176 dimethylglycoluril (21, R = Me) and paraformaldehyde in 5.4M HCl for 24h (Scheme 5, Method A). The structure of 177 BU[6] was confirmed by X-ray analysis of microcrystals (formed by slow evaporation of BU[6] in EtOH-CHCl₃-178 179 tetrabutylammonium chloride) and indicated a cyclic structure exhibiting a hyperbolic conformation (3, Figure 1) with the chloride anion situated within the 6.4 Å cavity and the tetrabutylammonium cation located outside the 180 macrocycle. 181



185

Synthesis of bambus[n]urils (n = 4, 6, 12). Method A (reflux, 24h); Method B (reflux, 24h); Method C Scheme 5. (reflux, 22h); Method D (200W, 75 °C, 72h); Method E (200W, 110 °C, 4h) 186

187

Subsequent to this report a number of bambus[n]urils were synthesised, from their prerequisite 2,4-disubstituted-188 glycoluril (**20**, R = Pr, Bn) derivatives [18], using either: (i) an equimolar mixture of paraformaldehyde and p-189 toluenesulfonic acid (pTSA) in refluxing chloroform (Scheme 5, Method B) or (ii) an equimolar mixture of 190 paraformaldehyde, p-toluenesulfonic acid and tetrabutylammonium iodide (TBAI) in refluxing toluene (Scheme 5, 191 Method C). Anion-free bambus[n]urils were originally prepared by sequential displacement of the chloride anion 192 within the cavity with a suitable solvent (e.g. MeCN), however, more robust methods involving displacement of the 193 194 chloride anion with an iodide ion and the concomitant removal via oxidation (with hydrogen peroxide in the absence of light; 92% yield) or photocatalytically (using flash photolysis and titanium dioxide; 84% yield) have been recently 195 reported [19]. 196

197

Bambus[n]urils exhibit solubility in both organic and aqueous media and are primarily capable of forming stable 198 complexes with various anions (e.g. Cl⁻, Br⁻, l⁻, BF₄⁻, NO₃⁻ and CN⁻) with significant selectivity [8, 17 – 19]. A number of 199 theoretical studies of the affinity of BU[6] with various cations (e.g. H⁺, Cs⁺) and anions (e.g. F⁻, CN⁻) using quantum 200 mechanical density functional theory [20, 21] have also been reported. 201

202

The application of microwave-mediated synthetic chemistry has grown in significance over the last two decades and 203 it is now routinely used in the chemical and pharmaceutical industries for parallel and combinatorial synthesis. Since 204 microwave energy (operating at a frequency of 2.45 GHz) can be focussed directly into the reactive components [22, 205 23], rather than being transferred from the bulk solvent by convection/conduction (as occurs in regular hotplate/oil 206 bath reactions), microwave irradiation is able to greatly enhance the rate and overall yield of many reactions [22]. It 207 is common for microwave-assisted processes to be significantly faster than 'conventional' methods; taking minutes 208

instead of days and/or seconds instead of hours [22, 23]. Microwave-mediated processes are, therefore, more
 economic in terms of overall energy efficiency, but also because fewer side-products are (in general) formed during
 these processes leading to simple work-up and purification of the desired targets [22, 23].

212

213 The application of microwave-assisted synthesis for the production of cucurbit[n]urils (Scheme 1, Method C) was first patented by Kim et al. using an equimolar mixture of (5, R = H) and paraformaldehyde in 9M sulphuric acid. 214 Microwave irradiation for 45s gave a mixture of CB[5] (15%), CB[6] (45%), CB[7] (20%) and CB[8] (15%) respectively 215 [24]. Sutcliffe and Wheate reinvestigated and optimised the application of microwave-assisted synthesis in the 216 production of CB[n]s and by varying the reaction time (1 - 10 mins), temperature $(100 - 200 \degree \text{C})$ and the type of acid 217 used to promote the cyclocondensation reaction. The study indicated that after irradiation (160 °C) a mixture of 218 CB[n]s was produced using either 12M hydrochloric (Scheme 6, Method B) or 98% sulfuric acids (Scheme 6, Method 219 C) [25]. Hydrochloric acid, after 10 minutes, was found to produce a mixture of CB[5] (23%), CB[6] (58%), CB[7] (13%) 220 221 and CB[8] (6%) which approximated the distribution reported by Kim (Scheme 6, Method A), whilst sulfuric acid, after 3 minutes, produced mostly CB[6] (circa. 90%) with small amounts of CB[5] and CB[7]. 222

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Scheme 6.

Microwave-assisted synthesis of cucurbit[n]urils (n = 5, 6, 7, 8 or 10).

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The type of the acid used was a critical factor as phosphoric, formic, acetic and trifluoroacetic (TFA) acids failed to produce any of the desired CB[*n*]s and nitric acid was unsuitable due to a violent off-gassing that occurred during dissolution of the starting materials. Mixtures of strong and weak acids (e.g. H₂SO₄:TFA in ratios between 20:80 and 80:20) produce CB[*n*] less efficiently and with CB[6] as the prominent macrocycle. Scale-up of the reaction by 2-4x was shown to be viable with the distribution of the CB[*n*]s produced remaining unchanged.

Though the application of microwave-assisted synthesis has not, to date, been applied to the preparation of 234 hemicucurbit[n]urils, Sutcliffe and Wheate's method for cucurbit[n]urils has been successfully applied to the 235 synthesis of novel bambus[n]urils by Heck et al. (Scheme 5, Methods D and E) [26]. Irradiation of an equimolar 236 mixture of 2,4-allylglycoluril (21, R = Allyl), paraformaldehyde and catalytic p-toluenesulfonic acid in chloroform (75 237 °C, 72h, Scheme 5, Method D) afforded allyl₈BU[4] (23, n = 4, 85% yield) and irradiation of an equimolar mixture of 238 239 paraformaldehyde, p-TSA and TBAI in toluene (110 °C, 4h, Scheme 5, Method E) gave the hexamer, allyl₁₂BU[6] (23, n = 6, R = R_1 = allyl), in 60% yield – which was vastly superior to the classical route (*circa*. 20% yield). Subsequent 240 manipulation of allyl₈BU[4] (23, n = 4, $R = R_1 = allyl$) was achieved through cross-metathesis with Hovyeda-Grubbs 241 242 catalyst (90% yield) and concomitant hydrogenation with Pd-C (100% yield) providing access to three further examples of this emerging class of macrocycles: propen₈BU[4] (**23**, n = 4, $R = R_1 = CH=CH_3$); allyl₇HepBU[4] (**24**, n = 1, 243 4, R = allyl, $R_1 = CH_2-CH=CH-(CH_2)_3CH_3$) and $Pr_8BU[4]$ (23, n = 4, R = $R_1 = propyl$). 244

245

246 3. Applications

The high binding affinity and selectivity of cucurbit [n] urils has attracted significant attention, as such, CB[n]s have 247 demonstrated enormous potential in a range of applications, including microelectronics and nanomachine 248 components, chromatography and waste remediation, drug delivery, biosensors, and self-assembly and reaction 249 250 catalysis [1 – 7]. The construction of self-assembling molecular machines, which can switch between two different 251 states via a chemical, photochemical or electrochemical stimulus, is currently of interest and the CB[n] family is ideally suited for such applications due to their recognition properties. A detailed discussion of these applications is 252 outside the scope of this chapter, so we recommend the extensive reviews by Lagona et al. [1], Kim [3] and Isaacs [4]. 253 Functionalised cucurbit[n]urils have also used for a number biomimetic applications as artificial biological structures 254 and processes in living cells such as: selective ion channels, vesicles for targeted drug delivery and even gene 255 transfection [1, 3, 5, 7]. Hemicucurbit[n]urils have limited applications but have been utilised as supramolecular 256 catalysts in the esterification of 4-methoxy-4-oxobut-2-enoic, acrylic and benzoic acids using methanol [27] and 257 258 patented as potential cosmetic agents [28]. In contrast to the cucurbit [n] uril and hemicucurbit [n] uril families, the detailed recognition properties and practical applications of bambus[n]urils remain to be explored. 259

260

261 4. Future Perspective

262 Many of the early concerns with the cucurbit[*n*]uril family, such as poor solubility, have been circumvented, either by 263 the use of salts or by chemical modification of the macrocycle itself. Since the turn of the century, the scientific 264 literature has expanded greatly, aided by the commercial availability of many of the CB[*n*]s and new synthetic 265 procedures for the functionalization of these materials, which has widened their use greatly. The utility of CB[*n*]s in 266 an assortment of chemical (microelectronics, chromatography, waste remediation and self-assembly or reaction

catalysts) and pharmaceutical (nanomachines, drug delivery and biosensors) applications have been consistently 267 demonstrated in vitro and in a few in vivo examples. Their progress from "bench to bedside", however, is reliant on 268 the future development of functionalized CB[n]s and reproducibility of structurally consistent CB[n]s and their host-269 270 guest complexes. Cucurbit[n]urils have already been revealed to prevent the degradation of some drugs and diminish their toxicity. The development of new CB[n]s with suitable functional groups (e.g. amine and carboxylic acids) will 271 272 allow the growth of targeted drug-delivery vehicles through the attachment of biologically relevant substrates, antibodies, peptides or aptamers to the CB[n]s. To date, few biologically useful functionalized CB[n]s have been 273 274 reported and the application of novel and efficient synthetic methodologies (for example: microwave-assisted organic synthesis [MAOI]) may provide instrumental in the expansion of the CB[n]s family. 275

276 A requirement of pharmaceutical drug approval is to demonstrate the ability to produce new drugs (or, in this case, CB[n]-based drug-delivery systems) in a single form both robustly and reproducibly. It is well documented that CB[n]s 277 produce a range of different crystal polymorphs (i.e. the ability of a solid material to exist in more than one 278 crystalline form) based on CB[n]-drug interactions and the crystal's hydration state [25]. As such, the crystallization 279 280 processes of CB[n]s and their host-guest complexes needs to be scrutinized in great detail, in order to determine the number of different polymorphs which may form, the physicochemical processes that drive their crystallization and 281 solid-state packing, as well as methods to produce CB[n]-drug formulations suitable for clinical use (e.g., oral, liquid 282 or solid dosage forms). Though the application of hemicucurbit[n]urils and bambus[n]urils has, to-date, not been 283 284 exploited as much as their cucurbit[n]uril cousins their potential in biological, pharmaceutical and chemical science, especially in light of new derivatives being reported in the literature, is far from exhausted. 285

286

287 Executive Summary

- Cucurbit[n]urils (CB[n]s) are macrocycles made from the acid-catalyzed condensation reaction of glycoluril
 and formaldehyde.
- Hemicucurbit[n]urils (HemCB[n]s) are macrocycles made from the acid-catalyzed condensation reaction of
 ethyleneureas and formaldehyde.
- Bambus[*n*]urils (BU[*n*]s) are macrocycles made from the acid-catalyzed condensation reaction of 2,4 disubstituted glycoluril subunits and formaldehyde.
- CB[*n*]s have applications in many chemical, pharmaceutical and biomimetic fields, however, the application
 hemCB[*n*]s and BU[*n*]s have, to-date, not been exploited to their full potential.
- Microwave synthesis can applied as an efficient and low cost method on the industrial scale for the
 production of CB[n]s and BU[n]s for a variety of applications.

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