From dendrimers to fractal polymers and beyond

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The advent of dendritic chemistry has facilitated materials research by allowing precise control of functional component placement in macromolecular architecture. The iterative synthetic protocols used for dendrimer construction were developed based on the desire to craft highly branched, high molecular weight, molecules with exact mass and tailored functionality. Arborols, inspired by trees and precursors of the utilitarian macromolecules known as dendrimers today, were the first examples to employ predesigned, $1 \rightarrow 3$ $C$-branched, building blocks; physical characteristics of the arborols, including their globular shapes, excellent solubilities, and demonstrated aggregation, combined to reveal the inherent supramolecular potential (e.g., the unimolecular micelle) of these unique species. The architecture that is a characteristic of dendritic materials also exhibits fractal qualities based on self-similar, repetitive, branched frameworks. Thus, the fractal design and supramolecular aspects of these constructs are suggestive of a larger field of fractal materials that incorporates repeating geometries and are derived by complementary building block recognition and assembly. Use of terpyridine-$M^3$-terpyridine (where, $M = Ru, Zn, Fe$, etc.) connectivity in concert with mathematical algorithms, such as forms the basis for the Sierpinski gasket, has allowed the beginning exploration of fractal materials construction. The propensity of the fractal molecules to self-assemble into higher order architectures adds another dimension to this new arena of materials and composite construction.

Uniterms: Arborol. Dendrimer. Supramolecular. Terpyridine. Self-assembly.

The article discusses the transition from dendritic chemistry to fractal-based materials design. It highlights the iterative synthetic protocols used for dendrimer construction and explores the properties of arborols, which are inspired by trees. These materials are noted for their globular shapes, excellent solubilities, and ability to aggregate. The fractal design of dendritic materials is discussed, along with the use of terpyridine-$M^3$-terpyridine connectivity in self-assembly processes. The potential of these constructs is suggested as a basis for new classes of materials and composite designs.

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INTRODUCTION

Building block architectures are dependent on the physical and chemical properties of the desired material under construction; although at times, the properties of the new materials are not readily apparent based solely on the building block(s) design. This leads to an evolution in materials’ synthesis and utility, as well as an element of excitement and anticipation with regard to the discovery of unknown local and aggregate attributes. Such has been the paradigm of a class of polymers known as dendrimers (Newkome et al., 2001; Newkome, Shreiner, 2010; Moorefield et al., 2012).

From their first published reports in 1985, dendrimers and dendritic materials, characterized by a regularly repeating, branched motif that may be simply described as tree-like, have captured the attention of research groups around the globe. This is due in-part to several factors, including the potential to integrate the dendritic constructs with other materials in order to instill the desired physical or structural properties. This generates a tailored molecular assembly possessing enhanced physical and/or structural characteristics for a utilitarian outcome. For example, high molecular weight species can easily be made soluble in aqueous or non-aqueous media, based on the incorporated functionality (Yu et al., 2013). As well, and perhaps most importantly, with the advent of dendritic synthetic strategies, chemists now have better insight to incorporate precise control over the placement of all the components of macromolecular materials that could be fully and unequivocally characterized.

Dendrimers can be envisioned (Figure 1) by considering a core possessing multiple arms, where each arm or branch is subsequently connected to a preconstructed branched building block (or dendron, adopted from the synthon terminology) giving rise to an initial layer or generation. Repetition of the building block layering produces the next higher generation dendrimer. Continued elaboration produces higher generations. Starting with a core possessing 4 arms and using a 1 → 3 branched dendron will ideally afford dendrimers possessing 12, 36, 108, 324, and 972 termini or surface groups for generations 1, 2, 3, 4, and 5, respectively. Design aspects of dendrimers or dendrons encompass branching centers (e.g., C, N, aryl, etc.), arm length and rigidity (e.g., aryl, acetylenic, alkyl, etc.), terminal moiety (CO₂H, NH₂, OH, etc.). These parameters combine to influence the ultimate utilitarian physical properties, such as: void volume, internal hydrophobicity/lipophilicity, dielectric potential, and porosity, to mention but a few.

Notably, dendrimer construction has been conceived and developed in many variations. Building from the outside inward complements construction from the inside outward (convergent vs. divergent protocols). Dendron attachment has been effected using a wide variety of covalent and non-covalent protocols. As well, dendrimers can be constructed using linear building blocks, where branching centers are created during generational growth. Dendrimers have been examined in a numerous applications, including chemical (Kumar et al., 2008; Park et al., 2009) and electrical sensing, (Koo et al., 2001) micellar host-guest ability (Boisselier et al., 2010), coatings and polymer additives (Hartmann-Thompson et al., 2009), and drug delivery vehicles (Cheng et al., 2011).

Dendrimers have further been demonstrated to be examples of molecular fractals (Mandelbrot, 1982; Farin, Avnir, 1991; Cho et al., 2006) based on their repeating architectural motif at differing size scales. It is the fractal quality of the dendrimer that has given rise to new types of material in our laboratories that we have coined - fractal polymers.

As witnessed over the past 50 years, chaos science has introduced us to a methodology that permits us to observe order and patterns that were noted earlier as the simply the complexity in nature. Thus, the elements of chaos, fractals and dynamics have been introduced to mathematically address the complexity of nature (Peitgen et al., 2004). Notably, dendrimers are but a small avenue in the world of fractals. Molecular architectures exhibit self-similarity on differing scales that allow incorporation of a balance of interrelated attributes and structural compo-
nents that are ideally suited for interdependent component positioning. Towards these directions, we are working to design and assemble nanoscale homo- and heteronuclear materials possessing a non-branched morphology in which metal center incorporation and the structural architecture join with supramolecular concepts to facilitate material applications. The use of ligand-metal-ligand connectivity has thus served to expand our directed and self-assembly work into the novel, utilitarian “fractal” macro- and nanomolecular architecture arena (Newkome et al., 2006).

A fractal polymer with the attributes of a Sierpinski triangle can be envisioned geometrically by considering the transformation upon proceeding from a 3-directional, 2D dendrimer to a triangular, 1 → 3 branched motif that can be overlaid onto a Sierpinski gasket design, which was mathematically envisioned in 1916. The dendritic branches bisect the collective triangle sides’ at all midway points; way points; and hence, form the basis for the construction of a Sierpinski triangle (Figure 2).

The use of ligand-metal-ligand connectivity in these constructs is anticipated to lead to the development of materials with demonstrated potential as energy storage and release devices, based on stable oxidized and reduced metal states, as components of molecular devices, such as in new photovoltaic cells and organic light emitting diodes (OLEDs), based on their photo- and electroluminescence properties (Chen et al., 2010) and as molecular switches and optical display components (Yoon et al., 2010) based on their low spin-high spin characteristics. These attributes coupled with the potential to self-assemble the fractal materials into nanotube and nanofiber structures make them ideal candidates for bio-pharma applications. This review is intended to give a brief overview of past and on-going work, in our laboratories, as well as introduce the reader the the molecular world via traditional mathematics and the beauty of Mother Nature’s fractal patterns that are all around each of us.

**EARLY WORK**

Initial work in our laboratories (1982-1985) resulted in the first example of a one-directional, 1 → 3 C-branched dendrimer (Newkome et al., 1985) (1, Figure 3), that was described as an arborol for its likeness to tree architecture. Inspiration for the terminology and the molecular architecture was derived from the Leeuwenberg model for a tree, (Hallé et al., 1982; Tomlinson1983) which branched 1 → 3 where the branch juncture was suggestive of a tetrahedral, tetravalent carbon. The critical building blocks chosen to facilitate the construction of the 1 → 3 branched motif included triethyl methanetricarboxylate (Newkome, Baker, 1986) that could be easily attached to an alkyl halide by nucleophilic substitution and tris(hydroxymethyl)aminomethane, which would undergo facile amidation of the ester moieties. Each of these simple dendrons possessed the requisite tetrahedral carbon branching center to allow maximum branching in a carbon framework and notably the amide connectivity was highly stable and not prone to retro-reactions. Its spherical shape and aqueous solubility properties led to the suggestion in that paper that these highly branched structures would possess unimolecular micelle character.

This arborol was closely followed in time by the construction of another [9] alcohol-terminated arborol (Newkome et al., 1986b) using benzene as the core (2, where [9] denotes the number of hydroxyl groups at the arms ends). Its synthesis provided more insight into highly branched tree-like, architectures. Transmission electron micrograph images revealed globular microcrystalline regions, whereby molecular modeling calculations indi-

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**FIGURE 2** - Conceptual progression from branched dendrimer architecture to interrelated, non-tree-like fractal motifs.
cated the ~200 Å diameter microcrystals were comprised of ca. 40 molecules; subsequently, the potential for the arborols to aggregate revealed their micellar behavior and the potential for host-guest encapsulation, reinforcing our concept of unimolecular micellar properties. Light scattering experiments confirmed this behavior by revealing a 2.02 mM critical micelle concentration.

Shortly after the [9]arborol report, a new class of arborol (3) was developed and communicated. The two-directional [9]-n-[9] and [6]-n-[6] arborol series (Newkome et al., 1986a; Newkome et al., 1990) that each formed aqueous gels at concentrations less than 2 wt % (where [9] or [6] denote the number of end group hydroxyl moieties and n denotes the number of methylene groups in the alkyl chain connecting the two hydroxyl-terminated, dendritic units). Access to these new arborols was achieved in two steps by reacting either the Na+ salt of the methanetricarboxylate or malonate esters with an appropriate α,ω-dibromoalkane to give the corresponding nona- or hexa-esters, followed by treatment under basic conditions with the tris(hydroxymethyl)aminomethane dendron to give the desired arborol series.

The gels derived from these two-directional dendritic constructs exhibited birefringence when viewed between cross-polarized and plane-polarized light filters indicative of extended ordered structure. This was confirmed by transmission electron microscopy (TEM) that revealed linear, fiber-like strands with diameters of 35 Å, which corresponded well with a calculated CPK length of ca. 30 Å. It was postulated that these dumbbell-like macro-molecules stacked in a perpendicular manner to produce the aggregates possessing lipophilic interiors and hydrophilic exteriors. Fluorescence of chlortetracycline in the presence of the gel provided support for the proposed aggregation based on the dyes dependence on a lipophilic environment for emission.

UNIMOLECULAR MICELLES

The unimolecular micelle concept came to fruition with the synthesis of a dendrimer possessing an all-saturated hydrocarbon framework termed a Micellenoic Acid (Newkome et al., 1991a; Newkome et al., 1991b) (4, Figure 4) – the first example of an alkyl-based unimolecular micelle. Key building blocks leading to its construction were accessed by employing a radical initiated nitro group substitution with acrylonitrile at a tertiary carbon. This provided a critical three carbon homologation of a functional group from the requisite quaternary branching center, thereby allowing for easy manipulation. Transformation of the newly incorporated nitrile group to a terminal alkyne provided for facile attachment of the building block to an alkyl halide. Employing an alkyl tetrabromide as the core and the 1 → 3 C-branched, monoalkyne tris-benzyl ether dendron allowed construction of the 1st and...
2nd generation acid-terminated dendrimers by converting the surface groups to carboxylic acids via benzyl deprotection and subsequent hydroxyl oxidation.

Aggregation of the dendrimers in solution, promoted by carboxylic acid-based $H$-bonding, was prevented by ion exchange of the Na$^+$ salt to afford the tetraalkylammonium carboxylate. Confirmation of the exchange was evidenced in the electron micrographs of 4 that individually exhibited 30 Å molecular diameters and that compared favorably to the calculated values (Newkome et al., 1991b).

Using diphenylhexatriene (DPH), as a molecular probe, fluorescence lifetime and anisotropy decay values observed in phase resolved anisotropy experiments corresponded well to that reported with DPH in phosphatidylycholine vesicles, thereby demonstrating the micellar host-guest relationship in an aqueous environment. Additional molecular probes used to explore the micellar properties of these dendrimers included chlortetracycline (fluorescence in a lipophilic environment), phenol blue, and naphthalene (UV absorbance and solubilization by the dendrimer in an aqueous medium), and pinacyanol chloride (color change in the dendritic lipophilic interior).

Another unique family of dendrimers (e.g., 5) (Young et al., 1994) that has been shown to possess micellar properties was derived from a novel aminotriester, commonly known as Behera’s amine after the first person to synthesize it; prepared in two-steps starting with the treatment of MeNO$_2$ with tert-butyl acrylate, followed by reduction of the nitro group to the amine). This dendron facilitates branched growth by peptide-type coupling to a carboxylic acid end group followed by removal of the tert-butyl moieties to generate new acid termini for continued elaboration. Dendrimers up to the 5th generation, theoretically possessing 972 surface ester or acid functional groups, have been prepared using this amide-based, iterative technology.

Systematic evaluation of the effect of pH on their hydrodynamic radii using 2-dimensional, diffusion ordered $^1$H NMR spectroscopy (DOSY NMR) revealed the carboxylic acid series to be largest, or in an expanded state, in neutral and basic pH; whereas, the corresponding amine-terminated constructs exhibited contraction in basic media. Hydroxyl-terminated dendrimers exhibited a constant hydrodynamic radius over the range from basic to acidic pH. Environmental effects can thus be employed as containment and release mechanisms for encapsulated drugs.

As well, the amide-based, acid-terminated dendrimers were shown to be useful as a micellar substitute in the analytical technique of electrokinetic capillary chromatography. (Kuzdzal et al., 1994, 1997) A series of parabens and in a separate experiment, the components of cough syrup, were separated, thereby demonstrating the sensitivity of the method. The incorporation of 2,6-diamidopyridine units onto the internal arms of the dendrimer (6, Figure 5) placed four, hydrogen-bonding urea-type
receptor groups within the framework (Newkome et al., 1996).

The *H*-bonding encapsulation of the cancer fighting drug AZT (3’-azido-3’-deoxythimidine) was demonstrated with this dendrimer and later, their use as sensors for the detection of the presence of organic solvents was exploited (Miller et al., 1998).

We have employed the polyalkyne dendritic precursors of the micellanoic acid series as scaffolding for the site specific incorporation of decaborane (B$_{10}$H$_{14}$) within a dendritic framework due to interest in Boron Neutron Cancer Therapy (BNCT) (Newkome et al., 1994). Thus, the reaction of decaborane with the 1$^\text{st}$ and 2$^\text{nd}$ generation alkyne dendrimers afforded the poly(1,2-dicarba-closo-dodecaboranes) 7 and 8, respectively. The alcohol termini of the o-carborane superclusters were rendered water soluble by transformation to the corresponding polysulfate analogs.

**MOLECULAR FRACTAL DESIGN AND UTILITY**

The past several years have witnessed increased interest in the general topic of metallo-fractal materials, particularly in relation to their future nanotechnological applications. Thus, projected research directions will build on the melding of classical synthetic strategies with materials science construction and characterization protocols to fine-tune bulk and localized supramolecular properties to specific tasks, and to assemble macromolecular infrastructures capable of functioning alone or in concert within materials at composite interfaces.

The art of macromolecular construction has evolved to include utilitarian composite materials; whereby, interdependent components are precisely matched for the desired physicochemical properties. Targeted materials and properties, derived from minimum-assembly protocols, are therefore of interest from various perspectives that include the ease of architecturally complex construction and commercial viability. The preparation of new monomers with bonding metal center(s) leading to new predesigned macro- and nano-scale constructs includes: branching multiplicity and choice of branching moieties (e.g. mono- or polyatomic), length and flexibility of connector units, coupled with localized and aggregate supramolecular properties. Research efforts in our laboratories suggest that fundamental supramacromolecular properties can be affected by subtle changes in these design parameters.

These fractal constructs have led to the development of materials with demonstrated potential as: molecular batteries, switches, and optical display devices, to name but a few (Figure 6) (Schubert et al., 2006; Winter et al., 2011a,b; Friebe et al., 2012). Hence, our research directions necessarily encompass studies on terpyridine-metal-terpyridine “<tpy-M-tpy>” connectivity employing preconstructed synthons to facilitate the one-step construction (as opposed to the multi-step construction of dendrimers) of nano- and macroscopic, precisely positioned, polymetal arrays giving rise to new precise multicomponent macromolecular systems.

Initial work (Newkome et al., 1999) resulted in the single-step, high yield, self-assembly of a hexaruthenium macrocycle (9, Figure 7) possessing the ubiquitous benzeneoid architecture derived from the 120° juxtaposed ditopic ligand, 3,5-*bis*(terpyridinyl)toluene. The resultant metallo-macrocycle supports a 12$^+$ charge and was isolated as the
Cl\(^{-}\); ion exchange to the PF\(_6\)\(^{-}\) salt facilitated characterization and solubility in common organic solvents. Structural confirmation of this original hexameric, multi-metal array was supported by TEM imaging.

From the observation that these multimetal arrays readily aggregate into larger assemblies that may present opportunities to prepare new materials with unique properties, we have investigated the ion-promoted, stoichiometric self-assembly of nanoscale composite fibers. Combining a structurally rigid hexameric macrocycle \([\text{10}^{12+}]\)\((\text{PF}_6)^{-}\)\(_n\) and a flexible, dodecarboxylate-terminated, first generation, \(1 \rightarrow 3\) branched dendrimer \([\text{11}^{12-}]\)\((\text{Na}^{+})\)\(_n\); (see Figure 1 and the accompanying text for an introduction to these materials] gave rigid fibers (Wang et al., 2008) \([\text{10}^{12+}]\)\((\text{11}^{12-})\)\(_n\) as revealed by TEM images (12, Figure 8).

The hydrodynamic diameter for the dendrimer of 23.6 Å determined under basic pH conditions using 2D diffusion ordered spectroscopy (DOSY) NMR experiments (Young et al., 1994) is larger than the internal open area diameter (17.5 Å) of the hexamer. This suggests that an ordered, molecular packing of the two components could be based on the dendrimer fitting above (and below) the cavity of the hexamer, and only partially into it.

Insight into the composites molecular ordering was obtained with selected area electron diffraction (SAED) of the fibers (i.e., 12). Two strong electron diffractions perpendicular to the fibers long axis were observed with a \(d\)-spacing of 3.85 nm; diffused higher diffractions were also observed. As well, a \(d\)-spacing of 1.92 nm was obtained from a pair of intense diffraction patterns parallel to the fiber’s long axis. This suggests that the in-plane direction of \((\text{10}^{12+})\) is perpendicular to the long axis and that self-assembly results in alternating stacking of the two components. With the impetus to explore the properties of triangular metallacycles, (Schultz et al., 2011) three new ligands (13 - 15; Figure 9) possessing 60° direction-
ality were prepared by Suzuki coupling with known aryl 1,2-dihalides and 4-terpyridinylphenylboronic acid. Treatment of the 60°-based bisteryridines with either Cd\(^{2+}\) or Zn\(^{2+}\) generated a series of triangular materials (16 - 21) that were observed to form fibers following counterion exchange with benzenehexacarboxylate. The linear, hair-like fibers were obtained using a mixed solvent system prepared from dilute solutions (~1 mM) of metallocone in MeCN and sodium hexaborate in water. TEM images revealed large bundled structures with cross-sections of ca. 300 nm comprised of narrower fiberous strands of ca. 2.0 nm.

Rigorous characterization of these molecular triangles included electrospray ionization mass spectrometry (ESI-MS) in concert with traveling-wave ion mobility mass spectrometry (TWIM-MS), along with gradient tandem mass spectrometry (gMS\(^2\)) that corroborated the stability of the triangles observed in the TWIM spectra.

Powder X-ray diffraction (XRD) data were obtained from microcrystalline regions within the fibers. Combined with SAXD data orthorhombic unit cell dimensions of 39.82, 8.38, and 49.14 Å for a, b, and c, respectively, were determined and a packing model for the triangles in the fiber was suggested using computer generated molecular modeling (Figure 10).

In general, self-assembly techniques lead to the most thermodynamically stable product(s), while kinetic control is required to access the less stable products. For example, kinetic control can be affected by utilizing non-reversible connectivity to achieve the desired target or by the removal of the potential for the formation of the enthalpically favored product.

Whereas, we have demonstrated the high yield synthesis of trimeric metallocones by the self-assembly of 60°-based bisteryridines, the dimerization of these

![FIGURE 9](image_url) - Synthesis of a series of triangular metallocones and a TEM image of fibers readily formed when the molecular triangles are paired with benzenehexacarboxylate [Schultz, A.; Cao, Y.; Huang, M.; Cheng, S. Z. D.; Li, X.; Moorefield, C. N.; Wesdemiotis, C.; Newkome, G. R. *Dalton Trans.*, v.41, p.11573-11575, 2012. – Reproduced by permission of The Royal Society of Chemistry].

![FIGURE 10](image_url) - Proposed fiber packing model for triangle 20 [Schultz, A.; Cao, Y.; Huang, M.; Cheng, S. Z. D.; Li, X.; Moorefield, C. N.; Wesdemiotis, C.; Newkome, G. R. *Dalton Trans.*, v.41, p.11573-11575, 2012. – Reproduced by permission of The Royal Society of Chemistry].
bisligands using a strongly binding metal that renders the complex non-reversible leading to the formation of folded molecular squares (Schultz et al., 2012b) upon self-assembly with itself.

Starting with ligand 15 (Figure 9) dimerization was achieved by a stoichiometrically controlled reaction with RuCl₂(DMSO), in CHCl₃:MeOH (1:1 v/v) to generate the new bisligand 22 (Figure 11).

Self-assembly with Ru²⁺, Fe²⁺, Zn²⁺, and Cd²⁺ gave the corresponding tetramers 23 – 26, respectively. For the Cd²⁺ and Zn²⁺ products, purification was achieved by simple counterion exchange with NH₄PF₆; whereas, the Fe²⁺ and Ru²⁺ tetramers required column chromatography. Notably, a one-pot reaction of ligand 15 with FeCl₂·4H₂O afforded (8%) tetramer 24, along with the expected trimer in 29% yield after chromatography. In addition to NMR, ESI-MS, TWIM-MS, and photophysical characterization, gradient tandem MS (gMS²) revealed an order of stability for stability order of <tpy-M²⁺-tpy> as Cd²⁺ < Zn²⁺ < Fe²⁺ < Ru²⁺, as expected. Owing to the folded shape of the tetramers (Figure 12), the descriptive term “Dondorff” rings was coined after the first person to describe the giant manta ray.

In an earlier report, we delineated the concept of larger, dimerized, bisterpyridine ligands to afford a series of heteronuclear hexameric, octameric, and decameric metallo-cycles (Chan et al., 2011a) via stepwise self-assembly procedures. In this case, the new bisligand (27, Figure 13) was constructed using a 120°-based building block architecture. As in the case of the 60°-juxtaposed starting ligand, the formation of smaller metallocycles is circumvented by using a longer building block, as the precursor. Traveling wave ion mobility mass spectrometry (TWIM-MS) and molecular modeling provided insight into their unique sizes and conformational flexibility.

Dimer 27 was isolated (29%) following column chromatography from a single-pot reaction of the cor-

**FIGURE 11** - Synthesis of a series of square-shaped metallocycles using a modified bisterpyridine to avoid trimer formation [Schultz, A.; Li, X.; McCusker, J. K.; Moorefield, C. N.; Castellano, F. N.; Wesdemiotis, C.; Newkome, G. R. Chem. Eur. J., v.18, p.11569-11572, 2012. – Reproduced by permission of Wiley-VCH].

**FIGURE 12** – Side view of a computer generated model of a tetrameric Dondorff ring showing the folded motif [Schultz, A.; Li, X.; McCusker, J. K.; Moorefield, C. N.; Castellano, F. N.; Wesdemiotis, C.; Newkome, G. R. Chem. Eur. J., v.18, p.11569-11572, 2012. – Reproduced by permission of Wiley-VCH].
responding 120°-based bis(terpyridine) ligand with 0.5 equivalent of RuCl$_2$(DMSO)$_4$. Treatment of 27 with 1.05 equivalents of FeCl$_2$·4H$_2$O in MeOH afforded hexamer 28, octamer 29, and decamer 30 in 36, 9, and 2% yields, respectively, after column chromatography.

The specific geometries of the different macrocycles and their cross-sectional areas were investigated using molecular modeling and annealing simulations. The obtained collision cross-section vs. relative energy plots exhibited three distinct areas, corresponding to three major regions of conformations, circular, twisted stretched, twisted folded. Whereas, when rigid ligands are used to construct the macrocycles, the overall flexibility of the larger polycomplexes increases with size, leading to a wider range of possible geometries. Flexibility is thus greatest for decamer 30 (Figure 14), where several different conformers are possible.

Expanding on the concept of enlarging bisligand building blocks for the construction of otherwise not easily accessible macrocycles, a trimeric ligand 31 has been...
conceived and isolated. (Chan et al., 2011b) Following a two-step reaction sequence, whereby bisterpyridine 32 was treated with 2 equivalents of RuCl₂ to give the paramagnetic bisRu³⁺ adduct, which upon addition of 2 equivalents of 31 under reductive conditions afforded the desired trimer (Figure 15). Incorporating bisterpyridine 32 into the same two-step sequence gave (80%) the desired hexaRu²⁺ metallocycle 33. Reaction of trimer 32 with FeCl₂ in refluxing CH₃OH for 18 h generated the heteronuclear hexamer 34 and nonamer 35 in 49 and 14% yields, respectively. Along with characterization by ESI-TWIM-MS, ¹H and ¹³C NMR, absorption spectroscopy and molecular modeling, 2D DOSY (diffusion ordered spectroscopy) supported the similar ring sizes of the two hexamers, as well as the larger nonamer; diffusion coefficients of 4.57 × 10⁻¹⁰, 4.51 × 10⁻¹⁰, and 3.31 × 10⁻¹⁰ m²/s were recorded for the homo- and hetero-nuclear hexamers and heteronuclear nonamer, respectively (Figure 16).

FIGURE 15 – Construction of a trimeric building block facilitated the synthesis of a large, nine-membered ring and homo- and hetero-nuclear hexamers in good yields [Chan, Y.-T.; Li, X.; Yu, J.; Carri, G. A.; Moorefield, C. N.; Newkome, G. R. J. Am. Chem. Soc., v.133, p.11967-11976, 2011. – Reproduced by permission of The American Chemical Society].

Efforts aimed at the construction of pentameric metallomacrocycles have utilized carbazole derivatives to introduce the requisite angle (Hwang et al., 2005). Thus, employing a 3,6-disubstituted carbazole unit between two ligands gave a 105° angle between two the terpyridine ligands (i.e., 36, Figure 17). Treatment of the carbazole-based bisterpyridine with Ru²⁺, Fe²⁺ or Zn²⁺ gave the corresponding metall-o-cycles 37 – 39.

Employed as sensitizer materials for solar cell devices, discharge experiments were conducted with these materials. A similar fill factor (ff) was observed for the three electrodes studied, while both the short circuit photocurrent (j_sc) and the open circuit photopotential (V_oc) exhibited the best results for the Ru²⁺ metallopentacycle. As well, photoconversion efficiency (η) using light covering the visible region of the spectrum again showed that the Ru²⁺ pentamer 37 possessed the highest value.

FIGURE 16 – 2D Diffusion Ordered Spectroscopy (DOSY) NMR confirmed ring size [Chan, Y.-T.; Li, X.; Yu, J.; Carri, G. A.; Moorefield, C. N.; Newkome, G. R. J. Am. Chem. Soc., v.133, p.11967-11976, 2011. – Reproduced by permission of The American Chemical Society].

After modifying a 120°-based bisterpyridine ligand with an acetyl protected sugar moiety with the expectation of crafting a peripherally functionalized, hexameric metallomacrocycle, we isolated (4%) a unique pentameric construct, (Chan et al., 2010) along with the anticipated hexameric species (12%). Thus, the Fe²⁺-based pentamer 40 - characterized unequivocally by 2D COSY NMR and ESI-MS - represents the first example of a smaller ring being obtained from the self-assembly of rigid, 120°-juxtaposed terpyridine ligands. A noticeable difference in the ¹H NMR of the pentamer in contrast to the hexamer was the upfield shift of the absorption attributed to the 4ArH, that is oriented towards the interior of the ring; a slightly enhanced crowding effect for the five-membered specie was postulated. All other NMR absorptions for both rings was similar.
Both the five- and six-membered rings formed fibers when hexane was allowed to diffuse into a solution of the macrocycle in a CHCl₃:MeOH:MeCN mixture (8:3:1 v/v). In each case, fiber lengths and diameters of 10-80 nm and 20-30 nm were observed, respectively. Molecular modeling of the rings showed interior void regions of 1.3 nm for the 5-membered specie and 1.8 nm for the larger ring.

During 2009, our lab was searching for better thermodynamic control over the self-assembly of the metallomacrocycles; the use of Cd²⁺ to facilitate the terpyridine-terpyridine connectivity provided that kinetic vs thermodynamic balance. Thus, treatment of tert-butyl-modified bisterpyridine 41 with a stoichiometric amount of Cd(NO₃)₂·4H₂O in CHCl₃:MeOH (3:2 v/v) quantitatively generated the desired hexacadmium (Chan et al., 2009) macrocycle 42 (Figure 18). Due to the lability of the Cd-terpyridine coordination, the thermodynamically most stable product was formed; no other species (e.g., 5-, 7-, or 9-membered rings) were detected. The absence of oligomeric products in the reaction to form the macrocycles was verified by the preparation of a linear bisterpyridine building block, oligomerization with Cd, and ESI-TWIM-MS analysis for comparison to patterns and drift times observed for hexamer 42.

Ions selected for ion mobility separation included [6L+6Cd]⁺⁺ and [3L+3Cd]²⁺, based on the small number of ligand-to-metal possible combinations. Ion mobility separation signals for the cyclic and linear fragments generated from the cyclic material occurred at shorter drift times (i.e., 10.30, 6.45, and 5.13 ms for the [3L+3Cd]²⁺ ion and linear and cyclic [6L+6Cd]⁺⁺ ions, respectively) then the signals for same m/z 1187 amu-based fragments observed in the case of the linear construct (i.e., 11.55 and 6.59 ms for linear [3L+3Cd]²⁺ and the [6L+6Cd]⁺⁺ ions respectively). Drift times are longer for the linear building block material and no signals were seen near the lower drift time of 5.13 ms, which confirmed that the 5.13 peak corresponds to the cyclic [6L+6Cd]⁺⁺ species and the 6.45 ms peak corresponds to the linear [6L+6Cd]⁺⁺ species. Thus, the linear fragments generated from the cyclic material in the mass spectroscopy instrument arise solely from the cyclic specie.

In a dramatic example of the synthetic potential that complementary geometric and thermodynamic control can achieve, the first terpyridine-based, nanoscale, supramolecular spoked wheel (Wang et al., 2011) was constructed in our laboratory. A three-component, single-step, coordination-driven, self-assembly with tris- and hexakis-terpyridine polyligands, along with Zn²⁺ or Cd²⁺ ions reacted in a stoichiometric ratio, generated the D₆h symmetric product that was isolated in >94% yield. Characterization of the product included TWIM-MS, diverse NMR techniques, and TEM imaging.

Tristerpyridine 43 and hexakisterpyridine 44 (Figure 19) were each prepared using Suzuki-type, cross-
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Coupling reactions with 4-terpyridylphenylboronic acid and the corresponding tri- or hexabromobenzene. Subsequent reaction of the two polyterpyridine ligands with $\text{M(NO}_3\text{)}_2\cdot6\text{H}_2\text{O}$ (where $\text{M} = \text{Cd}^{2+}$ or $\text{Zn}^{2+}$) in a $\text{CHCl}_3/\text{MeOH}$ solution at 25 °C for 30 min in a 1:6:12 ratio, respectively, afforded the dodeca$\text{M}^{2+}$ complex 45. The polynitrate form was converted to the $24\ \text{PF}_6^-$ salt to facilitate characterization.

The TEM supported the size and geometry of individual macromolecules revealing hexagonal-shaped, shadow-like images possessing average diameters of $6.5 \pm 1.0 \text{ nm}$; this diameter corresponds well with the molecular modeling calculations showing a $5.6 \text{ nm}$ diameter for the optimized structure.

The ability to create new materials with terpyridine-based self-assembly is limited only by the building blocks that are available and the imagination of the artist. In a logical extension of the polyterpyridines used in the construction of the spoked wheel, other polybromides were sought that would allow access to unique starting materials. Ready availability of two additional tetrabromoaryl synthons facilitated the creation, via high yield Suzuki coupling reaction, of two isomeric tetra$k$is terpyridines (i.e., 46 and 47, Figure 20).

With these new terpyridines added to our molecular quiver, two polycyclic, macromolecular, constitutional isomers (Schultz et al., 2012a) were constructed (Figure 21). As in the case of synthesizing large ring structures, use of the dimeric terpyridine 48 restricted the degrees-
of-freedom of reaction and subsequently facilitated optimum conditions for isolation of either molecular bowtie 49 when reacted with tetrakisterpyridine 47 or the isomeric molecular butterfly motif 50 when treated with tetrakisterpyridine 46. Structural characterization of both constitutional isomers was facilitated by ESI-MS, TWIM-MS, $^1$H NMR and $^{13}$C NMR. Of particular interest are the notably different drift times observed in the ion mobility separation that unequivocally demonstrated the different sizes and morphologies of the isomers at high charge states.

In an effort to examine the outcome(s) of reacting building blocks with differing geometries (Lu et al., 2012) we constructed two new bisterpyridine ligands 51 and 52 (120°- and 60°-oriented, respectively). Reaction of these two building blocks with Zn(NO$_3$)$_2$·6H$_2$O in a precise 2:1:1 ratio in MeOH for 30 min at 25 °C afforded rhomboidal complex 53 (Figure 22). Analysis by ESI-MS revealed the presence of the triangular species 55, which can also be prepared in high yields by reaction of building block 52 with a metal. Performing the reaction under refluxing conditions for 24 h gave the same mixture. And, replacing Zn(NO$_3$)$_2$·6H$_2$O with Cd(NO$_3$)$_2$·4H$_2$O gave the Cd$^{2+}$-based product mixture of rhombus and triangle, as revealed by ESI-MS.

To further probe the self-assembly a rhomboidal structures, tristerpyridine 43 was prepared; notably, it is a structural hybrid of both the 60°- and 120°-based bisligands. Upon treatment of monomers 52 and 43 and Zn(NO$_3$)$_2$·6H$_2$O in a ratio of 2:2:5 in MeOH at 25 °C for 30 min, followed by ion exchange with NH$_4$PF$_6$, a slightly yellow precipitate (i.e., rhombus 54) was obtained in 95% yield. Unequivocal characterization was obtained with 2D COSY and NOESY NMR, along with ESI-TWIM-MS, which showed the complete absence of the triangle-based by-products (i.e., 55). It is also worth noting that the Zn-complex of 54 show remarkable stability in that it was observed to remain intact when subjected to MALDI-ToF-MS analysis in the linear mode. Normally, only Ru-based constructs have survived these conditions.

FIGURE 20 – Two new tetrakisterpyridine building blocks that facilitated the synthesis of constitutional isomers.

FIGURE 21 – Self-assembly of isomeric bowtie and butterfly bicyclic metallomacrocycles 49 and 50, respectively [Schultz, A.; Li, X.; Barkakaty, B.; Moorefield, C. N.; Wesdemiotis, C.; Newkome, G. R. J. Am. Chem. Soc., v.134, p.7672-7675, 2012. – Reproduced by permission of The American Chemical Society].
FIGURE 22 – Preparation of rhombohedral structures [Lu, X.; Li, X.; Wang, J.-L.; Moorefield, C. N.; Wesdemiotis, C.; Newkome, G. R. *Chem. Commun.*, v.48, p.9873-9875, 2012. – Reproduced by permission of The Royal Society of Chemistry].

CONCLUSIONS

“Prediction is difficult, especially of the future” (Niels Bohr). The transformation of nature’s chaos as seen in the eyes of mathematics to the design and construction of novel fractal macromolecular assemblies that can possess utilitarian applications in daily life is tomorrow’s challenge.

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REFERENCES

BOISSELIER, E.; LIANG, L.; DALKO-CSIBA, M.; RUIZ, J.; ASTRUC, D. Interactions and Encapsulation of Vitamins C, B<sub>3</sub>, and B<sub>6</sub> with Dendrimers in Water. *Chem. Eur. J.*, v.16, p.6056-6068, 2010.

CHAN, Y.-T.; LI, X.; MOOREFIELD, C.N.; WESDEMIOTIS, C.; NEWKOME, G.R. Towards larger polygonal architectures: synthesis and characterization of iron(II)- and Ru(II)-Bisterpyridine metallomacrocycles. *Chem. Eur. J.*, v.17, p.7750-7754, 2011a.
CHAN, Y.-T.; LI, X.; SOLER, M.; WANG, J.-L.; WESDEMIOTIS, C.; NEWKOME, G.R. Self-Assembly and traveling wave ion mobility mass spectrometry analysis of hexacadium macrocycles. *J. Am. Chem. Soc.*, v.131, p.16395-16397, 2009.

CHAN, Y.-T.; LI, X.; YU, J.; CARRI, G.A.; MOOREFIELD, C.N.; NEWKOME, G.R. Design, synthesis, and traveling wave ion mobility mass spectrometry characterization of Iron(II)- and Ruthenium(II)-Terpyridine macrocycles. *J. Am. Chem. Soc.*, v.133, p.11967-11976, 2011b.

CHAN, Y.-T.; MOOREFIELD, C.N.; SOLER, M.; NEWKOME, G.R. Unexpected isolation of a pentameric metallomacrocycle from an Fe(II)-Mediated complexation of 120º juxtaposed 2,2':6',2''-Terpyridine Ligands. *Chem. Eur. J.*, v.16, p.1768-1771, 2010.

CHEN, X.; DING, Y.; CHENG, Y.; WANG, L. Synthesis, spectroscopy and electroluminescence of cadmium(II) polypyridyl complexes. *Synth. Met.*, v.160, p.625-630, 2010.

CHENG, Y.; ZHAO, L.; LI, Y.; XU, T. Design of biocompatible dendrimers for cancer diagnosis and therapy: current status and future perspectives. *Chem. Soc. Rev.*, v.40, p.2673-2703, 2011.

CHO, T.J.; MOOREFIELD, C.N.; WANG, P.; NEWKOME, G.R. Metallodendrimers: fractals and photonics. *ACS Symp. Ser.*, v.921, p.186-204, 2006.

FARIN, D.; AVNIR, D. Surface fractality of dendrimers. *Angew. Chem. Int. Ed. Engl.*, v.30, p.1379-1380, 1991.

FRIEBE, C.; HAGER, M.D.; WINTER, A.; SCHUBERT, U.S. Metal-containing polymers via electropolymerization. *Adv. Mater.*, v.24, p.332-345, 2012.

HALLÉ, F.; OLDEMANN, R.A.A.; TOMLINSON, P.B. Tropical trees and forests: an architectural analysis. Berlim: Ed. Springer, 1982. 441 p.

HARTMANN-THOMPSON, C.; KEELEY, D.L.; ROUSSEAU, J.R.; DVORNIC, P.R. Fluorescent dendritic polymers and nanostructured coatings for the detection of chemical warfare agents and other analytes. *J. Polym. Sci. Part A: Polym. Chem.*, v.47, p.5101-5115, 2009.

HWANG, S.-H.; WANG, P.; MOOREFIELD, C.N.; GODÍNEZ, L.A.; MANRÍQUEZ, J.; BUSTOS, E.; NEWKOME, G.R. Design, self-assembly, and photophysical properties of pentameric metallomacrocycles: [M₅(N-hexyl[1,2-bis(2,2',6',2''-terpyridin-4-yl)carbazole])₅][M = Fe(II), Ru(II), and Zn(II)]. *Chem. Commun.*, v.37, p.4672-4674, 2005.

KOO, B.W.; SONG, C.K.; KIM, C. CO gas sensor based on a conducting dendrimer. *Sens. Actuators B*, v.77, p.432-436, 2001.

KUZDZAL, S.A.; MONNIG, C.A.; NEWKOME, G.R.; MOOREFIELD, C.N. Dendrimer electrokineic capillary chromatography: unimolecular micellar behaviour of carboxylic acid terminated cascade macromolecules. *J. Chem. Soc., Chem. Commun.*, p.2139-2140, 1994.

KUZDZAL, S.A.; MONNIG, C.A.; NEWKOME, G.R.; MOOREFIELD, C.N. A Study of Dendrimer-solute interactions via electrokineic chromatography. *J. Am. Chem. Soc.*, v.119, p.2255-2261, 1997.

LU, X.; LI, X.; WANG, J.-L.; MOOREFIELD, C.N.; WESDEMIOTIS, C.; NEWKOME, G.R. From supramolecular triangle to heteroleptic rhombus: a simple bridge can make a difference. *Chem. Commun.*, v.48, p.9873-9875, 2012.

MANDELBROT, B.B. *The fractal geometry of nature*. New York: W.H. Freeman and Company, 1982.

MILLER, L.L.; KUNUGI, Y.; CANAVESI, A.; RIGAUT, S.; MOOREFIELD, C.N.; NEWKOME, G.R. “Vapoconductivity” sorption of organic vapors causes large increases in the conductivity of a dendrimer. *Chem. Mater.*, v.10, p.1751-1754, 1998.

MOOREFIELD, C.N.; PERERA, S.; NEWKOME, G.R. *Dendrimer chemistry*: supramolecular perspectives and applications. Hoboken: John Wiley & Sons, 2012. p.1-54.

NEWKOME, G.R.; BAKER, G.R. The chemistry of methanetricarboxylic esters: a review. *Org. Prep. Proced. Int.*, v.18, p.117-144, 1986.
NEWKOME, G.R.; BAKER, G.R.; ARAI, S.; SAUNDERS, M.J.; RUSSO, P.S.; THERIOT, K.J.; MOOREFIELD, C.N.; ROGERS, L.E.; MILLER, J.E.; LIEUX, T.R.; MURRAY, M.E.; PHILLIPS, B.; PASCAL, L. Synthesis and characterization of two-directional cascade molecules and formation of aqueous gels. *J. Am. Chem. Soc.*, v.112, p.8458-8465, 1990.

NEWKOME, G.R.; BAKER, G.R.; SAUNDERS, M.J.; RUSSO, P.S.; GUPTA, V.K.; YAO, Z.; MILLER, J.E.; BOUILLION, K. Two-directional cascade molecules: synthesis and characterization of [9]-n-[9] arborols. *J. Chem. Soc., Chem. Commun.*, p.752-753, 1986a.

NEWKOME, G.R.; CHO, T.J.; MOOREFIELD, C.N.; BAKER, G.R.; SAUNDERS, M.J.; CUSH, R.; RUSSO, P.S. Self- and direct-assembly of hexaruthenium macrocycles. *Angew. Chem. Int. Ed.*, v.38, p.3717-3721, 1999.

NEWKOME, G.R.; MOOREFIELD, C.N.; BAKER, G.R.; JOHNSON, A.L.; BEHERA, R.K. Alkane cascade polymers possessing micellar topology: micellanoic acid derivatives. *Angew. Chem., Int. Ed. Engl.*, v.30, p.1176-1178, 1991b.

NEWKOME, G.R.; MOOREFIELD, C.N.; KEITH, J.M.; BAKER, G.R.; ESCAMILLA, G.H. Chemistry within a unimolecular micelle precursor: boron superclusters by site- and depth-specific transformations of dendrimers. *Angew. Chem., Int. Ed. Engl.*, v.33, p.666-668, 1994.

NEWKOME, G.R.; MOOREFIELD, C.N.; VÖGTL, F. *Dendrimers and dendrons*: concepts, syntheses, applications. Weinheim: Wiley-VCH, 2001.

NEWKOME, G.R.; SHREINER, C.D. Dendrimers derived from 1 - 3 branching motifs. *Chem. Rev.*, v.110, p.6338-6442, 2010.

NEWKOME, G.R.; WANG, P.; MOOREFIELD, C.N.; CHO, T.J.; MOHAPATRA, P.; LI, S.; HWANG, S.-H.; LUKOYANOVA, O.; ECHEGOYEN, L.; PALAGALLO, J.A.; IANCU, V.; HLA, S.-W. Nanoassembly of a fractal polymer: a molecular “Sierpinski Hexagonal Gasket”. *Science*, v.312, p.1782-1785, 2006.

NEWKOME, G.R.; WOOSLEY, B.D.; HE, E.; MOOREFIELD, C.N.; GÜTHER, R.; BAKER, G.R.; ESCAMILLA, G.H.; MERRILL, J.; LUFTMANN, H. Supramolecular chemistry of flexible, dendritic-based structures employing molecular recognition. *Chem. Commun.*, p.2737-2738, 1996.

NEWKOME, G.R.; YAO, Z.; BAKER, G.R.; GUPTA, V.K. Cascade molecules: a new approach to micelles. A [27]-Arborol. *J. Org. Chem.*, v.50, p.2003-2004, 1985.

NEWKOME, G.R.; YAO, Z.; BAKER, G.R.; GUPTA, V.K.; RUSSO, P.S.; SAUNDERS, M.J. Cascade molecules: synthesis and characterization of a Benzene[9]-Arborol. *J. Am. Chem. Soc.*, v.108, p.849-850, 1986b.

PARK, C.; IM, M.S.; LEE, S.; LIM, J.; KIM, C. Tunable fluorescent dendron-cyclodextrin nanotubes for hybridization with metal nanoparticles and their biosensory function. *Angew. Chem. Int. Ed.*, v.47, p.9922-9926, 2009.

PEITGEN, H.-O.; JÜRGENS, H.; SAUPE, D. *Chaos and fractals*: new frontiers of science. New York: Springer-Verlag, 2004.

SCHUBERT, U.S.; HOFMEIER, H.; NEWKOME, G.R. *Modern terpyridine chemistry*. Weinheim: Wiley-VCH, 2006.

SCHULTZ, A.; CAO, Y.; HUANG, M.; CHENG, S.Z.D.; LI, X.; MOOREFIELD, C.N.; WESDEMIOTIS, C.; NEWKOME, G.R. Stable, trinuclear Zn(II)- and Cd(II)-metalocycles: TWIM-MS, photophysical properties, and nanofiber formation. *Dalton Trans.*, v.41, p.11573-11575, 2012.

SCHULTZ, A.; LI, X.; BARKAKATY, B.; MOOREFIELD, C.N.; WESDEMIOTIS, C.; NEWKOME, G.R. Stoichiometric self-assembly of isomeric, shape-persistent, supramacromolecular bowtie and butterfly structures. *J. Am. Chem. Soc.*, v.134, p.7672-7675, 2012a.

SCHULTZ, A.; LI, X.; MCCUSKER, J.K.; MOOREFIELD, C.N.; CASTELLANO, F.N.; WESDEMIOTIS, C.; NEWKOME, G.R. Dondorff rings: synthesis, isolation, and properties of 60°-directed bisterpyridine-based folded tetramers. *Chem. Eur. J.*, v.18, p.11569-11572, 2012b.

TOMLINSON, P.B. *Tree architecture*. *Am. Sci.*, v.71, p.141-149, 1983.
WANG, J.-L.; LI, X.; LU, X.; HSIEH, I.-F.; CAO, Y.; MOOREFIELD, C.N.; WESDEMIOTIS, C.; CHENG, S.Z.D.; NEWKOME, G.R. Stoichiometric self-assembly of shape-persistent 2D complexes: a facile route to a symmetric supramacromolecular spoked wheel. *J. Am. Chem. Soc.*, v.133, p.11450-11453, 2011.

WANG, P.; MOOREFIELD, C.N.; JEONG, K.-U.; HWANG, S.-H.; LI, S.; CHENG, S.Z.D.; NEWKOME, G.R. Dendrimer-metallomacrocycle composites: nanofiber formation by multi-ion pairing. *Adv. Mater.*, v.20, p.1381-1385, 2008.

WINTER, A.; HAGER, M.D.; NEWKOME, G.R.; SCHUBERT, U.S. The marriage of terpyridine and inorganic nanoparticles: synthetic aspects, characterization techniques, and potential applications. *Adv. Mater.*, v.23, p.5728-5748, 2011a.

WINTER, A.; NEWKOME, G.R.; SCHUBERT, U.S. Catalytic applications of terpyridines and their transition metal complexes. *Chem. Cat. Chem.*, v.3, p.1384-1406, 2011b.

YOON, Z.S.; CHAN, Y.-T.; LI, S.; NEWKOME, G.R.; GOODSON, T.G. III ultrafast time-resolved spectroscopy of self-assembled cyclic Fe(II)-Bisterpyridine complexes. *J. Phys. Chem. B*, v.114, p.11731-11736, 2010.

YOUNG, J.K.; BAKER, G.R.; NEWKOME, G.R.; MORRIS, K.F.; JOHNSON JR, C.S. ‘Smart’ cascade polymers. Modular syntheses of four-directional dendritic macromolecules with acidic, neutral, or basic terminal groups and the effect of pH changes on their hydrodynamic radii. *Macromolecules*, v.27, p.3464-3471, 1994.

YU, X.; YUE, K.; HSIEH, I.-F.; LI, Y.; DONG, X.-H.; LIU, C.; XIN, Y.; WANG, H.-F.; SHI, A.-C.; NEWKOME, G.R.; HO, R.-M.; CHEN E.-Q.; ZHANG, W.-B.; CHENG, S.Z.D. Giant surfactants provide a versatile platform for sub-10-nm nanostructure engineering. *P. Natl. Acad. Sci. USA*, v.110, p.10078-10083, 2013.