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A convergent growth approach to electroactive ferrocene rich carbosilane- and siloxane-based dendrons, dendrimers, and dendronized polymers†‡

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The construction of the first members of a novel family of structurally well-defined, ferrocenyl rich, dendritic macromolecules based on carbosilane skeletons and siloxane linkages has been achieved via a convergent growth approach. Starting from triferrocenylvinylsilane $\text{Fc}_3\text{SiCH}=\text{CH}_2$ ($\text{Fc} = \text{Fe}(\eta^5\text{-C}_5\text{H}_4)(\eta^5\text{-C}_5\text{H}_5)$) as the key monomer, the sequential utilization of platinum-catalyzed hydrosilylation and alkenylation steps with Grignard reagents (allylmagnesium bromide) can be applied to prepare three different branched structures: multiferrocenyl-terminated dendrons **2** and **3**, dendrimers **4** and **5**, and dendronized polymers **7_n–9_n**. All of the dendritic metallomacromolecules have been thoroughly characterized using a combination of elemental analysis, multinuclear (^1H , ^{13}C , ^{29}Si) NMR spectroscopy, FT-IR and MALDI-TOF mass spectrometry, to establish their chemical structures and properties. The molecular structures of **G1**-dendron **3** and dendrimer **4**, containing six and nine ferrocenyl units, respectively, have been successfully determined by single-crystal X-ray analysis, compound **4** being the branched multiferrocenyl-containing siloxane with the highest number of Fc substituents whose structure has been reported so far. Electrochemical studies (using cyclic voltammetry (CV) and square wave voltammetry (SWV) performed in dichloromethane solution with $[\text{PF}_6]^-$ and $[\text{B}(\text{C}_6\text{F}_5)]^{4-}$ as supporting electrolyte anions of different coordinating abilities) reveal that all the macromolecular compounds obtained exhibit a three-wave redox pattern, suggesting appreciable electronic interactions between the silicon-bridged triferrocenyl moieties as they are successively oxidized. In addition, dendrimer **5** and dendronized polymers **7_n–9_n**, with 12 and $4 < n < 14$ ferrocenyl units, respectively, linked in threes around the periphery, undergo remarkable oxidative precipitation in $\text{CH}_2\text{Cl}_2/[n\text{-Bu}_4\text{N}][\text{PF}_6]$ and are able to form chemically modified electrodes with stable electroactive films.

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Introduction

Dendrons, dendrimers, and dendronized polymers are among the most widespread, well-defined, branched macromolecular

structures.¹ Specifically, dendrimers constitute a unique class of macromolecular architectures that differ from all other synthetic macromolecules in their perfectly branched topology, which is constructed, in an exceptionally challenging step-wise manner, by a repetitive reaction sequence from a central core and expands to the periphery that becomes denser with increasing generation number. Many synthetic approaches have been developed for dendrimers since the seminal examples reported during the 80s and 90s,² the divergent^{2a–c} and convergent^{2g–i} growth being the two most widely used strategies.

In the divergent approach, dendrimers are grown from the inside to the outside, starting from a multifunctional core to which several layers of branching units are attached progressively. As a different strategy, in the convergent approach the construction of dendrimers starts at what will ultimately become the outer surface of the dendrimer and progresses inwardly. This approach first requires the synthesis of functionalized, progressively larger, dendritic wedges or dendrons, having a single reactive focal point, which is then covalently

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attached to a polyfunctional core to yield the desired dendrimer. Furthermore, dendrimer approaches imply other accelerated synthetic strategies that can minimize the number of reaction steps in dendrimer synthesis.³ Remarkably, the click reaction concept has proved to be a powerful synthetic tool for the design of highly sophisticated dendritic architectures, for instance to prepare dendrons and dendrimers *via* thiol-ene chemistry,⁴ to assemble dendrons in order to construct siloxane dendrimers⁵ or to prepare liquid-crystalline Janus dendrimers, merging two types of dendromesogens.⁶

Among the fascinating architectural features of dendrimers, their high overall symmetries and their well-defined nanometer sizes and internal cavities have allowed the rapid development of their new and improved properties. In fact, they are materials with valuable and promising applications in a wide range of fields, such as catalysis, cancer therapy, biosensors for diagnostics, light harvesting scaffolds, controlled drug delivery, medical diagnostics or nanoelectronics, to name just a few examples.⁷

One of the most active research areas in dendrimer chemistry involves the integration of transition metals into dendritic structures to create metalodendrimers.⁸ Since the first transition-metal containing dendrimers were reported in the early 1990s, advances in the synthesis and chemistry of these molecules have not ceased to blossom. For instance, the incorporation of electroactive organometallic units into dendrimer structures is an especially attractive target area because such highly branched macromolecules are good multielectron-transfer mediators in electrocatalytic processes of biological and industrial importance, and this remains a topical challenge. In particular, covalent incorporation of ferrocene units into dendrimers,⁹ polymers,¹⁰ and macrocycles¹¹ can greatly modify their physical and chemical properties. Ferrocene is an ideal donor moiety and the ease of functionalization, the chemical stability, and the diamagnetism in the neutral state (allowing NMR characterization) make this metallocene one of the mostly used building blocks to construct metallomacromolecular architectures.¹² In addition, this organometallic sandwich molecule exhibits attractive electrochemical behavior, a fast electron-transfer rate, low oxidation potential, and two stable redox states (neutral ferrocene and oxidized ferrocenium). In fact, the achievements in the synthesis of ferrocene-containing organosilicon polymers and oligomers,¹³ and well-defined branched molecules of different nature, including ferrocene dendrimers, have attracted significant attention over the years due to their beauty, synthetic complexity, and valuable applications.

Likewise, in order to expand the dendrimer chemistry, silicon chemistry has played a dominant role in the field of dendritic polymers, enabling the synthesis of a robust and diverse family of silicon-based dendritic macromolecules.^{14,15} As a result, silicon-containing dendritic skeletons based on carbosilane (Si–C), siloxane (Si–O–Si) or carbosiloxane (Si–O–C) linkages are among the most widely used ones for the synthesis of transition metal-containing dendrimers, because of their kinetic and thermodynamic stability, chemical inertness,

and accessibility. Regarding this point, our research group has also contributed to the design, construction and electrochemical studies of dendritic macromolecules based on silicon, mainly carbosilanes and siloxanes, functionalized with electroactive organometallic units of a very diverse nature.¹⁶

On the other hand, dendronized polymers have become an important class of macromolecules, which merges the concepts of dendrimers and linear polymers.^{17,18} They are composed of a linear polymer backbone to which dendrons of increasing size (*i.e.*, generations G1, G2, G3...) are appended. At high degrees of dendronization, dendronized polymers gain stiffness and adopt a cylindrical shape. This exclusive architecture makes dendronized polymers unique macromolecules and interesting candidates for a variety of nanoscale applications, such as molecular wires, self-adaptive materials, and functional scaffolds for catalysis. The vast majority of dendron-branched polymers present an organic nature. Few examples of branched siloxanes have been published so far.¹⁹ Furthermore, in spite of the fact that numerous metalodendrimers have been reported, only very few studies have addressed the synthesis of dendronized polymers containing metallic moieties.²⁰

Herein, we merge the convergent approach, the silicon chemistry and the unique redox properties and ease of functionalization of the ferrocene unit and present full details of the synthesis, characterization (including the solid-state structures of some macromolecules), thermal properties, and electrochemical behavior of first- and second generation, ferrocene-rich and siloxane-based dendrons, dendrimers, and dendronized polymers. Remarkably, the presence of triferrrocenyl end moieties, bridged by a silicon atom, has led us to explore the electrochemical behavior and electronic interactions between the redox-active metal sites. Furthermore, we have investigated in detail the chemically modified electrodes formed with some ferrocenyl-dendritic molecules, highlighting their ability to form robustly adsorbed films onto metallic electrode surfaces.

Experimental

General procedures

Materials and equipment. All reactions and compound manipulations were performed in an oxygen- and moisture-free Ar atmosphere using standard Schlenk techniques. Solvents were dried by standard procedures over the appropriate drying agents and distilled under argon, immediately prior to use. The vinyl-functionalized precursor $\text{Fc}_3\text{Si}-\text{CH}=\text{CH}_2$ (**1**) was synthesized from monolithioferrocene (generated *in situ* from the reaction between ferrocene and *t*-BuLi, at $-10\text{ }^\circ\text{C}$) according to the procedure already described.²¹ The platinum-divinyltetramethyldisiloxane complex in xylene (3–3.5% Pt concentration) (Karstedt's catalyst), chlorophenylsilane, allyl-magnesium bromide (1 M in diethyl ether) and poly(methylhydrosiloxane) ($(\text{Me}_3\text{SiO})(\text{MeSiHO})_n(\text{SiMe}_3)$, M_n 1700–3200) (available from Merck-Sigma-Aldrich), and tetrakis(dimethylsiloxy)silane (ABCR) were used as received. Silica gel (70–230 mesh) (Merck-Sigma-Aldrich) was used for column



chromatography purifications. Infrared spectra were recorded on a PerkinElmer 100 FT-IR spectrometer. Elemental analyses were performed on a LECO CHNS-932 elemental analyzer, equipped with a MX5 Mettler Toledo microbalance. All NMR spectra were recorded on Bruker Avance III-HD NANOBAY 300 MHz and Bruker DRX 500 MHz spectrometers. Chemical shifts were reported in parts per million (δ) with reference to CDCl_3 residual solvent resonances for ^1H (δ 7.26 ppm) and ^{13}C (δ 77.2 ppm). ^{29}Si NMR resonances were recorded with inverse-gated proton decoupling to minimize nuclear Overhauser effects and were referenced externally to tetramethylsilane. For the types of silicon atoms, depending on the number of bonded oxygen atoms, the terminologies M, D, T and Q represent the structures $\text{R}_3\text{SiO}_{0.5}$, R_2SiO , $\text{RSiO}_{1.5}$ and SiO_2 , respectively (see Scheme S1 in the ESI†). MALDI-TOF mass spectra were recorded using a Bruker-Ultraflex III TOF/TOF mass spectrometer equipped with a nitrogen laser emitting at 337 nm. Dichloromethane solutions of the matrix (dithranol, 10 mg mL^{-1}) and CH_2Cl_2 solutions of the corresponding compound (1 mg mL^{-1}) were mixed in a ratio of 20:5. Then, 0.5–1 μL of the mixture was deposited on the target plate using the dried droplet method. The positive ion and the reflectron mode were used for these analyses. Thermogravimetric analyses were performed using a TGA Q-500 instrument coupled with an EGA oven. The samples (5–10 mg) were loaded in platinum pans. The measurements were carried out under N_2 (90 mL min^{-1}) with a heating rate of 10 $^{\circ}\text{C min}^{-1}$. The morphology of the ceramic residues was investigated with SEM using a Philips XL30 instrument coupled with an EDAX DX4i analyzer.

Electrochemical measurements. Cyclic voltammetry (CV) and square wave voltammetry (SWV) experiments were performed on Bioanalytical Systems BAS CV-50 W and Autolab PGSTAT302F potentiostats. CH_2Cl_2 and CH_3CN (SDS, spectro-grade) were freshly distilled from calcium hydride under Ar. The supporting electrolytes used were tetra-*n*-butylammonium hexafluorophosphate (Alfa-Aesar), which was purified by recrystallization from ethanol and dried in a vacuum at 60 $^{\circ}\text{C}$, and tetra-*n*-butylammonium tetrakis(pentafluorophenyl) borate, which was synthesized as described in the literature,²² by metathesis of $[\text{n-Bu}_4\text{N}]\text{Br}$ with $\text{Li}[\text{B}(\text{C}_6\text{F}_5)_4]\cdot(\text{nOEt}_2)$ (Boulder Scientific Company) in methanol and recrystallized twice from $\text{CH}_2\text{Cl}_2/n$ -hexane. The supporting electrolyte concentration was 0.1 M. A conventional three-electrode cell connected to an atmosphere of prepurified nitrogen was used. The counter electrode was a coiled Pt wire, and the reference electrode was a BAS saturated calomel electrode (SCE). All cyclic voltammetric experiments were performed using a Pt-disk working electrode ($A = 0.020 \text{ cm}^2$) (BAS). The working electrode was polished on a Buehler polishing cloth with Metadi II diamond paste for about 3 min followed by sonication in absolute ethanol, rinsed thoroughly with purified water and acetone, and allowed to dry. Under our conditions, the ferrocene redox couple $[\text{FeCp}_2]^{0/+}$ is + 0.462, and the decamethylferrocene redox couple $[\text{FeCp}_2^*]^{0/+}$ is −0.056 V vs. the SCE in $\text{CH}_2\text{Cl}_2/0.1 \text{ M} [\text{n-Bu}_4\text{N}]\text{[PF}_6]$. The solutions were, typically, 10^{-4} M in the

redox-active species and were purged with nitrogen and kept under an inert atmosphere throughout the measurements. No IR compensation was used. SWV was performed using frequencies of 10 Hz.

The preparation of electrode surfaces modified with films of dendrimer 5 and polymers 7_n–9_n was accomplished by electrodeposition of the corresponding species on Pt electrodes. The electroactive films were prepared by cyclically scanning the potential (between +0.0 and +1.3 V vs the SCE) in degassed CH_2Cl_2 solutions of the corresponding dendritic species at different numbers of times (10 or 20 scans for dendrimer 5 and 10 scans for polymers 7_n–9_n). The electrochemical behavior of the films was then studied by CV in multiferrocene-free CH_2Cl_2 or CH_3CN solution containing only the supporting electrolyte ($[\text{n-Bu}_4\text{N}]\text{[PF}_6]$ or $[\text{n-Bu}_4\text{N}]\text{[B}(\text{C}_6\text{F}_5)_4]$). From the CVs of the modified electrodes, the surface coverages, Γ (mol cm^{-2}), of the ferrocenyl sites were calculated from the charge, Q , under the voltammetric current peaks, using $\Gamma = Q/nFA$.

X-ray crystal structure determination. Suitable orange crystals of 3 and 4 were isolated and coated with mineral oil, and mounted on Mitegen MicroMounts. Diffraction data were collected using a Bruker D8 KAPPA series II diffractometer with graphite monochromated Mo K α radiation ($\lambda = 0.71073 \text{ \AA}$). Full details of the data collection and refinement can be found in the ESI (see Section 2†). The good redundancy in data allowed empirical absorption corrections (SADABS)²³ to be applied using multiple measurements of symmetry-equivalent reflections. Raw intensity data frames were integrated with the SAINT program,²⁴ which also applied corrections for Lorentz and polarization effects. SHELXTL was used for space group determination, structure solution, and refinement.²⁵ The space group determination was based on a check of the Laue symmetry and systematic absences were confirmed using the structure solution. The structures were solved by direct methods (SHELXS-97), completed with difference Fourier synthesis, and refined with full-matrix least-squares using S minimizing $\omega(F_o^2 - F_c^2)^2$.^{26,27} Weighted R factors (R_w) and all goodness of fit S are based on F^2 ; conventional R factors (R) are based on F . All non-hydrogen atoms were refined with anisotropic displacement parameters. All scattering factors and anomalous dispersion factors are present in the SHELXTL 6.10 program library. The crystal structures of compounds 3 and 4 have been deposited at the Cambridge Crystallographic Data Centre with deposit numbers CCDC 2105207 and 2105208, respectively.†

Synthesis

Synthesis of first generation dendrons: $(\text{C}_6\text{H}_5)\text{SiCl}\{(\text{CH}_2)_2\text{SiFc}_3\}_2$ (2) and $(\text{C}_6\text{H}_5)\text{Si}(\text{CH}_2\text{CH}=\text{CH}_2)\{(\text{CH}_2)_2\text{SiFc}_3\}_2$ (3). To a solution of dendron 1 (1.00 g, 1.64 mmol) in 25 mL of dry and degassed toluene, Karstedt's catalyst (60 μL) was added under argon. The mixture was stirred for 20 min before a toluene solution of chlorophenylsilane (110 μL , 0.82 mmol) was added dropwise. The reaction mixture was heated at reflux temperature until the ^1H NMR experiments showed the com-



plete disappearance of the Si–H signal of the starting $(C_6H_5)Si(H)_2Cl$ (18 h). At this moment, the reaction mixture was allowed to cool to room temperature, and the solvent was removed under vacuum. The remaining orange oil $(C_6H_5)SiCl\{(CH_2)_2SiFc_3\}_2$ (2) was immediately used in the next reaction step.

To a three-necked, round-bottomed flask equipped with a gas inlet, a pressure equalizing addition funnel, an Allihn condenser topped with a gas inlet and bubbler, and a Teflon-covered magnetic stir bar, freshly distilled and degassed diethyl ether (30 mL) was added. A solution of recently prepared 2 in 30 mL of diethyl ether was added dropwise, with vigorous stirring. Simultaneously, allylmagnesium bromide (0.9 mL, 0.9 mmol, 1 M in diethyl ether) was added dropwise. The resulting yellow-orange solution was refluxed for 20 h, cooled to 0 °C with an ice bath, and then hydrolyzed with aqueous NH_4Cl (10%). The organic layer was separated, washed three times with water, and dried over anhydrous $MgSO_4$. After filtration, the solvent was removed under vacuum yielding an orange-yellow oily product, which was purified by column chromatography on silica gel (3 cm × 10 cm). A first band was observed corresponding to unreacted trifluorocenylvinylsilane 1 which was eluted with *n*-hexane/ CH_2Cl_2 (10 : 2), and subsequently, a second major orange band was observed with *n*-hexane/ CH_2Cl_2 (10 : 5). Solvent removal afforded the desired grown dendron 3, carrying six ferrocenyl units, which was obtained as an analytically pure, air-stable, orange crystalline solid.

Yield: 0.48 g (43%). **Anal.** Calcd for $C_{73}H_{72}Si_3Fe_6$: C 64.06; H 5.30. Found: C 64.10; H 5.21. 1H NMR ($CDCl_3$, 300 MHz, ppm): δ 1.26 (s, 8H, Si– CH_2 – CH_2), 2.11 (m, 2H, – CH_2 CH=CH₂), 3.98 (s, 30H, C_5H_5), 4.24, 4.37 (m, 12H + 12H, C_5H_4), 5.03 (m, 2H, – CH_2 CH=CH₂), 5.98 (m, 1H, – CH_2 CH=CH₂), 7.38 (m, 3H, Ph), 7.60 (m, 2H, Ph). $^{13}C\{^1H\}$ NMR ($CDCl_3$, 75 MHz, ppm): δ 5.2, 8.3 (Si– CH_2 – CH_2), 19.5 (– CH_2 CH=CH₂), 68.5 (C_5H_5), 69.8 (*ipso*-Fc), 70.5, 73.8 (C_5H_4), 114.1 (– CH_2 CH=CH₂), 128.0 (m–Ph), 129.3 (*p*-Ph), 134.3 (*o*-Ph), 134.7 (– CH_2 CH=CH₂), 136.3 (*ipso*-Ph). $^{29}Si\{^1H\}$ RMN ($CDCl_3$, 59 MHz, ppm): δ –8.3 (Fc–Si), –0.1 (Si– CH_2 CH=CH₂). **IR** (KBr, cm^{-1}): δ (Si–C) 1258, ν (Si–O–Si) 1033–1105, δ (Si–H) 890, ν (Si–C) 818. **MS** (MALDI-TOF): *m/z* 1368.0 [M^+].

General hydrosilylation reaction procedure. All the hydrosilylation reactions were performed under an oxygen- and moisture-free atmosphere (Ar) using standard Schlenk techniques. A typical experimental procedure was as follows. In a three-necked, round-bottomed flask equipped with a gas inlet, an Allihn condenser topped with a gas inlet and bubbler, and a Teflon-covered magnetic stir bar, trimetallic compound 1 was dissolved in freshly distilled toluene. Into the resulting orange solution, Karstedt's catalyst was injected using a Hamilton precision syringe, under a flow of argon, and the mixture was stirred at room temperature during 30 min. Subsequently the corresponding siloxane was slowly added dropwise. The hydrosilylation mixture was heated, and the progress of the reaction was monitored by 1H NMR from the progressive disappearance of the Si–H signal of the corresponding precursor. After the appropriate time, the reaction was

stopped, the mixture was allowed to cool to room temperature and the solvent was removed under vacuum.

Synthesis of dendrimers 4 and 5. Dendrimer 4 was obtained from the reaction of 1 (0.3 g, 0.49 mmol) in 6 mL of toluene, with 30 μ L of Karstedt's catalyst, and 41.78 μ L of tetrakis(dimethylsiloxy)silane I (0.11 mmol). After 6 h at 70 °C, the orange-brown, oily residue was purified by column chromatography on silica gel (2 cm × 20 cm). A first band was observed corresponding to the excess of compound 1 which was eluted with *n*-hexane/ CH_2Cl_2 (10 : 2). Then, upon eluting with *n*-hexane/ CH_2Cl_2 (10 : 6), a second major band was observed. Solvent removal afforded dendrimer 4, with nine terminal ferrocenes, as an analytically pure, air-stable, orange, crystalline solid.

Dendrimer 4. **Yield:** 0.09 g (38%). **Anal.** Calcd for $C_{104}H_{118}Si_8Fe_9O_4$: C 57.85; H 5.51. Found: C 58.09; H 5.48. 1H NMR ($CDCl_3$, 500 MHz, ppm): δ 0.28 (s, 24H, CH_3), 0.98, 1.22 (m, 6H + 6H, CH_2), 4.01 (s, 45H, C_5H_5), 4.26, 4.37 (m, 36H, C_5H_4), 4.86 (sep, 1H, Si–H). $^{13}C\{^1H\}$ NMR ($CDCl_3$, 125 MHz, ppm): δ –0.1 (CH_3), 1.2 (CH_3SiH), 7.8, 11.2 (CH_2), 68.6 (C_5H_5), 70.2 (*ipso*-Fc), 70.6, 74.1 (C_5H_4). $\{^1H\}^{29}Si$ HMQC ($CDCl_3$, 500, 99 MHz, ppm): δ –5.3 (Si–H). $\{^1H\}^{29}Si$ HMBC ($CDCl_3$, 500, 99 MHz, ppm): δ –103.0 (SiO_4), –8.2 (Fc–Si), –5.3 (Si–H), 10.0 (Si(CH_3)₂). **IR** (KBr, cm^{-1}): ν (Si–H) 2137, δ (Si–C) 1258, ν_{as} (Si–O–Si) 1033–1105, δ (Si–H) 890, ν (Si–C) 818. **MS** (MALDI-TOF): *m/z* 2159.1 [M^+].

Dendrimer 5 was obtained by the reaction of 1 (0.3 g, 0.49 mmol) in 6 mL of toluene, with 39 μ L of tetrakis(dimethylsiloxy)silane I (0.10 mmol) in the presence of 30 μ L of Karstedt's catalyst. The reaction mixture was stirred at 70 °C for 17 h. The oily residue was purified by column chromatography on silica gel (2 cm × 20 cm). After isolating the excess of compound 1 with *n*-hexane/ CH_2Cl_2 (10 : 2), a second major band was eluted with *n*-hexane/ CH_2Cl_2 (1 : 1), which afforded the desired dendrimer 5 as an analytically pure, air-stable, orange, crystalline solid.

Dendrimer 5. **Yield:** 0.20 g (74%). **Anal.** Calcd for $C_{136}H_{148}Si_9Fe_{12}O_4$: C 58.98; H 5.39. Found: C 59.22; H 5.29. 1H NMR ($CDCl_3$, 300 MHz, ppm): δ 0.31 (s, 24H, CH_3), 0.99, 1.24 (m, 16H, CH_2), 4.01 (s, 60H, C_5H_5), 4.26, 4.37 (m, 24H + 24H, C_5H_4). $^{13}C\{^1H\}$ NMR ($CDCl_3$, 75 MHz, ppm): δ –0.1 (CH_3), 7.8, 11.2 (CH_2), 68.6 (C_5H_5), 70.2 (*ipso*-Fc), 70.6, 74.1 (C_5H_4). $^{29}Si\{^1H\}$ RMN ($CDCl_3$, 59 MHz, ppm): δ –103.1 (SiO_4), –8.3 (Fc–Si), 9.8 (Si(CH_3)₂). **IR** (KBr, cm^{-1}): δ (Si–C) 1249, ν_{as} (Si–O–Si) 1028–1107, ν (Si–C) 820. **MS** (MALDI-TOF): *m/z* 2770.1 [M^+].

Synthesis of dendronized polymers 7_n–9_n. Compound 1 (0.5 g, 0.82 mmol), Karstedt's catalyst (30 μ L), and poly(methylhydrosiloxane) II (42 μ L, 0.02 mmol) were mixed in 6 mL of toluene. After 45 h of stirring at 100 °C, the oily reaction product was dissolved in a small amount of CH_2Cl_2 and precipitated twice into methanol.

Yield: 0.38 g. 1H NMR ($CDCl_3$, 300 MHz, ppm): δ 0.20–0.26 (m, CH_3), 0.88, 1.26 (m, CH_2), 3.99 (s, C_5H_5), 4.24, 4.34 (m, C_5H_4). $^{13}C\{^1H\}$ NMR ($CDCl_3$, 75 MHz, ppm): δ 0.2, 2.7 (CH_3), 7.6, 10.7 (CH_2), 68.8 (C_5H_5), 70.1 (*ipso*-Fc), 70.7, 74.1 (C_5H_4). $^{29}Si\{^1H\}$ RMN ($CDCl_3$, 59 MHz, ppm): δ –21.9 (SiO_2) –8.2 (Fc–



Si), 7.5 (*SiO*). IR (KBr, cm^{-1}): $\delta(\text{Si}-\text{C})$ 1260, $\nu_{\text{as}}(\text{Si}-\text{O}-\text{Si})$ 1031–1105, $\nu(\text{Si}-\text{C})$ 815, 799. MS (MALDI-TOF). 7_n: *m/z* 2844 [M^+ , *n* = 4], 3514 [M^+ , *n* = 5], 4184 [M^+ , *n* = 6], 4854 [M^+ , *n* = 7], 5524 [M^+ , *n* = 8], 6194 [M^+ , *n* = 9], 6864 [M^+ , *n* = 10], 7534 [M^+ , *n* = 11], 8204 [M^+ , *n* = 12], 8874 [M^+ , *n* = 13], 9544 [M^+ , *n* = 14]. 8_n: *m/z* 2681 [M^+ , *n* = 4], 3351 [M^+ , *n* = 5] γ 4021 [M^+ , *n* = 6]. 9_n: 2904 [M^+ , *n* = 4], 3574 [M^+ , *n* = 5], 4244 [M^+ , *n* = 6], 4914 [M^+ , *n* = 7], 5584 [M^+ , *n* = 8], 6254 [M^+ , *n* = 9], 6924 [M^+ , *n* = 10], 7594 [M^+ , *n* = 11], 8264 [M^+ , *n* = 12], 8934 [M^+ , *n* = 13], 9604 [M^+ , *n* = 14].

Attempts to synthesize dendrimer **6** and the second generation dendronized polymer from dendron **3** are explained in the ESI.‡

Results and discussion

Convergent growth approach from triferrocenylvinylsilane **1**: synthesis of dendrons **2** and **3**

The convergent approach^{1g,2g-i} has been selected for the synthesis of a new family of ferrocene rich, silicon-containing, dendrons and dendrimers. We first constructed small ferrocene-based monofunctional dendrons, bearing a single carbon–carbon double bond at the focal point, and subsequently we used them in hydrosilylation chemistry to synthesise either dendrons of higher generations or final dendrimer molecules.

Prior to initiating dendrimer synthesis, triferrocenylvinylsilane^{21,28} $\text{Fc}_3\text{Si}-\text{CH}=\text{CH}_2$ (**1**) ($\text{Fc} = \text{Fe}(\eta^5\text{-C}_5\text{H}_4)(\eta^5\text{-C}_5\text{H}_5)$) was selected as the key monomer for the synthesis of higher generation dendrons. Triferrocenylvinylsilane **1** fulfils several fundamental requirements and shows features for being used in a convergent methodology: (1) it is experimentally accessible; (2) it bears a single vinylsilyl functional group that can act as the reactive point for Si–H functional backbones through hydrosilylation reactions; (3) from an electronic point of view, the $-\text{CH}=\text{CH}_2$ group of **1**, linked to three electron-donating ferrocenyl moieties, is particularly electron rich and, accordingly, is highly reactive toward hydrosilylation;²⁹ (4) it has a very stable chemical nature, which implies high tolerance to the reaction conditions of carbosilane chemistry; (5) it shows good solubility in common organic solvents (such as THF, CH_2Cl_2 and *n*-hexane), and finally, (6) **1** is an

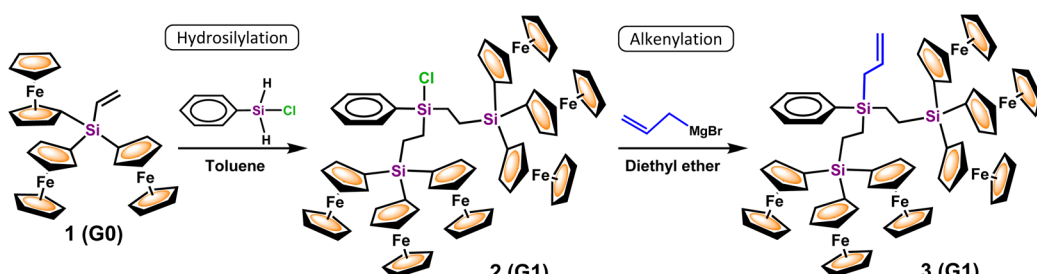
organometallic molecule with remarkable thermal and air stability, which can be obtained and stored for a long period of time (several years) under an air atmosphere at room temperature without noticeable decomposition (¹H NMR spectroscopy).

Beforehand, perhaps the only structural challenge that monofunctional **1** could offer for a convergent dendrimer synthesis is the sterically hindered environment of the reactive $\text{Si}-\text{CH}=\text{CH}_2$ group, surrounded by three bulky ferrocenyl moieties that might well affect the effectiveness of hydrosilylation reactions. However, triferrocenylvinylsilane **1** has already been employed in successful hydrosilylation reactions,²⁹ which led to the preparation of cyclic and cubic macromolecules peripherally decorated with as many interacting ferrocenyl units as chemically possible (24 ferrocenyl moieties). In addition, the presence of three ferrocenes, linked through the same silicon atom, enables an electronic interaction between metal centers and provides strong motivation to investigate their electrochemical processes.

Therefore, by using triferrocenylvinylsilane **1**(G0) as the dendritic precursor, the first generation (G1) dendrons **2** and **3** were obtained, by means of a two-step convergent growth methodology (Scheme 1). The reaction sequence began with the hydrosilylation of **1** and $(\text{C}_6\text{H}_5)\text{Si}(\text{H})_2\text{Cl}$. Platinum-catalyzed hydrosilylation was chosen since it has been shown to be a highly efficient process, and Karstedt's catalyst was selected because it is widely used in hydrosilylation chemistry.³⁰

Dendron **2** was obtained as an orange dense oil and was immediately used in the subsequent reaction, avoiding its isolation to prevent any possible decomposition by hydrolysis. In the second step, **2** was successfully alkenylated with allylmagnesium bromide in diethyl ether. A subsequent hydrolytic workup and careful purification by column chromatography afforded an air-stable, orange and crystalline solid (in 43% isolated yield), which was characterized as the desired grown dendron **3** (Scheme 1), carrying six ferrocenyl units linked in three ways through a silicon atom. The coordination sphere of the external silicon atom was completed by a phenyl ring, and an allyl reactive group to allow further dendritic growth.

On the basis of the ¹H NMR spectra of both dendrons **2** and **3**, it was determined that only the β -isomers formed under the described conditions, and consequently hydrosilylation was anti-Markovnikov. The structural identity of novel dendron



Scheme 1 Dendritic growth from $\text{Fc}_3\text{Si}-\text{CH}=\text{CH}_2$ (**1**): synthesis of dendritic wedges with $\text{Si}-\text{Cl}$ (**2**) and $\text{Si}-\text{CH}_2-\text{CH}=\text{CH}_2$ (**3**) reactive focal points.

3 was confirmed by IR, multinuclear ^1H , ^{13}C and ^{29}Si NMR spectroscopy, MALDI-TOF mass spectrometry and elemental analysis. As can be seen in the ^1H NMR spectra of precursor **1** and grown dendron **3** (Fig. 1), the mono-substituted ferrocenyl areas, around δ 4 ppm, are almost identical for both compounds. The allylic group of **3** appears as three resonances: a multiplet at δ 5.98 ppm for the CH unit, another multiplet at δ 5.03 ppm for the terminal $-\text{CH}_2$ group, and a doublet at δ 2.11 ppm assigned to the internal $-\text{CH}_2-$ protons. The hydrogen signals of the two $-\text{CH}_2-\text{CH}_2-$ bridges are observed at δ 1.26 ppm under a broad singlet, and those of C_6H_5 as two multiplets are centered at δ 7.60 and 7.38 ppm.

The same occurs with the ^{13}C NMR experiments. The vinyl resonances at about δ 135 ppm of **1** are replaced with the allyl signals at δ 19.5, 114.1 and 134.7 ppm in **3** (Fig. S3‡). In the ^{13}C NMR spectrum of **3** the corresponding resonances of the phenyl unit (at δ 128.0, 129.3, 134.3 and 136.3 ppm) and of the asymmetric $-\text{CH}_2-\text{CH}_2-$ bridges (at δ 5.2 and 8.3 ppm) can also be observed. The ^{29}Si NMR spectrum of the resulting dendron **3** shows two peaks centered at δ -0.1 ppm (for the $-\text{Si}-\text{CH}_2-\text{CH}=\text{CH}_2$ unit) and at δ -8.3 (for the $(\text{Fc})_3\text{Si}-\text{CH}_2\text{CH}_2$ group). The latter is considerably shifted downfield in comparison with the same signal of precursor **1** (δ -17 ppm), reflecting the change in the electronic environment of the Si units due to the transformation of the adjacent C=C bond into a $-\text{Si}-\text{CH}_2-\text{CH}_2-$ bridge. The MALDI-TOF mass experiment shows a peak at m/z 1368.0 for **3** corresponding to the molecular ion M^+ , showing an excellent agreement between the experimental and the calculated isotopic patterns (Fig. S7‡). Finally, the solid-state structure of hexaferrocenyl dendron **3** was proved by single-crystal X-ray diffraction, as it will be commented below.

Incorporation of dendrons **1** and **3** into Si-H polyfunctional siloxane backbones: synthesis of dendrimers **4**, **5** and **6** and dendronized polymers 7_n-9_n

The final step of a dendritic molecular construction, following the convergent approach, consists of integrating the previously

synthesized monofunctional dendritic wedges into a multifunctional core.^{1g,2g-i} In this case, the availability of free olefinic substituents, at the focal point of dendrons **1** and **3**, facilitates the incorporation of interacting organometallic redox-active centers in dendritic structures through hydrosilylation chemistry with Si-H polyfunctional cores. Particularly, two different hydrosiloxanes were carefully selected as multifunctional cores (Fig. 2): star shaped tetrakis(dimethylsiloxy)silane, $\text{Si}[\text{OSi}(\text{CH}_3)_2\text{H}]_4$ (**I**), which will act as a four-functional core allowing the formation of dendrimers; Si-H-rich linear poly(methylhydrosiloxane), $[(\text{Me}_3\text{SiO})(\text{MeSiHO})_n(\text{SiMe}_3)]$, M_n 1700–3200, **II**], which will allow the formation of dendronized polymers. Based on our previous synthesis experience, the choice of core units, in which the reactive Si-H sites are located at the ends of extended flexible siloxane spacer chains, is important to reduce steric congestion in the desired final macromolecules.

For the synthesis of the dendrimeric molecules, hydrosilylation assembly reactions of **1** and **3** (in excess) with tetrakis(dimethylsiloxy)silane **I** were performed at 70 °C, in toluene solutions and in the presence of Karstedt's catalyst (Schemes 2 and 3). Completion of the anchoring reactions was easily monitored by ^1H NMR spectroscopy.

From the first hydrosilylation reaction between **1** (in stoichiometric excess) and $\text{Si}[\text{OSi}(\text{CH}_3)_2\text{H}]_4$ (**I**) instead of the target dendrimer **5**, only the nonmetallic dendritic molecule **4** was formed and isolated (38% yield, Scheme 2), with three of the four Si-H bonds being functionalized with $-(\text{CH}_2)_2\text{SiFc}_3$ units. This result is reproducible under the same reaction conditions (6 hours at 70 °C) and no other functionalization degrees, such as dendrimers with one, two or four $-(\text{CH}_2)_2\text{SiFc}_3$ units, were detected by MALDI-TOF experiments of the crude reaction mixtures. It is interesting to note here that the presence of the free single reactive Si-H functionality in molecule **4** enables further elaboration through, for example, thiol–ene reactions. Therefore, **4** could serve as an excellent candidate to construct two-faced “Janus dendrimers”, which are molecules composed of two dendrimeric wedges (dendrons) and terminated by two different but complementary functionalities.³¹

We first reasoned that this “structural dendritic defect” was probably caused by a low hydrosilylation time (6 hours of reaction). To test this hypothesis, and to ensure the full substitution of the four Si-H bonds, this reaction was repeated using a larger stoichiometric excess of triferrrocenylsilane **1** and it

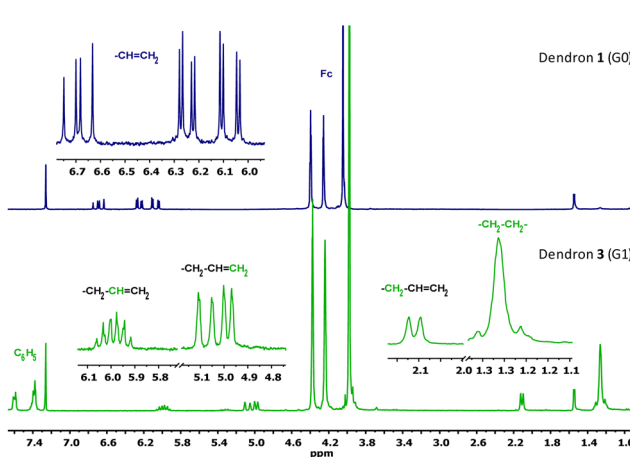


Fig. 1 ^1H NMR spectra of dendrons **1** (G0) and **3** (G1) (recorded in CDCl_3 , 300 MHz).

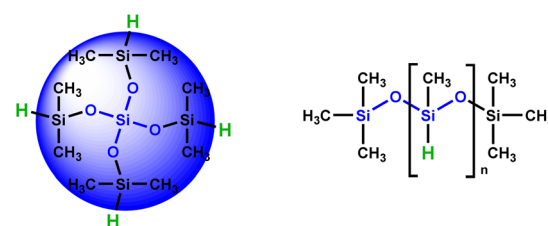
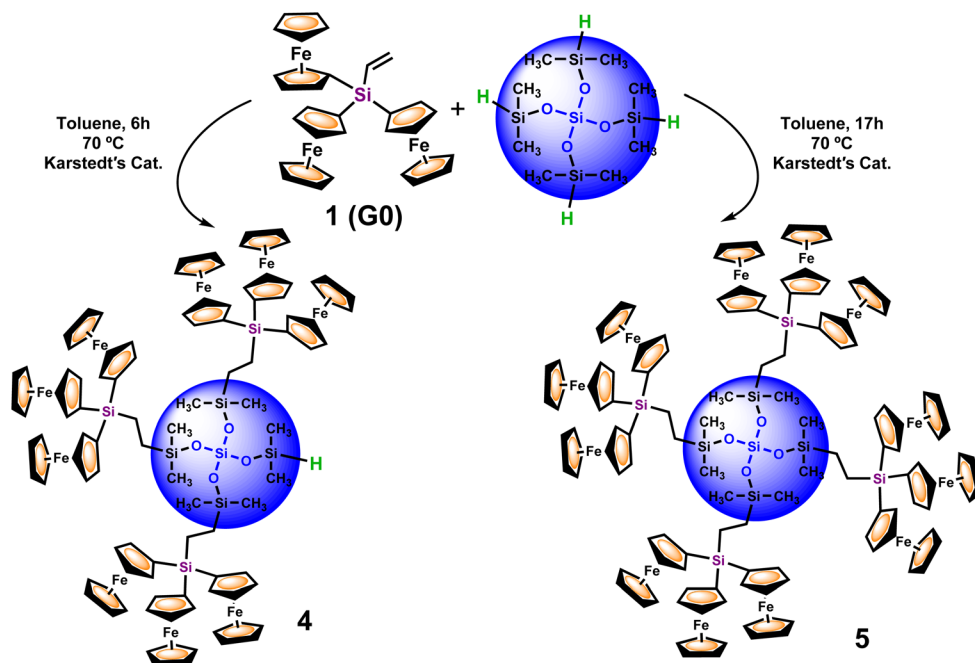
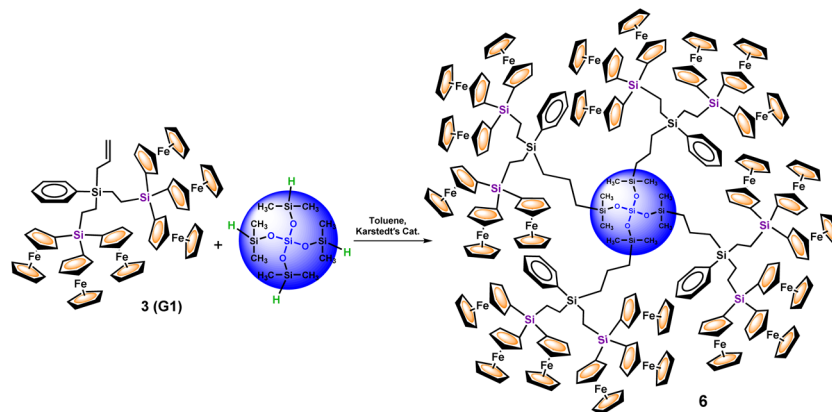


Fig. 2 Selected Si-H containing siloxane backbones for the growth of ferrocenyl-rich dendrimers and dendronized polymers.





Scheme 2 Hydrosilylation reactions of **1** with tetrakis(dimethylsiloxy)silane **I** leading to the first-generation dendrimers **4** and **5**.



Scheme 3 Hydrosilylation between the first generation dendron **3** and the tetra-directional core **I**.

was allowed to react for 17 hours. After appropriate purification, the target dendrimer **5** was isolated as an air-stable, crystalline, and orange solid (74% yield). The ^1H NMR and MALDI-TOF experiments confirm its identity. Particularly, there are no other peaks in the mass spectrum, allowing us to exclude the formation of, for instance, other dendrimers with structural defects. In addition, the ^1H NMR spectrum of the hydrosilylated dendrimer product proved that only the β -isomer was formed. This ensured a regular dendritic growth and the generation of molecules of maximum symmetry.

The next task was then focused on the hydrosilylation of **G1**-dendron **3** with the siloxane core $\text{Si}[\text{OSi}(\text{CH}_3)_2\text{H}]_4$ (**I**), in order to prepare larger generation dendrimers. The reaction was carried out for 24 hours (Scheme 3), and the orange oil

obtained was dissolved in a small amount of CH_2Cl_2 and precipitated into *n*-hexane. Both the solution and the oily residue obtained were studied by MALDI-TOF mass spectrometry, showing similar results (see Fig. 3). The most intense peak (at m/z 1369.1) matches with unreacted dendron **3**, but additional peaks that correspond to partially functionalized dendrimers can also be observed. Therefore, the peak at m/z 4450.0 matches with the dendrimer containing three functionalized arms and the fourth one ends in a $\text{Si}-\text{OH}$ group. The one at m/z 3080.2 corresponds to the bifunctionalized dendrimer and two unfunctionalized branches (one $\text{Si}-\text{H}$ and one $\text{Si}-\text{OH}$). This peak is followed by another one at m/z 3096.2 of the bifunctionalized dendrimer with two terminal $\text{Si}-\text{OH}$ groups. Finally, the peak at m/z 1726.1 correlates with the monofunc-

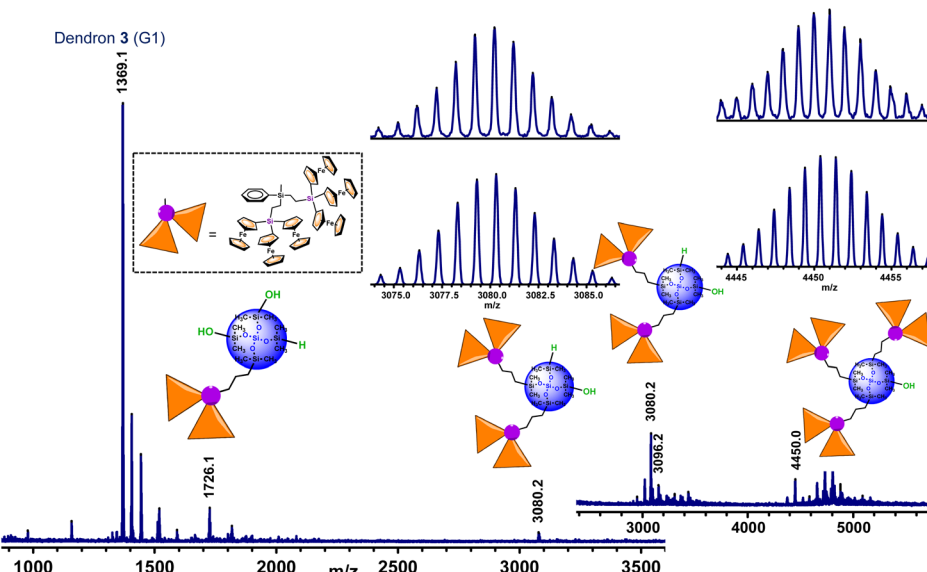


Fig. 3 MALDI-TOF mass spectrum of the reaction product resulting from the hydrosilylation of **3** and $\text{Si}[\text{OSi}(\text{CH}_3)_2\text{H}]_4$ (**I**). The insets show the experimental and calculated isotopic patterns of the peaks at m/z 3080.2 and 4450.0.

tionalized dendrimer, two Si–OH and one Si–H groups (the peak of the monofunctionalized dendrimer with three Si–OH groups appears at m/z 1742.1).

As can be observed in Fig. 3, the peak of the target dendrimer **6** was not detected by the MALDI-TOF technique. On the basis of the good results obtained with the reactions of dendron **1** (G0), this last hydrosilylation was repeated with a larger excess of **3** and the increased reaction time (72 hours) and temperature (100 °C), but it was not possible to fully react the four available Si–H groups. It is evident that there is a noteworthy difference in the reactivity of the first generation dendron **3** in comparison with its corresponding precursor **1**. Steric congestion appears to be responsible for the inability of the four Si–H groups of the tetradirectional core $\text{Si}[\text{OSi}(\text{CH}_3)_2\text{H}]_4$ (**I**) to react with G1-dendron **3**, which is formed by six ferrocenyl units. Even though the fourth branched core of **I** is selected for its extended silicon-containing arms, and dendron **3** is functionalized with an allyl group (to generate a long $-\text{CH}_2-\text{CH}_2-\text{CH}_2-$ chain), steric hindrance imposed by the sterically demanding organometallic units seems to be the key factor that dominates this reaction, preventing full functionalization and leading to dendrimers with structural defects. However, it is important to emphasize that the MALDI-TOF technique tends to show considerable molecular weight discrimination for polydisperse samples.³² This means that, although peaks at higher m/z values are not observed in this spectrum (Fig. 3), the formation of higher molecular weight species cannot be ruled out.

Our attention was then turned to the synthesis of dendronized polymers by polyhydrosilylation reactions of **1** and **3** with poly(methylhydrosiloxane) **II**. The reaction with dendron **1** (G0) as the starting material was accomplished using the methodology mentioned before (see the top of Fig. 4). After

45 hours at 100 °C, analysis of the reaction mixture by ^1H NMR experiments confirmed the complete consumption of the Si–H protons of $(\text{Me}_3\text{SiO})(\text{MeSiHO})_n(\text{SiMe}_3)$ (**II**) at δ 4.72 ppm. New resonances at δ 0.8–1.3 ppm, corresponding to the new hydrosilylated product with CH_2-CH_2 linkages, were also observed indicating the quantitative transformation of the Si–H reactive groups.

The oily reaction mixture was dissolved in CH_2Cl_2 and precipitated twice into methanol. The obtained precipitate was then analyzed by MALDI-TOF mass spectrometry (Fig. 4), showing three different groups of peaks. First of all, the peaks of the expected linear oligomeric species 7_n ($n = 4-12$) appear in the range of m/z 2844–8204 with an interval between the peaks at m/z 670 (this value corresponds to the mass of the repeated fragment $-\text{MeSi}[-\text{CH}_2\text{CH}_2-\text{Si}-\text{Fc}_3]\text{O}-$). Then, there are peaks corresponding to cyclic species 8_n ($n = 4-6$) without terminal groups, at m/z 2681, 3351 and 4021, also separated by m/z 670. Finally, a group of more intense peaks appears between m/z 2903 and 8264 (equally spaced at m/z 670). These signals that differ from the values of 7_n at m/z 60 (mass of one unreacted $-\text{H}_3\text{C}-\text{Si}(\text{H})\text{O}-$ fragment) (see Fig. 4) form a new family of dendronized oligomers and polymers, 9_n ($n = 4-12$). The isotopic masses of all these peaks (see Fig. S27 in the ESI[†]) agree well with the calculated ones, confirming their precisely defined linear or cyclic structures.

In an attempt to obtain a fraction of ferrocenyl-bearing polymers with higher molecular weights, the first precipitate was dissolved again in a small amount of CH_2Cl_2 and precipitated into methanol. The MALDI-TOF spectrum (Fig. S28[†]) proved to follow an identical pattern, when compared to the previous one. However, when the matrix deflection high voltage is set at m/z 2000, species 7_n and 9_n with n values up to 14 (m/z 9604) can be observed. As shown before, although

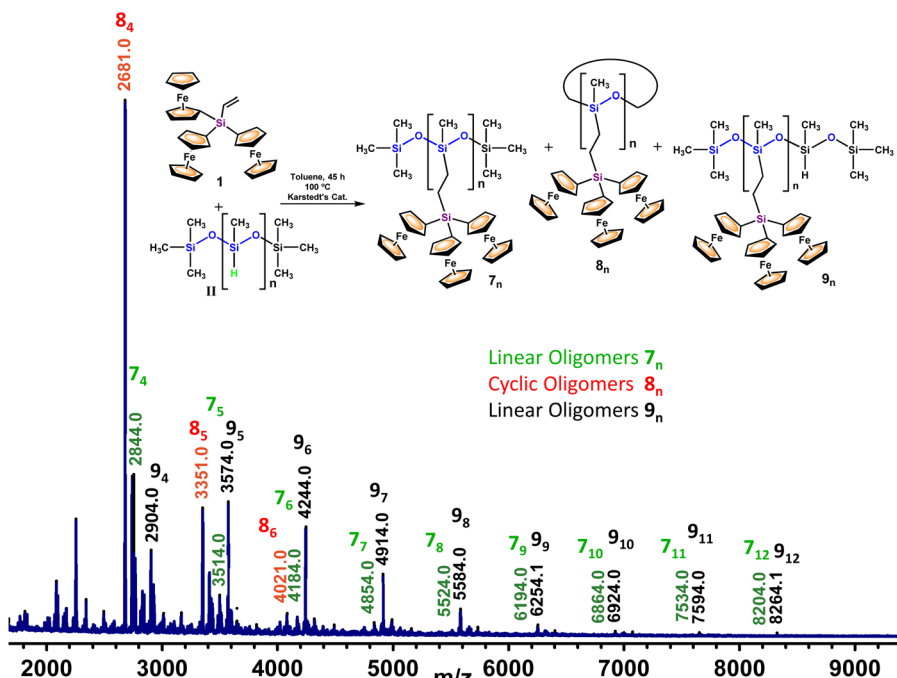


Fig. 4 Structures of the dendritic products 7_n , 8_n and 9_n formed by platinum-catalyzed hydrosilylation of dendron **1** (G0) onto the multifunctional silicone backbone $(\text{Me}_3\text{SiO})(\text{MeSiHO})_n(\text{SiMe}_3)$ (**II**) (top) and MALDI-TOF mass spectrum (bottom).

peaks at higher m/z values are not observed, the formation of higher molecular weight species cannot be ruled out, since MALDI-TOF tends to show a considerable molecular weight discrimination for polydisperse samples.³²

The last step in our proposed dendritic growth approach consisted of incorporating the first generation dendron **3** into the polymeric siloxane backbone **II**. After 72 hours of the hydrosilylation reaction at 100 °C, the oily product was treated, as before, and the precipitates were analyzed by MALDI-TOF. In all the recorded spectra, a major peak was observed at m/z 1369.1, and corresponds to the starting dendron **3**, but we were not able to identify any other species. As observed with dendrimer **6**, this result can be due to the steric demand of **G1**-dendron **3**, which is responsible for the failure of the $\text{Si}-\text{CH}_2-\text{CH}=\text{CH}_2$ groups to react. Nonetheless, given the proved efficiency of the hydrosilylation processes with **G0**-dendron **1**, we are convinced that both dendrimers and dendronized polymers can be chemically prepared with **G1**-dendrons of type **3**. Obviously, reducing the steric congestion proves to be critical, and this can be achieved by adding spacer units to the focal point in the **G1**-dendron **3** and/or in the $-\text{Si}-\text{H}$ arms of the siloxane cores. Indeed, in closely related $\text{Si}-\text{H}$ rich polysiloxane backbones, it was suggested that steric constraints were responsible for the inability of all $\text{Si}-\text{H}$ groups to react with the dendrons.^{19a}

Multinuclear NMR (^1H , ^{13}C , and ^{29}Si) experiments of dendrimers **4** and **5** and dendronized polymers 7_n – 9_n support their assigned structures. On the basis of their ^1H and ^{13}C NMR studies it was determined that the addition of $\text{Si}-\text{H}$ functionality to the vinyl group of **1** proceeds selectively according

to the anti-Markovnikov rule. While the ^1H NMR spectrum of dendrimer **4** (Fig. S8 \ddagger) shows an informative septuplet at δ 4.86 ppm, confirming the presence of one unreacted $\text{Si}-\text{H}$ functionality, the ^1H NMR spectrum of dendrimer **5** (Fig. S14 \ddagger) proves its fully functionalized structure, as no signal appears in this region. The ^1H NMR spectrum of the mixture of dendronized polymers 7_n – 9_n (Fig. S21 \ddagger) shows broader signals, due to the complex mixture of the species and their large size, but with quite similar chemical shift values as the ones of the already reported dendrimers. The ^{13}C NMR spectrum of dendrimer **4** (Fig. S9 \ddagger) shows a representative signal at δ 1.2 ppm for the $\text{H}-\text{Si}-(\text{CH}_3)_2$ group.

Regarding the ^{29}Si NMR experiments, the spectra of dendrimers **4** and **5** and polymers 7_n – 9_n display a peak at around δ –8.3 ppm, corresponding to the $(\text{Fc})_3\text{Si}-\text{CH}_2\text{CH}_2$ group, which supports the incorporation of the trifluorocenylsilyl moieties around the dendritic siloxane frameworks *via* hydrosilylation reactions. Moreover, these spectra show resonances at δ –103.0 (for **4**) and –103.1 ppm (for **5**) which are consistent with the Q-type silicon atoms (Si_QO_4), at δ –21.9 ppm (for 7_n – 9_n) of the D-type silicon atoms (Si_DO_2), and at δ 9.8 (for **5**), 10.1 (for **4**) and 7.5 ppm (for 7_n – 9_n) of the M-type silicon atoms (Si_MO). In the case of **4** an additional resonance can be observed at δ –5.3 ppm corresponding to the unreacted $\text{Si}-\text{H}$ bond. This last signal was corroborated by a $\{^1\text{H}-^{29}\text{Si}\}$ HMQC experiment (see Fig. S10 \ddagger).

MALDI-TOF mass spectrometry provided further evidence for the formation of the new dendritic macromolecules. The mass spectra of **4** and **5** show the most intense peaks at m/z 2159.1 and 2770.1, respectively, which correspond to the M^+

ions. Their isotopic masses agree well with those of the calculated ones, confirming their precisely defined star-shaped dendritic structures and the presence of 9 and 12 pendant ferrocenyl moieties, grouped in three dimensions at the end of each arm. In addition, the IR spectra of dendrimers **4** and **5** and the dendronized polymers **7_n–9_n** are very similar to those of a predominant band corresponding to the $\nu(\text{Si}-\text{O}-\text{Si})$ of the siloxane backbone. The main difference can be observed in the dendrimer **4** spectrum, in which two bands appear at 2137 cm^{-1} ($\nu(\text{Si}-\text{H})$) and at 890 cm^{-1} ($\delta(\text{Si}-\text{H})$) (see Fig. S12‡).

Investigations of the properties of the obtained ferrocenyl-rich dendritic molecules were complemented by the evaluation of their thermal stability by thermogravimetric analysis (TGA). We have found that the mixture of dendronized polymers **7_n–9_n** shows a reasonably high thermal stability (Fig. 5). This TGA experiment was performed under nitrogen and was compared with the TGA results previously obtained by our research group²⁹ for the related octasilsesquioxanes (OS) molecules **10** and **11**, as also shown in Fig. 5. Ferrocenyl-functionalized cubic macromolecules **10** and **11**, peripherally decorated with 24 ferrocenyl units, have also shown good thermal stability.²⁹

As can be seen in Fig. 5, the three multiferrocenyl macromolecules (**7_n–9_n**, **10** and **11**) show similar weight losses when heated between 0 and $1000\text{ }^{\circ}\text{C}$. In particular, the mixture of dendronized polymers **7_n–9_n** displays two weight losses. The first loss, beginning near $200\text{ }^{\circ}\text{C}$, can be attributed to the cleavage of the organic components of the polymeric chains,

whereas the second and major weight loss, between 370 and $650\text{ }^{\circ}\text{C}$, can be ascribed to the breakdown of the inorganic structure. Above this temperature, the TGA curve is almost leveled off and the ceramization yield of **7_n–9_n**, at $1000\text{ }^{\circ}\text{C}$, is 64 wt\% (50 wt\% for **10** and 66 wt\% for **11**). This comparison clearly highlights the great thermal stability of the three ferrocenyl-containing siloxane backbones, regardless of their precisely siloxane nature.

We further examined the residues obtained *via* pyrolysis of **7_n–9_n** under a nitrogen stream at $1000\text{ }^{\circ}\text{C}$, by SEM and energy-dispersive X-ray analysis (Fig. S30‡). The pyrolyzed material obtained from **7_n–9_n** in the ceramization process contains iron particles (Fig. S31–S33‡) and can be attracted to a bar magnet, revealing that polymers **7_n–9_n** could be promising precursors for ceramic materials.

Solid-state structures of G1-dendron **3** and dendrimer **4**

A key challenge in the synthesis of highly branched molecules is to obtain good single crystals, suitable for SCXRD analysis. Since the dendrons and dendrimers reported here are large and complex molecules, they were very hard to crystallize, due to the difficulty in organizing the molecules in the crystalline network. Fortunately, we were able to obtain suitable crystals for hexametallic dendron **3** (**G1**) and nonametallic dendrimer **4** by crystallization in $\text{CH}_2\text{Cl}_2/n\text{-hexane}$ ($1:2$) at $-4\text{ }^{\circ}\text{C}$. The resulting structures are shown in Fig. 6 and complete structural information is collected in the ESI.‡ Both compounds crystallize in the monoclinic space group $P2_1/n$ with one molecule per asymmetric unit. The cyclopentadienyl rings in **3** and **4** are essentially parallel and slightly staggered, and the molecular packing in both cases is achieved by C–H··· π interactions involving the cyclopentadienyl rings.

The structure of hexaferrocenyl dendron **3** (**G1**) shows that the two largest $(\text{CH}_2)_2\text{SiFc}_3$ substituents of the central silicon atom are located as separated as possible (Fig. 6A). The three silicon atoms in the molecule show a nearly tetrahedral environment, with the C–Si–C bond angles close to 109° , similar to the ones reported for **1**.²⁸ The metal atoms of the ferrocenyl substituents attached to the same silicon center are separated by distances ranging from $5.401(1)$ to $6.164(1)$ \AA , while the Fe centers of the ferrocenyl units attached to different silicon atoms are separated by longer distances (from $6.967(1)$ to $12.730(1)$ \AA), as can be seen in Table S5.‡

Dendrimer **4** provides a unique opportunity to examine the solid-state structure of a branched siloxane with nine ferrocenyl units. We consider it important to highlight here that a bibliographic search in the CSD has revealed dendrimer **4** as the branched multiferrocenyl-containing siloxane with the highest number of ferrocenes fully characterized by single-crystal X-ray diffraction. The central tetrahedral SiO_4 unit displays three large $(\text{CH}_3)_2\text{Si}-(\text{CH}_2)_2-\text{SiFc}_3$ substituents and a smaller $-\text{Si}(\text{CH}_3)_2\text{H}$ one. This shorter moiety shows a statistical disorder that was modelled with two sets of positions for the $-\text{OSi}(\text{CH}_3)_2\text{H}$ fragment, with $60\text{--}40\%$ occupations. The presence of a remaining Si–H reactive group in dendrimer **4** is not only a simple structural defect, but makes it a genuine func-

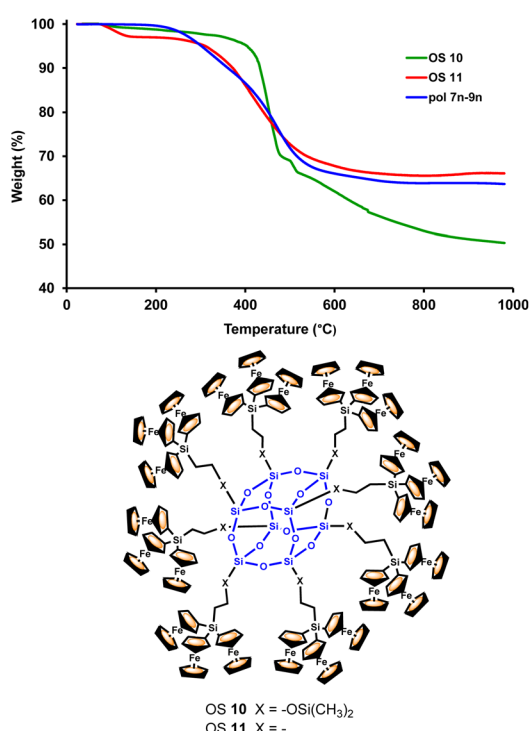


Fig. 5 Comparison of the TGA thermogram of dendronized polymers **7_n–9_n**, under N_2 at a heating rate of $10\text{ }^{\circ}\text{C min}^{-1}$, and the TGA thermograms of ferrocenyl-functionalized octasilsesquioxanes **10** and **11** under the same experimental conditions.



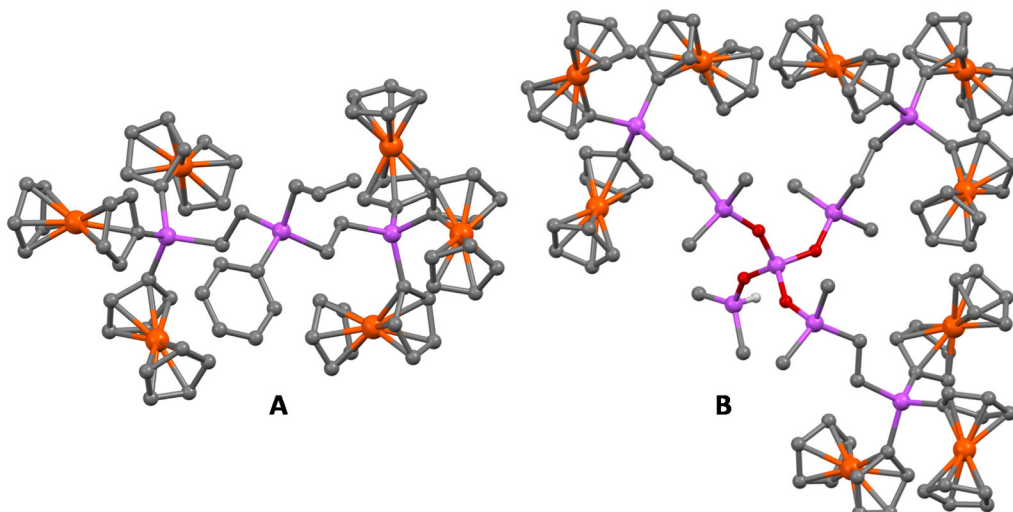


Fig. 6 Crystal structures of G1-dendron 3 (A) and dendrimer 4 (B). Hydrogen atoms are omitted for clarity.

tional core, being capable to build new types of asymmetric dendrimers by hydrosilylation reactions.

The eight silicon atoms in the molecule display the expected tetrahedral environment with all the C–Si–C, O–Si–O and O–Si–C bond angles close to 109° . The metal atoms of the ferrocenyl substituents attached to the same silicon center are located at distances ranging from 5.806(2) to 6.160(2) Å, and the iron centers of the Fc units attached to different silicon atoms are separated by distances ranging from 6.224(2) to 18.487(2) Å (see Table S5†). It is remarkable that the large number of ferrocenyl units in this molecule packs them closer to each other, and consequently, the difference between the largest distance for the iron atoms from Fc attached to the same Si atom and the shortest Fe–Fe distance from the metals in adjacent SiFc_3 groups is only 0.064 Å.

Electrochemical studies of 1 (G0) and 3 (G1), dendrimers 4 and 5 and dendronized polymers 7_n – 9_n

Due to the ferrocenyl-rich nature of the silicon-based dendritic molecules obtained here, a significant property to consider is their electroactivity. The solution electrochemical behavior of isolated ferrocenyl-rich dendritic macromolecules 3–5 and the mixture of dendronized polymers 7_n – 9_n was examined by cyclic (CV) and square wave (SWV) voltammetry and was compared to the redox behavior of the precursor molecule, triferro-

cenylvinylsilane $\text{Fc}_3\text{SiCH}=\text{CH}_2$ (1) (G0). The electrochemical responses are shown in Fig. 8 and 9, and the electrochemical data are summarized in Table 1.

It is worth noting that the solution redox behavior of this multiferrocenyl dendritic family is sensitive to the dendrimer generation and to the number of ferrocenyl moieties. In order to check the evolution of the dendritic effect on the voltammetric response, cyclic voltammograms (CVs) were recorded using a Pt-disk working electrode, in dichloromethane solution with 0.1 M of tetra-*n*-butylammonium hexafluorophosphate ($[\text{n-Bu}_4\text{N}]^+[\text{PF}_6]^-$) as the supporting electrolyte, and containing 10^{-4} M of the redox-active species, in all cases.

Fig. 7 shows the representative CV responses of dendron 3 and dendrimer 5, as examples, together with the CV of triferrocenyl precursor 1, for comparative purposes. Qualitatively similar responses were observed for dendrimer 4 and dendronized polymers 7_n – 9_n (see Fig. S38 and S43,‡ respectively). As can be seen, the solubility of all the studied multimetallic dendritic systems is drastically affected by the oxidation and reduction steps. Therefore, the voltammetric response deviates from the typical CV wave shape, characteristic of freely diffusing soluble species undergoing reversible transfer processes.³³

Specifically, it is evident from Fig. 7 that for dendritic precursor 3 (bearing 6 ferrocenyl (Fc) units) and dendrimer 5 (12 Fc), the anodic and cathodic peak currents are dissimilar. The

Table 1 Electrochemical data for dendritic species 11, 3–5 and 7_n – 9_n

| Compound ^a | ${}^1E_{1/2}$ | ${}^2E_{1/2}$ | ${}^3E_{1/2}$ | $\Delta E ({}^2E_{1/2} - {}^1E_{1/2})^b$ | $\Delta E ({}^3E_{1/2} - {}^2E_{1/2})^b$ | $\Delta E ({}^3E_{1/2} - {}^1E_{1/2})^b$ |
|-----------------------|---------------|---------------|---------------|------------------------------------------|------------------------------------------|------------------------------------------|
| 1 | 0.436 (0.400) | 0.736 (0.568) | 1.048 (0.704) | 300 (168) | 312 (136) | 612 (304) |
| 3 | 0.400 (0.440) | 0.720 (0.604) | 1.068 (0.736) | 320 (164) | 348 (132) | 668 (296) |
| 4 | 0.392 (0.444) | 0.684 (0.620) | 0.896 (0.756) | 292 (176) | 212 (136) | 504 (312) |
| 5 | 0.456 (0.452) | 0.696 (0.624) | 0.819 (0.753) | 240 (172) | 123 (129) | 363 (301) |
| 7_n – 9_n | (0.452) | (0.645) | (0.736) | (193) | (91) | (284) |

^a $E_{1/2}$ for V vs. the SCE, determined by square wave voltammetry in CH_2Cl_2 solution with $[\text{n-Bu}_4\text{N}]^+[\text{B}(\text{C}_6\text{F}_5)_4]^-$, or $[\text{n-Bu}_4\text{N}]^+[\text{PF}_6]^-$ with $\text{CH}_2\text{Cl}_2/\text{CH}_3\text{CN}$ (3 : 0.5) for values indicated in parentheses. ^b Peak potential separation values, ΔE , are given in mV.



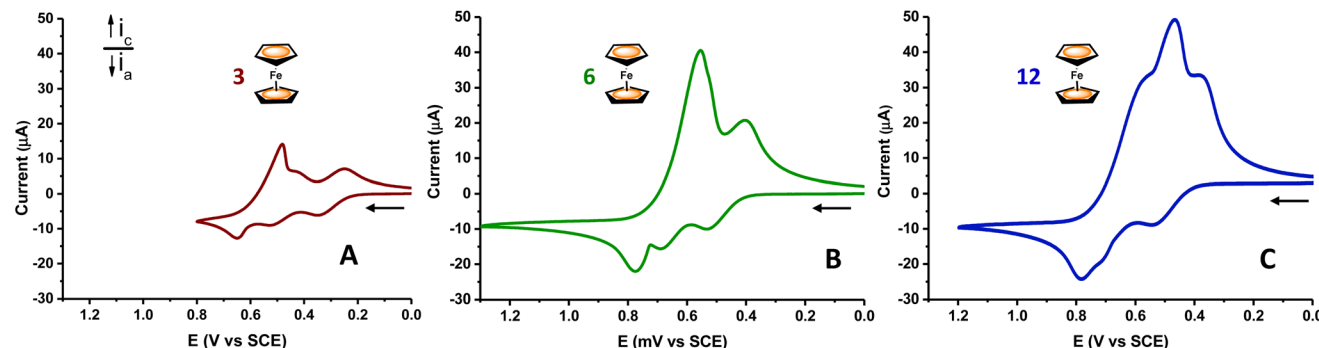


Fig. 7 Cyclic voltammograms of multiferrocenyl dendritic molecules **1** (A), **3** (B), and **5** (C) measured in $\text{CH}_2\text{Cl}_2/[n\text{-Bu}_4\text{N}][\text{PF}_6]$ at 100 mV s^{-1} .

anodic waves of macromolecules **3–5** and $7_n\text{--}9_n$ occur by three poorly resolved redox processes, when compared to those of precursor **1**. In addition, the cathodic scan gives rise to extensive precipitation of the oxidized, highly charged species $[3^{6+}][\text{PF}_6^-]_6$, $[4^{9+}][\text{PF}_6^-]_9$, $[5^{12+}][\text{PF}_6^-]_{12}$ and $[7_n\text{--}9_n]^{n+}[\text{PF}_6^-]_n$ on the electrode surface, which results in distorted, very sharp peaks. The sharpness of these peaks is strongly related to the stripping nature of the electrochemical processes. As the dendritic material is deposited onto the electrode surface, it is redissolved back in the solution, as a result of the cathodic electron-transfer reaction. Nevertheless, not all the electrodeposited dendritic molecules are stripped from the surface in a single scan. In fact, there are some strong precipitated and/or adsorbed dendrimer molecules remaining on the electrode surface upon reduction.

Likewise, as the number of ferrocenyl moieties per dendritic molecule increases (that is, as the generation starts), the cathodic stripping wave increases in magnitude (see Fig. 7), meaning that the electrogenerated polycationic macro-molecules $[3^{6+}][\text{PF}_6^-]_6$ and $[5^{9+}][\text{PF}_6^-]_9$ become less soluble in this solvent/electrolyte medium.

Taking dendrimer **5** as a representative example, Fig. 8 shows how by enriching the solvent mixture with acetonitrile, the cathodic stripping peaks progressively tend to disappear. Thus, the CVs of dendron **3**, dendrimers **4** and **5**, and the CV of the mixture $7_n\text{--}9_n$, show three well-resolved anodic waves, which are indicative of appreciable electronic interactions between the silicon-bridged triferrrocenyl moieties as they are successively oxidized. Under these conditions, the $E_{1/2}$ values

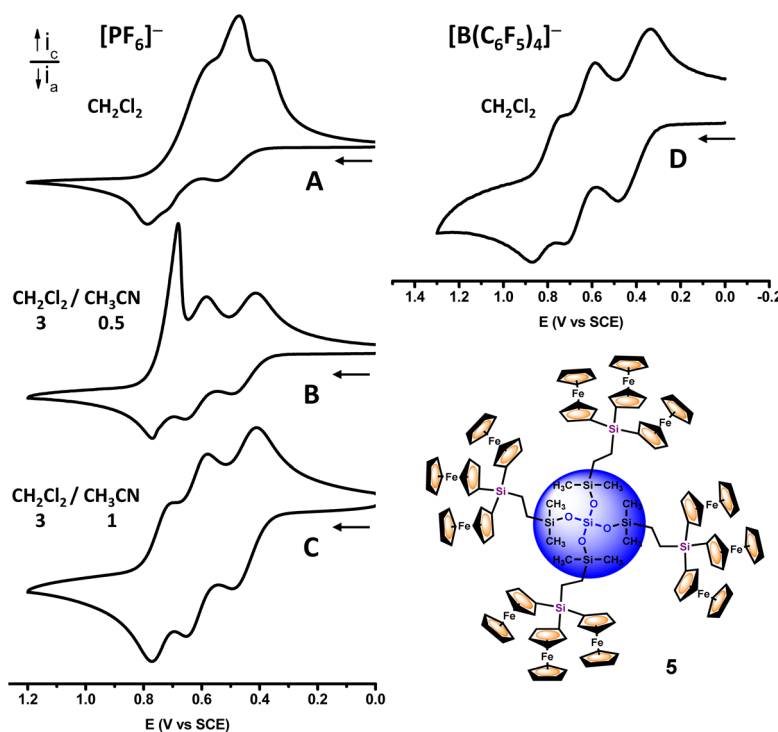


Fig. 8 Cyclic voltammograms of dendrimer **5**, measured under the conditions indicated therein, at 100 mV s^{-1} .



of the three redox processes can be measured for all the compounds and are summarized in Table 1.

Fig. 8 also shows the remarkable improvement for the CV response of dendrimer 5 when CH_2Cl_2 and tetrakis(pentafluorophenyl)borate ($[\text{n-Bu}_4\text{N}][\text{B}(\text{C}_6\text{F}_5)_4]$) are employed as the solvent/supporting electrolyte. A similar qualitative behavior is observed with dendritic species 3 and 4 and dendronized polymers $7_{\text{n}}\text{--}9_{\text{n}}$ (see Fig. S37, S39 and S41 \ddagger). The weak coordinating counter-anion $[\text{B}(\text{C}_6\text{F}_5)_4]^-$ is known to stabilize the highly charged species in solution. Furthermore, the low ion-pairing capabilities lead to a poor shielding of the electrostatic interactions among the redox active sites, resulting in an increase of the observed redox splitting. 34 Consequently, in the presence of the less coordinating electrolyte anion $[\text{B}(\text{C}_6\text{F}_5)_4]^-$, a better resolution, improved electrochemical reversibility, and very little electrode adsorption are observed, which is indicative of a better solubility of the multiferrocenium electrogenerated dendritic species 3^{6+} , 4^{9+} , 5^{12+} and $7_{\text{n}}\text{--}9_{\text{n}}^{n+}$ in this solvent/electrolyte medium. Clearly, in agreement with our previous observations, 21,29 and with the seminal results reported by Geiger and co-workers, 34 the combination of CH_2Cl_2 and $[\text{n-Bu}_4\text{N}][\text{B}(\text{C}_6\text{F}_5)_4]$ as the solvent/electrolyte medium provides more favorable conditions for the electrochemical studies of multiferrocenyl compounds, minimizing the ion-pairing interactions between the fluoroarylborate anion and the polycationic species generated in the oxidation processes.

In addition to the electrochemical studies of the dendrimers in a homogeneous solution, we have explored the redox properties of the ferrocenyl-containing dendritic molecules confined to electrode surfaces, where the films of the dendritic molecules serve as electrode modifiers. It is well known that the deliberate and controlled modification of electrode surfaces with films of electroactive molecules can produce electrodes with new and interesting properties that may form the basis of new electrochemical applications and novel devices. In addition, such modified electrodes can provide a better insight into the nature of charge transfer and charge transport processes in the films. Therefore, chemically modified electrodes are important due to their potential applications in the areas of photo- and electro-catalysis, ion recognition, biosensing and information storage. 33a,35

Remarkably, a valuable feature of the new dendritic macromolecules with the highest number of ferrocenes, dendrimer 5 and dendronized polymers $7_{\text{n}}\text{--}9_{\text{n}}$, is their noteworthy tendency to modify electrodes, resulting in dendrimer-modified electrode surfaces that remain persistently attached to the electrodes. Electrodes modified with the films of these multiferrocenyl-containing dendritic molecules were prepared using a 10^{-4} M solution of the corresponding macromolecule in a 10^{-1} M $\text{CH}_2\text{Cl}_2/[\text{n-Bu}_4\text{N}][\text{PF}_6]$ mixture (see the Experimental section). The electrochemical deposition could be observed onto the Pt electrodes by continuous increases in both the anodic and cathodic peak currents during consecutive cyclic voltammetric scans. We have evaluated the stability of the films of macromolecules 5 and $7_{\text{n}}\text{--}9_{\text{n}}$, by transferring the modified Pt electrodes into fresh CH_2Cl_2 or CH_3CN /electrolyte solutions. As a

representative example, Fig. 9 shows the CV responses of an electrodeposited film of dendronized polymers $7_{\text{n}}\text{--}9_{\text{n}}$, in both solvents. When CH_2Cl_2 is used for the fresh solution, three successive well-resolved, reversible oxidation–reduction waves are observed, with formal potential values of ${}^1E_{1/2} = 0.459$, ${}^2E_{1/2} = 0.600$ and ${}^3E_{1/2} = 0.705$ V, vs. the SCE. These values are similar to the ones observed for the dendronized polymers $7_{\text{n}}\text{--}9_{\text{n}}$ in solution (see Table 1). The surface-confined nature of the electroactive $7_{\text{n}}\text{--}9_{\text{n}}$ films is proved by the linear dependences of the anodic and cathodic peak currents, of the three waves, on the potential sweep rate v (see the top of Fig. 9A). 35,36

As expected, the extent of multiferrocenyl-dendritic molecule adsorption, Γ , is strongly dependent on the number of scans performed (see Fig. S42 and S44 \ddagger). The surface coverages, Γ (mol cm^{-2}), of the ferrocenyl sites can be estimated from the charge, Q , under the ferrocene voltammetric current

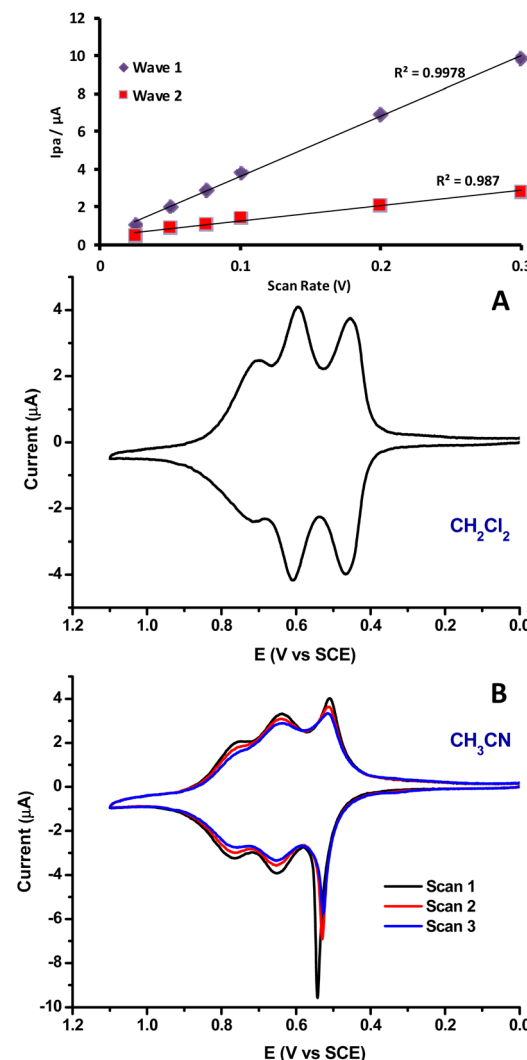


Fig. 9 CVs of a Pt-disk electrode modified with a film of $7_{\text{n}}\text{--}9_{\text{n}}$ (after 10 successive scans) measured in: (A) $\text{CH}_2\text{Cl}_2/[\text{n-Bu}_4\text{N}][\text{PF}_6]$ ($\Gamma = 1.20 \times 10^{-9} \text{ mol Fe cm}^{-2}$) and (B) $\text{CH}_3\text{CN}/[\text{n-Bu}_4\text{N}][\text{PF}_6]$ ($1.10 \times 10^{-9} \text{ mol Fe cm}^{-2}$). Inset (top): plot of the peak currents versus the sweep rate for the CV shown in (A).

peaks, in the ferrocenyl dendritic molecule-free electrolyte solution, using the following relationship: $\Gamma = Q/nFA$, where Q is the integrated charge of the redox peak, n is the number of electrons involved in the redox reaction, F is the Faraday constant, and A is the real surface area of the electrode. In particular, the surface coverage of the electroactive ferrocenyl sites in the film for the studied example of dendronized polymers **7_n–9_n** shown in Fig. 9A is 1.20×10^{-9} mol Fe per cm², and corresponds to a modified electrode prepared by cyclically scanning the potential (between +0.0 and +1.3 V vs. SCE), in degassed CH₂Cl₂ solutions of the corresponding dendritic molecule, during 10 scans.

For comparison purposes, the surface coverages obtained from the electrochemical responses of the dendrimer **5** film ($\Gamma = 4.19 \times 10^{-11}$ mol Fe per cm², see Fig. S42[‡]) and the octaferrocenyl silsesquioxane **10** film ($\Gamma = 6.05 \times 10^{-12}$ mol Fe per cm²)²⁹ measured under identical conditions (CH₂Cl₂/[*n*-Bu₄N][PF₆]) and prepared with the same Pt-disk electrode and with identical number of scans (10 scan), are considerably lower. These differences in the values of the surface coverages of the modified electrodes could be related to the attachment mode of the electroactive triferrrocenyl groups to the corresponding siloxane backbone. For dendronized polymers **7_n–9_n** the electroactive silicon-bridged triferrrocenyl units ($-\text{SiFc}_3$) append from the main linear and flexible siloxane polymeric chain (pendant ferrocenyl units). In contrast, in dendrimer **5** and octaferrocenyl silsesquioxane **10** (see Fig. 5), the $-\text{SiFc}_3$ units are attached to more rigid spherical and cubic cores. Perhaps, the increased flexibility of the $-\text{Si}-\text{O}-\text{Si}-$ siloxane chain in the polymer backbone affects the redox behavior of dendronized polymers **7_n–9_n**, allowing more effectiveness during the multiferrocenyl-dendritic molecule electrodeposition and/or electroprecipitation process onto the Pt electrode, resulting in modified electrodes with considerably higher values of the surface coverages, Γ (mol cm⁻²). It therefore appears from these data that dendronized polymers **7_n–9_n** is a better electrode modifier than dendrimer **5** and octasilsesquioxane **10**.

On the other hand, the Pt electrode modified with dendronized polymers **7_n–9_n** shows a very different response in a fresh CH₃CN/[*n*-Bu₄N][PF₆] solution. As can be seen in Fig. 9B, the first anodic peak in this CV is atypically sharp and narrow, something that is not observed in CH₂Cl₂ solution. This peak can be detected in the first four scans and corresponds to the process mentioned before, known as break-in. This means that the electrodeposited film is activated during the first scans, as it breaks to allow the flow of both the solvent and the [PF₆]⁻ anions.³⁴ This behavior is typical of redox-active films into which the diffusion of the solvent molecules and supporting electrolyte ions is slow, as observed for thick redox-active films or films that do not swell easily in the solvent employed for the electrochemical study. The anodic peak current decreases gradually with each successive scan. After the “break-in” period of four scans, the film reaches a steady state, at which stable and reproducible, and well-defined reversible current–potential curves are obtained.

Finally, the electrodeposited films of dendrimer **5** and dendronized polymers **7_n–9_n** are quite robust, persisting after

rinsing or soaking the electrode in CH₂Cl₂ (with or without a dissolved electrolyte). The stability of these electroactive films was further demonstrated by their nearly quantitative persistence after continuous CV scans (for more than 4 h) in ferrocenyl-macromolecule-free CH₂Cl₂ solutions with [PF₆]⁻ as the electrolyte anion.

Conclusions

In summary, we have designed and successfully synthesized, *via* a convergent approach, three different multiferrocenyl-terminated branched molecules: dendrons, dendrimers, and dendronized polymers, starting from triferrrocenylvinylsilane (**1**) as the key monomer, and using Pt-catalyzed hydrosilylation and alkenylation with Grignard reagents, as consecutive reaction steps. Remarkably, just by varying the reaction conditions between **1** and the tetradirectional siloxane core Si[OSi(CH₃)₂H]₄, the fully functionalized dendrimer **5** (with 12 Fc moieties) or the dendrimer **4** (with 9 Fc units) might be prepared. This last one is a certainly outstanding achievement, because one of its branch-ends is a Si–H group, making it a reactive macromolecule for further convergent growth. Unfortunately, all the attempts to use **G1**-dendron **3** in this convergent methodology, to prepare larger generation dendrimers and dendronized polymers, were hampered by its high steric hindrance. However, dendrimers with structural defects were detected by MALDI-TOF experiments, suggesting that by adding spacer units to the focal point in the **G1**-dendron **3** and/or in the $-\text{Si}-\text{H}$ arms of the siloxane cores, the desired macromolecules may be obtained. Despite the complex nature of these macromolecules, the obtention of good quality crystals has allowed the detailed structure solution of **G1**-dendron **3** and dendrimer **4** by SCXRD and consequently, a unique insight and a better understanding of the stereochemistry of these remarkable species. In particular, dendrimer **4**, peripherally decorated with nine Fc units, has been revealed as the branched multiferrocenyl-containing siloxane with the highest number of ferrocenes fully characterized by SCXRD. Furthermore, solution electrochemical studies have confirmed that the ferrocenyl units in all the obtained dendrons, dendrimers, and dendronized polymers show electronic interaction in threes through a silicon atom. Finally, we have demonstrated the feasibility of modifying Pt-disk electrode surfaces with dendritic molecules such as dendrimer **5**, bearing a predetermined number of redox centers, and a mixture of dendronized polymers **7_n–9_n**. A noteworthy feature of platinum electrodes modified with films of such multiferrocenyl dendritic molecules is that they are extremely durable and reproducible.

Author contributions

Conceptualization, S. B. and I. C.; methodology, S. B., I. C., and J. P.; X-ray analysis, J. P.; electrochemical studies, S. B. and I. C.; formal analysis, S. B., I. C., and J. P.; investigation and



data curation, S. B., I. C., and J. P.; writing, original draft preparation, and editing, S. B., I. C., and J. P.; supervision, S. B.; project administration and funding acquisition, I. C.

Conflicts of interest

There are no conflicts to declare.

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