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# Dendrimer crown-ether tethered multi-wall carbon nanotubes support methyltrioxorhenium in the selective oxidation of olefins to epoxides†

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Benzo-15-crown-5 ether supported on multi-wall carbon nanotubes (MWCNTs) by tethered poly(amidoamine) (PAMAM) dendrimers efficiently coordinated methyltrioxorhenium in the selective oxidation of olefins to epoxides. Environmentally friendly hydrogen peroxide was used as a primary oxidant. Up to first and second generation dendrimer aggregates were prepared by applying a divergent PAMAM methodology. FT-IR, XRD and ICP-MS analyses confirmed the effective coordination of methyltrioxorhenium by the benzo-15-crown-5 ether moiety after immobilization on MWCNTs. The novel catalysts converted olefins to the corresponding epoxides in high yield without the use of Lewis base additives, or anhydrous hydrogen peroxide, the catalyst being stable for more than six oxidative runs. In the absence of the PAMAM structure, the synthesis of diols largely prevailed.

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#### Introduction

Multi-wall carbon nanotubes (MWCNTs) are efficient supports for the immobilization of reactive organometallic species thanks to their high surface area to volume ratio value, chemical stability, and unique redox, rheological and structural properties. <sup>1,2</sup> They show surface defect sites for the anchoring of spacer chains and selective recognition sites, as for example for crown ethers. <sup>3</sup> Different applications of MWCNTs-crown ether functionalized systems are reported, <sup>4</sup> including dispersive microsolid phase extraction procedures, <sup>5</sup> exchange reaction in quasi-solid-state processes, <sup>6</sup> absorption of metals and organic pollutants, <sup>7</sup> and ion-detection. <sup>8</sup> In these latter cases, MWCNT-

crown ether tethered poly(amidoamine) (PAMAM) dendrimers received increasing attention. These supramolecular aggregates are characterized by recognition sites on both the dendrimer core and the end-termination of the system. The divergent methodology for the preparation of PAMAM dendrimers encompasses the grafting of 1,3-diamino propane intermediates with methyl acrylate and ethylenediamine, followed by linkage of the crown ether moiety. The chemical complexity of PAMAM dendrimers (number of dendrimer sidechains and type of ramification) depends on the successive methyl acrylate/ethylenediamine treatments, and controls the distribution of the recognition sites, the leaching of the active species, 12-14 and the dispersibility of MWCNTs in organic solvents. In organic solvents.

Crown ethers are coordination sites for methyltrioxorhenium (MTO),16,17 a potent catalyst for the activation of hydrogen peroxide (H2O2) by formation of two reactive intermediates, namely monoperoxo [MeRe(O<sub>2</sub>)O<sub>2</sub>] and bis(peroxo) [MeRe(O<sub>2</sub>)O<sub>2</sub>] η2-metal complexes.<sup>18</sup> The chemistry of MTO is largely reviewed, 19 including σ C-H oxygen atom insertion 20 and oxy-functionalization,21 oxidative nucleophilic substitution,22 oxidative desulfurization (ODS),23 synthesis of quinone derivatives,24 Baeyer-Villiger rearrangement,25 metathesis,26 methanolysis of glycals,27 and epoxidation of olefins in traditional28 and non-conventional solvents, such as chiral aliphatic amines,24 fluorinated solvent29 and ionic liquids.30 The formation of diols is the major drawback in the epoxidation of olefins, as a consequence of the acidic property and high affinity of Re(vII) for the oxygen atom. Lewis bases such as pyridine, pyridine derivatives,31,32 and bipyridines,33,34 have been used as

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<sup>†</sup> Electronic supplementary information (ESI) available: SI#1 FT-IR of intermediates II A-D, III A-D and catalysts IV A-D and V E-H; SI#2: difference method of neutron diffraction and isotopic substitution SI#3: mass spectrometry profiles of compounds 1-21; SI#4: nuclear magnetic resonance spectra of compounds 1-21, SI#5 far IR of IV D and V G compared to the reference MTO-1,2-cyclohexanediamine complex and SI#6 transmission electron microscopy (TEM) of V F and V H. See DOI: 10.1039/d0ra02785e

additives to inhibit this side-reaction, their efficacy depending from the stability of the corresponding MTO complex. As an alternative, anhydrous H<sub>2</sub>O<sub>2</sub> and urea-H<sub>2</sub>O<sub>2</sub> adduct (UHP) have been also applied.35 The immobilization of MTO on solid supports can solve the problem of selectivity, stability and recyclability of the catalyst. Immobilization procedures have been reported including organic polymers bearing Lewis bases, such as chitosan, <sup>36</sup> poly(4-vinylpyridine), <sup>37</sup> poly(4-vinylpyridine-N-oxide),38 and MTO/nitrogen ligand complexes,39 and in alternative different inorganic supports.40-42 More recently, MWCNTs wrapped on poly(4-vinylpyridine)/MTO showed high reactivity in the desulfurization of synthetic diesel fuel. 43 Lewisbase sites generally enhance the formation of epoxides, 44,45 but drawbacks relative to toxicity of the Lewis base additives (e.g. pyridine and pyridine derivatives), partial selectivity, low reactivity and leaching of rhenium have yet to be completely solved.

We report here the preparation and structural characterization of novel heterogeneous catalysts based on the immobilization of MTO on MWCNTs functionalized with benzo-15crown-5 ether (B15C5) in the presence of tethered PAMAM dendrimers. Up to first and second generation of B15C5 PAMAM aggregates were obtained by applying the divergent methodology. The variety of catalysts was increased by the use of alkyl-diamino linkers of various length (from C-2 to C-6 atoms, respectively). FT-IR, ICP-MS and XRD analyses confirmed the coordination of MTO by the B15C5 moiety after immobilization on MWCNTs. The novel catalysts were applied in the epoxidation of a large panel of olefins using H<sub>2</sub>O<sub>2</sub> (35% water solution) as green oxidant in the absence of Lewis bases additives to yield epoxides in high yield and selectivity. As a general trend, alkyl-diamino linkers with longer carbon atom chain afforded the most reactive systems in PAMAM aggregates. Diols were obtained as the main reaction products in the absence of the dendrimer structure, highlighting the role of PAMAM in the selectivity of the reaction.

#### Results and discussion

#### Preparation of MWCNTs-B15C5/MTO catalysts IV A-D

MWCNTs-B15C5/MTO catalysts **IV A–D** were prepared by immobilization of MTO on previously functionalized MWCNTs-B15C5 support. Briefly, MWCNTs **I** were treated with different alkyl-diamino linkers, ethylene diamine (EDA), propylene diamine (PDA), cadaverine (CV), and hexamethylene diamine (HAD) (from C-2 to C-6 atoms, respectively), in the presence of *N*,*N*-di-isopropylcarbodiimide (DIC), 1-hydroxybenzotriazole (HOBt), and *N*,*N*-diisopropyl ethylamine (DIPEA) in DMF at

25 °C for 24 h, to yield intermediates II A-D.46 The coupling procedure was repeated with 4'-carboxy-benzo-15-crown-5 (B15C5) as electrophilic reagent, to afford MWCNTs-B15C5 III A-D (Scheme 1). The FT-IR analysis of intermediates II A-D and III A-D showed the shift of the peak at 1730 cm<sup>-1</sup> (stretching vibration of the carboxylic group in native MWCNTs I) at 1637 cm<sup>-1</sup> (SI#1, Fig. S1 and S2†), confirming the formation of the amide bond. 47 Moreover, intermediates III A-D showed the characteristic peaks at 1120 cm<sup>-1</sup> and 1210 cm<sup>-1</sup>, corresponding to C-O stretching vibrational mode of B15C5 (SI#1, Fig. S2†).8 MWCNTs-B15C5 III A-D were successively treated with MTO in THF under gentle stirring at 25 °C for 6 h, to afford MWCNTs-B15C5/MTO catalysts IV A-D.43 Catalysts IV A-D showed the typical Re=O stretching vibrational mode at 953 cm<sup>-1</sup> (SI#1, Fig. S3†). Scanning Electron Microscopy (SEM) analysis of IV A-D (Fig. 1) confirmed the integrity of MWCNTs after the functionalization procedure. Powder X-ray analysis (XRD) was applied to evaluate the coordination geometry of MTO in representative intermediate III D and catalyst IV D.48-50 The X-ray data of III D and IV D were treated by the difference method of neutron diffraction and isotopic substitution (NDIS)37,51-53 in order to solve the contemporary presence of highly structured MWCNTs and amorphous B15C5 (generally indicated as support in the next discussion). Assuming that the general structure of III D was not significantly altered during the MTO loading, structural data for MTO/B15C15 complex were

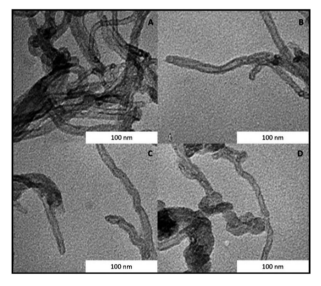
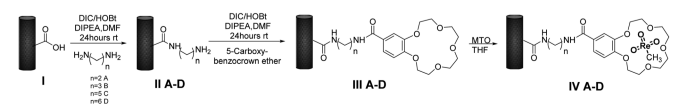


Fig. 1 SEM images of catalyst IV A–D. The structure of MWCNTs is not modified after the functionalization procedure with B15C5.



Scheme 1 Synthesis of catalysts IV A-D.

obtained by subtracting the radial distribution D(r) function of **III D** to that of **IV D** (Fig. 2, panels A and B) (see SI#2† for a detailed description of the difference method technique).<sup>54</sup> The structure and distribution functions of **III D** and **IV D** are reported in SI#2 (Fig. S5 and S6,† respectively).

Note that the oscillation pattern in the D(r) function of the two samples remain almost the same, confirming the hypothesis that the MTO loading didn't alter the structural order of the support. The D(r) function showed three peaks, set at about 1.80, 2.45, and 3.00 Å, respectively (Fig. 2, panel C). The first coordination sphere of the Re atom consists of six interactions assuming a quite octahedral conformation around the metal. The first peak was fitted by three metal-oxygen interaction lengths of exactly 1.80 Å and one metal-carbon interaction of 1.75 Å, respectively. The second peak at 2.45 Å is due to the other two interactions Re-O necessary to complete the octahedral coordination sphere. These interactions are also responsible for the peak at 3.00 Å (mutual interactions). The carbon atoms of the crown ether moiety might also contribute to this peak. The successive signals in the D(r) function were not further considered, since they derived from long-range MTO-MWCNTs interactions. In Fig. 2 (panel C) the function is compared with the theoretical peak shape function calculated for the octahedral model represented in Fig. 2 (panel D). The Re-O and Re-C bond distance values are quite in accord with the mean values previously reported for MTO/Ln complexes.<sup>55</sup> As shown by the model (Fig. 2, panel D), the Re atom is coordinated by two adjacent oxygen atoms of the B15C5 ring. The model refers to the couple of oxygen atoms bonded to the aromatic ring, but the possibility that another couple of adjacent oxygen atoms is involved in the interaction with the metal cannot be completely ruled-out.

Moreover, the first Re coordination sphere analysis does not allow to determine the relative position of the methyl group with respect to the crown-ether oxygen atoms, due to the small electron differences between this group and the oxygen ones.

The distances between Re atom and ligands in the octahedral arrangement are reported in Table 1.

#### Preparation of MWCNTs-B15C5/PAMAM/MTO catalysts V E-H

We prepared first and second-generation dendrimer catalysts **V** E-F and **V** G-H, respectively. As a general procedure, intermediates **II A** and **II D** were treated with methyl acrylate in MeOH at 25 °C for three days, followed by addition of alkyl-diamino linkers EDA and HAD, to yield **III** E-F, respectively (Scheme 2, pathway A). The reaction of **III** E-F with B15C5 in the presence of DIC, HOBT, and DIPEA, in DMF at 25 °C for 24 h, afforded **IV** E-F. These latter intermediates were further converted to first-generation MWCNTs-B15C5/PAMAM/MTO catalysts **V** E-F by treatment with MTO in THF at 25 °C. In a similar way, two successive treatments of **II A** and **II D** with methyl acrylate and alkyl-diamino linkers afforded **III** G-H (Scheme 2, pathway B),

Table 1 Distances between Re atom and ligands in the octahedral arrangement of the complex

Atoms	$r^a$ , Å
Re-O1	1.80
Re-O2	1.80
Re-O3	1.80
Re-O4	2.45
Re-O5	2.45
Re-C	1.75

<sup>&</sup>lt;sup>a</sup> The mean square deviation was  $\sigma_{ij} = 0.06$ .

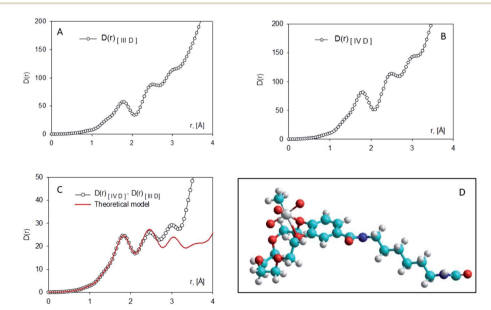
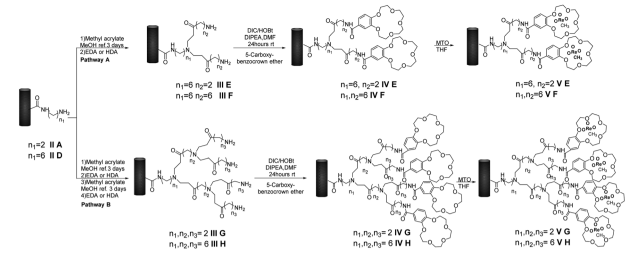


Fig. 2 Red line represents the theoretical model. Panel (A) weighted experimental radial distribution function D(r) of III D. Panel (B) experimental radial distribution function D(r) of IV D. Panel (C) difference curve for the two radial distribution functions, D(r)[IV D] D(r)[III D] D

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Scheme 2 Synthesis of MWCNTs-B15C5/PAMAM/MTO catalysts V E-H. Pathway A: preparation of the first-generation of dendrimer hybrids V E-F. Pathway B: preparation of the second-generation of dendrimer hybrids V G-H.

successively converted into IV G-H under previously reported experimental conditions.

Treatment of IV G-H with MTO in THF at 25 °C yielded the second-generation MWCNTs-B15C5/PAMAM/MTO catalysts IV G-H. FT-IR analyses of catalysts V E-H are reported in SI#1 (Fig. S4†).

Irrespective from the sample, the characteristic Re=O stretching vibrational mode at 950 cm<sup>-1</sup>, and the CONH stretching vibrational mode at 1630 cm<sup>-1</sup>, were always observed, besides to B15C5 C-O stretching signals at 1120, 1210, and 1250 cm<sup>-1</sup>, respectively. The integrity of MWCNTs in catalysts V E-H after the PAMAM functionalization was confirmed by SEM analysis (Fig. 3). Moreover, some sections of the MWCNTs in catalysts V F and V H, as representative examples, showed the presence of the PAMAM coating in TEM images (SI#6†). The loading factor of catalysts IV A-D and V E-H was evaluated by Inductively Coupled Plasma Mass-Spectrometry (ICP-MS) analysis (Table 2, entries 1-8).

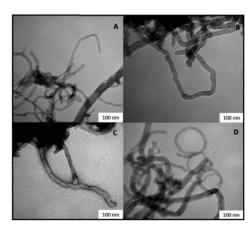


Fig. 3 Scanning electron microscopy images of catalysts V E-H. Panel (A) catalyst V E. Panel (B) catalyst V F. Panel (C) catalyst V G. Panel (D) catalyst V H.

Catalysts V E-F showed loading factors of the same order of magnitude than IV A-D (Table 2, entries 5-6 versus entries 1-4). As a general trend, a slight decrease of the loading factor was observed for catalysts V G-H (Table 2, entries 7-8). Note that, among PAMAM catalysts, the higher value of the loading factor was obtained in the presence of the longest linker in terminal position (Table 2, entry 6 versus entry 5, and entry 8 versus entry 7).

Examples of the decreased accessibility to coordinative sites into higher generation dendrimer systems have been reported as a consequence of conformational changes, steric hindrance and encapsulation processes.56

#### Epoxidation of olefins with catalysts IV A-D and V E-H

Catalysts IV A-D and V E-H have been evaluated for the epoxidation of a large panel of aliphatic and aromatic olefins, including 1-hexene 1, 1-octene 2, trans-3-octene 3, vinylcyclohexane 4, styrene 5, cis-stilbene 6, and allylbenzene 7. The oxidation was performed by treating the selected olefin (0.05 mmol) with H<sub>2</sub>O<sub>2</sub> (35% water solution, 0.5 mmol) and the

Table 2 ICP-MS analyses of catalysts IV A-D and V E-H

Entry	Catalyst	$\operatorname{Re}^{a}\left(\%\right)$	Loading factor <sup>b</sup>
1	IV A $(n=2)^c$	$29.9 \pm 0.3$	1.6
2	IV B $(n=3)$	$16.4\pm0.3$	0.9
3	IV C $(n=5)$	$30.6\pm0.4$	1.7
4	IV D $(n=6)$	$8.3\pm0.3$	0.5
5	<b>V E</b> $(n_1 = 6, n_2 = 2)$	$22.3\pm0.5$	1.2
6	$\mathbf{V} \mathbf{F} (n_1 = 6, n_2 = 6)$	$26.4\pm0.4$	1.4
7	$V G (n_1 = n_2 = n_3 = 2)$	$3.7\pm0.2$	0.2
8	$V H (n_1 = n_2 = n_3 = 6)$	$\textbf{4.1} \pm \textbf{0.3}$	0.3

a Values are expresses as wt/% with respect to the total weight of the sample. Measurements were repeated in triplicate. b Amount of mg of MTO per gram of support.  $^{c}$  The n value represents the number of carbon atoms in the linker as reported in Scheme 2.

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Scheme 3 Schematic representation for the MTO/ $H_2O_2$  epoxidation of olefins. The reaction proceeds through the formation of monoperoxo [MeRe( $O_2$ ) $O_2$ ] and bis(peroxo) [MeRe( $O_2$ ) $O_2$ ]  $\eta$ 2-metal complexes.<sup>18</sup>

appropriate amount of catalyst (containing 0.0025 mmol of MTO) at 25  $^{\circ}$ C, in the absence of Lewis base additives as resumed in Scheme 3.

The reaction products have been detected by gaschromatography associated to mass-spectrometry (GC-MS) and nuclear magnetic resonance (NMR) analysis after chromatographic purification (SI#3 and SI#4,† respectively). The oxidation with MTO alone was used as reference. Irrespective from the experimental conditions, the oxidation of 1 with MTO and catalysts IV A-D afforded the diol 9 as the only recovered product (Table 3, entries 1-5), IV B being the most efficient catalyst (Table 3, entry 3) The corresponding epoxide was not detected in the reaction mixture, confirming the detrimental effect of the Lewis acidity of MTO in the ring-opening of the oxirane ring, even in the presence of B15C5. As a general trend, the catalytic activity was improved by increasing the number of carbon atoms in the alkyl-diamino spacer, as in the case of IV C-D (Table 3, entries 4 and 5), the lowest reactivity being observed with IV A, characterized by a linker with only two carbon atoms. MWCNTs-B15C5/PAMAM/MTO catalysts V E-H showed a different selectivity in the oxidation of compound 1, the epoxide 8 being obtained as the only recovered product in quantitative yield and high conversion of the substrate (Table 3, entries 6-9). In particular, the second-generation of PAMAM catalysts V G-H was more reactive than the first-generation counterpart V E-F (Table 3, entries 6-7 versus entries 8-9). It is reasonable to suggest that the high selectivity toward the synthesis of the epoxide in the oxidation of 1 with PAMAM catalysts may be associated to the occurrence of additional interactions between MTO and nitrogen atoms in the dendrimer structure, acting as internal Lewis base sites. In accordance with this hypothesis, the vibrational mode at 492 cm<sup>-1</sup>, corresponding to stretching of the Re-N group, was detected for V G, as representative example, by Far Infrared analysis (FIR) (SI#5;† MTO/1,2-cyclohexanediamine complex was used as reference). The ligand properties of tertiary amine atoms in PAMAM dendrimer are reported.<sup>57</sup> For example, octahedral Co(II)-PAMAM dendrimer complexes showed a slight distorted octahedral coordination of the metal atom by the polymeric

matrix.<sup>58,59</sup> Moreover, atomistic molecular dynamics (MD) simulations on ethylenediamine (EDA) PAMAM dendrimer wrapped on MWCNTs (up to 11th generation) showed the high mobility and the low distance occurring between tertiary nitrogen atoms and the terminal amino groups of the structure, where the metal coordinative sites are expected to be located.<sup>58</sup> The oxidation of aliphatic olefins 1-octene 2, *trans*-3-octene 3 and vinyl-cyclohexane 4 proceeded in a similar way. The epoxides 10, 12 and 14 were obtained in quantitative yield in the presence of catalysts V E-H (Table 3, entries 15–18, 24–27, and 33–36, respectively), while diols 11, 13 and 15 were isolated as the main recovered products with MTO alone and catalysts IV A-D (Table 3). In this latter case, only small amounts of epoxides were detected.

The second generation of PAMAM catalysts was slightly more reactive than the first-generation counterpart in the conversion of substrate. Catalysts V E-H efficiently catalyzed the epoxidation of low reactive aromatic olefins, styrene 5, cis-stilbene 6, and allylbenzene 7, affording the corresponding epoxides 16, 18, and 20, as the only recovered products in quantitative yield, from acceptable (52%) to high (89%) conversion of substrate (Table 3, entries 42-45, 51-54, 60-63, respectively). Instead, diols 17, 19, and 21, largely prevailed in the oxidation performed with MTO and catalysts IV A-D. The recyclability of catalysts V F and V H in the oxidation of compound 1 was studied as representative example (Table 4). After the first run, the catalyst was recovered by filtration, washed with CH<sub>2</sub>Cl<sub>2</sub>, and used without any further purification. Catalysts V F and V H showed a slight decrease in the conversion of substrate (Table 4, entries 2-6 and entries 7-12, respectively) yielding the epoxide 8 in quantitative yield for at least six successive reactions. In accordance with these data, a low decrease of the loading factor of the catalyst was observed by ICP-MS analysis of the sample at the end of any run (Table 4).

# **Experimental section**

#### **Materials**

1-Hexene 1, 1-octene 2, trans-3-octene 3, vinyl-cyclohexane 4, styrene 5, cis-stilbene 6, allylbenzene 7, organic solvents, MTO, methyl acrylate, ethylene diamine (EDA), propylene diamine (PDA), cadaverine (CV), hexamethylene diamine (HDA), diisopropylcarbodiimide (DIC), hydroxybenzotriazole (HOBt), N,Ndiisopropyl ethylamine (DIPEA), 4'-carboxy-benzo-15-crown-5, and GH polypro membrane filters were purchased from Sigma-Aldrich-Merck (Germany) and used without further purification. Gas chromatography associated to mass spectrometry (GC-MS) was recorded on a Varian 410 GC-320 MS (Palo Alto, CA, USA) using a VF-5 ms column (30 m, 0.25 mm, 0.25 µm), and an electron beam of 70 eV. All experiments were done in triplicate. Ultrapure HNO3 and HCl obtained from a sub-boiling system (DuoPUR, Milestone, Bergamo, Italy) and ultrapure 18.2 M $\Omega$  water from a Milli-Q (Millipore, Burlington, MA, USA) were used for the sample dissolution. <sup>1</sup>H NMR and <sup>13</sup>C NMR spectra were recorded on a Bruker (400 MHz) spectrometer.

Table 3 Oxidation of olefins 1–6 with catalysts IV A–D and V E– $H^a$ 

Entry	Substrate	Catalyst	Product	Conversion (%)	Yield of epoxide <sup>b</sup> (%)	Yield of $diol^b$ (%)	TON
1		МТО		94	n.d. <sup>c</sup>	>98	
2		IV A	H <sub>3</sub> C	16	n.d.	>98	_
3	CH <sub>2</sub>	IV B	8	84	n.d.	>98	_
Į.	H <sub>3</sub> C	IV C		68	n.d.	>98	_
5	1	IV D	H <sub>3</sub> C OH	58	n.d	>98	
<u>,</u>		V E	9 OH	89	>98	n.d.	178
7		V F	•	91	>98	n.d.	182
3		V G		>98	>98	n.d.	200
)		V H		>98	>98	n.d.	200
.0		MTO		80	n.d	>98	_
.1		IV A	^ ^ 1	12	n.d	>98	_
.2		IV B	H <sub>3</sub> C \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	83	4	96	8
3	LI COCH2	IV C	10	64	11	87	22
4	H <sub>3</sub> C <b>2</b>	IV D	H <sub>3</sub> C OH	58	n.d.	n.d.	_
15		VE	H <sub>3</sub> C OH OH	80	>98	n.d.	160
.6		VF	ОН	77	>98	n.d.	154
.7		V G		>98	>98	n.d.	200
.8		VH		>98	>98	n.d.	200
.9		MTO	211	84	n.d.	>98	0
20		IV A	$H_3C$ $CH_3$	18	3	97	6
21		IV B	12	79	12	85	24
22	CH <sub>3</sub>	IV B	12	69	12	88	24
3	$H_3C$	IV D	ÓН	62	>98	n.d	124
.3 24	•	V E	Λ ↓ Λ .CH₂	85	>98		
			H <sub>3</sub> C Y Y			n.d	170
25		VF	OH 13	89	>98	n.d	178
6		V G		>98	>98	n.d	200
27		VH		>98	>98	n.d	200
8		MTO	$\sim$	88	n.d.	>98	_
9		IV A	( ) 6	16	n.d. <sup>b</sup>	>98	_
0	CH <sub>2</sub>	IV B	<b>14</b> QH	83	14	85	28
1		IV C		64	3	97	6
2	4	IV D	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	58	n.d. <sup>b</sup>	>98	_
3		V E		93	>98	n.d	186
4		VF	15	90	>98	n.d	180
5		V G		>98	>98	n.d	200
6		V H		>98	>98	n.d	200
57		MTO	$\wedge$	70	n.d.	>98	_
8		IV A	[ ] 6	31	$\text{n.d.}^b$	>98	_
9	CH <sub>2</sub>	IV B	16 OH	48	5	90	10
0		IV C	16 OH	36	8	92	16
1	_	IV D	OH	29	11	89	22
2	5	V E		64	>98	n.d	128
3		V F	17	67	>98	n.d	134
4		V G		72	>98	n.d	144
5		V H		74	>98	n.d	148
6		MTO		80	n.d. <sup>b</sup>	>98	
7		IV A	<b>⟨                                    </b>	18	n.d. <sup>b</sup>	>98	_
8		IV B		35	5	95	10
.9		IV C	_ 8	36	7	93	14
0	\_/ \_/	IV D	18	34	4	96	8
1	6	VE		65	>98	n.d	130
2	v	VF		61	>98	n.d	122
3		V G	но́ он <b>19</b>	74	>98	n.d	148
4		V G V H		77	>98	n.d	154
/ <del>-</del> ±		V 11		//	r 30	11.U	134

Table 3 (Contd.)

Entry	Substrate	Catalyst	Product	Conversion (%)	Yield of epoxide <sup>b</sup> (%)	Yield of diol <sup>b</sup> (%)	TON <sup>c</sup>
55		МТО		69	n.d. <sup>b</sup>	>98	_
56		IV A		21	n.d. <sup>b</sup>	>98	_
57	CH <sub>2</sub>	IV B	20 0	40	10	90	20
58		IV C	20	37	7	93	14
59	<u> </u>	IV D		39	9	91	18
60	7	V E	но́ он	52	>98	n.d	104
61		VF	21	48	>98	n.d	96
62		V G		62	>98	n.d	124
63		VН		89	>98	n.d	178

<sup>&</sup>lt;sup>a</sup> The reaction was performed by treating the selected olefin (0.5 mmol) with a stoichiometric amount of  $H_2O_2$  (35%, water solution) in the presence of the appropriate catalyst (0.5 mol/% with respect to loaded MTO) in  $CH_2Cl_2$  (5.0 mL) at 25 °C for 5 h. Experiments were performed in triplicate. As ethylene diamine  $NH_2(CH_2)_2NH_2$ . B: propylene diamine  $NH_2(CH_2)_3NH_2$ . C: cadaverine  $NH_2(CH_2)_5NH_2$ . D: hexamethylene diamine  $NH_2(CH_2)_6NH_2$ . The product yield is calculated with respect to converted substrate. N.d. not determined. <sup>c</sup> Turn over number expressed as the ratio between the mmol of epoxide *versus* the mmol of MTO in the heterogeneous catalyst.

#### Preparation of oxidized MWCNTs I

In a round-bottomed flask, MWCNTs (1.0 g) and sulphonitric mixture  $[H_2SO_4-HNO_3 \ (3:1),\ 500\ mL]$  were kept under magnetic stirring for 4.0 h at 25 °C, sonicated for 1 h and heated for 12 h at 40 °C. The reaction mixture was cooled down to 25 °C and the suspension was washed with cold  $H_2O$  (400 mL). The supernatant was removed by centrifugation (4000  $\times$  g rpm, 30 min). The residue was washed with deionized  $H_2O$  (200 mL) and filtered using GH polypro membrane filters 0.2 ( $\mu$ m). The resulting MWCNTs I were dried under reduced pressure and used without further purification.

#### Preparation of MWCNTs-B15C5/MTO catalysts IV A-D

MWCNTs I (400 mg) were suspended in dimethyl formamide (500 mL) and treated with N,N-diisopropylcarbodiimide (DIC; 250 mg, 2.0 mmol), hydroxybenzotriazole (HOBt; 270 mg, 2.0 mmol), and N,N-diisopropyl ethylamine (DIPEA; 700  $\mu$ L, 4.0 mmol) in a round-bottomed flask with an egg-shaped magnetic stirring bar for 15 min at 25 °C. Thereafter, the appropriate diamine (ethylene diamine for II A, propylene diamine for II B,

cadaverine for II C, and hexamethylene diamine for II D; 2.0 mmol) was added to the solution under gentle stirring for 12 h at 30 °C. The resulting solution was washed with DMF (5  $\times$  5.0 mL), centrifugated (4000  $\times$  g rpm, 20 min), and filtered using GH polypro membrane filters (0.2 µm). The resulting intermediates II A-D were dried under reduced pressure. Successively, 4'-carboxy-benzo-15-crown-5 (1.5 g, 6.0 mmol) was dissolved in DMF (250 mL), followed by treatment with DIC (790 mg, 6.0 mmol) and DIPEA (2.1 mL, 12 mmol), in a 500 mL roundbottomed flask under stirring for 4 h at 25 °C. Thereafter II A-**D** (200 mg) were added to the solution and the mixture stirred for 8 h at 30 °C. The resulting intermediates III A-D were washed with DMF and  $H_2O$ , centrifugated (4000  $\times$  g rpm, 20 min), and filtered using GH polypro membrane filters (0.2 μm). Intermediates III A-D (125 mg) were suspended in THF (50 mL) in a round-bottomed flask, and MTO (25 mg) in THF (25 mL) was added dropwise. After 1 h the suspension was concentrated under reduced pressure, washed with THF (5  $\times$  5.0 mL) and filtered using GH polypro membrane filters (0.2 µm) to afford MTO catalysts IV A-D.

Table 4 Recyclability and stability of catalysts V E and V H in the oxidation of compound 1

Run	Substrate	Catalyst	Conversion (%)	Yield of epoxide <sup>a</sup> (%)	$\operatorname{Re}^{b}\left(\%\right)$
1	1	V F	91	>98	$26.4 \pm 0.4$
2			90	>98	$26.1\pm0.3$
3			90	>98	$26.0 \pm 0.2$
4			85	>98	$24.3\pm0.4$
5			84	>98	$23.9 \pm 0.3$
6			82	>98	$21.6 \pm 0.2$
7		V H	>98	>98	$4.1\pm0.3$
8			>98	>98	$4.0\pm0.2$
9			>98	>98	$4.0\pm0.2$
10			97	>98	$4.0\pm0.3$
11			95	>98	$3.5\pm0.2$
12			92	>98	$3.3\pm0.1$

<sup>&</sup>lt;sup>a</sup> The reactions were performed by treating the olefin 1 (0.5 mmol) with  $H_2O_2$  (0.5 mmol) and the appropriate quantity of MTO catalyst (containing 0.0025 mmol of active rhenium species) calculated with respect to the MTO loading value. After each reaction, the catalyst was recovered by filtration, washed with  $CH_2Cl_2$ , and used without any further purification for the successive run. <sup>b</sup> Values are expresses as wt/% with respect to the total weight of the sample. Measurements were repeated in triplicate.

#### Preparation of MWCNTs-B15C5/PAMAM/MTO catalysts V E-H

The grafting of MWCNTs I with PAMAM was performed by applying a slightly modified procedure from the literature.8 Methyl acrylate (10 mL) was added to a suspension of the appropriate intermediate II D in methanol (10 mL), and the resulting mixture was stirred at 80 °C for 3 days. Thereafter, ethylene diamine (EDA, 0.15 mol, 400 mg), and in alternative, hexamethylene diamine (HAD, 0.15 mol, 775 mg), was added to the suspension in order to prepare the first-generation PAMAM intermediates III E-F, respectively. Two successive treatments of II A and II D with methyl acrylate and the appropriate alkyldiamino linker afforded III G-H, respectively. Next, the appropriate PAMAM intermediate III E-H (100 mg) in DMF (50 mL) was added to a mixture of 4'-carboxybenzo-15-crown-5 (2.0 mmol, 624 mg), DIC (2.0 mmol, 406 mg), and HOBt (2.0 mmol, 270 mg) in dry CH<sub>2</sub>Cl<sub>2</sub> (20 mL). The reaction mixture was stirred for 48 h at room temperature and then filtered on GH polypro membrane filters (0.2 µm) to afford PAMAM intermediates IV E-H, respectively. The black solid was washed several times with DMF (3  $\times$  2.0 mL) and CH<sub>2</sub>Cl<sub>2</sub> (5  $\times$  10 mL) and dried under vacuum. PAMAM intermediates IV E-H (100 mg) were than suspended in THF (25 mL) and a solution of MTO (25 mg) in THF (25 mL) was added dropwise. After 1 h the suspension was concentrated under reduced pressure, washed with THF (5  $\times$  5.0 mL) and filtered using GH polypro membrane filters (0.2 µm) to afford catalysts V E-H.

#### FT-IR analysis

FT-IR analyses were performed, at room temperature, with a PerkinElmer Spectrum One and a BRUKER VERTEX 70V spectrometers, respectively, both equipped with an ATR unit. Mid Infrared spectra were recorded by averaging 32 scans, with a resolution of 4 cm<sup>-1</sup>. In the case of FIR (Far Infrared) configuration, spectra were registered within the (600–50 cm<sup>-1</sup>) interval.

# Scanning electron microscopy (SEM) and transmission electron microscopy (TEM)

For the scanning electron microscopy (SEM), the sample suspensions (50 L) were let to adsorb onto carbon tape attached to aluminum stubs and air dried at 25 °C. The observation was made by a JEOL JSM 6010LA electron microscope (Waltham, MA, USA) using Scanning Electron (SE) and Back Scattered Electrons (BSE) detectors. Energy Dispersive Spectroscopy (EDS) analysis was carried out to reveal the chemical elements. Transmission electron microscopy (TEM) analysis was carried out by JEM-1200 EXII Transmission Electron Microscope. For this aim one drop of nanocomposite was put on copper lace coated with carbon and dried at 25 °C.

#### XRD analysis

The X-ray diffraction data were collected by a Bruker D8 Advance with DaVinci design located at CNIS – La Sapienza University of Rome diffractometer with a molybdenum source. The angle dispersive instrument is equipped with a Mo  $K\alpha$  X-ray

tube ( $\lambda=0.7107$  Å), whose radiation was focused onto the sample with Göbel mirrors. The  $2\theta$  angle range available was  $2.75-142^{\circ}$  with a step of  $0.25^{\circ}$  within Bragg-Brentano *para*focusing geometry. The scattered intensity was collected with the Lynxeye XE Energy-Dispersive 1-D detector.

# Inductively coupled plasma mass-spectrometry (ICP-MS) analysis

The samples were weighed (from 1.0 mg to 10 mg) and transferred in fluorinated ethylene propylene (FEP) vials, previously washed to avoid any kind of external contamination. Regia solution was chosen for the mineralization. In particular, 750  $\mu L$  of HCl and 150  $\mu L$  HNO $_3$  were added and the solution was heated to 80 °C for 3 hours. The volume was adjusted to 5.0 mL and then diluted another 10 times before the ICP-MS analysis. The analysis was performed with ultrahigh vacuum PHI 1257 system and an Agilent 7500 ICP-MS instrument under clean room ISO6 (Santa Clara, CA, USA), respectively.

#### Oxidation of olefins 1-7

A 5.0 mL flask equipped with a magnetic bar was charged with CH<sub>2</sub>Cl<sub>2</sub> (5.0 mL), the olefins 1-7 (0.5 mmol), the appropriate amount of selected MTO catalyst (0.0025 mmol of MTO calculated with respect to the MTO loading value) and H<sub>2</sub>O<sub>2</sub> (35% water solution, 50 µL, 0.5 mmol). The two-phase mixture was stirred vigorously at room temperature for 5 h. The progress of the reaction was monitored at half-hour intervals by gas chromatography associated to mass spectrometry (GC-MS) analysis of small aliquots of the organic phase. The conversion of olefins and yield of corresponding products were determined by GC-MS using n-octane as internal standard and by comparison with commercially available standards. GC-MS was recorded on a Varian 410 GC-320 MS (Palo Alto, CA, USA) using a VF-5 ms column (30 m, 0.25 mm, 0.25 m), and an electron beam of 70 eV. Helium was used as the carrier gas at a constant flow of 1.5 mL min $^{-1}$ . The GC oven temperature was started at 50  $^{\circ}$ C (1 min hold) ramped at 20 °C min<sup>-1</sup> to 115 °C (10 min hold), then ramped 5 °C min<sup>-1</sup> to 280 °C (10 min hold).

#### Conclusions

MTO was efficiently supported on B15C5 and PAMAM functionalized MWCNTs. The novel catalysts showed a different selectivity in the oxidation of olefins depending on the chemical complexity of the system. Epoxides were synthesized in quantitative yield and high conversion of substrate in the presence of PAMAM catalysts, while diols largely prevailed when B15C5 was directly linked to the surface of MWCNTs. XRD analysis showed that the Re atom was coordinated by an octahedral structure through interaction with a couple of oxygen atoms of the B15C5 ring. The high selectivity observed in the synthesis of epoxides with PAMAM catalysts was probably due to the Lewis base property of the nitrogen atoms present in the dendrimer aggregate in buffering the Lewis acidic character of Re(vII) atom. The reactivity of the dendrimer-based catalysts increased by increasing the order of complexity of the structure, the PAMAM

of second-generation affording higher yield of epoxide with respect to the first-generation counterpart. In the PAMAM series, terminal alkyl-diamino spacer bearing carbon atoms (cadaverine) afforded the most reactive catalysts. Note that PAMAM catalysts **V** E-**H** afforded styrene epoxide in yield higher than poly(4-vinylpyridine) and microencapsulated polystyrene MTO based systems,<sup>37</sup> highlighting the beneficial role of the dendrimer structure in the selectivity of the reaction. In the absence of PAMAM, the Lewis acidity of MTO was responsible for the ring-opening of epoxides to diols. In this latter case, the reactivity decreased in the presence of alkyl-diamino spacer bearing more than three carbon atoms (propyl diamine). Finally, catalyst **V H** retained the catalytic activity for at least six

successive runs without any appreciable leaching of the metal.

#### Conflicts of interest

Paper

There are no conflicts to declare.

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