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3D Ruthenium Nanoparticle Covalent Assemblies from Polymantane Ligands for Confined Catalysis

Yuanyuan Min,[†] Houssein Nasrallah[‡] Didier Poinsot,[‡] Pierre Lecante,[§] Yann Tison,[#] Hervé Martinez,[#] Pierre Roblin,[#] Andrea Falqui,^{*} Romuald Poteau,⁴ Iker del Rosal,⁴ Iann C. Gerber,^{*,4} Jean-Cyrille Hierso,^{*,‡} M. Rosa Axet^{*,†} and Philippe Serp^{*,†}.

[†] LCC-CNRS, INPT, 205 route de Narbonne, 31077 Toulouse Cedex 4, France

[‡] Institut de Chimie Moléculaire de l'Université de Bourgogne (ICMUB - UMR CNRS 6302), Université de Bourgogne Franche-Comté (UBFC), 9 avenue Alain Savary, 21078 Dijon Cedex, France

[§] Centre d'élaboration des matériaux et d'études structurales UPR CNRS 8011, 29 Rue Jeanne-Marvig, BP 4347, 31055 Toulouse, France

∥Universite de Pau et des Pays de l'Adour, e2s UPPA, IPREM CNRS UMR 5254, 2 Avenue du président Angot, Pau, 64053, France.

Laboratoire de Génie Chimique and Fédération de Recherche FERMAT, 4 allée Emile Monso, 31030 Toulouse, France

[∗] King Abdullah University of Science and Technology (KAUST), Biological and Environmental Sciences and Engineering (BESE) Division, NABLA Lab, 23955 Thuwal, Saudi Arabia

^{*d} INSA–CNRS–UPS, LPCNO, Université Fédérale de Toulouse Midi-Pyrénées, 135 Avenue de Rangueil, F-31077 Toulouse, France*</sup>

31 ABSTRACT: The synthesis of metal nanoparticle (NP) assemblies stabilized by functional molecules is an important research topic in nanoscience, and the ability to control interparticle distances and positions in NP assemblies is one of the major 32 challenges in designing and understanding functional nanostructures. Here, two series of functionalized adamantanes, bis-33 adamantanes and diamantanes bearing carboxylic acid or amine functional groups have been used as building blocks to 34 produce, via a straightforward method, networks of ruthenium NP. Both the nature of the ligand and the Ru/ligand ratio affect 35 the interparticle distance in the assemblies. The use of 1,3-adamantanedicarboxylic acid allows the synthesis of 3D networks 36 of 1.7-1.9 nm Ru NP presenting interparticle distance of 2.5-2.7 nm. The surface interaction between Ru NP and the ligands 37 were investigated spectroscopically using a ¹³C labeled ligand, as well as theoretically with Density Functional Theory (DFT) 38 calculations. We found that Ru species formed during the NP assembly are able to partially decarbonylate carboxylic acid 39 ligands at room temperature. Decarbonylation of a carboxylic acid at room temperature in the presence of dihydrogen usually 40 occurs on catalysts at much higher temperature and pressure. This result reveals a very high reactivity of ruthenium species 41 formed during network assembling. The Ru NP networks were found active catalysts for the selective hydrogenation of phenyl 42 acetylene, reaching good selectivity towards styrene. Overall, we demonstrated that catalyst activity, selectivity, and NP 43 network stability are significantly affected by Ru NP interparticle distance, and electronic ligand effects. As such, these 44 materials constitute a unique set that should allow a better understanding of the complex surface chemistry in carbon-45 supported metal catalysts. 46

INTRODUCTION

The self-assembly of molecules, clusters or nanostructures is
a promising avenue for the development of organized
nanostructures with peculiar properties and broad spectrum
of emerging applications.¹⁻⁴ In the case of metallic
nanoparticles (NP), a combination of self- and directed

assembly processes involving interparticle forces can be used to produce the desired materials.⁴ These processes generally involve non-covalent interactions between NP,⁴ resulting in assemblies of poor stability, which can be detrimental for some applications. To obtain stable NP assemblies, molecular mediators that can induce a covalent bond between the NP have to be used. Such molecules enable interparticle distance and interaction tuning in the assemblies, allowing studying not only the properties of the individual NP, but also the collective effects of the twodimensional (2D) and three-dimensional (3D) networks of NP.⁵⁻⁷ Networks of gold⁸⁻¹⁰ and ruthenium¹¹ NP have already been described; these networks are assembled by means of di- or polytopic ligands. For instance, thiol-amine¹² and disulfide employed as ligands^{8, 9} lead to 2D assemblies of Au NP, which are organized in a hexagonal array. In general, the interparticle distance can be modulated by changing the length of the alkyl chain of the ligands, and in that context, ligand rigidity should be considered. The use of multi-carboxylate ligands, such as the C₆₆(COOH)₁₂ hexa-adduct, allowed producing 3D networks of Ru NP (1.6 nm) showing an interparticle distance of 2.85 nm.¹¹

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15 For applications in catalysis, such networks of metal NP -16 provided they are sufficiently robust-13 could combine 17 several advantages compared to traditional catalysts, such 18 as: i) a narrow particle size distribution, ii) a controlled 19 interparticle distance, iii) a well-defined chemical 20 environment, and iii) the possibility to take advantage of 21 confinement effects if porous. Whenever built from organic 22 ligands, NP networks constitute ideal model catalysts to 23 study carbon-supported metal catalysts and their intricate 24 surface chemistry.^{14, 15} Indeed, such catalysts generally 25 consist in metal NP deposited on *sp²/sp³* hybridized carbon 26 materials presenting surface functional groups that contain 27 heteroatoms such as oxygen or nitrogen.¹⁵⁻¹⁷ Although the 28 nature of the interaction between surface functional groups 29 and metal NP has been identified as essential for catalysis, since it can affect both metal dispersion and metal-support 30 31 interaction, it is not yet well understood.¹⁷ Thus, in the case of oxygen- and nitrogen-functionalized carbon nanotubes 32 (CNT), it has been shown that: i) Pd NP supported on N-33 CNT showed a narrower size distribution than those on O-34 CNT; and ii) Pd/N-CNT catalyst showed better catalytic 35 performances in selective olefin hydrogenation in terms of 36 activity and selectivity compared to the Pd/O-CNT 37 catalyst.¹⁸ Additionally, in such catalysts, the control of 38 fundamental characteristics for catalysis such as particle 39 size, interparticle distance and metal-support interaction is 40 not achieved. 41

In this context, we have used bi-functional polymantanes 42 (i.e. rigid sp3-hybridized diamondoids) as ligands to build, 43 via a straightforward and up-scalable method, Ru NP 44 networks presenting uniform particle size and interparticle 45 distance, as well as a well-defined chemical environment. 46 Commercially available adamantane and diamantane have 47 mechanically rigid and thermodynamically extremely stable 48 structures,19, 20 which are easily amenable to synthetic 49 modifications,21-23 and have been used with success to 50 assemble various innovative functional materials.²⁴⁻²⁶ Thus, 51 we employed adamantane, bis-adamantane and diamantane 52 as platform for synthesizing dicarboxylic and diamine 53 ligands, which are used for the first time to produce a set of 54 Ru NP networks with well-defined characteristics (NP size, 55

interparticle distance, electronic environment) of interests for catalysis. These materials were investigated for the selective hydrogenation of phenylacetylene into styrene. The nature of the metal-ligand interface that mimics surface oxygen or nitrogen groups is shown to affect catalyst activity, selectivity but also NP network stability. A significant increase in TOF by decreasing interparticle distance is also observed.

MATERIALS AND METHODS

The solvents we used in this work were purified with a MBraun SPS-800 solvent purification system and degassed cycles. [Ru(COD)(COT)] by freezing [(1,5cyclooctadiene)(1,3,5-cyclooctatriene)ruthenium] was purchased from Nanomeps Toulouse. 1-Adamantanecarboxylic acid (AdCOOH, 1) and 1-Adamantylamine (AdNH₂, **5**), phenylacetylene and decane were purchased from Sigma-Aldrich and used as received. The synthesis of the other polymantanes ligands is given in the Supporting Information.

Synthesis of Ru nanoparticles. In a typical experiment [Ru(COD)(COT)] complex and the desired ligand were introduced in a Fisher-Porter bottle, solubilized with THF, and stirred 1h under argon. The resulting solution was pressurized with 3 bar of H₂. The solution, which turned black after few minutes of reaction, was kept under stirring overnight at room temperature. After this period of time, excess of H₂ was eliminated and the volume of solvent was reduced under vacuum. Pentane was then added to the colloidal suspension to precipitate the Ru nanoparticles. After filtration under argon with a cannula, the black solid powder was washed twice with pentane and filtrated again before drying under vacuum. For each ratio studied, the quantities of reactants and the obtained yields are detailed in the Supporting Information.

Synthesis of Ru complexes. The synthesis and characterizations of [Ru(adamantane carboxylate)($\eta^{4-1,5-COD}$)($\eta^{3-1,2,3-COD$)] – **Ru**₁, [Ru₂(adamantane dicarboxylate)($\eta^{4-1,5-COD}$)₂($\eta^{3-1,2,3-COD$)₂] – **Ru**₂, and [Ru₂(μ -adamantane carboxylate)₂(μ - η^{-3} η^{2-} $\eta^{2-1,2,3-COD$)($\eta^{3-1,2,3-COD$)(THF)] – **Ru**₃ is given in the Supporting Information.

Synthesis of Ru NP on functionalized carbon nanotubes. 9.6 mg [Ru(COD)(COT)] dissolved in 20mL THF, was added to 150 mg of dried functionalized carbon nanotubes (O-CNT or N-CNT). After stirred under Ar atmosphere overnight, the resulted mixture was filtered and washed with pentane to obtain a black solid, which was reduced under H₂/Ar gas flow (flow ratio of Ar:H₂= 4:1) at 200 °C for 1h. The rate of increasing temperature is 7 °C/min. The final products were kept in inert atmosphere for further catalysis. ICP analyses have shown a Ru loading of 1.6% w/w for Ru/O-CNT and 0.8% for Ru/N-CNT.

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Catalytic hydrogenation of phenylacetylene. In a typical catalytic reaction, a mixture of phenylacetylene (4 mmol, 412 mg), decane (0.5 mmol, 71 mg) and Ru NP catalyst (Ru atom amount 0.02 mmol) were dispersed in MeOH (25 mL) in a magnetically stirred autoclave with Teflon inner cylinder. The autoclave was purged three times with H_2 . The autoclave was pressurized with 5 bar of H₂ at room temperature and stirred at 1500 rpm. Under such conditions, external mass transfer limitation should not be relevant (see ESI, S1). The suspension was continuously stirred until the 10 end of the reaction. Gas chromatography (GC) was used to 11 identify the products. After catalysis, samples were taken for 12 TEM analyses, also, the % of Ru on the filtered catalytic 13 solution was ascertained by ICP. The TOF were calculated 14 according to the total surface Ru content (see ESI, S1).

15 Characterization. The details of ICP, TEM, NMR, XRD, 16 infrared, WAXS, XPS and SAXS characterization are given in 17 the Supporting Information.

18 Computational details. The DFT calculations were 19 performed using the Vienna *ab initio* simulation package 20 VASP,²⁷⁻³⁰ based on the full-potential projector augmented 21 wave framework,^{31,32} Exchange-correlation effects have been 22 approximated using the spin-polarized version of PBE 23 functional.33 A kinetic-energy cutoff of 400 eV was found to 24 be sufficient to achieve a total-energy convergence within 25 several meV, considering the k-point sampling in Gamma-26 point only calculations for isolated molecules and 27 complexes, in conjunction with a Gaussian smearing with a 28 width of 0.05 eV. During geometry optimization runs, all the 29 atoms were fully relaxed until forces on individual atoms 30 were smaller than 0.01 eV/Å. Calculation cells for isolated 31 molecules and complexes were (25x26x27) Å³, to avoid 32 spurious interactions between periodic images. Figures of 33 the different geometries were produced thanks to the 3D 34 visualization program VESTA.³⁴ Bader charge analyses were 35 performed using Henkelmann's group code.35 The optimal 36 geometries upon H₂ adsorption were constructed following 37 the results of Ref.,³⁶ *i. e.* all available μ_3 sites were occupied 38 and then the top sites and, if needed, some bridge sites were used to build the starting geometries. Reaction barriers were 39 estimated by the climbing image nudge elastic band (CI-40 NEB) method³⁷⁻³⁹ with a spring force between images of 5 41 eV/Å² and a force tolerance of 0.05 eV/Å. The harmonic 42 vibrational modes were systematically calculated to 43 distinguish minima and saddle points using dynamical 44 matrix extraction scheme of VTST tools. 45

RESULTS

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Synthesis and characterization of Ru NP. We have used 49 a set of carboxylic acid and amine ligands bearing the *sp*³-C 50 hybridized platforms adamantane (Ad), bis-adamantane 51 (BAd), and diamantane (DAd) (Scheme 1). 1,3-52 diaminoadamantane ($\underline{6}$), Ad-(NH₂)₂, was prepared from 1,3-53 dibromoadamantane (28% yield) following a reported 54 method.⁴⁰ The related compound BAd-(NH₂)₂ 7 was 55

obtained from BAd, which was itself prepared from 1bromoadamantane homocoupling by metallic sodium.40, 41 From commercial diamantane, DAd-(NH₂)₂ **8** was obtained after selective dibromination of 4,9-C-H apical positions.42 Ad-COOH 1 and Ad-(COOH)2 2 are commercially available, and DAd-(COOH)₂ 4 was prepared based on literature methods and isolated (23% yield) starting from 4,9dihydroxy-diamantane.43, 44 BAd-(COOH)₂ 3, was also synthesized (see SI); however, its insolubility in common organic solvents obviated its use as stabilizer of Ru NP.



Scheme 1. Polymantane ligands based on adamantane, bisadamantane, and diamantane backbone used for the stabilization of discrete Ru NPs and/or networks.

The discrete NP and NP networks reported herein were conveniently obtained by reaction of (1,5cyclooctadiene)(1,3,5-cyclooctatriene)ruthenium(o),

[Ru(COD)(COT)], at room temperature in THF under 3 bar of H₂ in the presence of the corresponding stabilizing polymantane ligand (see Scheme 2 for a representative example with ligands 1 and 2).

Different amounts of ligand were employed for the synthesis of some of the Ru NP, specifically 0.2, 0.1 and 0.05 molar equiv., with the aim to obtain different sizes of nanoparticles and as such a lever to control reactivity. The size of ruthenium NP was determined by Transmission Electron Microscopy (TEM) and correlated with Wide-Angle X-ray Scattering (WAXS). In Ru NP network samples, the interparticle distances were measured by Small Angle X-ray Scattering (SAXS). The results of these analyses, as well as the Ru metal content, are reported on Table 1. Adamantane carboxylic acid 1 has already been used to stabilize FeCo, NP45 and CdSe quantum dots.46

To the best of our knowledge, the adamantane amine **5** has not been used before as ligand for surface stabilization in NP synthesis. The decomposition of [Ru(COD)(COT)] in the presence of 1 or 5 (Ru/ligand molar ratio = 10) leads to the formation of isolated Ru NP with size of 1.8 and 1.6 nm

(TEM), respectively (see Table 1 and Figure 1). Similar NP sizes were reported for Ru NP prepared from [Ru(COD)(COT)] and amines,⁴⁷ and carboxylic acid ligands.48 The coherence length (2.2 nm, Ru NP produced from 1) indicated by the Radial Distribution Function (RDF) resultant from WAXS analyses (Figure S1) is slightly longer than the mean diameter determined by TEM observation. This feature, which was common to many of the samples prepared (see Table 1), was already reported for ligand stabilized metal NP.49 For amine ligands, for which a facile reversible binding leads to dynamic exchanges between free and coordinated species, Ru NP coalescence has been reported.⁵⁰ Consistently, the NP produced from the amine ligand 5 are much more agglomerated than the ones produced from the carboxylic acid ligand 1. Adsorption energy of -60 kcalmol⁻¹ was obtained for 1 on a bare Ru₁₃ cluster, presenting a syn-syn bridging coordination with two Ru–O distances of 2.0 and 2.1 Å after the spontaneous O-H bond dissociation on the cluster, see Figure S2.

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Scheme 2. Synthesis of ruthenium NP and NP networks using polymantane ligands as stabilizers.

 Table 1. Mean size distributions, interparticle distances and ruthenium content of the synthesised Ru NP and Ru NP networks.

Ligand	Ru/L (molar)	Ru loading (%)	Ru NP size (nm) ^a	Ru NP size (nm) ^b	Interparticle distance (nm) ^c
	5/1	77.9	1.5±0.6	1.5	-
1	10/1	65.5	1.8 ± 0.5	2.2	-
	20/1	78.7	1.9 ± 0.6	2.5	-
	5/1	50.2	1.7 ± 0.7	1.7	2.2
2	10/1	61.8	1.8 ± 0.8	2.0	2.7
	20/1	70.1	1.9 ± 0.7	2.2	2.7
4	10/1	67.7	1.8 ± 0.6	2.1	2.9
5	10/1	82.5	1.6 ± 0.7	1.5	-
6	10/1	80.1	1.8 ± 0.7	1.9	2.4
7	10/1	65.7	1.7 ± 0.6	1.2	2.5
8	10/1	70.2	1.7 ± 0.7	2.1	2.7
		CANC .			

^a From TEM. ^b From WAXS. ^c From SAXS.

38 Similar energies have been obtained in a previous work 39 involving the coordination of acetic acid on Ru55.51 A 40 significant charge transfer from the cluster to the ligand (-41 o.84 e⁻, as a result of the oxygen lone pair sigma donation 42 and π^* backdonation), with 0.27 e⁻ given to the hydride) has 43 been estimated by Bader charge analysis. Slightly smaller 44 adsorption energy on a bare Ru₅₅ NP was calculated: -48 45 kcal.mol⁻¹, with very similar charge transfer. In the case of **5**, 46 in agreement with previous results involving the 47 coordination of amines on RuNP,⁵² the adsorption energy is 48 reduced to -30 kcal/mol. This difference can be associated to a top coordination, with a single Ru-N bond-length of 2.2 49 Å. In the same way, a lower charge transfer $(0.12 e^{-})$ from the 50 ligand to the metal is calculated. From these fundamental 51 data, it can be anticipated that these two types of ligands 52 53 (carboxylic acid vs amine) should induce different surface states for Ru NP of similar size, and possibly also different 54

network assembly, and as a consequence, different catalytic performances.

As expected, the use of both the bis-carboxylic acid $(\underline{2}-\underline{4})$ and bis-amino $(\underline{6}-\underline{8})$ derived bi-functional polymantanes induces the formation of networks of Ru NP (Figure 1). In these networks, the mean Ru NP size is relatively homogeneous (1.7-1.9 nm from TEM). Pleasingly, this synthetic method allows for the production of networks of metal NP with a narrow particle size distribution compared to conventional catalytic materials prepared on carbon support. Whatever the ligand used, we did not find any correlation between the amounts of adsorbed ligand (Table S1) and the Ru NP size for the isolated NP or the NP networks.

SAXS analyses were performed to evaluate the interparticle distances in the assemblies. The scattering intensity profiles (Figure 2a for ligand <u>2</u>, and Figure S₃ for the other samples) show a general increase of the scattering intensity towards

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small q values, which is coherent with a system constituted of Ru NP.^{11, 53} At higher values of q ($q_{max} = 0.2-0.3$ Å⁻¹) we observed a peak interpreted as a correlation distance between the metallic NP (center-to-center distance between Ru NP in the superstructure).^{11, 54} It is worth mentioning that SAWS analyses do not allow calculating the distance distribution. These correlation distances were found to vary between 2.2 and 2.9 nm, according to the nature of the ligand, and the Ru/L molar ratio. Slightly larger correlation distances were measured in the case of dicarboxylic ligands 10 compared to diamine ones displaying the same backbone. 11 This is consistent with the dimensions of these stabilizing 12 species. The self-assembly process can be directed to tune 13 the interparticle spacing by using BAd and DAd ligands; and 14 with ligands 4, 7 and 8, larger correlation distances were 15 measured compared to ligands <u>2</u> and <u>6</u>, respectively.

16 For a given ligand, the increase of the Ru/L molar ratio 17 contributes to enhance the spatial distribution of the NP, 18 creating a more homogeneous particle network. Indeed, for 19 ligand 2 the SAXS curves shows at intermediate angles a 20 correlation peak larger for the Ru/L ratio 5/1 (Figure 2a), 21 reflecting a change in distribution of interparticle distance 22 with an increase of the polydispersity.

23 In the case of ligands $\underline{2}$ and $\underline{6}$, DFT calculations have 24 confirmed larger interactions of the carboxylate groups 25 compared to the diamine ones with both Ru₁₃ and Ru₅₅ NP 26 models. With a single ligand 2, Ru-O interactions are strong 27 enough to avoid the coalescence of two Ru₁₃ or Ru₅₅ clusters. 28 For ligands **6**, larger clusters are obtained due to coalescence 29 upon geometry optimizations. See Figure 2b for the 30 coordination mode of ligand <u>2</u> bridging two Ru₅₅ clusters. In 31 this case, the Ru NP-Ru NP distance is 1.8 nm, measured 32 from both NP centers.

33 To confirm the short-range order between the Ru NP, we 34 performed an electron tomography (ET) analysis on a typical 35 aggregate obtained from ligand 2 at a Ru/L molar ratio of 10 36 (Figure 2c). After performing ET on large NP assemblies, it 37 was possible to determine the distribution of their distance 38 (Figure S4) and to reconstruct the 3D shape of some of them, 39 indicating a Ru NP mean size of 1.9 ± 0.3 nm and an 40 interparticle distance of 2.65 nm. We have correlated these 41 results to a higher binding energy (calculated by DFT, see 42 Electronic Supporting Information for computational 43 details) of 1 compared to 5 to the Ru NP. The Ru NP mean 44 size determined by ET for this sample is in agreement with the distances evaluated by TEM and WAXS data (Table 1), 45 and the interparticle distance is consistent with the one 46 47 determined by SAXS analysis (Table 1, 2.7 nm).



Figure 1. TEM micrographs of Ru NP and NP networks obtained at a Ru/L molar ratio of 10 for ligands: a) 1 (scale bar = 50 nm); b) $\underline{2}$ (scale bar = 50 nm); c) $\underline{4}$ (scale bar = 50 nm); d) 5 (scale bar = 100 nm); e) 6 (scale bar = 50 nm); f) 7 (scale bar = 50 nm); g) 8 (scale bar = 50 nm); and h) HRTEM of Ru@2 (scale bar = 20 nm).

We have shown by DFT that the use of ligands 1 and 5 leads to different charge transfer between the ligand and the Ru NP. We have also investigated possible charge transfer in the case of the ligands Ad-(COOH)₂, 2, and Ad-(NH₂)₂, 6. DFT calculations, XPS and IR analyses show a similar tendency: the charge transfer is from the ligand to the metal with the

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amine ligand, and from the metal to the ligand with the carboxylic ligand. For instance, when a single ligand $\underline{2}$ is bridging two Ru₅₅ NP, the charge transfer corresponds to a loss of around o.8 e⁻ for each NP. We used carbon monoxide adsorption on the Ru NP produced from ligands 2 (Ru@2) and $\underline{6}$ (Ru@6) to probe the charge transfer by infrared spectroscopy.

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Prior to CO adsorption, the Ru NP were first outgassed under vacuum to eliminate the CO ligands arising from the decomposition reaction (vide infra). After CO adsorption at room temperature, the v_{CO} stretching frequency were measured at 2050-1940 and 1980-1920 cm⁻¹ for Ru@2 and Ru@6, respectively (see Figure S5 for the corresponding spectra). The use of the donor ligand 6, contributes to increase the electronic density on Ru NP, and consequently to a significant Ru \rightarrow CO π -donation.



40 The increase of the electronic density in the CO π^* causes 41 the C-O bond length to increase and thus the v_{CO} to 42 decrease. In the case of ligand 2, the charge transfer occurs 43 from the metal to the ligand, and consequently the v_{CO} 44 shifted to higher wavenumber. XPS analyses support the 45 charge repartition deduced from DFT/IR. The details of the analyses of Ru@2 and Ru@6 at a Ru/L molar ratio of 10 are 46 47 given in Tables S₂ and S₃ and Figure S6. As far as the Ru (3d)and (3p) core levels are concerned, the data obtained for the 48 Ru $3d_{5/2}$, Ru $3d_{3/2}$ and Ru $3p_{3/2}$ peaks are consistent with 49 metallic Ru.55 A slight difference in the binding energy of the 50 Ru $_{3p_{3/2}}$ peaks between the product obtained from ligands 2 (461.3 eV) and 6 (461.2 eV) is consistent with Ru NP that are 52 more electro-deficient in the case of ligand 2. 53

54 Surface coordination chemistry. The coordination of 55 primary amine⁵⁶ and carboxylic acid ligands⁴⁹ to Ru NP

Figure 2. a) SAXS patterns of Ru NP assembly produced from ligand 2 at different Ru/L molar ratio 5/1, 10/1 and 20/1. The experimental SAXS curves are plotted in Log I(q) fct Log q representation in black line and the fitting curves corresponding to the computed SAXS curves from model are plotted in red line for the Ru/L molar ratio 5/1, in yellow for the molar ratio 10/1 and in green for the molar ratio 20/1. For the plot molar ratio 5/1, the contributions of the different components of the model (surface description, elementary object and distance between particle), are shown in dashed line with the principal corresponding fitting parameters. The sum of these tree contributions gives the red fitting curve. b) Optimized structure of the (Ru₅₅)₂@Ad-(COO)₂ species; and c) 3D manual reconstruction of some Ru NP from the corresponding tomogram. A slice of the latter was used just as a visual background for the NP. It should be noticed that not all the NP imaged by ET are reported, but just few of them, properly chosen in order to make it clearly distinguishable. Their size is calculated to be 1.9 ± 0.3 nm.

prepared by the decomposition of [Ru(COD)(COT)] has been studied: NMR contributed to demonstrate the coordination of octylamine to the surface of Ru NP.57 Herein, the coordination of $\underline{6}$ by the amine has been confirmed experimentally by XPS, IR and SS-NMR. In XPS, the N 1s spectrum of **Ru@6** shows a peak at 398.6 eV (see Table S3 and Figure S6), which corresponds to a surface-attached nitrogen atom transferring electron density to the Ru surface via its lone pair. Consistently, it was also reported that the coordination of primary amines such as hexadecylamine to iron oxide NP gives a binding energy (BE) at 399.7 eV, whereas the corresponding free amine gives a BE value at 397.2 eV.57 Figure S7 shows the IR and SS-NMR spectra of the free ligand 6 and of Ru@6. The IR spectrum of Ru@6 shows bands at 3275-3205 and 2903-2852 cm⁻¹ corresponding to the N-H and C-H stretching bands of

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the ligand, respectively; confirming the presence of amine 1 ligands near the surface of the particles. The ¹³C SS-NMR 2 spectrum of Ru@6 shows broad bands at 30-50 ppm 3 characteristic of the sp^3 carbons of ligand <u>6</u>. In the case of 4 acetic acid ligands, spectroscopic data and DFT calculations 5 have shown that the acetic acid lies on the metal surface as 6 acetate, together with hydrides.⁴⁹ For the adamantane 7 ligand 2, DFT calculations have shown that the most stable 8 structure on a Ru₅₅ cluster involves the coordination of the 9 two carboxylate moieties on a large facet of the cluster after 10 spontaneous O-H bond activation, with an adsorption 11 energy of -50 and -63 kcal.mol-1 for the mono- and bi-12 anchoring coordination mode of the carboxylate group, 13 respectively, see Figure S8 for the latter geometry. However, 14 we assume that a mono-anchoring mode is favored 15 thermodynamically since in that case, more ligands can be 16 adsorbed on the NP surface. For ligands 6, only the mono-17 anchoring mode is allowed due to the strong repulsion 18 between surface Ru and the closest hydrogen atoms lying 19 between the two amine groups.

20 Surface modeling from the analysis of coordination 21 complexes. In order to obtain better insight on the 22 coordination mode of carboxylic acid ligands onto the Ru 23 NP surface we achieved the reaction of [Ru(COD)(COT)] 24 with 1 and 2. Reports concerning the reactivity of 25 [Ru(COD)(COT)] with carboxylic acids have previously 26 shown that reaction with 9-anthracenecarboxylic gives the 27 complex [Ru(9-anthracenecarboxylate)₂(COD)],⁵⁸ and with 28 acetic acid gives the Ru complex [Ru(CH₃COO)(η⁴-1,4-29 COD)(η^{5-1} , 5-COT)].⁵⁹ [Ru(COD)(COT)] reacts smoothly 30 with 1 at room temperature to give $[Ru(\kappa-AdCOO)(\eta^{4}-$ 31 C_8H_{12} (η^3 - C_8H_{11})] **Ru**₁, isolated as a pale yellow solid in 64% yield and characterized by NMR, IR and X-ray 32 crystallography (see S.1 in the SI). 33

34 Ru, results from the coordination of the carboxylic acid (see 35 Scheme 3), through oxidative addition of the O-H bond 36 inducing the reduction of cyclooctatriene to form an η³-allyl 37 system. Likewise, the coordination of ligand 2 with 38 [Ru(COD)(COT)] leads to the O-H bond cleavage of the 39 ligand to produce a carboxylate Ru complex. In this case, the presence of two carboxylic acids on the ligand allowed 40 synthesizing a bimetallic complex in which 2 in the 41 carboxylate form acts as a bridge. Ru₂ was isolated as a 42 yellow powder in 52% yield and characterized by NMR, IR, 43 and elemental analyses (see S.1 in the SI). Similar reactivity, 44 i.e. oxidative addition followed by cyclooctatriene reduction, 45 has been reported with Si-H bonds, in which the reduced 46 cyclooctratriene is coordinated to the Ru center trough the 47 Surface coordination chemistry of carboxylic ligands. 48 In the case of the use of carboxylic stabilizers for NP 49 synthesis, different coordination modes on metallic surface 50 have been reported: i) monodentate; ii) chelating bidentate; 51 and iii) bridging bidentate.⁶⁴ As far as Ru NP stabilized by 52 carboxylate ligands are concerned, Poteau et al. reported a 53 bridging bidentate coordination mode ($\Delta v = 145 \text{ cm}^{-1}$),⁴⁹ in 54 accordance with data reported for molecular (or polymeric) 55

three carbons in the allyl form and the two carbons of the double bond.60, 61 Upon heating these carboxylate Ru complexes in the presence of ligands, the unsaturated ligands of the Ru center are generally removed. We experimented a similar procedure, *i.e.* heating **Ru**, at 55 °C in THF in an attempt to reproduce the formation of Ru NP by loss of the unsaturated ligands. A mixture of 1, Ru, and other species was obtained. Crystals grown from a saturated solution of the mixture in THF, which correspond to a bimetallic complex Ru₂. Single-crystal XRD analysis showed that Ru₃ contains two ligand 1 coordinated, chelating both metallic centers with a THF molecule attached to one Ru atom (see S.1 in the SI). This finding is in line with the formation of Ru NP by the coordination of ligand 1 to the Ru NP surface. In the two isolated Ru complexes bearing ligands 1 and 2, the carboxylic acids are easily deprotonated and the carboxylate ligands are chelating bidentate. Complexes **Ru**, and **Ru**, show in their IR spectra the typical $v_{as}(COO^{-})$ and $v_{s}(COO^{-})$ bands of the carboxylate ligand at 1497/1443 and 1502/1430 cm⁻¹, respectively. In IR, the difference in $v_{as}(COO^{-})$ and $v_{s}(COO^{-})$ (Δv), is generally used to determine the coordination mode of the carboxylate group. In that case, the Δv are 54 and 72 cm⁻¹ for **Ru**, and \mathbf{Ru}_{2} , respectively, which is in accordance with a chelating bidentate coordination ($\Delta v < 110$ cm⁻¹).⁶² The ¹³C NMR spectra of these complexes show singlets at 190.3 and 189.5 ppm for Ru1 and Ru2, respectively, attributed to the chelating bidentate carboxylates, which are consistent with chemical shifts reported in the literature for Ru complexes bearing chelating bidentate acetate ligands.63



Scheme 3. Structure of complexes \mathbf{Ru}_{1-3} isolated from reaction of [Ru(COD)(COT)] with ligands $\underline{1}$ and $\underline{2}$.

Ru species bearing such ligands.⁶⁵⁻⁶⁸ In Poteau's work, the ¹³C NMR spectrum of the NP show a singlet at 186 ppm attributed to the acetate. Interestingly, a low intensity band at 1950 cm⁻¹ is also observed, attributed to a v_{CO} band; the CO ligand presumably arising from THF (the reaction solvent) decomposition. XPS, IR, and SS-NMR analyses were performed on the final compound **Ru@2** prepared from ligand <u>2</u>. The O1s XPS spectrum of **Ru@2** (Ru/ligand ratio =

10, Figure S6) shows a main peak at 532.2 eV, which can be assigned to deprotonated carboxylic acids.⁶⁹

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2 The IR spectra of **Ru@2** materials prepared at different Ru/L 3 4 ratio are shown in Figure S9 and compared to the one of the free ligand (Figure 3a shows the spectrum obtained for a 5 Ru/ligand ratio of 10). First, it is clear that like in the 6 coordination complexes, the ligand 2 has been deprotonated 7 since no carboxylic acid (-C=OOH) stretching vibration 8 bands are visible at 1710 cm⁻¹. Whatever the Ru/L ratio, 9 several bands are visible in the 1600-1300 cm⁻¹ region. First, 10 as in the case of Ru NP stabilized by acetate ligands,49 we 11 identified bands at 1560/1390 cm⁻¹ ($\Delta v = 170$ cm⁻¹), which are 12 consistent with bridging bidentate carboxylate ligands. DFT 13 calculations performed on a Ru₁₃H₁₈@(Ad-COOH-COO⁻) 14 model predicted bands at 1473 and 1340 cm⁻¹ ($\Delta v = 133$ cm⁻¹). 15 The frequency of the $v_{as}(COO^{-})$ mode is underestimated in 16 our calculation, similarly to reported elsewhere.49 Other 17 bands are visible at 1450, 1361, 1337 and 1312 cm⁻¹. The band at 18 1450 cm⁻¹ is characteristic of CH, scissoring. These bands are 19 not consistent with the presence of monodentate 20 carboxylate species, since monodentate species should give 21 a Δv > 200 cm⁻¹. DFT calculations performed on a 22 Ru₁₃H₁₈@Ad(COOH)₂ model predicted bands at 1590 and 23 1309 cm⁻¹ ($\Delta v = 281$ cm⁻¹). Additionally, monodentate 24 carboxylate Ru complexes give bands for the $v_{as}(COO^{-})$ 25 mode at around 1600 cm^{-1.64} Although it is difficult at that 26 point to safely assign these bands, we can propose the 27 presence of chelate bidentate species. Beside to the bands 28 attributed to carboxylate species, a band at 1940-1960 cm⁻¹ 29 (according to the Ru/ligand ratio) is also observed, as in the 30 work of Poteau et al.49 In our case, however the intensity of this v_{CO} band is very high, and depends on the Ru/ligand 31 ratio. At a Ru/ligand ratio of 5, two bands are visible at 2040 32 and 1942 cm⁻¹; for a ratio of 10 a single broad band at 1962 33 cm⁻¹ is observed (Figure 3b), and at a ratio of 20 the band at 34 1957 cm⁻¹ is very broad with a shoulder at 1800 cm⁻¹ (Figure 35 S9), which could correspond to bridged CO.68 The shift of 36 the main band as well as the appearance of new bands could 37 be due to an influence of the CO coverage.70-73 The origin of 38 CO in the medium will be discussed in the next section. The 39 ¹³C SS-NMR spectrum of Ru@2 (Figure 3c) has been 40 recorded on a compound prepared with ¹³C labeled 2 (Ad-41 (¹³COOH)₂) in order to obtain clear signals of the *sp*² carbons 42 of carbonyl groups. Beside the peak at 27.1 ppm 43 corresponding to the *sp*³ carbons of the adamantane cage, 44 carbonyl species are detected at 178.3, 193.6 and 220.1 ppm, 45 which have been attributed to the bridging bidentate 46 carboxylate ligands (186.0 ppm in the work of Poteau et al.49 47 and 196.5 ppm in \mathbf{Ru}_3), whereas the peak at 178.3 ppm could 48 be tentatively attributed to chelate bidentate species (a peak 49 at 185.2 ppm was also observed in the mixture of Ru, 50 synthesis). ¹³C-NMR studies on Ru complexes bearing 51 chelate bidentate carboxylate ligands have reported 52 We recorded the infrared spectra for which a shift should be 53 observed in the v_{CO} band if the ¹³CO comes from the ligand. 54 Figure S11 show the spectra obtained with unlabeled and 55

chemical shifts (182-186 ppm), which are consistent with this attribution.⁶⁴

Room temperature carboxylic acid decarbonylation induced by reactive Ru species. The presence of CO ligands on the Ru NP was rather surprising regarding the intensity of the CO band and the origin of their formation. Chaudret *et al.*^{74, 75} and Poteau *et al.*⁴⁹ have also observed the presence of CO ligands on Ru NP that were formed in THF solvent from [Ru(COD)(COT)]. They have attributed this CO formation to the decomposition of THF into propane and CO. Herein, CO could arise either from THF decomposition or from the decarbonylation of <u>2</u>. To determine whether the CO arises from ligand <u>2</u> or THF, we performed the reaction with labeled <u>2</u> using Ad-(¹³COOH)₂, with a Ru/ligand ratio = 10.



Figure 3. a) IR spectra of <u>2</u> and **Ru@2** (Ru/ligand ratio = 10), and b) ¹³C SS-NMR spectrum of **Ru@2** (Ru/ligand ratio = 10) obtained with a ¹³C labeled 2 (Ad-(¹³COOH),).

labeled <u>2</u>. The CO band for the unlabeled <u>2</u> is observed at 1962 cm⁻¹, whereas for the labeled <u>2</u> the v_{CO} band is found at 1934 cm⁻¹. When ${}^{12}C{}^{16}O$ is replaced by ${}^{13}C{}^{16}O$ (R = 0.978), the

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differences between observed and calculated frequencies are 1 generally low. In our case, the calculated frequency should 2 be at 1918 cm⁻¹, so the difference (16 cm⁻¹) is relatively large. 3 This could be because, in addition to ligand 2 4 decarbonylation, we may also observe some THF 5 decomposition. The large envelops at 1934 cm⁻¹ could 6 encompass two bands (Ru(13CO)(12CO)); the Ru(13CO) 7 arising from 2 decarbonylation and the Ru(12CO) from THF 8 decomposition. The fact that THF decomposition also 9 occurs during the reaction of [Ru(COD)(COT)] with H₂ and 10 polymantane ligands is supported by a v_{CO} band at 1914-1923 11 cm⁻¹ that was also observed in the case of the diamine ligands 12 5, 6, 7 and 8 (Figure S12). Decarbonylation of a carboxylic 13 acid at room temperature in the presence of dihydrogen is 14 very surprising since this reaction usually occurs on catalysts 15 at much higher temperature. Accordingly, the 16 decarbonylation of propanoic acid in the presence of H₂ on 17 Ru catalyst has been observed above 180 °C,^{76, 77} in processes 18 rationalized by the energy barrier necessary to pass for C-C 19 bond breaking, as an essential step to produce CO.77 This 20 decarbonylation, operated at room temperature, suggests an 21 unanticipated high reactivity of under-coordinated Ru 22 species that are formed during the decomposition of 23 [Ru(COD)(COT)] in the presence of H₂ and <u>2</u>. A DFT study was performed on a $Ru_{12}H_{16}$ model, as representative species 24 of a small-size reactive intermediate under realistic 25 conditions, where the ligand 2 was replaced by CH₃COOH 26 for simplicity. Several reaction pathways were considered 27 (Figure 4), and the lowest energy routes are discussed below. 28 29 The reference initial state is given by those two species 30 infinitely separated. The reaction begins by the 31 thermodynamically favorable coordination of CH₃COOH (-23 kcal.mol⁻¹). We show below that in an energetically 32 favorable process CO@Ru13H16 may form with the assistance 33 of an extra H_2 , and the release of one H_2O and one CH_4 (-46 34 kcal.mol⁻¹). This pathway is reasonable in terms of kinetics 35 and thermodynamics. In this case, the reaction occurs first 36 through the facile (10 kcal.mol⁻¹) C-OH bond dissociation, 37 leading to the formation of CH₃CO* and OH* on the NP. A 38 C–C bond breaking with a lower barrier height (5 kcal.mol⁻¹) 39 follows this step. Yet, this (CH₃)*(CO)*(OH)* intermediate 40 is slightly less stable than the previous one (-35 vs. -43 41 kcal.mol⁻¹). Owing to the experimental conditions, under H₂ 42 pressure, one H₂ can easily react on the surface, and act as a 43 thermodynamic driving force leading to the release of CH₄ 44 and H₂O yielding the final adsorbed CO product (-46 45 kcal.mol⁻¹). Another plausible pathway is a direct C-C 46 dissociation, assuming that mobile surface hydrides allow 47 for the clearance of a facet composed by three Ru atoms. The 48 corresponding energy barrier is 29 kcal.mol⁻¹, when the 49 (CH₃)*(COOH)*@Ru₁₃H₁₆ final state remaining high in 50 energy (-1 kcal.mol⁻¹). A second possible pathway is a further 51 dissociation of the carboxylic group (almost barrierless) that 52 yields a very deep well in the potential energy landscape 53 ((CH₃COO)*(H)*: -52 kcal.mol⁻¹). This value is slightly 54 lowered compared to the one reported above for 1@Ru₁₂. 55 This is mainly due to the surface saturation by hydrides. This 56

intermediate corresponds to a thermodynamic trap since neither a further C-C nor C-O dissociation could provide a more stable state. The question thus arising is which principle reaction favor the CO formation herein. Some reported semi-quantitative clues based on previous phase diagrams calculations are to be considered.49, 78 These diagrams show that under realistic conditions close to the ones we used, a Ru₅₅ NP in the presence of acetic acid and pressure can respectively accommodate CO 16 $(CH_2COO)^*(H)^*$ and up to 63 $(CO)^*$. Owing to the average adsorption energy per ligand (-35 kcal.mol⁻¹ per (CH₃COO)*(H)* and -40 kcal.mol⁻¹ per (CO)*), the final decomposition of a large amount of acids into CO can be considered as the strongest driving force. It is however worth noting that to turn this unexpected reactivity into a catalytic reaction at low temperature, the system has to face the very strong CO adsorption at the Ru NP surface, which may block the catalytic activity of the Ru centers.

Overall, the results from structural characterization and surface coordination chemistry studies demonstrate, for the first time, that bifunctional rigid polymantane ligands with well-defined functionalization (functions and their position) promote the synthesis of unprecedented Ru NP networks in which are well controlled both Ru NP size uniformity and interparticle distances. Because of the relatively low cost and good availability of Ru among precious metals, we investigated the Ru NP network catalytic performances for an atom-economic hydrogenation reaction, with the additional aims of providing high catalytic selectivity and evidencing structure/performance relationship.

Ru NP network activity in catalytic hydrogenation of **phenylacetylene.** The selective hydrogenation of alkynes to alkenes is a particularly valuable reaction in synthetic organic chemistry. The hydrogenation of phenylacetylene in liquid phase is a benchmark method for assessing catalytic activity and selectivity of Pd NP.79 Although there is a need for selective hydrogenation catalysts that are less expensive than conventional Pd-based catalysts, Ru NP have been little studied.⁸⁰⁻⁸⁷ For this reaction, the catalytic properties of the transition metal strongly depend on its electronic structure. The catalytic behavior of Ru NP prepared with the different polymantane ligands was studied in the hydrogenation of phenylacetylene into styrene at room temperature and a constant H₂ pressure of 5 bar (Table 2, entries 1-11). The evolution of activity and selectivity over time (Figure S13) shows that for most of the catalysts, at the exception of the less active **Ru@4**, the styrene selectivity decreases abruptly at high conversions due to the formation of ethylbenzene and ethyl cyclohexane, resulting from the complete hydrogenation of the phenyl ring. Product distribution over time is shown on Figure S14 for Ru@1, Ru@2, Ru@5 and **Ru@6** (Ru/ligand = 10). The hydrogenation of arenes under mild reaction conditions using (supported or colloidal) Ru NP as catalysts has already been reported,⁸⁸⁻⁹¹ and is not surprising, even at room temperature.

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Figure 4. DFT investigation of decarbonylation's reaction mechanism. Blue values correspond to total energy differences with respect to the initial state, when red ones give energy barrier height.

Figure 5a shows the calculated TOF according to the nature 33 of the ligand. For each series of ligands (acids and amines), 34 the use of discrete Ru NP (from 1 and 5, adamantane 35 monoacid and monoamine, respectively) provides higher 36 TOF than the use of Ru NP networks (from 2 and 6, 37 adamantanes, and 4 and 8, diamantanes, diacids and 38 diamines, respectively). We independently checked that this 39 result cannot be rationalized by the amount of adsorbed 40 polymantane ligands (Table S1) on the NP surface. 41 Considering that diffusion limitations should not be limiting 42 in our case since catalytic activity is low (in the h⁻¹ range) 43 and the catalytic particles are small (in the sub-micrometric 44 range), this result can rely on an important difference 45 between these materials, which is the presence of significant 46 amount of adsorbed CO on the Ru NP networks. This 47 adsorbed CO poisons the Ru NP in the network, which are 48 finally less active that the discrete NP. Additional 49 experiments were performed to validate this hypothesis. We 50 tested as catalyst a Ru NP network (Ru*@2 - 10/1) that have 51 been vacuum treated to remove the adsorbed CO, and 52 compared the TOF at 100% conversion of Ru@1 - 10/1, Ru@2 53 10/1 and **Ru***@2 - 10/1. The TOF obtained were respectively 54 113.8, 101.9, and 117.4 h⁻¹. From Figure 5a, it is discernible that 55

the use of donating amino ligands (5-8) allows reaching higher TOF than with the carboxylic ligands. The fact that amine ligands, which insure a higher electronic density on Ru, allow reaching higher TOF for phenylacetylene hydrogenation is in accordance with results reported in the literature. Thus, it was reported that for the hydrogenation of various unsaturated compounds on stabilized Ru NPs, the use of σ -donor ligands increases the activity of the Ru NP.^{92,} 93 Similarly, Ru/N-doped carbon catalysts were reported to be very efficient catalysts for arene hydrogenation.94

(COO)*@Ru₁₃H₁₆

-24 kcal.mol⁻¹

(CO)*(O)*@Ru13H16

- H₂O

-47 kcal.mol⁻¹

(CO)*@Ru13H16

-46 kcal.mol⁻¹

We assume that the amine as σ -donor ligand increases the electron density at the NP surface, favoring thus the π -back donation from the *d* Ru orbitals to the antibonding π^* orbital of the alkyne/alkene, which facilitates rapid hydrogenation. Accordingly, a slightly higher selectivity towards styrene was observed when using carboxylic acid ligands (Figure 5b). Indeed, in that case, the charge transfer from the metal to the ligand results in electro-deficient Ru NP that provide less π -back donation to the alkene, which weakens Rualkene bonding and favors its quick desorption, thus avoiding overhydrogenation to produce ethylbenzene.



Figure 5. a) Activity, and b) styrene selectivity at 20% conversion for catalysts **Ru@1**, **Ru@2**, **Ru@4**, **Ru@5**, **Ru@6** and **Ru@8** (Ru/ligand = 10/1). Red horizontal lines = carboxylic ligands; and blue vertical lines = amine ligands.



Figure 6. a) Activity (TOF calculated according to surface Ru content) as a function of the interparticle distance, and b) styrene selectivity at 20% conversion as a function of the v_{CO} frequency of CO adsorbed on the catalysts (blue spheres amine ligands, and red squares acid ligands). The Ru/ligand ratio in these catalysts is 10/1.

 Table 2. Hydrogenation of phenylacetylene catalyzed by Ru catalysts.^{a)}

К н-	[Ru] H ₂ , 5 bar r.t., MeOH	+	
Phenylacetylene	Styrene	Ethylbenzene	Ethylcyclohexane

	Phenyl	acetylene	Styrene	Ethylbenzene	Ethylcyclohexane	
Entry	Catalysts	Time (h)	Conv. (%)	S20% (%)	S _{60%} (%)	TOF $(h^{-1})^a$
1	Ru@1 - 5/1	2	84.4	62.3	58.3	61.6
2	Ru@1 - 10/1	3	73.1	64.5	61.5	112.9
3	Ru@1 - 20/1	4	73.2	63.5	61.7	45.6
4	Ru@2 - 5/1	4	66.7	68.9	66.7	30.3
5	Ru@2 - 10/1	3	71.6	66.7	63.6	64.9
5	Ru@2 - 20/1	3	79.1	66.7	63.6	74.2
7	Ru@4 - 10/1	5	62.1	71.3	65.4	18.9
3	Ru@5 - 10/1	2	72.7	64.3	59.6	98.7
9	Ru@6 - 10/1	2	65.9	62.5	58.9	101.4
10	Ru@7 - 10/1	3	72.6	65.2	64.9	66.6
1	Ru@8 - 10/1	3	66.8	60.5	56.3	74.9
12	1.6%Ru/O-CNT	8	10.6	67.8^{b}	-	-
13	o.8%Ru/N-CNT	4	9.0	67.4^{b}	-	-

Reaction conditions: 0.02 mmol Ru, 412 mg (4.00 mmol) phenylacetylene, 71 mg (0.50 mmol) decane (internal standard), room temperature, 5 bar H_2 , 25 mL MeOH. a TOF calculated according to surface Ru content at 1 h reaction. b Selectivity given at 10% of conversion.

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In a series of ligand (acid or amines) providing Ru NP network, some differences can also be noted. The use of bulkier ligands such as BAd and DAd is detrimental to the activity. We also observed a positive confinement effect in relation with shorter interparticle distances. Notably, the interparticle distance is not affected by changes in Ru NP size since all the catalysts present very similar size (1.7-1.9 nm). Whatever the type of ligand a correlation between the interparticle distance and the catalyst activity is indicated by the analysis of Figure 6a: the closer the particles the higher the activity. The effects of confinement in carbon materials have been identified as possible levers to modify catalytic performances.95 It has been shown that soft confinement effects, such as reactant enrichment or rapid diffusion, will directly affect the kinetics of the reaction. Hard effects, like the charge transfer between the metal and the support, will also affect activity/selectivity and stability. In our case, the lower is the interparticle distance, thus giving more confined Ru NP, higher is the activity. A correlation can also be drawn between the interparticle distance and selectivity (Figure 6b) considering both amine and acid as ligands, which is reverse to the correlation observed between interparticle distance and activity. Larger distance induces a decrease of the activity; in that case styrene could be desorbed before further hydrogenation, leading to higher selectivity. Considering that reactant concentration enrichment (as well as high diffusion rates) have been regularly reported as explanation of "soft" confinement effects, we preferred to rely on this explanation. It should nevertheless be noted that the search for a detailed explanation of the observed confinement effects would require a study in itself, which falls outside the scope of our study. Additionally, in acid or amines series no evident correlation was found between the NP size and their activity. Considering the surface electronic density, which was probed via the v_{CO} frequency of CO adsorbed on the catalysts after their synthesis and theoretically, we find a correlation for the acid series as far as activity and selectivity a concerned (Figure S15). As expected, we can noticed an increase of the selectivity and a decrease of the activity when the electronic density on the Ru NP is decreased (higher v_{CO} frequency). For the σ -donor amine ligands, which provide a higher electronic density on the Ru NP, the selectivity is lower. However, differences of v_{CO} shifts between the amine ligands are too close to allow drawing any general correlation (Figure S15). It is nevertheless important to note that in this series appears the ligand z, which presents a more flexible backbone than the other ligands, which could also explain this lack of correlation. Indeed, if this ligand is disregarded, the same type of tendency as for the ligands of the acid series is observed.

We also compared the stability of catalysts prepared from amine and carboxylic ligands. The Figure S16 shows TEM micrographs of the fresh and spent catalysts. It is clear from this figure that the networks prepared from acid ligands provide more robust catalysts, as expected from the DFT calculations (Figure S₂). While an influence of the solvent for adsorption (and ligand displacement) or swelling of the networks in the course of catalytic reactions could not be completely ruled out, Ru NP networks were not found to be significantly modified after catalysis suggesting a rather robust/rigid framework.

Finally, to better delineate the performances of the Ru NP and NP networks prepared from the functionalized polymantane ligands we compared their reactivity with the ones of individual unprotected Ru NP deposited on functionalized carbon nanotubes (CNT). Two types of CNT were used, CNT functionalized with nitric acid (O-CNT) that contains significant amounts of carboxylic groups to stabilize Ru NP,47 and CNT prepared from acetonitrile, which contains different types of nitrogen groups (N-CNT), such as pyridinic or amines.96 Both catalysts present similar Ru NP size (1-2 nm) and show much lower conversions (Table 2, entries 12 and 13) than the NP networks. However, the same tendency as the one observed for the Ru NP networks was observed, i.e. a higher activity is obtained with the catalyst containing nitrogen functionalities. As for the Ru NP networks at low conversion (Figure S14), these two Ru/CNT catalysts show similar selectivity. The higher activity obtained with Ru NP networks nicely illustrate the interest provided from rigid polymantane ligands in Rucatalyzed hydrogenation. These also hold promises concerning the use of NP networks in relation with inedited confinement effects in which short-range interactions between NP may greatly improve their catalytic activity.

CONCLUSIONS

We have successfully prepared catalytic materials based on well-defined small nanometric-sized Ru NP (< 2 nm) stabilized by mono or difunctionalized adamantanes, bisadamantanes and diamantanes, bearing either carboxylic acid or amine functional groups. Unprecedented assembling of networks based on polymantanes are described. We have shown that the interparticle distances can be tuned by a proper choice of the ligand and the Ru/ligand ratio. We have also shown that the choice of the ligand is crucial to tune the electronic properties of the metal. In a systematic approach from molecular regime to NP and networks, the coordination chemistry of these species has been investigated, and it was shown that highly reactive Ru clusters or colloidal species are formed during NP network synthesis, which were experimentally shown to be able to decarbonylate carboxylic acids at room temperature in the presence of 3 bar of H₂. DFT theoretical investigation, which supported this high reactivity with small clusters, also allowed proposing a preferential low energy pathway for this innovative decarbonylation reaction.

These new nanoparticle networks constitute a unique set to investigate structural and electronic effects in

heterogeneous catalysis with flexible assembly of nanoobjects, especially confinement effects were revealed that exalt catalytic activity. In the hydrogenation of phenyl acetylene we evidenced that Ru NP interparticle distance and electronic effects control the catalyst activity; whereas electronic effects mainly govern the catalyst selectivity.

ASSOCIATED CONTENT

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Supporting Information. This material is available free of charge via the internet at <u>http://pubs.acs.org</u>.

It contains, materials synthesis and characterization by ICP, XPS, IR, SAXS, WAXS, optimized geometry from DFT, and data related to phenyl acetylene hydrogenation.

AUTHOR INFORMATION

Corresponding Author

*E-mail: Philippe.serp@lcc-toulouse.fr.

Author Contributions

19 Y. M. synthesized and participated to the characterization of 20 the 3D ruthenium nanoparticle covalent assemblies and 21 performed the catalytic studies. H.N. designed the global 22 synthetic pathways of polymantane ligands and achieved 23 most of the late-stage functionalization for polymantane 24 ligands. D.P. synthesized and characterized the precursors for 25 functionalized polymantane ligands.P.L. performed the 26 WAXS analyses and interpreted the results. Y. T. and H. M. 27 performed the XPS analyses and interpreted the results. P. R. 28 performed the SAXS analyses and interpreted the results. A.F. performed the electron tomography characterization. R. P., I. 29 del R. and I. Gerber performed the DFT calculations. J.C. H. 30 supervised the work on polymantane ligands synthesis and 31 contributed to catalysis discussion. M. R. A. participated to 32 the supervision of the work on 3D ruthenium nanoparticle 33 covalent assemblies and catalytic studies. P. S. conceived the 34 idea of 3D ruthenium nanoparticle covalent assembly 35 synthesis for catalytic applications and coordinated the entire 36 work. 37

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Notes

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ABBREVIATIONS

CCR2, CC chemokine receptor 2; CCL2, CC chemokine ligand 2; CCR5, CC chemokine receptor 5; TLC, thin layer chromatography.

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