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An overall one-pot strategy for the planning of mono and bimetallic nanoparticles upheld on carbon nanotubes.

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Abstract

The disintegration of organometallic metal buildings within the sight of stoichiometric measures of ligands have devil strated to be a compelling strategy for the readiness of all around characterized metal nanoparticles under gentle conditions.1 Recently, another system for incorporating little and very much characterized NHC-settled NiNPs has been developed.2 This approach per-mitted the arrangement of colloidal nanoparticles as well as the immediate immobilization on carbon nanotubes by a basic "one-pot" technique. The upheld Ni NPs/CNTs uncovered to be productive impetuses in the particular hydrogenation of inward alkynes into the comparing (Z)- alkenes. In the current con-tribution, the "one-pot" strategy was reached out for the planning of monometallic (Cu and Pd) and bimetallic nanoimpetuses (NiCu and PdCu) settled by a N-heterocyclic carbene ligand. Both colloidal and upheld nanoparticles (NPs) on carbon nanotubes (CNTs) showed welldefined control on their size, morphology and organization and were assessed in the particular hydrogenation of alkynes and alkynols. PdCu/CNTs uncovered to be a productive reactant framework giving exceptionally se-lectivity in the hydrogenations of terminal and inward alkynes. Besides, this impetus was tried in the semihydrogenation of acetylene in modernly significant acetylene/ethylene-rich model gas takes care of showing phenomenal soundness even after 40 h of response.

A facile and straightforward methodology for the preparation of monometallic (copper and palladium) and bimetallic nanocatalysts (NiCu and PdCu) stabilized by a N-heterocyclic carbene ligand is reported. Both colloidal and supported nanoparticles (NPs) on carbon nanotubes (CNTs) were prepared in a one-pot synthesis with outstanding control on their size, morphology and composition. These catalysts were evaluated in the selective hydrogenation of alkynes and alkynols. PdCu/CNTs revealed an efficient catalytic system providing high selectivity in the hydrogenation of terminal and internal alkynes. Moreover, this catalyst was tested in the semi-hydrogenation of acetylene in industrially relevant acetylene/ethylene-rich model gas feeds and showed excellent stability even after 40 h of reaction.

Bimetallic nanoparticles represent attractive catalytic systems thanks to the synergy between both partners at the atomic lev-

el, mainly induced by electronic effects which in turn are associated with the corresponding structures (alloy, core-shell, hetero-dimer). This type of engineered material can trigger changes in the kinetics of catalyzed processes by variations on the electrophilicity/nucleophilicity of the metal centers involved and also promote cooperative effects to foster organic transformations, including multi-component and multi-step processes. Solvents become a crucial factor in the conception of catalytic processes, not only due to their environmental impact, but also because they can preserve the bimetallic structure during the catalytic reaction and therefore increase the catalyst life-time. In this frame, the present review focuses on the recent works described in the literature concerning the synthesis of bimetallic nanoparticles in non-conventional solvents, i.e., other than common volatile compounds, for catalytic applications.

Furthermore, one can envisage the concomitant synthesis of both BMNPs and the solid support, such as silica, by in situ methodologies from metal salts and organosilicates. As a highly valuable and sophisticated impregnation method, Zhang et al. reported the synthesis of CuNi@SiO2 from the reverse-microemulsion of Cu(NO3)2•3H2O and Ni(NO3)2•6H2O in a mixture of Brij[®]58 (polyethylene glycol hexadecyl ether) and cyclohexane, followed by aqueous ammonia treatment and tetraethyl orthosilicate (TEOS) hydrolysis at the oil/water interface to trigger the formation of a 3-D network structure Cu(II)/Ni(II)@SiO2. After drying, Cu(OH)2/Ni(OH)2@SiO2 was generated, followed by CuO/NiO@SiO2 after calcination at 500 °C. Finally, CuO and NiO were reduced to CuNi@SiO2 under H2 atmosphere at a high temperature (350 °C). The size and composition of these as-prepared BMNPs were characterized by FT-IR (Fast Transform-InfraRed spectroscopy), TEM, XPS, XRD, and ICP-OES (Inductively Coupled Plasma Optical Emission Spectrometry). As a catalyst for the reduction of p-nitrophenol in water by NaBH4, the calculated activation energy of Cu54Ni46@SiO2 (CuNi BMNPs showed a mean size of 5.7 nm) was much lower (16.6 kJ/mol) than that of monometallic Cu@SiO2 (29.0 kJ/mol) and Ni@SiO2 (39.5 kJ/mol). This CuNi@SiO2 nanocatalyst was recycled up to 10 times whilst preserving its reactivity (>90% conversion for 10th run).

Fe-Ni BMNPs supported on montmorillonite (MMT, soft phyllosilicate mineral) have recently been developed by Chikate

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and coworkers. Thus, an aqueous solution of FeSO4•7H2O and NiSO4•6H2O was mixed with a suspension of Na+-MMT in deionized water, before being reduced by the dropwise addition of NaBH4 under N2 atmosphere. The as-prepared composites were characterized by XRD, TEM, and NH3-TPD (Temperature Programmed Desorption) and were then applied as catalysts for the reductive cyclization of levulinic acid (LA) to γ -valerolactone (GLV). The reduction of LA took place on the bimetallic sites, while the subsequent cyclization to GLV was favored on the strong acidic sites of MMT. Despite the high reactivity (conversion of 90% after 1 h), substantial leaching of iron was observed when using water as the solvent. Another example of a co-reduction methodology by Yamauchid et al. yielded AuPd alloys supported on graphene oxide (GO) nanosheets by the treatment of a GO water suspension with an aqueous solution of H[AuCl4] and PdCl2, using ascorbic acid as the reducing agent. The resulting material was employed as an efficient photocatalyst for the degradation of phenolic compounds in water and it could be recycled up to five times. Long et al. reported the preparation of FePd BMNPs supported on chelating resin from Fe2(SO4)3 and PdCl2 using NaBH4 as the reducing agent in water; these catalysts were used for the selective reduction of nitrate into nitrogen.