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Rational Design of Non-precious Metal Oxide Catalysts by Means of Advanced Synthetic and Promotional Routes

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Message from the Guest Editors

Nowadays the catalytic performance of transition metal oxides, such as ceria-based mixed oxides, perovskites, hexaaluminates, hydrotalcites, and spinels can be considerably enhanced by tailoring the local surface structure (e.g., work function, reducibility, oxygen vacancies) and interfacial phenomena (e.g., metal-support interactions). The latter can be accomplished via the following strategic approaches that could be applied independently or in synergy: (i) employment of state-of-the art nano-synthesis routes towards engineering particle's size and shape (e.g., nanocubes, nanorods), ii) use of structural/surface promoters (e.g., alkali, graphene oxide) towards the optimization of structural and electronic properties, iii) employment of special pretreatment protocols towards the regulation of surface chemistry and metal-support interactions. This holistic approach in conjunction to the fundamental understanding of metalsupport interactions (either geometric or electronic) is expected to lead to the development of low cost (NMsfree), highly active and stable catalysts, for real life applications in the energy and environmental sector.



