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A compact catalytic foam reactor for decomposition of ammonia by the Joule-heating mechanism

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ABSTRACT

Ammonia (NH₃) is a viable hydrogen (H₂) carrier that allows storage and transport of H₂ using well-established infrastructure while maintaining high H₂ storage density. However, cracking NH₃ into H₂ is energy-intensive. Herein, direct Joule-heating of the NiCrAl foam catalyst support is suggested and demonstrated, to minimize heat transfer scale for lower reactor volume, higher efficiency and power density than previously reported reformers. The power density of 128 W/cm³_{Reactor} is achieved based on the lower heating value of H₂: this is 90% higher than previously reported microreactors. Also, even in a small-scale demonstration with a low internal volume of 7.7 cm³ and a high surface-area-to-volume ratio of 5.7 cm⁻¹, a high reforming efficiency of 69.2% is achieved with low catalyst loadings, showing the feasibility of the concept. The as-proposed reactor concept offers a strong prospect for facile adoption of the power-to-X scheme for numerous applications including H₂-fueled islanded networks, and decarbonized energy conversion.

1. Introduction

As the detrimental effects of using hydrocarbon fuels are becoming rapidly tangible in daily life, campaigns for reducing the use of fossil fuels and replacing them with clean energy sources are surging. Hydrogen (H₂) is one of the promising alternatives as a chemical energy carrier. However, despite its high gravimetric energy density, *i.e.*, high heating value (HHV) of 141.91 MJ/kg [1], its low volumetric energy density, *i.e.*, 0.01 MJ/L at 0 °C and 1 atm [2], poses a challenge for H₂ to fully reach its potential as a prominent energy carrier. H₂ can be safely stored and transported via the available chemical H₂ carriers such as ammonia (NH₃), methanol [3], liquid organic hydrogen carriers (LOHCs) [4], formic acid [5], and formate [6]. Among these, NH₃ has the highest gravimetric and volumetric hydrogen storage density (ca. 17.8 wt%, 108 g/L at 20 °C and 8.6 bar [7]) while offering zero carbon emissions during dehydrogenation (*i.e.*, H₂ release by chemical reaction) and no side reactions. Annually, NH₃ is produced by millions of tons (estimated 150 Mt in 2019 [8]) and transported for numerous industries including agriculture, chemicals, and pharmaceuticals. Hence, the

infrastructure hitherto exists, and portable use of NH₃ as an H₂ carrier is feasible. Conversion of NH₃ to H₂, however, is highly endothermic, and providing an efficient heat source for this reaction requires advances in both the material and reactor designs. Therefore, studies have been conducted to address these challenges. For example, with advances in catalyst design, a decomposition temperature of 500 °C for > 99.9% NH₃ conversion has been reported [9]. Nevertheless, the application of sustainable heat sources and delivering the required thermal energy even at this temperature is still a restricting challenge for the use of NH₃ as a reliable H₂ carrier. Therefore, devising a mechanism under the power-to-X scheme [10], in which surplus electric power can be used for other types of energy storage and conversion, should be of main concern for the development of viable NH₃-driven energy industry.

Some methods have been proposed and investigated for efficient delivery of heat via the autothermal design of reactors. Autothermal design refers to the thermally-coupled decomposition-oxidation reactors. Related reports are mostly inclined towards thermal coupling of NH₃ decomposition reactor with oxidation reactors fueled with NH₃ [11–13] and/or hydrocarbons [14,15]. However, heat loss resulted from these designs is a major issue. In some reports, process intensification

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