Hybrid Technology for DeNOxing by LNT-SCR System for Efficient Diesel Emission Control: Influence of Operation Parameters in H₂O + CO₂ Atmosphere

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**Abstract:** The behavior and operation parameters were analyzed for the hybrid LNT-SCR (Lean NOₓ-Trap–Selective Catalytic Reduction) system with advanced catalyst formulations. Pt-Ba-K/Al₂O₃ was used as an NSR (NOₓ Storage and Reduction) or LNT catalyst effective in NOₓ and soot simultaneous removal whereas Cu-SAPO-34 with 2 wt.% of copper inside the structure was the small pore zeolite employed as the SCR catalyst. Under alternating and cyclic wet conditions, feeding volumetric concentrations of 1000 ppm of NO, 3% of O₂, 1.5% of water, 0.3% of CO₂, and H₂ as a reductant, the NOₓ-conversion values were above 95% and a complete mineralization to nitrogen was registered using θ ≤ 3 (20 s of regeneration) and a hydrogen content between 10,000 and 2000 ppm in the whole temperature range tested. An excess of hydrogen fed (above 1% v/v) during the rich phase is unnecessary. In addition, in the low temperature range below 250 °C, the effect is more noticeable due to the further ammonia production and its possible slip. These results open the way to the scale up of the coupled catalytic technologies for its use in real conditions while controlling the influence of the operation map.

**Keywords:** NSR-SCR hybrid catalysts; coupled aftertreatment technologies; DeNOxing; fixed double bed; NOₓ emission control

1. Introduction

The increase in nitrogen monoxide emissions with the use of new biofuels has been proven in a motor bench [1]. The way of driving modifies the production of NOₓ and efficient catalytic after-treatment technologies are required to reduce the levels below the limits imposed by the Euro 6 regulation. At present, NOₓ Storage and Reduction (NSR) technology is the most efficient in the removal of nitrogen oxides from diesel exhaust gases. However, it is not enough to achieve zero emissions at the outlet of the vehicles, due to the formation of an NH₃ slip that is strongly dependent on the catalyst formulation and the operation conditions [2–5]. The Selective Catalytic Reduction (SCR) process, which is based on the mineralization of NOₓ to N₂ and water with ammonia [6–9], requires the injection of an external reducing agent. Therefore, the coupling of SCR and NSR technologies is being widely studied to solve these drawbacks [10–15].

The idea consists of placing an SCR catalytic bed downstream of the NSR catalyst, so that the ammonia produced during the lean phase in the LNT-catalyst (Lean NOₓ Traps) can be stored in the SCR catalyst and, then, it would react with the unconverted NOₓ in the oxidant phase, which increases the NOₓ conversion and the nitrogen selectivity. Commercially, the system was implemented in the Mercedes E320 Blue-Tech vehicle in 2007 [16]. The main advantages of this system compared to the
single NSR catalytic bed are: the increase in NO\textsubscript{x} conversion values, the ammonia-slip reduction, a wider operating temperature window, the improvement of the catalyst durability, and the reduction of unburned hydrocarbon production due to a reaction in the SCR catalyst [10,17,18].

It is necessary to delve into the study of coupling technologies and their possible application in real conditions. Several authors have studied the coupling (double-bed) of standard Pt-Ba/Al\textsubscript{2}O\textsubscript{3} and Fe-ZSM\textsubscript{5} zeolite as an NSR and SCR catalyst, respectively, in powder form [13,19,20] by detecting higher nitrogen production in the overall cycle. Similar results have been obtained by other authors when the Cu-ZSM-5 was placed downstream of a Pt-Rh-Ba/Al\textsubscript{2}O\textsubscript{3} [21] or when Cu-ZSM-5 is substituted for Co/Al\textsubscript{2}O\textsubscript{3} [22]. In the double-bed configuration, a higher level of nitrogen is detected in the lean phase when compared to a single bed due to the reaction of the nitrogen oxides with the ammonia that had been stored in the system in the previous phase.

The physical mixture of the catalysts produces a lower amount of nitrogen in the rich phase because one of the nitrogen production routes uses the ammonia as an intermediate and, with the zeolite being so close, the ammonia formed could be retained on the zeolite and, thus, the nitrogen formation would decrease. Therefore, the use of two independent catalysts without a physical mixture seems to be the most appropriate configuration [13]. Although other authors [23] have also proposed dual-layer monolithic catalysts, with both materials over the same wall, the higher conversions of NO\textsubscript{x} to nitrogen are, however, achieved using sequential monoliths.

As SCR catalysts, some authors [24] have developed materials based on modified alumina and prepared by a sol-gel method with the required acidic and redox properties by introducing metal ions such as cerium (for oxygen mobility), titanium, zirconium, and silica. Although, these materials are effective in the NH\textsubscript{3}-SCR reaction together with the NH\textsubscript{3}-SCO (Selective Catalytic Oxidation). The zeolite with the incorporation of copper is the most effective material that can be used in this technology up to now [25,26]. As far as copper zeolites are concerned, materials prepared by ion exchange or impregnation have been studied in granular and monolithic form [12,23,27]. In addition, large and small pore zeolites are used, and a higher hydrothermal stability and a wider operation temperature window were observed when the small pore zeolites were analysed. In all cases, the ammonia slip was negligible and N\textsubscript{2}O selectivity decreased with the use of double-bed configuration. Among them, SAPO-34 with a chabazite structure has been reported as one of the most active small pore zeolite used as an SCR catalytic bed related to a high substitution capacity by disperse and isolated copper species [28–30].

Regarding the operation parameters, the time of the reduction phase or the ratio between oxidation/reduction period times is another open key point in the behaviour of these materials [31–33]. A decrease in the rich pulse length provokes a reduction in the NO\textsubscript{x} removal efficiency because insufficient time was available for the completely regeneration of the catalysts [22,34]. These parameters influence the cost of the process.

It is important to establish the relationship between new improved formulations able to cushion NO\textsubscript{x} emissions in a wet atmosphere with the operation conditions of diesel engines for a suitable and real penetration of the zero emission technologies in the automotive sector.

Therefore, this work is focused on the study of the behaviour and the operation conditions of the sequential Pt-Ba-K/Al\textsubscript{2}O\textsubscript{3} and Cu-SAPO-34 catalysts, as an LNT-catalyst and SCR material, respectively, in powder form and in the presence of water and CO\textsubscript{2}, which has never been addressed in literature.

2. Results and Discussion

2.1. Evaluation of V\textsubscript{NSR}/V\textsubscript{SCR} Ratio in Dry He-Atmosphere

Several cyclic experiments up to 50 cycles (1000 ppm of NO + 3% of O\textsubscript{2} and 2000 ppm of H\textsubscript{2}) were carried out using transient response method (TRM) in an He-atmosphere (dry conditions) at each temperature (200 or 350 °C) up to the stabilization of the catalyst behaviour. A low temperature value of the operation window of SCR and a representative temperature of the NSR catalysts were
selected. Therefore, in Figure 1, the influence of \( V_{\text{NSR}}/V_{\text{SCR}} \) in DeNOX performance expressed as NO\(_x\) conversion and selectivity to N\(_2\), NH\(_3\), and N\(_2\)O are presented.

\[ \begin{array}{c}
1 \text{ NSR: 0 SCR} & 1 \text{ NSR: 1 SCR} & 1 \text{ NSR: 0.5 SCR} \\
1 \text{ V}_{\text{NSR}}/V_{\text{SCR}} & 1 \text{ V}_{\text{NSR}}/V_{\text{SCR}} & 1 \text{ V}_{\text{NSR}}/V_{\text{SCR}} \\
X_{\text{NO}_x} & S_{\text{N}_2} & S_{\text{NH}_3} & S_{\text{N}_2\text{O}} \\
\end{array} \]

\( \text{Figure 1.} \) Conversion and selectivity values for different NSR/SCR volume ratios in experiments with NSR conditions (1000 ppm NO + 3% O\(_2\)/2000 ppm H\(_2\) in He, 15–15 min) at (a) 200°C and (b) 350°C.

An increase in the NO\(_x\) removed can be noticed compared to the values obtained with the single NSR bed-catalyst, Pt-Ba/K/Al\(_2\)O\(_3\), as can be verified in Reference [35], and was also observed and reported by other authors for both Pt-(Rh)-Ba/Al\(_2\)O\(_3\) catalysts and Cu-zeolites, such as ZSM5, BETA, or CHA [10, 22, 27] working in cyclic conditions. For 1 NSR: 0 SCR configuration, i.e., with only a Pt-Ba-K/Al\(_2\)O\(_3\) catalyst, high NO\(_x\) conversion to N\(_2\) is detected with low ammonia production and a slight amount of N\(_2\)O formation only at low temperatures. Ammonia was observed after nitrogen production in the middle of the rich phase, whereas an intense signal of N\(_2\)O was detected at the beginning of the step with a maximum of 70 ppm and was related to the low efficiency of the oxygen-covered Pt sites during the surface regeneration phase by H\(_2\), which prevents the NO dissociation and leads to N\(_2\)O formation [36]. This does not take place at a high temperature since the Pt is kept in the reduced form [18]. The placing of the SCR catalyst after the NSR catalytic bed completely removes the ammonia production in all the volume proportions presented in this scenario. For 1 NSR: 1 SCR, besides the improvement in the NO\(_x\) conversion and in the ammonia slip, a decrease in the N\(_2\)O, which is a highly polluting greenhouse gas, emission from 0.7% to 0.1% is also observed, as a consequence of the copper-zeolite bed-catalyst placed downstream. Nevertheless, the decrease of the SCR catalyst-volume, 1 NSR: 0.5 SCR, in spite of reducing the NO\(_x\) and NH\(_3\) production, negatively affects the N\(_2\)O decrease, as other authors have observed for the coupling of an NSR catalyst based on noble metals and an Fe-zeolite [37] with commercial catalysts and similar space velocity values. A plausible explanation of this effect is that the increase in the NSR weight with respect to the SCR produces a higher amount of N\(_2\)O and the SCR in this condition is not able to reduce all the gas produced. In this way, a reduction of N-containing products via the synergy of LNT-SCR coupled catalysts facilitated a major NO\(_x\) conversion. Considering the NO\(_x\) conversion, NH\(_3\) slip, and N\(_2\)O production evaluated for three different NSR-SCR catalytic ratios in a dry He-atmosphere, the most suitable volume ratio of catalysts was 1:1 for the NSR-SCR hybrid system, since it keeps the NO\(_x\) conversion, prevents the ammonia slip, and reduces the N\(_2\)O emission.

2.2. Influence of H\(_2\)O and CO\(_2\) Presence over NSR-SCR Systems

Hybrid LNT-SCR double-bed catalysts with \( V_{\text{NSR}}/V_{\text{SCR}} = 1/1 \) was also studied in CO\(_2\) wet-atmosphere (H\(_2\)O + CO\(_2\)), and, although the NSR catalyst presents higher selectivity values, this...
is a more realistic condition that influences the deNOxing performance. In Figure 2, the profiles of the outlet gas distribution at 350 °C (1000 ppm of NO + 3% of O2/2000 ppm of H2) are represented for a cycle in which the catalysts are working in a stable way, in helium (Figure 2a), and water + CO2 (Figure 2b) atmospheres, respectively.

![Figure 2. Storage-reduction stationary cycle (15-15 min) for Pt-Ba-K/Al2O3 + 2Cu-SAPO-34 at 350 °C in (a) He and (b) H2O + CO2.](image)

Under He atmosphere in the lean phase, the typical NOx saturation curve was appreciated, as a consequence of the storage of nitrites/nitrates, and the reaction with the adsorbed ammonia producing nitrogen and water, as is shown. NO and NOx curves overlapped since NO2 was not detected at the outlet due to the combination of the reaction with ammonia and the adsorption over both catalytic surfaces. In the rich phase, the profile of hydrogen is representative of the consumption by the reduction of the NOx adsorbed on the catalyst surface up to the entire regeneration of the catalyst with the corresponding nitrogen and water formation. However, during this phase (step), ammonia presence was undetectable due to the net storage in the SCR catalyst placed downstream. Moreover, the formation of N2O is almost insignificant due to the capacity of the copper modified-zeolite for the N2O decomposition, which other authors have reported with different copper-based materials [38–41] since copper ionic pairs or clusters and a chabazite structure are able to decompose N2O.

On the other hand, in the presence of water and CO2 (Figure 2b), besides the same gas profiles, an additional consumption of CO2 was detected with CO formation (data not shown) due to the reverse reaction of the Water Gas Shift [14]. The obtained profiles both in dry and wet conditions in Helium and 1.5% H2O + 0.3% CO2 presence, respectively, are similar to those reported [19] for the coupled Pt-Ba and Fe-ZSM5 catalysts, in terms of nitrogen and water production and the shape and trend of the NOx and H2 saturation curves.

In Table 1, the representative activity parameters are shown and compared with those obtained for the single NSR catalytic bed. Similar trends were observed for both systems. However, the difference between the analyzed parameters should be pointed out. For all the experiments, as was expected, the amount of NOx removed increased with the incorporation of an SCR bed-downstream, with increased values similar to those previously reported [13,19,20]. Nevertheless, the effect depended on the temperature and the reaction atmosphere.
Table 1. Reactivity data calculated during the storage-reduction of NO\textsubscript{x} experiments from isothermal TRM profiles at 200 \degree C and 350 \degree C for Pt-Ba-K/Al\textsubscript{2}O\textsubscript{3} (NSR) and Pt-Ba-K/Al\textsubscript{2}O\textsubscript{3} + 2Cu-SAPO-34 (NSR-SCR) systems.

| Temperature (\degree C) | Atmosphere   | System         | Removed NO\textsubscript{x} (10\textsuperscript{3} mol NO\textsubscript{x} g\textsuperscript{-1} NSR\textsuperscript{-1}) | X\textsubscript{NOx} (%) | S\textsubscript{N2} (%) | S\textsubscript{NH3} (%) | S\textsubscript{N2O} (%) |
|-------------------------|--------------|----------------|-------------------------------------------------------------------------------------------------|--------------------------|--------------------------|--------------------------|--------------------------|
| 200                     | He           | NSR            | 0.39                                                                                             | 91                       | 94.1                     | 5.2                      | 0.7                      |
|                         |              | NSR-SCR        | 0.51                                                                                             | 99                       | 99.9                     | 0                        | 0.1                      |
| H\textsubscript{2}O + CO\textsubscript{2} | NSR            | 0.24                                                                                             | 83                       | 90.5                     | 9.5                      | 0                        |
|                         |              | NSR-SCR        | 0.35                                                                                             | 84                       | 100                      | 0                        | 0                        |
| 350                     | He           | NSR            | 0.42                                                                                             | 88                       | 97                       | 3                        | 0                        |
|                         |              | NSR-SCR        | 0.53                                                                                             | 91                       | 100                      | 0                        | 0                        |
| H\textsubscript{2}O + CO\textsubscript{2} | NSR            | 0.41                                                                                             | 86                       | 100                      | 0                        | 0                        |
|                         |              | NSR-SCR        | 0.52                                                                                             | 92                       | 100                      | 0                        | 0                        |

At low temperatures, the impact was more noticeable because of the higher ammonia production that can be used in the SCR process and the initially, more unfavourable conditions. The higher the amount of ammonia adsorbed, the more efficient the configuration since the SCR catalyst begins to work when a considerable amount of ammonia is stored in the catalyst. The conversion values also increased as shown by U. De La Torre et al. [12] who observed an increment in the presence of copper-exchanged zeolites. In this case, the initial values registered are high, almost 90\%, and the effect is less remarkable with increments below 10\%.

At 200 \degree C and in an H\textsubscript{2}O + CO\textsubscript{2} atmosphere, it seems that a balanced-cooperated effect causes a somewhat different behaviour. On the one hand, the Pt-Ba-K/Al\textsubscript{2}O\textsubscript{3} catalyst in the presence of water and CO\textsubscript{2} reduced the NO\textsubscript{x} adsorption capacity and conversion, which is in agreement with other authors [42,43], due to the interaction of the carbonates that compete with the nitrates for the same active sites [44]. The NO\textsubscript{2} release is difficult in the presence of water and in the negative impact in the regeneration step when water and CO\textsubscript{2} are co-fed because of the poisoning of the Pt sites by CO\textsubscript{2} formation associated with a reverse water gas shift reaction [45]. On the other hand, the presence of water and CO\textsubscript{2} in the SCR process over the 2 Cu-SAPO-34 improves the performance in terms of conversion and selectivity values since it inhibits the ammonia oxidation and the nitrate decomposition and modifies the copper environment by obtaining more active species [46]. Therefore, in spite of the improvement provided by the SCR catalyst in these conditions, the negative effect of the reaction atmosphere over the NSR catalyst is stronger and, as a consequence, the conversion value is not considerably increased.

At 350 \degree C, the calculated parameters referred to the single NSR catalyst and obtained from the mathematical treatment of the curves represented in Figure 2, were high, with conversion and N\textsubscript{2} selectivity values of around 90\% and higher than 95\%, respectively, and the influence of placing an SCR catalyst downstream is less noteworthy. Nevertheless, conversion and the removed NO\textsubscript{x} increased due to the reduction with the adsorbed ammonia. The CO\textsubscript{2} inhibition impact decreases with the temperature [47]. An increase in the conversion value can also be obtained because, in the reduction phase, the hydrogen can react with an initial ammonia formation and this ammonia could interact with the unconverted NO\textsubscript{x} forming nitrogen and water by selective catalytic reduction. L. Castoldi et al. [13] have observed the same effects caused by the presence of water and CO\textsubscript{2} for the coupling of Pt-Ba/Al\textsubscript{2}O\textsubscript{3} and Fe-ZSM5 with an increase in the selectivity and conversion with a rise in temperature or compared to the single NSR catalyst. However, the overall NO\textsubscript{x} conversion of the hybrid system presented in this case is higher than data found in the literature.

As far as selectivity is concerned, almost complete nitrogen selectivity was observed at 200 and 350 \degree C. Thus, the double bed configuration allows the adsorption of the ammonia that is produced in the NSR-catalyst and the reaction with the nitrogen oxides of the following cycle. In addition,
the decrease of the low amount of N\textsubscript{2}O, only formed at a low temperature in dry conditions, was performed by the zeolite because that is able to decompose the N\textsubscript{2}O producing nitrogen.

### 2.3. Influence of the Duration of Lean-Rich Cycles and the Concentration of H\textsubscript{2}

The system with \(V_{\text{NSR}}/V_{\text{SCR}}\) equal to one was selected due to the NO\textsubscript{x} conversion level reached, ammonia slip prevention, and N\textsubscript{2}O emission reduction. Besides temperature, the duration of lean-rich cycles and the concentration of the reducing agent (currently H\textsubscript{2}) in the respective lean and rich phases are very critical parameters [16]. A series of experiments were carried out in order to analyze the influence of these other operation parameters in the catalytic activity of the NSR-SCR system. For this purpose and with the intention of comparing the results with other authors, the dimensionless parameter \(\theta\) has been defined as the ratio between the oxidation and reduction times and different values have been used, while the space velocity, the weight ratio between catalytic beds, and the feed composition were maintained. The optimization of lean and rich periods in order to ensure stable lean–rich cycles is essential for the engine operation. In addition, the concentration of the reducing agent (H\textsubscript{2}) was also modified. All the operation parameters were summarized in Table 2.

**Table 2. Flow conditions used in the TRM experiments over NSR-SCR systems.**

|                        | Lean Phase | Rich Phase |
|------------------------|------------|------------|
| NO (ppm)               | 1000       | -          |
| O\textsubscript{2} (v/v %) | 3         | -          |
| H\textsubscript{2} (v/v %) | -         | 0.2, 1, 2.5 |
| H\textsubscript{2}O (v/v %) | 1.5       | 1.5        |
| CO\textsubscript{2} (v/v %) | 0.3       | 0.3        |
| He                     | Balance    | Balance    |
| GHSV (h\textsuperscript{-1}) | 3 \times 10\textsuperscript{4} | 3 \times 10\textsuperscript{4} |
| Temperature (°C)       | 200, 350   | 200, 350   |
| \(\theta = 1\)         | 20, 60 s   | 20, 60 s   |
| \(\theta = 3\)         | 60 s       | 20 s       |
| \(\theta = 7.5\)       | 150 s      | 20 s       |

\(\theta = \) ratio between the oxidation and reduction times.

In Figure 3, the profiles throughout the cycles for the experiments over the hybrid system (\(V_{\text{NSR}}/V_{\text{SCR}} = 1 @350 \, ^{\circ}\text{C}\)) when the parameter \(\theta\) is set at 7.5 are shown, varying the amount of H\textsubscript{2} as a reductant from 0.2 to 2.5 v/v % and under an H\textsubscript{2}O + CO\textsubscript{2} atmosphere. As can be observed, the behavior in the storage and reduction of NO\textsubscript{x} is completely dependent on the reductant concentration. At hydrogen contents close to the stoichiometric value (2000 ppm, Figure 3a), the NO\textsubscript{x} amount at the outlet increases throughout the cycles up to stabilization, which indicates that the amount of the reducing agent is not enough for the complete regeneration of the catalyst. This effect is similar to that observed in the transient experiments carried out in the thermobalance where the baseline increased at the end of the experiment due to an incomplete regeneration of the catalyst in the LNT condition with Pt-Ba-K/Al\textsubscript{2}O\textsubscript{3} catalyst, which we reported in a previous work [35]. Hydrogen was not detected in the outlet stream during the cycles due to the total consumption by reaction with the adsorbed nitrates.
Figure 3. Storage-reduction cycles for Pt–Ba-K/Al₂O₃ + 2Cu-SAPO-34 (V_{NSR}/V_{SCR} = 1) at 350 °C with lean-rich periods of 150–20 s (θ = 7.5) with a hydrogen concentration of (a) 0.2, (b) 1, and (c) 2.5 v/v % in H₂O + CO₂.
At hydrogen content of 1% (Figure 3b), the amount of NO\textsubscript{x} detected at the outlet was lower than in the previous experiments, even though it also increased with time, and hydrogen was not observed during the experiment, which means that, again, this concentration is not enough for the whole regeneration of the LNT-catalyst’s surface.

For a reducing agent concentration of 2.5% (Figure 3c), the product distribution was completely different. The profile of NO\textsubscript{x} was identical for all the cycles performed, which means that the nitrates adsorbed onto the catalyst are fully removed by their reaction with the hydrogen in each cycle. In addition, H\textsubscript{2} was detected during the rich step since an excess of the reductant was being fed into the reactor in these conditions. In terms of the product distribution and selectivity, nitrogen was the only compound observed at the outlet. Neither ammonia nor N\textsubscript{2}O were detected. However, the nitrogen production was higher as the hydrogen content increased, in consequence with the regeneration process explained above. In addition, if an overestimated amount of hydrogen is used, the production of ammonia would be too high and the SCR activity would decrease, which is the effect identical to the use of long, rich periods \[14,48\]. However, the N\textsubscript{2}O formation decreases with the amount of reductant agent \[49\].

In Table 3, the TRM-parameters calculated from the experiments with lean-rich time variations as a function of hydrogen percentage and temperature is summarized. NO\textsubscript{x} stored and NO\textsubscript{x} conversion values follow the trend described above from Figure 3. The influence of hydrogen concentration on the catalytic behaviour can be confirmed for the Pt-Ba-K/Al\textsubscript{2}O\textsubscript{3} + Cu-SAPO-34 mixture.

| H\textsubscript{2} (ppm) | T (°C) | 60 s–20 s (θ = 3) | 150 s–20 s (θ = 7.5) |
|--------------------------|--------|-----------------|-----------------|
|                          |        | mmolNO\textsubscript{x}·g\textsubscript{NSR}\textsuperscript{−1} | X\textsubscript{NOX} (%) | mmolNO\textsubscript{x}·g\textsubscript{NSR}\textsuperscript{−1} | X\textsubscript{NOX} (%) |
| 2000                     | 200    | 0.44            | 78.6            | 0.61            | 76.3            |
|                          | 350    | 0.46            | 82.5            | 0.62            | 81.3            |
| 10,000                   | 200    | 0.69            | 97.3            | 1.2             | 94.3            |
|                          | 350    | 0.71            | 99.3            | 1.3             | 95.6            |
| 25,000                   | 200    | 0.69            | 97.7            | 1.6             | 97.7            |
|                          | 350    | 0.71            | 99.4            | 1.6             | 97.9            |

The operation temperature was another parameter to control since the amount of nitrogen oxides that can be stored on the surface depends on the temperature as well as the selectivity. The improvement of the activity values at a low temperature is more noteworthy than at higher temperatures \[32\]. As can be seen, the influence of the reaction temperature was less noticeable than the hydrogen concentration. Nevertheless, the amount of nitrogen produced as well as the NO\textsubscript{x} at the outlet were slightly different depending on the temperature. Higher values of the activity parameters were obtained in the experiments at high temperatures, which occurs in the single NSR catalytic bed. However, the improvement of the coupling of both technologies in the hybrid system was more remarkable at low temperatures since the initial conditions were less favorable.

In Figure 4, the profiles throughout the cycles for the experiments over the hybrid system (V\textsubscript{NSR}/V\textsubscript{SCR} = 1, @350 °C) when the parameter θ is set at 3 are shown, which varies the amount of H\textsubscript{2} as a reductant from 2000 to 25,000 ppm and under the H\textsubscript{2}O + CO\textsubscript{2} atmosphere. The decrease of the θ value below 3 modifies the behavior of the hybrid system. For the runs performed with 0.2% of hydrogen (Figure 4a), it is observed that the NO\textsubscript{x} concentration at the outlet increased throughout the cycles, but 1 v/v % of H\textsubscript{2} is enough for the complete reaction with the adsorbed nitrates. The conversion values obtained in these conditions (Table 3) indicated that total regeneration is possible when working with reducing agent content around five times the stoichiometric value, which is consistent with other authors \[50\] for higher flowrates and dual-layer systems.
Figure 4. Storage-reduction cycles for Pt-Ba-K/Al₂O₃ + 2Cu-SAPO-34 (V_{NSR}/V_{SCR} = 1) at 350 °C with lean-rich periods of 60–20 s (θ = 3) with a hydrogen concentration of (a) 0.2, (b) 1, and (c) 2.5 v/v % in H₂O + CO₂.
For the runs performed with 2.5 v/v % of H₂ (Figure 4c), the amount of H₂ was overestimated for the regeneration of the catalyst and a minimum amount of NOₓ was detected (as can be observed in the inset). Zero emissions of nitrogen pollutant compounds were appreciated in these conditions. However, the amount of hydrogen at the outlet was high and this would imply the use of an unnecessary excess of reductant. Regarding the product distribution, nitrogen was the only product detected during the experiments, which was observed for higher values of θ. Nevertheless, the amount of nitrogen for lower θ values was higher since the time of reduction was the same whereas the time of oxidation was lower.

From all the results presented, it can be assumed that, in the whole operation range, the amount of nitrogen oxides removed and the conversion and nitrogen selectivity values (Table 3) were higher than in the experiments with the single Pt-Ba-K/Al₂O₃ catalyst. Particularly, they increased around 30% and 5%, respectively. The amount of removed NOₓ decreased with the reduction of hydrogen concentration, as a consequence of the incomplete regeneration of the catalyst explained above. The increase in the oxidation phase time caused an increment of almost double (between 38% and 100%) the amount of removed NOₓ, at a constant removal rate. At high temperatures, the effect of the reductant content is less significant provided that it is higher than the stoichiometric concentration since the conversion values are above 90% and the nitrogen selectivity is high.

This means that such high amounts of hydrogen are not necessary, and the reducing agent consumption can be reduced as well as the chemical-equivalent input of a reducing agent. Nevertheless, although all the conditions are highly effective, hydrogen content around the stoichiometric seems to not be suitable for the removal of NOₓ in cyclic periods when the time of oxidation is higher than the reduction time. In addition, hydrogen levels below 0.8% were not recommendable to work with a θ value around 7, due to the fact that the reductant supply would defected and, as consequence, the LNT-catalyst could not be completely regenerated, which decreases the NOₓ removal efficiency [11], as has been verified. The θ decrease allows the regeneration of the catalyst, high NOₓ removal, and remarkable conversion values with less hydrogen in the whole operation range. Additionally, the reduction in θ implies the decrease in the necessary content of the reducing agent for the regeneration of the catalyst and obtaining high conversion values that prevent catalyst saturation. If the ratio between oxidant and reduction times is above 1, the amount of hydrogen used in the regeneration phase has to be higher than the stoichiometric value. Nevertheless, as has been observed in all the experiments up to saturation with the same time for both phases, the regeneration of the catalysts was complete.

Therefore, it is worth considering the influence of the period time when the value of θ is set to 1. For that, experiments with the same time for both steps were carried out without modifying the θ value, and the profiles of the product distribution for the experiments with 60–60 s and 20–20 s using 0.2% and 1% of hydrogen in the H₂O + CO₂ atmosphere, V<sub>NSR</sub>/V<sub>SCR</sub> = 1, and 350 °C, are represented in Figure 5.

As can be observed in Figure 5b,d, hydrogen concentrations above the stoichiometric, which is 1% in this case and higher than the exact content for the reduction of the nitrates. This would involve an unnecessary fuel overfeeding and cost. However, the total removal of the nitrogen oxides and the complete selectivity to nitrogen can be ensured from the first cycles. On the other hand, the use of H₂ in the stoichiometric value presented a different behavior depending on the length of each period. In these conditions, the LNT-catalyst was being saturated slowly, even though the saturation time is modified by the period time. The saturation of the catalyst is reached before when longer times are used. Comparing Figure 5a,c, in the experiment with 60 s for each phase, the catalyst is saturated faster than using 20 s for each phase. This is due to the fact that the regeneration rate is different from the adsorption rate. The removed amount percentage and the adsorption rate are identical in each cycle because they depend on the temperature.
The changes detected were similar to the kinetic and geometric effects observed in the TG (thermogravimetric) experiments in a previous work [35] when the NSR catalyst was not completely regenerated and, consequently, 2000 ppm of hydrogen is the exact stoichiometric value and the total reduction of the adsorbed species requires much longer time periods.

Therefore, stoichiometric hydrogen levels in over accumulation conditions cause the incomplete regeneration of a part of the catalyst and the saturation of the surface throughout the cycles. The operation conditions should ensure the availability of the catalyst for the following cycle. Therefore, hydrogen concentrations above 1.5 times the stoichiometric value are needed in order to obtain zero emissions and the regeneration of the catalyst when the θ value is around 1.

Additional experiments were carried out in order to obtain a response surface modifying the parameters $X = f (\% H_2, \theta)$. The distribution of the NO\textsubscript{x} conversion data obtained after 20 cycles, while keeping a constant rich phase time of 20 s, is represented in Figure 6. The operation regions are displayed with different colors, according to the conversion values, and isoconversion lines limit them.
As explained above, the increment in the hydrogen content increases the conversion and removal of nitrogen oxides, even though the hydrogen content should not be in excess, which produces unnecessary cost and unburned hydrocarbon emission. At low time ratios, with hydrogen content above the stoichiometric value, more than 90% of NO\textsubscript{x} conversion is achieved. When increasing the $\theta$ value, an ascendant trend with a low slope can be observed because the increment of the NO\textsubscript{x} adsorbed required a higher amount of reductant agent to regenerate the surface and convert the NO\textsubscript{x}. Yet, high conversion values are obtained in all the operation ranges. These formulations, keeping a $V_{\text{NSR}}/V_{\text{SCR}} = 1/1$, are suitable to incorporate into a cordierite monolithic structure (400 cpsi), with an Al\textsubscript{2}O\textsubscript{3} wash-coating (12–15 $\mu$m), as layers below 5 $\mu$m of LNT or SCR catalysts. The macroscopic cubic structure of Cu-SAPO-34 material limits the thickness of the layer to keep the homogenous structure along the channel. Therefore, the hybrid system can operate in a bench engine.

3. Materials and Methods

3.1. Synthesis of Catalysts

Pt-Ba-K/Al\textsubscript{2}O\textsubscript{3} was employed as an NSR catalyst due to its excellent properties in NO\textsubscript{x} and soot removal. The multi-metallic supported catalyst was prepared by co-impregnation with the following metal contents 0.4 at—Pt, 3.5 at—Ba, and 1.5 at—K nm\textsuperscript{-2} using $\gamma$-Al\textsubscript{2}O\textsubscript{3} TH, with 144 m\textsuperscript{2}\cdot g\textsuperscript{-1} of surface area, which is equivalent to 1.9, 11.5, and 1.4 wt.% of Pt, Ba, and K, respectively, with the same procedure described in Reference [35] where an optimization of the metal loading compared to the standard Pt-Ba model catalyst was demonstrated. Cu-SAPO-34 was used as an SCR catalytic bed because of the combination of its catalytic activity and structural properties [30,46]. The catalyst was synthesized by a hydrothermal method assisted by ultrasound with 2 wt. % of copper in the molar gel composition: 2 DEA (diethylamine): 0.6 SiO\textsubscript{2}: 1 Al\textsubscript{2}O\textsubscript{3}: 0.2 P\textsubscript{2}O\textsubscript{5}: 50 H\textsubscript{2}O. The synthesis procedure of these materials and their characterization and activity in the individualized technologies have been previously reported [30,35,51].

3.2. Catalytic Activity

Experiments were carried out in a quartz U-tube reactor connected to the QMS 200 Mass Spectrometer (Pfeiffer Vacuum Prisma\textsuperscript{TM}, Málaga, Spain) and the Gas Chromatograph $\mu$-GC 3000A (Agilent, Santa Clara, CA, USA) with a data sampling interval of 4 s and 3 min, respectively. The
zeolitic material was placed downstream in the reactor and both catalytic beds were separated by a quartz wool layer. The total gas flow was 100 cm$^3$·min$^{-1}$ using 60 mg of each catalytic bed in powder form, where the catalysts were mixed with quartz in order to obtain different ratios. Experiments were performed alternating lean-rich cycles as a TRM (Transient Response Method) procedure with 1000 ppm of NO + 3% of O$_2$ and 2000 ppm of hydrogen as a reductant up to saturation and complete regeneration, i.e., 15 min for each phase for the experiments in stoichiometric conditions. The hydrogen content and the time of each period were also modified. In Table 2, the reaction conditions used in the experiments are summarized.

As can be observed, although the LNT cyclic procedure was used, some parameters varied in order to study their effect, such as: the catalytic-bed volume ratio (V$_{NSR}$/V$_{SCR} = 1/0$, 1/1 and 1/0.5), the hydrogen content as a reductant agent (0.2, 1, 2.5% (v/v)), the oxidant/reductant phases time ratio ($\theta = 1, 3, 7.5$), and the impact of H$_2$O + CO$_2$ atmosphere. The Gas Hourly Space Velocity (GHSV) was maintained at 3·10$^4$ h$^{-1}$ with respect to each catalytic bed in order to obtain data that were comparable to those reported for the single bed. The amount of NO$_x$ removed per gram of catalyst was calculated as the difference in area between the inlet and the outlet in the lean phase, which is multiplied by the corresponding ratio between total flow (F) and molar volume (V$_m$) (Equation (1)).

$$\frac{\text{mmol NOx removed}}{g_{\text{cat}}} = \frac{(A_{\text{in NOx}} - A_{\text{out NOx}})_{\text{lean}} \cdot F}{w_{\text{cat}}}$$

(1)

The NO$_x$ conversion was calculated as the difference between the removed amount and the NO$_x$ at the outlet in the rich phase divided by the NO$_x$ removed (Equation (2)).

$$X_{\text{NOx}} = \frac{(A_{\text{in NOx}} - A_{\text{out NOx}})_{\text{lean}} - (A_{\text{out NOx}})_{\text{rich}}}{(A_{\text{in NOx}} - A_{\text{out NOx}})_{\text{lean}}}$$

(2)

4. Conclusions

In this study, the application of a combined LNT-SCR dual-bed catalytic system to improve NO$_x$ reduction efficiency together with the potential in deNOxing performance under simulated diesel exhaust conditions were demonstrated. This opens the way toward the use of a hybrid after-treatment catalytic system to reach zero NO$_x$-emission. Volume ratio of LNT and SCR catalysts and the operation parameters (reductant content and lean/rich time ratio) were established for the efficient and enhanced formulations of catalysts Pt-K-Ba/Al$_2$O$_3$ + Cu-SAPO-34.

The NSR-SCR hybrid system, with a volumetric ratio 1:1 working with $\theta$ values lower than 3 and a hydrogen content as a reducing agent between 2000 and 10,000 ppm, favored high levels of mineralization to nitrogen, complete regeneration of catalysts after the rich phase, and hindered the detection of N-intermediates as N$_2$O or NH$_3$ during the nearly 50 cycles by working with conversion average values close to 95%.

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