Influence of the nature of the promoter in NiO catalysts on the selectivity to olefin during the oxidative dehydrogenation of propane and ethane

by

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Abstract

A comparative study of the catalytic properties for the oxidation of C$_2$-C$_3$ alkanes and olefins has been carried out over unpromoted and M-promoted NiO catalysts (Me= K, La, Ce, al, Zr, Sn, Nb). The catalysts have been characterized by several physico-chemical techniques (UV Raman, Visible Raman, FTIR of adsorbed CO and XPS). The characteristics of promoter elements are of paramount importance, since they are able to modify both the nature of the active nickel and the concentration of electrophilic O$_2^−$/O$^-$ oxygen species. Thus, a relatively high acidity and valence of the promoter oxide (with oxidation state higher than +3) are necessary to achieve high selectivity to olefins during the oxidative dehydrogenation (ODH) of C$_2$-C$_3$ alkanes. In addition, an inverse correlation between the selectivity to the corresponding olefin and the concentration of electrophilic oxygen species has been observed, although the selectivity to propene during propane ODH is lower than the selectivity to ethylene achieved during ethane ODH. On the other hand, a very low influence of alkane conversion on the selectivity to the corresponding olefins is observed. This behaviour can be explained by considering that the reaction rate for olefin combustion are lower to the reaction rate for alkane oxidation. However, the comparative study of the oxidation of alkanes and olefins suggest that the differences observed between the ODH of propane and ethane are not related to the reactivity of olefins, but to the different number and reactivity of C-H bonds in both alkanes. A discussion on the importance of the concentration of active sites and the characteristics of the alkanes fed on the selectivity to olefin during the alkane ODH is also presented.

Keywords: propane; ethane; oxidative dehydrogenation; olefins; promoters; nickel oxide.
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INTRODUCTION

Short chain olefins, especially ethylene and propylene, are undoubtedly the most important raw materials for the petrochemical industry, and perhaps also for the chemical industry in general [1]. These olefins constitute the fundamental pieces from which a multitude of chemical compounds are obtained.

Propylene is a highly versatile compound since can be obtained through different ways and can be transformed into a large number of industrial products. It is mainly obtained by steam cracking or fluid catalytic cracking (FCC) [2], but also by propane dehydrogenation (PDH) [3] and metathesis [4]. Currently, the new trends in steam cracking, especially in US, indicate a growth in the use of ethane (obtained from natural gas) as a feedstock at the expense of naphtha. The use of ethane implies a higher ethylene production and a sharp diminution of propylene production. Therefore, in order to cope with this increasing demand of propylene (ca. 10% in US in the past 3 years) [5], other “on-purpose propylene” (OPP) technologies, such as the non-oxidative dehydrogenation of propane, olefin metathesis and methanol to olefins (MTO) processes, have been industrially implemented [6].

The oxidative dehydrogenation (ODH) of propane could be an interesting alternative to steam cracking and non-oxidative dehydrogenation of propane, since from an energetic point of view, ODH is a much more favorable process, due to its exothermic character [7]. Besides, the presence of molecular oxygen in the feed permits an in situ regeneration of the active sites and prevents coke formation [8].

In terms of selectivity to propylene, one of the most efficient materials reported for the ODH of propane are supported vanadium oxide catalysts (ca. 90 % selectivity to propylene at low propane conversion) [8-12]. Unfortunately, yields to propylene described in literature are generally lower than 30%, due to olefin deep oxidation into
carbon oxides at increasing propane conversions [9]. Importantly, the acid-basic character of the support has to be controlled, so that basic supports are preferred to favor the propylene desorption [12].

Supported vanadium oxide catalysts were reported to be also efficient for ODH of ethane to ethylene [13-16], although they present the same disadvantage, i.e. the olefin formed readily decomposes by deep oxidation. In contrast with the optimal catalysts in the ODH of propane, optimal supports for the ODH of ethane present a rather acidic character [12-16].

Interestingly, the use of optimized promoted NiO catalysts [17-22] in the ODH of ethane has allowed the minimization of the ethylene decomposition into carbon oxides, this way achieving high olefin yields (> 40%), especially after the first publication by Lemonidou et al. [17]. However, not every promoter improves the catalytic performance of nickel oxide, the nature of the dopant element being of paramount importance. Basic promoters lead to the predominant formation of carbon oxides, whereas those with an acidic character show high selectivity to ethylene [18].

Unlike vanadium oxide catalysts [8-16], there are very few studies related to propane ODH over NiO-based catalysts [23-28]. Moreover, reports comparing the catalytic performance in ethane and propane ODH for NiO-based catalysts are even more scarce [28, 29].

Overall, most of the works published with nickel oxide catalysts use cerium as a promoter due to its high oxygen storage capacity and concentration of oxygen vacancies. In this way, several authors propose Ni-Ce-O catalysts as relatively selective in ODH of propane [23-26]. The best catalytic results were obtained by a catalyst with an excess of cerium (Ni/Ce atomic ratio of 0.5) and the authors ascribed the optimal results to the presence of highly dispersed nickel oxide species and a high concentration of anionic vacancies.
Nevertheless, the study of the influence of the characteristics of dopants in promoted NiO catalysts has not been undertaken so far.

In addition, the selectivity to propylene during propane ODH over NiO-based catalysts is lower than that achieved during the ethane ODH. However, it is not clear what are the key aspects for ethane ODH that are not met for the propane ODH.

In the present article, we want to study if this positive behavior observed in the ODH of ethane with NiO-based materials can also take place in the ODH of propane. Thus, the objective of this work is: i) to analyze if a similar improvement in the selectivity to the olefin takes place when NiO is modified by the addition of specific promoters [18]; ii) to determine, for propane ODH, the nature of the optimal dopants; and, iii) to compare the reactivity of C$_2$ and C$_3$ alkanes and their corresponding olefins (ethylene and propylene), in order to evaluate the contribution of consecutive reactions to the selectivity profiles.

For this purpose, promoted NiO catalysts, considering a set of dopants with different physicochemical characteristics and acidities (i.e. K-, Ce-, Zr-, La-, Nb-, Sn-promoted catalysts) have been prepared, characterized and tested in the ODH of propane. For comparison the same catalysts have been tested in the ODH of ethane. The results are discussed in terms of the effect of promotors on the chemical nature of surface oxygen sites, and the derived catalytic consequences.

2. EXPERIMENTAL

2.1. Preparation of catalysts

Promoted NiO catalysts have been prepared through the evaporation of an ethanolic solution of nickel nitrate, a salt of the promoter, and oxalic acid using a (Ni+M)/oxalic acid molar ratio of 1 (the M-salts employed are shown in supporting
The paste obtained was dried in a furnace at 120ºC for 12 h and then calcined in air at 500ºC for 2 h. The characteristics of the catalysts are shown in Table 1.

The samples have been called as M-NiO, Me being the promoter (K⁺, Al³⁺, Ce³⁺, La³⁺, Zr⁴⁺, Sn⁴⁺ or Nb⁵⁺). The amount of promoter has been fixed in M/(M + Ni) atomic ratio of 0.08, as this is close to the optimal ratio in promoted NiO catalysts for ethane ODH [18]. For comparison, an unpromoted nickel oxide (i.e. uNiO) was also studied.

2.2. Characterization techniques

Catalyst surface areas were determined by multi-point N₂ adsorption at 77 K, in a Micromeritics ASAP 2000 instrument. The data were treated in accordance with the BET method.

Average chemical composition of catalysts was determined by inductively coupled plasma atomic emission spectroscopy (ICP-AES).

Powder X-ray diffraction (XRD) was used to identify the crystalline phases present in the catalysts. An Enraf Nonius FR590 sealed tube diffractometer, with a monochromatic CuKα₁ source operated at 40 kV and 30 mA was used. The particle sizes have estimated from XRD data using the Scherrer formula. This formula has been applied to peaks related to (101), (012), (113), and (024) crystal planes of NiO.

Raman spectra were collected in an inVia Renishaw spectrometer equipped with an Olympus microscope. The measurements were carried out at different wavelengths: 325 (UV-Raman) and 514 nm (Visible-Raman).

IR spectra of adsorbed CO were recorded at low temperature (-176 ºC) with a Nexus 8700 FTIR spectrometer, at a spectral resolution of 4 cm⁻¹, using a DTGS detector. An IR cell allowing in situ treatments in controlled atmospheres and temperatures from -176 ºC to 500 ºC has been connected to a vacuum system with gas dosing facility. Self-
supporting wafers were initially pre-treated at 250 °C/1.5 h (in an oxygen flow, 20 ml min⁻¹), followed by evacuation at 300°C/1h (at 10⁻⁴ mbar), and cooled down to -176°C (under dynamic vacuum). Finally, CO was adsorbed by CO dosing at increasing pressure (0.4-8.5 mbar), and IR spectra were recorded after each dosage.

X-ray photoelectron spectra were collected using a SPECS spectrometer with a MCD-9 detector and using a non-monochromatic AlKa (1486.6 eV) X-ray source. Spectra were recorded using analyzer pass energy of 50 eV, an X-ray power of 200 W, and under an operating pressure of 10⁻⁹ mbar. During data processing of the XPS spectra, binding energy (BE) values were referenced to C1s peak (284.5 eV). Spectra treatment has been performed using the CASA software.

2.3. Catalytic tests

The catalytic experiments were carried out at atmospheric pressure in a tubular isothermal fixed bed reactor, at atmospheric pressure, working in the 250-350 °C temperature range. Catalyst sample (0.3-0.5 mm particle size) were introduced in a quartz reactor diluted with silicon carbide in order to keep a constant volume. Typically, 0.1 g of catalyst and a flow rate of reactants of 50 ml/min were used, although both the catalyst amounts loaded and the total flows used were varied to achieve different hydrocarbons conversions at a fixed reaction temperature. The feed corresponds to a mixture consisting of C₃/O₂/He with a molar ratio of 10/10/80. For comparison, some experiments were undertaken in the oxidation of ethane, propylene or ethylene using a feed consisting of a mixture of hydrocarbon/O₂/He with a molar ratio of 10/10/80. Reactants and reaction products have been analysed using a gas chromatograph equipped with two packed columns: (i) molecular sieve 5 Å (2.5 m); and (ii) Porapak Q (3 m). Olefins (propylene or ethylene), water and carbon dioxide were the main reaction products detected from alkanes (propane
or ethane, respectively). CO could be identified in a couple of catalysts but with selectivity lower than 0.2%. Blank runs in the absence of catalyst were carried out until 450°C. Zero conversion was observed in both C₂-C₃ alkane ODH and C₂-C₃ olefin oxidation in blank experiments.

3. RESULTS AND DISCUSSION

3.1 Catalysts characterization

Table 1 shows some physicochemical and catalytic properties for propane oxidation of unpromoted (uNiO sample) and M-promoted nickel oxide catalysts (M-NiO; M= K, La, Ce, Al, Zr, Sn, Nb). XRD patterns of the catalysts show the only presence of cubic phase of NiO (JCPDS: 47-1049), although low intensity lines corresponding to other phases were observed in the case of Ce-, La- and Zr-promoted materials (Fig. S1, supporting information). On the other hand, the incorporation of promoters to NiO leads to a decrease of NiO crystallite size, and a subsequent increase in catalyst surface area (Table 1). This way, uNiO sample presents a mean crystallite size of NiO of 35 nm, while for promoted catalysts NiO crystallite size ranges from 9 to 17 nm.

The series of promoted NiO catalysts was characterized by Raman spectroscopy (Fig. 1). Figure 1A shows Raman spectra collected at an irradiation wavelength of 514 nm. Unpromoted NiO shows the characteristic vibrational features of nickel oxide, displaying a band centered at ca. 495 cm⁻¹ (Fig. 1A, spectrum a) [30]. Interestingly, depending on the nature of the promoter, two different trends are observed in the Raman spectra. Except for K⁺-promoted NiO catalysts (Fig. 1A, spectrum b), which shows a similar Raman profile than uNiO, band-broadening is observed for all promoted NiO catalysts (Fig. 1A, spectra c to d). This fact suggests that one common effect of adding high oxidation state dopants is to promote a loss of order in NiO, likely due to the presence
of defects. In addition, a shift to higher frequencies is observed for Ni-O modes in some promoted NiO catalysts (from 495 cm$^{-1}$ to 510-589 cm$^{-1}$, for Ce-, Sn-, Zr- and Nb-promoted materials), indicating changes in the Ni-O bond nature, likely due to the partial incorporation of $M^{m+}$ species within NiO crystal lattice (Fig. 1A, spectra d to g).

**Figure 1**

Figure 1. Raman spectra of NiO-based catalysts recorded at an irradiation wavelength of 514 nm (A) and 325 nm (B). Samples: a) uNiO; b) K-NiO; c) La-NiO; d) Ce-NiO; e) Zr-NiO; f) Sn-NiO; g) Nb-NiO. Characteristics of catalysts in Table 1.

In order to shed some light on the vibrational characteristics of the catalysts, the materials were further analyzed by UV Raman, by irradiating the samples with a UV-laser (i.e. 325 nm) (Fig 1B). This technique has been reported to be more sensitive to the surface features of solid materials, thus being more suitable for the study of heterogeneous catalysts [31-33]. All the catalysts show typical UV Raman features of NiO: one-phonon transversal optical (TO) and longitudinal optical (LO) modes at ca. 568 cm$^{-1}$ (1P); and two-phonon modes 2TO (at ca. 725 cm$^{-1}$), TO+LO (at ca. 897 cm$^{-1}$) and 2LO (at ca. 1110 cm$^{-1}$) (Fig. 1B) [34]. The relative intensity of these signals can provide information on the structural disorder and particle size of NiO. The lowest relative intensity of defect-induced one-phonon signal (1P, at ca. 568 cm$^{-1}$) is observed for unpromoted NiO and K$^+$-promoted catalysts (Fig. 1B, spectra a and b). On the contrary, NiO catalysts promoted with elements showing higher oxidation states (La$^{3+}$, Ce$^{3+/4+}$, Zr$^{4+}$ or Nb$^{5+}$) present a substantially higher intensity of this 1P mode (Fig. 1B, spectra c to g). This increase in the relative intensity of one-phonon mode observed by UV Raman can be explained in terms of the following features induced by the incorporation of promoter: i) a smaller NiO
particle size [35]; ii) the presence of a higher concentration of defects [36]; and, iii) a higher structural disorder [37].

FTIR of adsorbed CO have been also undertaken in order to study the Lewis acidity of the surface of the catalysts [38], in a manner that strong acid sites arise at high frequencies, whereas weak acid sites arise at low frequencies. Taking this into account, and according to the results of Table 1 (see also Fig. S2 in supporting information), the strength of acid sites in our catalysts increases in the order [18]: K-NiO << uNiO < La-NiO < Al-NiO < Zr-NiO < Sn-NiO < Nb-NiO. Similarly, the number of acid sites follows the order: Nb-NiO > Sn-NiO > Zr-NiO > La-NiO > Al-NiO > NiO, K-NiO.

A detailed XPS study was also conducted in order to get insights into surface features of NiO-based catalysts. As electrophilic O$_2^-$ and O$^-$ oxygen species can be directly involved in total oxidation reaction pathways, the nature of surface oxygen sites has been studied (Table 1). Figure 2 shows O1s core-level XPS spectra of promoted NiO catalysts. All the spectra can be fitted to three single peaks, centered at ca. 528, 531 and 533 eV, which can be assigned to lattice O$_2^-$, OH$^-$, and electrophilic (O$_2^-$ and O$^-$) oxygen species, respectively [39].

**Figure 2**

*Figure 2.* O1s XPS spectra of catalysts: a) uNiO; b) K-NiO; c) La-NiO; d) Ce-NiO; e) Zr-NiO; f) Sn-NiO; g) Nb-NiO.

Interestingly, differences in the relative intensity of these signals are clearly observed depending on the nature of the promoter. For instance, the incorporation of a low oxidation state promoter like K$^+$ leads to an increase in the relative intensity of the signal assigned to electrophilic O$^-$ species (Fig. 2, spectrum b) with respect to undoped
uNiO catalyst (Fig. 2, spectrum a). These electrophilic sites are those reported to be responsible for total oxidation of alkanes [40, 41]. However, the relative intensity of this signal decreases notably when metal cations with an oxidation state higher than $2^+$ are incorporated (i.e. Nb$^{5+}$, Zr$^{4+}$, La$^{3+}$, etc.) (Fig. 2, spectra c to g). The surface concentration of O$^-$ sites on promoted nickel oxide catalysts has been estimated (Table 1). Then it has been observed that the concentration of surface electrophilic O$^-$ sites decreases from ca. 22.9 at.% in K-NiO catalyst down to 6.6 at.% in Nb-NiO catalyst.

3.2. Catalytic results in the oxidative dehydrogenation of propane

The main reaction products observed in the oxidation of propane using NiO-based catalysts have been propylene and CO$_2$. Partially oxygenated hydrocarbons were not detected and CO was only observed as traces.

Figure 3 shows comparatively propane conversion and selectivity to propylene achieved during the ODH of propane for the different NiO-based catalysts at 325ºC. For comparison, the selectivity to propylene achieved with promoted NiO catalysts at a propane conversion of 10% is shown in Table 1. The most active catalyst was that with cerium, whereas that with potassium achieved the lowest activity. Thus, the order of catalytic activity observed is as follows: Ce-NiO > La-NiO > Zr-NiO > Al-NiO > uNiO > Nb-NiO > Sn-NiO >> K-NiO. Moreover, the selectivity to propylene follows a different trend: Nb-NiO > Zr-NiO, Sn-NiO > Al-NiO, La-NiO > Ce-NiO >> uNiO > K-NiO. Interestingly, promoted NiO catalysts present a selectivity to propylene higher than undoped (uNiO) or K-doped (K-NiO) catalysts; with Nb-promoted NiO presenting the highest values of selectivity to propylene (ca. 45%), whereas uNiO and K-NiO samples present the lowest ones (16 and 11% respectively).
3.3. Relationship between the nature of the promoter and the characteristics of the catalysts with the selectivity to propylene.

The acid characteristics of the NiO-based catalysts can have an important influence on the adsorption and desorption of reactants and reaction products, and therefore on the catalytic performance. Thus, with the aim of understanding the catalytic results over promoted catalysts, we have attempted to correlate the selectivity to propylene with the acid characteristics of the catalysts (Fig. 4), considering the results achieved by FTIR from adsorbed CO (Table 1 and Fig. S2). According to these results, the higher the amount and strength of Lewis acid sites on the NiO-catalysts, the higher the selectivity to propylene (obtained in the ODH of propane) (Fig. 4).

We must inform that a similar trend was observed during the ODH of propane and ethane on similar catalysts [18]. However, in the case of the ODH of ethane the selectivity to the olefin was substantially higher than in the ODH of propane (Fig. S3). Then, a selectivity to ethylene up to 85-90 % over Nb-NiO and Sn-NiO catalysts have been achieved.
Figure 5 displays the variation of the selectivity to the olefin during the ODH of propane or ethane as a function of the surface concentration of electrophilic O\textsuperscript{-} sites. A drop of the selectivity to the olefin is observed when the amount of electrophilic sites increases. Then, the proportion of electrophilic sites for the least selective catalyst (i.e. K-NiO) is 23\% whereas for the most selective catalyst (i.e. Nb-NiO) is only of 6.6 at.\%.

This fact suggests that it is possible to eliminate non-selective oxygen sites on NiO by the incorporation of promoters. Interestingly, such a NiO-promoter interaction can take place either by the incorporation of the promoter within NiO framework (as in Nb-NiO) [17-22, 42], or by intimate contact between separated oxide phases (as in Sn-NiO) [42].

**Figure 5**

**Figure 5.** Influence of amount of electrophilic O\textsuperscript{-} sites (determined by XPS) on the selectivity to propylene, during the ODH of propane (○), or on the selectivity to ethylene, during the ODH of ethane (●). Data at 10\% alkane conversion and a reaction temperature of 350ºC. Catalytic results for propane ODH (this study), using a C\textsubscript{3}/O\textsubscript{2}/He molar ratio of 10/10/80; Catalytic results for ethane ODH (from ref. 18), using a C\textsubscript{2}/O\textsubscript{2}/He molar ratio of 10/3.3/86.7.

It can be stated that, for NiO-based catalysts, high valence promoters (like Nb\textsuperscript{5+}, Sn\textsuperscript{4+} or Zr\textsuperscript{4+}) [17-20, 42-45], or the use of supports [46, 47], tend to remove unselective electrophilic O\textsuperscript{-} species. Thus, the highest selectivity to propylene from propane (but also ethylene from ethane) is observed with catalysts displaying the lowest amount of electrophilic species. These results go in line with previous reports on ethane ODH, which show that the selectivity to ethylene increases concomitantly with a decrease in the number of Ni neighbours in the first (Ni-O) and second coordination shell (Ni-Ni) for supported NiO catalysts [46]. In addition, it has been also suggested that the incorporation of promoters and supports/diluters leads to a decrease of the average oxidation state of Ni\textsuperscript{a+} sites, and to a decrease in the concentration of non-selective electrophilic oxygen
sites on the surface, likely to occur due to the elimination of non-stoichiometric Ni$^{3+}$ surface sites [42].

In this way, it can be interesting to indicate that the lowest relative intensity of defect-induced one-phonon (1P mode) signal determined by UV Raman is observed for unpromoted NiO and K$^+$-promoted catalysts (Fig. 1B, spectra a and b), which are those showing the lowest selectivity in ODH. On the contrary, the higher intensity of this 1P mode in the UV Raman (Fig. 1B, spectra c to g) is observed in NiO catalysts promoted with elements showing higher oxidation states (La$^{3+}$, Ce$^{3+/4+}$, Zr$^{4+}$ or Nb$^{5+}$), which present a higher selectivity to olefins in ODH reactions.

3.4. On the reactivity of alkanes and olefins in oxidation reactions over NiO-based catalysts

In the case of propane oxidation, only propylene and CO$_2$ are produced (Tables S1-S3), which can be considered as primary reaction products. This behaviour is similar than that observed for ethane oxidation in which ethylene and CO$_2$ are the main reaction products. CO was observed only as traces in both cases.

For this reason, and in order to evaluate the contribution of consecutive reactions, Figure 6 shows the variation of the selectivity to olefin during the ODH of propane (Fig. 6A) and ethane (Fig. 6B) at 350ºC for some representative catalysts, i.e. Nb- and Zr-promoted and unpromoted NiO catalysts (fixing the reaction temperature and modifying the contact time).

**Figure 6**
The selectivity to ethylene (during ethane ODH) is higher than the selectivity to propylene achieved during propane ODH. However, in both cases, a low influence of the alkane conversion on the selectivity to the corresponding olefin is observed over the studied catalysts. Thus, the consecutive deep oxidation of the olefins should be very low with respect to the formation of olefins during the ODH reaction. This behavior is completely different to that achieved over supported vanadium oxide catalysts [12-16, 48-50], in which it is well known that the alkane conversion has a strong influence on the selectivity to olefins.

To confirm this trend, a comparative study on the oxidation of C2 and C3 alkanes and their corresponding olefins (i.e. ethylene and propylene) has been carried out on three selected catalysts, i.e. uNiO, Zr-NiO and Nb-NiO, at the same reaction conditions (350°C, using a hydrocarbon/oxygen/helium molar ratio of 10/10/80 and a contact time, W/F, of 8.2 g_{cat} h/(mol_{hydroc})^{-1}) (Tables S1-S3). Figure 7 presents the reaction rate for alkane (r_{C2H6} or r_{C3H8}) or olefin (r_{C2H4} or r_{C3H6}) conversion as well as the reaction rate for the formation of the main reaction products (r_{CO2}, r_{CO2}, r_{CO2} and r_{CO}). In all cases, the reaction rate for hydrocarbon conversion decreases according to: Zr-NiO > uNiO > Nb-NiO. However, when considering the specific reaction rate (i.e. considering the surface area of catalysts) for ethane and propane oxidation, the catalytic activity decreases according to: uNiO > Zr-NiO > Nb-NiO. In addition, there exist common features in the reactivity when comparing the reaction rate for the conversion of the alkane and alkene for C2 and C3 substrates. Thus, in all cases, the reaction rate for the olefin transformation is lower than that for the corresponding alkane, being this difference greater in the case of C2 hydrocarbons than for C3 hydrocarbons. Indeed, the relative reaction rates for
ethylene/ethane is ca 0.5 (more specifically, 0.51 for uNiO; 0.53 for Zr-NiO and 0.42 for Nb-NiO), whereas the relative reaction rates for propylene/propane it is ca. 0.70 (more specifically, 0.69 for uNiO; 0.64 for Zr-NiO and 0.80 for Nb-NiO) (see Table 2). These values are very different to those reported for supported vanadium oxide catalysts, in which the ratio between olefin/alkane oxidation is between 4 to 10 [16]. Thus, the results presented here confirm the low influence of alkane conversion on the selectivity to olefin during ODH of ethane or propane on NiO-based catalysts. In addition, regardless of the catalyst (see Fig. 7, for C$_2$H$_4$ and C$_3$H$_6$), the activity for propylene oxidation is slightly higher than for ethylene oxidation.

On the other hand, no major differences are observed between the reaction rates for alkanes and olefins oxidation when comparing more selective (such as Nb-or Zr-promoted NiO) or less selective (unpromoted NiO) catalysts for alkane ODH. This suggests that: i) the differences in selectivity to olefins during the ODH of ethane and propane depend largely on the characteristics of the fed alkane and not on the reactivity of the olefins; and ii) the presence or absence of promoted in NiO-based catalysts has a very low influence on the deep oxidation of olefins.

**Figure 7**

**Figure 7.** Reaction rate for hydrocarbon conversion (r$_{C2H6}$, r$_{C3H8}$, r$_{C2H4}$ or r$_{C3H6}$) as well as the reaction rate for the formation of the main reaction products (r$_{C3H6}$ or r$_{C2H4}$, r$_{CO2}$ and r$_{CO}$) during the oxidation of ethane (a), ethylene (b), propane (c), and propylene (d) over NiO, Zr-NiO and Nb-NiO catalysts at 350ºC. Experimental conditions as in table S1-S3).

Accordingly, the differences observed for undoped and doped NiO catalysts during the propane and ethane ODH are much more related to changes in the ratio between the olefin formation and the CO$_2$ formation from each alkane (which probably is related to the
different C-H and C-C bond strengths in both alkanes) rather than changes in the olefin deep oxidation over each catalyst.

**General remarks and comparison between propane and ethane ODH**

The nature of the promoter highly affects the catalytic performance in the ODH of propane over NiO-based catalysts (in a parallel trend to that observed for ethane [18]). As observed previously for ethane ODH [17-22, 43-47], those promoters with high oxidation state and high acidity lead to the best catalytic results as this minimizes the presence of surface electrophilic O$_2^-$ and O$^-$ oxygen species [43], especially when using Nb$^{5+}$ as promoter [17, 19, 42-45].

In our case, niobium promoted NiO sample resulted to be the most efficient catalyst. The positive role of acidity in promoted NiO catalysts, especially for propane ODH, contrasts with the results observed for supported vanadium oxide catalysts in which the presence of acid sites has detrimental influence for obtaining high propylene yields [12, 16]. In fact, among vanadium based catalysts, K-doped VO$_x$/Al$_2$O$_3$ [12, 48] or V-Mg-O [49] have been reported as selective in propane ODH, whereas VO$_x$/Al$_2$O$_3$ [13,14, 48, 49] is more selective in ethane ODH.

Accordingly, the selectivity to olefins during C$_2$-C$_3$ alkane ODH seems to be mainly related to the nature of the active sites (and in the ratio between electrophilic and nucleophilic oxygen species), whereas the deep oxidation of olefins on NiO-based catalysts seems to be very low (and similar independently of the presence or absence of promoters) as concluded from results in Figure 7.

However, the selectivity to ethylene during the ethane ODH is remarkably higher than that observed for the selectivity to propylene during propane oxidation (Figures 5 and 6). Thus, on Nb-NiO catalyst, the selectivity to the olefin in the ODH of ethane is
over 80%, whereas in the ODH of propane is only of ca. 45%. Similarly, over unpromoted NiO catalyst the selectivity to ethylene is 40-45%, whereas that to propylene is ca. 16%.

In agreement to previous results [17-22, 43-47], a general reaction scheme for the oxidative dehydrogenation of propane and ethane on nickel oxide catalysts involves the direct transformation of propane or ethane into propylene or ethylene but also the direct formation of CO₂ from either propane or ethane (Fig. 8). Thus, according to the results shown in Fig. 7 and Table 2, the selectivity to olefins strongly depends on the $k_2/k_1$ ratio (i.e. parallel reactions which strongly depend on both the characteristics of catalyst an the alkane feed). However, in all cases, the consecutive reactions (with low $k_3/k_1$ ratio) has a low importance on the selectivity to olefins. On the other hand, the formation of CO is negligible from alkanes and very low from olefins.

A last question to be considered is the fact that the selectivity to olefin during the ODH of ethane is higher than that achieved during ODH of propane, when relatively small differences in the $k_3/k_1$ ratio are concluded. In fact, it has been observed that the ratio between the reaction rate for olefin and alkane oxidation is ca. 0.5 ($r_{-C_2H_4}/r_{-C_2H_6}$) and 0.7 ($r_{-C_3H_6}/r_{-C_3H_8}$) (Table 2), suggesting a low influence of the deep oxidation of olefins during the alkane ODH in both unpromoted and promoted NiO-based catalysts.

**Figure 8**

**Figure 8.** Reaction pattern of the ODH of propane and ethane on NiO based catalysts.

In fact, the reaction rate for deep oxidation of propylene would be expected to be greater than that of ethylene, since the stability of propylene (presenting weak C-H allylic bonds) is supposed to be lower than that of ethylene (presenting stronger C-H vinylic bonds). [51]. In addition, the adsorption constant of ethylene on NiO is considerably lower than
that of propylene [52, 53], which apparently would decrease the extent of the ethylene decomposition. However, and although this can be the case of the ODH over V-containing catalysts, this does not take place for NiO-based catalysts. Thus, unexpectedly, the higher olefin yields achieved in the ODH of ethane with respect to those achieved during propane ODH are not due to the differences in the stability of the olefins (since in both cases the reaction rate for olefin oxidation is lower than that for alkane oxidation), but to the tendency of propane to be more easily transformed into carbon dioxide.

Yao and Kreamer in 1973 [28] reported a catalytic behavior for the oxidation of C₂-C₄ hydrocarbons on pure NiO catalysts similar to the trend presented here. These authors observed that, unexpectedly, the catalytic activity for alkane oxidation was higher than for olefin oxidation. This is especially surprising in the case of propylene since, despite showing weaker allylic C-H bonds, it is less reactive than propane (with stronger primary and secondary C-H bonds), and even less reactive than a stable molecule as ethane. The interesting results presented here verify that this trend occurs not only for the unselective NiO (with low selectivity to olefins during the ODH of ethane and propane), but also on promoted NiO catalysts. Thus, NiO catalysts promoted with Nb⁵⁺ or with Zr⁴⁺ present high selectivity to olefin, especially during the ODH of ethane, and to a less extent during the ODH of propane, but both undoped and doped NiO catalysts present similar reaction rates for deep oxidation of C₂-C₃ olefins (Fig. 7, b and d).

The results obtained here suggest that the selective sites for ethane or propane ODH hardly intervene (their contributions is very low) in the combustion of olefins. Therefore, and in agreement with previous results [17, 42, 43], there must be two types of centers involved in the alkane ODH: i) selective sites for olefin formation, which are strongly related to the presence of the promoter (especially in the case of Nb); and ii) non-selective sites, favoring the formation of CO₂, which are majority in the case of pure NiO.
On the other hand, and in agreement to previous reported mechanism, the alkane ODH involves three different steps: i) for ethane ODH, in which the scission of C-H bond is the rate-determining step [43]; ii) ethane deep oxidation, in which the rate-determining step may be either the adsorption of ethane as C₂H₅ radical with breaking of the C-H bond if the slow step is the chemisorption of the ethane molecules, or the interaction between the adsorbed O₂ and either gas phase or physically adsorbed C₂H₆ through attack at the C-H bond [28]; iii) ethylene deep oxidation, minority on NiO-based catalysts, in which the rate of adsorption of ethylene or reaction between oxygen and the double bond of ethylene (adsorbed on the surface, with the C₂H₄ molecules adsorbed on the surface through the π bond) may be the rate-determining step [28].

Finally, the lower selectivity to propylene over promoted catalysts, when comparing with the catalytic results for ethane ODH, could be related to the different number and dissociation energies of C-H bonds present in ethane and propane molecules, as suggested by Iglesia et al. [14].

In the present article, it has been clearly observed for both ethane and propane ODH that the selectivity to carbon dioxide increases with the surface concentration of O- electrophilic sites. As the selectivity to olefins hardly falls with the alkane conversion, it indicates that these unselective sites are more reactive in the alkane oxidation than in the olefin oxidation. In the case of Nb-NiO catalyst 6.6% of the surface oxygen is O- electrophilic sites, which leads to an initial selectivity to CO₂ of 10-15% in the ODH of ethane and 45-50% in the ODH of propane. In the case of Zr-NiO catalyst 11% of O- surface electrophilic sites lead to an initial selectivity to CO₂ of 25-30% and ca. 60% for ODH of ethane and propane respectively. In the unpromoted NiO catalyst 16% of surface electrophilic sites lead to an initial selectivity to CO₂ of 45-50% and ca. 85% for ODH of ethane and propane respectively. All these data indicate that in all catalysts for each 1%
of O’ sites the selectivity to CO₂ increases by 2-3% in the ODH of ethane whereas increases by 5-7% in the ODH of propane. Accordingly, the combustion of propane takes place at a higher reaction rate than the combustion of ethane in the O’ surface electrophilic sites. It is possible that the higher size and the greater number of C-H bonds in propane (compared to ethane) gives rise to a better attack of the unselective O’ species to the propane molecule, with the consequent higher formation of CO₂.

Interestingly, the formation of CO is not detected, neither from ethane nor from propane, and this is likely related to the high reaction rate of the CO oxidation to CO₂ on nickel oxide based catalysts, which could take place at temperatures below 150°C [43], which are significantly lower than those required for the ODH of propane. Moreover, CO is usually a reaction product obtained as a result of the decomposition of the olefin and, for NiO-based catalysts, this step seems to be negligible, at least up to 20% alkane conversion.

Conclusions

Oxidative dehydrogenation of propane to propylene can be undertaken using promoted NiO catalysts achieving propylene yields of ca. 10% at low reaction temperatures (in the 300-350°C range). The catalytic behaviour of unpromoted NiO is poor but can be improved, as well as the physicochemical properties, if NiO is modified by the incorporation of a second metal oxide.

Overall, the oxidation state of the promoter determine the interaction NiO-promoter affecting the size of the NiO particles and the acidity of the catalyst, and importantly the concentration of surface O’ electrophilic species. A clear inverse relationship has been observed between the selectivity to the olefin (in both cases from propane or from ethane) and the concentration of surface O’ electrophilic species. Then, the higher the amount of
these unselective O species the highest is the extent of the direct alkane combustion whereas the extent of the olefin decomposition is hardly affected.

The optimal catalytic results have been obtained by the catalysts in which the promoter metal presents high valence, as Nb$^{5+}$. Generally speaking, selective catalysts present tiny NiO crystallites and high acidity although the most determining factor seems to be the removal of unselective electrophilic O’ species. A link of all these characteristics is very likely. Nb-promoted NiO catalysts resulted to be the most efficient for the ODH of propane whereas the unpromoted NiO and K-doped NiO are the least selective to propylene.

On the other hand, a parallelism between the selectivity to ethylene observed in the ODH of ethane and the selectivity to propylene in the ODH of propane has been observed (the electrophilic O’ species are related to the direct combustion of the alkane), although the selectivity to ethylene achieved is significantly higher than the selectivity to propylene. However, the reaction rate for olefin combustion (ethylene or propylene) is lower than the reaction rate for alkane transformation, with the consequent high stability of olefins during the alkane ODH.

The lower selectivity to propylene (referred to that achieved for ethylene) cannot be explained by the different reactivity of olefins (very low in both olefins when comparing to the corresponding alkanes). Thus, the greater number of C-H bond in propane could facilitate a better attack of unselective oxygen species, favouring a higher formation of carbon dioxide.

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Table 1. Physicochemical and catalytic properties of NiO containing catalysts.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>$S_{\text{BET}}$ (m$^2$ g$^{-1}$)</th>
<th>NiO crystal size (nm)$^a$</th>
<th>Concentration of O$^-$ (from XPS) at.%</th>
<th>FTIR of adsorbed CO$^b$</th>
<th>Catalytic activity $^c$</th>
<th>Selectivity to propylene (%) $^d$</th>
</tr>
</thead>
<tbody>
<tr>
<td>uNiO</td>
<td>15</td>
<td>35</td>
<td>15.8</td>
<td>2148</td>
<td>87</td>
<td>16</td>
</tr>
<tr>
<td>K-NiO</td>
<td>30</td>
<td>17</td>
<td>22.9</td>
<td>-</td>
<td>8.7</td>
<td>11$^e$</td>
</tr>
<tr>
<td>La-NiO</td>
<td>79</td>
<td>11</td>
<td>14.8</td>
<td>2151</td>
<td>251</td>
<td>28</td>
</tr>
<tr>
<td>Ce-NiO</td>
<td>84</td>
<td>15</td>
<td>11.8</td>
<td>n.d.</td>
<td>923</td>
<td>23</td>
</tr>
<tr>
<td>Zr-NiO</td>
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<td>9</td>
<td>10.5</td>
<td>2155</td>
<td>154</td>
<td>37</td>
</tr>
<tr>
<td>Sn-NiO</td>
<td>103</td>
<td>12</td>
<td>10.7</td>
<td>2165</td>
<td>35</td>
<td>36</td>
</tr>
<tr>
<td>Nb-NiO</td>
<td>115</td>
<td>9</td>
<td>6.6</td>
<td>2169</td>
<td>59</td>
<td>45</td>
</tr>
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</table>

$^a$ Mean NiO crystallite sized determined by XRD using the Scherrer equation; $^b$ $\nu$(CO) IR band associated to surface Lewis acid sites with acidity higher than that observed on NiO; $^c$ Catalytic activity in g$_{\text{C}_3\text{H}_8}$ reacted/(kg$_{\text{cat}}$ h), determined at 300°C and using a contact time, W/F, of 8.2 g$_{\text{cat}}$ h/mol$_{\text{C}_3\text{H}_8}$; $^d$ Selectivity to propylene at 300°C and 10% propane conversion (modifying the contact time depending on the catalyst); $^e$ At 3% conversion as higher conversion could not be achieved due to the low activity of this catalyst.
Table 2. Reaction rate for the oxidation of alkanes and olefins over NiO-based catalysts at 350°C

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Reaction ratea</th>
<th>r-C2H4/r-C2H6</th>
<th>Reaction ratea</th>
<th>r-C3H6/r-C3H8</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>r-C2H6</td>
<td>r-C2H4</td>
<td>r-C3H8</td>
<td>r-C3H6</td>
</tr>
<tr>
<td>uNiO</td>
<td>111</td>
<td>56.1</td>
<td>0.51</td>
<td>140</td>
</tr>
<tr>
<td>Zr-NiO</td>
<td>134</td>
<td>70.7</td>
<td>0.53</td>
<td>174</td>
</tr>
<tr>
<td>Nb-NiO</td>
<td>117</td>
<td>48.8</td>
<td>0.41</td>
<td>130</td>
</tr>
</tbody>
</table>

a) Reaction rate for hydrocarbon conversion, r_i, in 10^4 molhydroc h^{-1} (g_{cat})^{-1}.

Caption to figures

Figure 1. Raman spectra of NiO-based catalysts recorded at an irradiation wavelength of 514 nm (A) and 325 nm (B): a) NiO; b) K-NiO; c) La-NiO; d) Ce-NiO; e) Zr-NiO; f) Sn-NiO; g) Nb-NiO. Characteristics of catalysts in Table 1.

Figure 2. O1s XPS spectra of catalysts: a) NiO; b) K-NiO; c) La-NiO; d) Ce-NiO; e) Zr-NiO; f) Sn-NiO; g) Nb-NiO.

Figure 3. Propane conversion (■) and selectivity to propylene (■) for unpromoted and promoted NiO catalysts during the ODH of propane. Reaction conditions: 325°C, W/F = 8 g_{cat} h /mol_{C3}. Remaining conditions in the Experimental section. For Ce-NiO catalyst, W/F = 2 g_{cat} h /mol_{C3}. 

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Figure 4. Variation of the selectivity to propylene at 10% conversion and 350°C with the amount of adsorbed CO (FTIR). Note: Adsorbed CO total FTIR is the total number of surface acid sites calculated as the number of CO molecules adsorbed on each site.

Figure 5. Influence of amount of electrophilic O` sites detected by XPS on the selectivity to propylene, during the ODH of propane (○), or on the selectivity to ethylene, during the ODH of ethane (●). Data at 10% alkane conversion and a reaction temperature of 350°C. Catalytic results for propane ODH (this study), using a C3/O2/He molar ratio of 10/10/80; Catalytic results for ethane ODH (from ref. 14), using a C2/O2/He molar ratio of 10/3.3/86.7.

Figure 6. Variation of the selectivity to olefin with the alkane conversion at 350°C during the ODH of propane (A) or the ODH of ethane (B) on Ni-based catalysts: uNiO; Zr-NiO and Nb-NiO. Remaining reaction conditions in text.

Figure 7. Reaction rate for hydrocarbon conversion (rC2H6, rC3H8, rC2H4 or rC3H6, in 10^4 molCxHx gcat^{-1} h^{-1}) as well as the reaction rate for the formation of the main reaction products (rC3H6, rC2H4, rCO2 or rCO, in 10^4 molproduct gcat^{-1} h^{-1}) during the oxidation of ethane (a), ethylene (b), propane (c), and propylene (d) over NiO, Zr-NiO and Nb-NiO catalysts at 350°C. Experimental conditions as in Tables S1-S3.

Figure 8. Reaction pattern of the ODH of propane and ethane on NiO based catalysts.
Fig. 1

Fig. 2
Fig. 3
Fig. 4

Fig. 5
Figure 8. Reaction network of the ODH of propane or ethane on NiO based catalysts.