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1 **Bridging the gap between academic and industrial** 2 **hydrocracking: on catalyst and operating conditions effects**

3
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12 **Abstract**

13 This work aims at bridging the knowledge gap between the well-studied Pt/zeolite catalysts and
14 the industrially-employed NiMoS/(Al₂O₃+zeolite) ones. To do so, catalyst series based on HUSY
15 zeolite were evaluated in the hydroconversion of *n*-hexadecane under similar operating
16 conditions, but the industrially-relevant H₂S and NH₃ contaminants were added when evaluating
17 the latter. The intrinsic performance of Pt/HUSY and NiMoS/(Al₂O₃+HUSY) catalysts was
18 noticeably similar, when well-balanced, with the catalytic activity being however much lower in
19 the latter (temperature gap of 120 K). Ammonia inhibition of more than 99% of the protonic sites
20 was revealed to be at the origin of such low activity. On the other hand, the metal-acid balance
21 was only reached at sufficiently low reaction temperatures (ca. 593 K), i.e. when the NH₃
22 inhibiting effect increased the metal to acid sites ratio in the NiMoS/(Al₂O₃+HUSY) to values five
23 orders of magnitude larger than in the Pt/HUSY one. In addition to the understanding achieved
24 on the role of metal-acid balance in industrial-like catalysts and its key controlling parameters,
25 these findings also point to the need of developing better hydrogenating functions to improve
26 the efficiency, and consequently the sustainability, of liquid fuels production.

27 **Keywords:** industrial hydrocracking, metal-acid balance, HUSY zeolite, inhibition.

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1 Introduction

2 The actual energy context as well as recent forecasts ¹ sustain the preeminent role of
3 hydrocracking (HCK) in modern conversion of heavy low-quality feedstocks into lighter high-
4 quality middle distillates fuels, particularly jet fuel, and also base chemicals for industry. This
5 conversion occurs in presence of a bifunctional catalyst which plays thus a key role on the HCK
6 process performances. Therefore, adequate catalyst design and optimization can have major
7 impact in the sustainability of this process.

8 HCK processes are carried out under hydrogen pressure and in the presence of a
9 heterogeneous catalyst. Nowadays, the main feedstock for HCK is straight run distillation
10 vacuum gas oil (VGO) that may be mixed with other heavy feeds ². VGO is mainly composed by
11 aromatic and paraffinic compounds with a wide range of molecular weights. Other elements
12 besides carbon and hydrogen can also be found on VGO. Sulphur and nitrogen are commonly
13 presented in polyaromatic molecules, which can also contain metals in trace amounts ². To
14 remove such impurities, an initial hydrotreating step is required, generating a more
15 hydrogenated stream rich in H₂S and NH₃. In single-stage processes, this stream is directly fed
16 to the hydrocracking reactor without any intermediate separation ³.

17 The HCK catalysts are bifunctional catalysts, i.e. featuring two catalytic functions: a
18 hydrogenation-dehydrogenation (HDH) function and an acidic function ²⁻⁴. In presence of H₂S,
19 the HDH function is commonly granted by a transition metal sulfide (TMS) promoted with other
20 metals (commonly, Mo or W promoted with either Ni or Co), because noble metals, e.g.
21 platinum, are strongly inhibited by sulfur compounds ⁵. The acidic function can be provided by
22 amorphous silica-aluminas or zeolites, in particular the ultrastabilized Y zeolite (HUSY), usually
23 shaped with an alumina binder ^{2,3}. The selection of the acid function is governed by the desired
24 product carbon chain range and process type as well as the nitrogen content in the feed. Zeolite
25 based catalysts show a high activity (owing to high Brønsted acid strength), high
26 thermal/hydrothermal stability, good resistance to poisoning, and low coking rate resulting in a
27 good performance and long catalyst life ^{6,7}.

28 Most of the academic studies have been conducted over zeolite-supported platinum catalysts,
29 which are typically evaluated in the hydroconversion of pure *n*-alkanes. As a result, a panoply of
30 evermore tuneable acidic functions is available nowadays ⁷⁻¹⁰. More importantly, the
31 (bifunctional) reaction mechanism ^{11,12} in these simplified laboratorial bench systems has been
32 understood at its fundamental level. Particularly, for a given acidic function, feedstock, and
33 operating conditions, the catalytic behaviour is known to depend on the balance (i.e. ratio of
34 activity) ^{13,14} and intimacy (i.e. distance) ^{15,16} between the hydrogenating and Brønsted acid
35 sites. For selected feed and reaction conditions, catalyst performances are optimal when
36 catalyst functions are well-balanced and intimacy is adequate ^{11,16,17}. The activity and (e.g.

1 hydroisomerization) selectivity are then essentially dictated by the properties of the acid
2 function, i.e. are intrinsic to the selected acid function.

3 In contrast, studies with industrial-like catalysts and feedstocks, i.e. featuring promoted TMS
4 supported on shaped zeolites and feedstocks replicating industrial ones, are much less
5 frequent. More importantly, the focus is often held on the evaluation of industrial performances
6 ¹⁸, comparing different catalysts ¹⁹⁻²⁴, typically at optimal operating conditions, but rarely
7 providing fundamental insights into the industrial-like systems. The great complexity of industrial
8 feedstocks together with the lack of information (due to confidential clauses) on industrial
9 catalysts further hinders the understanding of these systems. In the same way, only a few
10 studies compare laboratorial and industrial catalysts, but the focus is in any case put on the
11 performances. For instance, the phenanthrene hydroisomerization selectivity over promoted
12 MoS₂/zeolite catalytic system was found to be poorer than Pt/zeolite ones, but the influence of
13 the metal-acid balance was not taken into account ²⁵. The same conclusion was reached via
14 kinetic modelling over a MoS₂/(HBEA+Al₂O₃) catalyst (in this case for *n*-hexadecane), but once
15 again no characterization of the active sites was performed ²⁶.

16 In brief, in-depth rationalization of industrial-like catalysts and their behaviour in hydrocracking is
17 still lacking in the open literature. In addition, the state-of-the-art understanding on
18 hydroconversion is based on investigations at laboratorial conditions. The aim of this work is
19 hence twofold. First, to characterize, at the active site level, and second to rationalize the
20 behavior of industrial-like catalysts in hydroconversion under operating conditions as
21 comparable as possible to those commonly used in industry. The ultimate goal is to unveil the
22 similarities and differences between the poorly-studied industrial-like catalytic systems and the
23 typical laboratorial catalytic systems. As the comprehension of individual phenomenon,
24 particularly at the active site level, under industrial-like conditions is virtually impossible, a
25 reduction of the complexity in the system under study is required. As the focus of this work is
26 hold at the catalytic system, the feedstock will be as simple as possible to be somehow
27 representative of reactions occurring in hydrocracking reactors. The simplest model molecules
28 among the ones present in VGO are long chain *n*-alkanes. A molecule of particular interest is
29 then *n*-hexadecane, as it is been widely studied in literature as well. Moreover, H₂S and NH₃ will
30 be present in the reaction medium, to reproduce of the effect of contaminants in industrial
31 reactors.

32 Concerning the catalysts formulation, HUSY zeolite was selected as acid function as this zeolite
33 is widely used in industrial hydrocracking catalysts. The laboratorial catalyst consist hence of
34 noble metal over HUSY powder labelled Pt/HUSY, which formulation was previously optimized
35 ¹³. The industrial-like catalysts feature a Ni-promoted molybdenum sulphide (NiMoS) phase as
36 HDH function loaded into extrudates of alumina-bound HUSY zeolite, labelled
37 NiMoS/(Al₂O₃+HUSY). A preliminary study on catalysts with different molybdenum loadings was
38 firstly performed and a catalyst was selected based on the observed performances.

1 2 Materials and Methods

2 2.1 Zeolite and catalyst preparation

3 Zeolite HUSY (CBV720) was supplied by *Zeolyst*. The framework was previously identified as of
4 type FAU and the global and framework silicon to aluminium molar ratio were determined to be
5 16.7 and 18.4, respectively ²⁷. Pt/HUSY catalyst was prepared by incipient wetness
6 impregnation of 0.7 wt.% of platinum according to a protocol published elsewhere ²⁷. The
7 impregnated material was dried overnight at 383 K. Calcination under continuous air flow (4 NL
8 h⁻¹ g⁻¹) consisted of three plateaux at 423, 523 and 623 K over 1 h each and a final plateau at
9 773 K over 2 h. Ex-situ reduction under hydrogen flow (4 NL h⁻¹ g⁻¹) included two plateaux: 393
10 K over 1 h and 723 K over 2 h. A heating rate of 5 K min⁻¹ was used in all cases.

11 The shaped support was prepared by mixing-extrusion of HUSY (CBV720) with boehmite
12 employing a procedure published elsewhere ²⁸. The metals were introduced in the calcined
13 support by incipient wetness impregnation with an aqueous solution of H₃PO₄, Ni(OH)₂ and
14 MoO₃. Catalysts with three different Mo contents were prepared, while the nickel and
15 phosphorous to molybdenum (atomic) ratios were kept constant at 0.40 and 0.48, respectively.
16 Further details on the preparation protocols can be found in a previous work ²⁹. After a
17 maturation step, the material was then dried overnight at 493 K and calcined under air flow (1.5
18 NL h⁻¹ g⁻¹) at 723 K over 2 h (heating at 5 K min⁻¹). Ex-situ sulphidation was carried out over a
19 part of the samples in order to characterize the NiMoS phase. The sulphidation took place at
20 623 K over 2 h (heating at 5 K min⁻¹) under a H₂/H₂S flow (15 vol.%) of 1.5 NL h⁻¹ g⁻¹ and the
21 samples were sealed under a N₂ atmosphere. Throughout this article, calcined samples will be
22 denoted NiMo and sulphided samples will be denoted NiMoS.

23 2.2 Materials characterization

24 Both the HUSY zeolite and the Pt/HUSY catalyst have been extensively characterized in
25 preceding works ^{13, 27, 30}. The HUSY shaped support has been also characterized in detail
26 elsewhere ²⁸. Concerning the NiMo-based catalysts, a summary of the techniques employed is
27 provided below, with the focus being held on the determination of the amount of active NiMoS
28 sites. Additional details are provided on the referred works and in section SI.3 of the Supporting
29 Information.

30 Quantitative elemental analysis was performed by x-Ray Fluorescence (XRF). Samples were
31 grinded and sieved into a granulometry under 200 µm. Measurements were carried out directly
32 over powder samples in a Thermofischer Scientific Advant-X instrument. The micrometric
33 elemental distribution was assessed via electron probe micro analysis (EPMA), carried out in a
34 JEOL JXA 8100 equipment. The detailed procedure can be found elsewhere ²⁹ (vide SI.3 for a

1 summary). The MoS₂ slab length was measured by transmission electron microscopy (TEM)²⁹,
2 using a JEOL JEM 2100F microscope operated at 200 kV.

3 X-ray photoelectron spectroscopy (XPS) was carried out in a ESCA KRATOS Axis Ultra
4 spectrometer following a previously published procedure²⁹. The atomic Ni/Mo ratio of MoS₂
5 slabs was calculated as the product of the bulk Ni fraction (by XRF) multiplied by the fraction of
6 Ni atoms involved in the NiMoS phase divided by the product of the bulk Mo fraction (by XRF)
7 multiplied by Mo atoms corresponding to MoS₂. The molar amount of mixed sites per gram of
8 sulfided catalyst (n_{NiMoS}) was calculated based on the weight fraction of Ni in the sample
9 measured by XRF (x_{Ni}) along with the fraction of Ni engaged in the NiMoS sites ($x_{NiNiMoS}$) as
10 estimated by XPS and the atomic mass (AM) of nickel (Eq. 1).

$$n_{NiMoS} = \frac{x_{Ni} \times x_{NiNiMoS}}{AM(Ni)} \quad \text{Equation (1)}$$

11 **2.3 Catalytic testing**

12 The hydroconversion of pure *n*-hexadecane was carried out in a catalytic test unit with a fixed-
13 bed downflow reactor (detailed in SI.1). The catalysts powder was shaped into 0.2-0.35 mm
14 pellets and pre-treated with H₂ (4 NL h⁻¹ g⁻¹ at 4.1 MPa) at 723 K over 1h (heating at 5 K min⁻¹).
15 The tests were conducted at 4.1 MPa of pressure and a H₂ to *n*-C₁₆ molar ratio of 12. The
16 conversion level was changed by either decreasing the reaction temperature in the 533-548 K
17 range or increasing the weight hourly space velocity in the 10-100 g_{nC16} g_{zeolite}⁻¹ h⁻¹ range.

18 The hydroconversion of *n*-hexadecane in presence of H₂S and NH₃ was carried out in two
19 catalytic test units with fixed-bed downflow reactors. The units differed on the operating space
20 velocities range: 1-4 g_{nC16} g_{catalyst}⁻¹ h⁻¹ for the “high-feed” unit and 0.2-1.0 g_{nC16} g_{catalyst}⁻¹ h⁻¹ for
21 the “low-feed” unit. Consequently, the range of reaction temperatures was also different: 613-
22 633 K and 583-613 K, respectively. Once again, both reaction temperature and space velocities
23 were modified to screen a wide conversion range. All employed procedures were similar. The
24 only difference was on the product separation and analysis, as detailed in SI.1. The shaped
25 catalysts were directly loaded into the reactors. The catalysts were sulphided *in situ* at 673 K
26 over 1 h (heating at 5 K min⁻¹) using a feedstock with increased H₂S partial pressure compared
27 to reaction one (vide SI.1) and a volume hourly space velocity of 4.0 mL_{feed} g_{catalyst}⁻¹ h⁻¹.

28 The operating conditions for *n*-hexadecane hydroconversion in presence of H₂S and NH₃ are
29 summarized in Table 1. The catalytic tests were carried out in presence of 11 kPa of H₂S
30 (through decomposition of dimethyl disulphide) and 3.8 kPa of NH₃ (through decomposition of
31 aniline), at 4.1 MPa of total pressure and a molar H₂ to *n*-C₁₆ ratio of 12. These operating
32 conditions are in line with the industrial ones (Table 1). The only exception is the total pressure
33 which was kept lower than the industrial one due to safety reasons.

Table 1: Comparison of the operating conditions used in this work and in industrial hydrocracking processes.

	Pressure (MPa)	Temperature (K)	H ₂ /HC (m ³ /m ³)	LHSV (h ⁻¹)	Sulphur (wt. ppm)	Nitrogen (wt. ppm)
This work	4.1	593-633	1000	0.2-4	10200	750
Industrial HCK ³¹	10-20	623-703	800-2000	0.2-2	1000-20000	300-2000

In the case of *n*-paraffins hydroconversion, it has been repeatedly confirmed that the overall reaction follows a first-order kinetics in the paraffin concentration ^{13, 32, 33}. The catalytic activity was thus calculated as the apparent kinetic constant for first-order consumption of *n*-hexadecane normalized per catalyst mass. The selectivity of a given product was calculated as the amount of carbon atoms from that product divided by total amount of carbon atoms in the products. Yield was calculated as a product of conversion and selectivity.

3 Academic case study: Pt/HUSY in pure *n*-hexadecane hydroconversion

As referred in the Introduction, the effect of metal-acid sites ratio in Pt/HUSY has been extensively study in the hydroconversion of *n*-alkanes. For the catalysts employed in this study, this effect has been even quantified in a previous work, establishing the well-balanced character of catalysts holding 0.4 and 0.7 wt.% of platinum ¹³. Hence, the latter catalyst was selected as the well-balanced Pt/HUSY catalyst to be tested at similar conditions to the NiMo-based ones in this study. It shall be noted that the preceding hydroconversion tests ¹³ were performed at a total pressure of 1.1 MPa whereas the tests herein were carried out at 4.1 MPa, to better reproduce industrial conditions (vide previous section).

3.1 Physico-chemical properties of the materials

Table 2 summarize the main features of both the HUSY zeolite and the Pt/HUSY catalyst. Most relevant for the purpose of this work are the concentrations of the active sites in the Pt/HUSY catalyst. The concentration of Brønsted acid sites amounted to 820 μmol g⁻¹, based on the concentration of Al^{IV} atoms (n_{Al}) ²⁷. The metal dispersion was ca. 52 %, resulting in a metal to acid sites molar ratio of 0.023. Concerning the location, Pt nanoparticles have been observed to be mainly located in the micropores of HUSY zeolite ³⁰.

Table 2: Physico-chemical, textural and structural characteristics of HUSY zeolite and corresponding Pt-containing bifunctional catalyst ^{13, 27}.

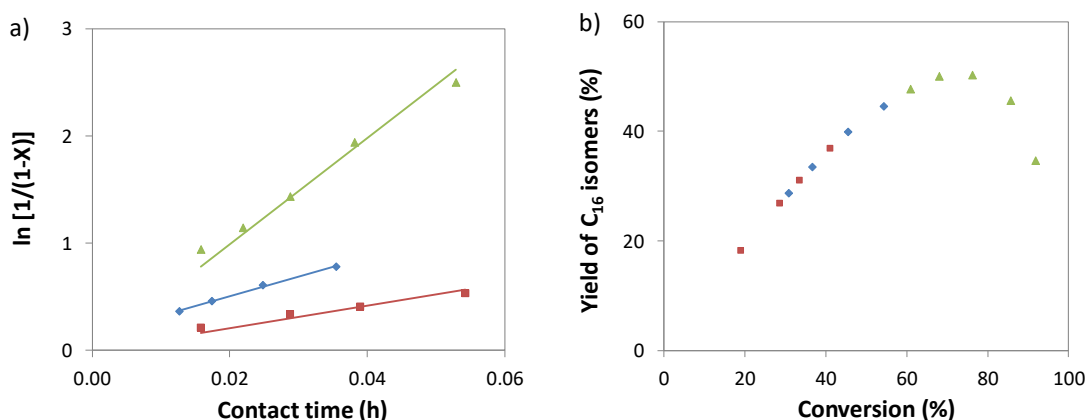
HUSY					Pt/HUSY		
V _{micro} (cm ³ g ⁻¹)	S _{ext} (m ² g ⁻¹)	(Si/Al) _{global}	(Si/Al) _{framework}	n _{Al} (μmol g ⁻¹)	Pt loading (wt.%)	Pt dispersion (%)	n _{Pt} /n _{Al} (mol mol ⁻¹)
0.33	101	16.7	18.4	820	0.7	52	0.023

1

2 3.2 Catalytic behaviour

3 Figure 1a) depicts the evolution of *n*-hexadecane conversion in terms of 1st-order kinetics as
4 function of contact time. A very good fit to the rate law can be observed, confirming once again
5 the results in literature for *n*-paraffins hydroconversion^{13, 32, 33}. This allowed the calculation of
6 both the catalyst activity and turnover frequency at the different temperatures (Table SI2). At
7 548K, for instance, the values were 61 $\mu\text{mol g}^{-1} \text{s}^{-1}$ and $75 \cdot 10^{-3} \text{s}^{-1}$, respectively. The apparent
8 activation energy was ca. 188 kJ mol^{-1} , in agreement with previous values reported (150-190 kJ
9 mol^{-1}) for pure *n*-hexadecane hydroconversion on Pt/zeolite catalyst^{13, 34}.

10 The yield of hexadecane isomers as a function of *n*-hexadecane conversion is represented in
11 Figure 1b). A unique curve, regardless of the reaction temperature, was obtained, as expected
12 for a well-balanced catalyst^{28, 35, 36}. The maximal isomer yield was about 50%.



13 **Figure 1:** a) 1st-order plot of *n*-hexadecane conversion (*X*) as function of contact time and b)
14 yield of C₁₆ isomers as function of conversion, over Pt/HUSY catalyst. Reaction temperatures:
15 528 (■), 538 (◆), and 548 K (▲). Symbols stand for experimental data and lines for the
16 corresponding linear regression.

17

18 4 Industrial case study: NiMoS/(Al₂O₃+HUSY) in *n*-hexadecane 19 hydroconversion in presence of H₂S and NH₃

20 Conversely to Pt/zeolite catalysts, the impact of the balance between the two catalytic functions
21 on the catalyst performances has been seldom studied over NiMoS-based shaped catalysts. To
22 investigate it, catalysts with different molybdenum loadings (from 9 to 17 wt.%) were firstly
23 synthesized and evaluated for *n*-hexadecane hydroconversion. Aiming at somewhat mimicking
24 the industrial reaction medium, both H₂S and NH₃ were present in this case. No noticeable
25 evolution of the activity with Mo loading and only a minor evolution on the maximal C₁₆ isomer

1 yield were observed (vide SI.4 for full results). Hence, a NiMoS/(Al₂O₃+HUSY) catalyst with
2 intermediate Mo content (12 wt.%) was selected to compare against the well-balanced
3 Pt/HUSY.

4 The comprehensive characterization results and discussion of all NiMoS/(Al₂O₃+HUSY)
5 catalysts can be found in SI (section SI.3). The focus herein has been put on the
6 12NiMo/(Al₂O₃+HUSY) sample.

7 4.1 Physico-chemical properties of the materials

8 The zeolitic trilobed extrudates comprised 83.5% of γ -alumina and 16.5% of HUSY zeolite,
9 based on the amount of silicon detected in shaped samples (**Erreur ! Source du renvoi**
10 **introuvable.**). In a previous study, the characterization of the shaped sample hinted at a full
11 preservation of the zeolite properties in the extrudates, in particular the zeolite porosity and
12 Brønsted acidity²⁸. The concentration of protonic Al^{IV} sites in the extrudates can be thus directly
13 estimated via that of the parent zeolite, taking into account the zeolite fraction in the shaped
14 support. The concentration of Brønsted acid sites is thus expected to be around 135 μmol
15 $\text{g}_{\text{support}}^{-1}$.

16 **Table 3:** Physico-chemical, textural and structural characteristics of HUSY shaped support²⁸
17 and NiMoS/(Al₂O₃+HUSY).

Al ₂ O ₃ +HUSY extrudates			NiMoS/(Al ₂ O ₃ +HUSY)		
Si (wt.%)	Zeolite (wt.%)	n _{Al} ($\mu\text{mol g}_{\text{support}}^{-1}$)	Mo loading (wt.%)	n _{NiMoS} /n _{Mo} (mol.%)	n _{NiMoS} /n _{Al} (mol mol ⁻¹)
7.3	16.5	135	12.0	16	2.9

18

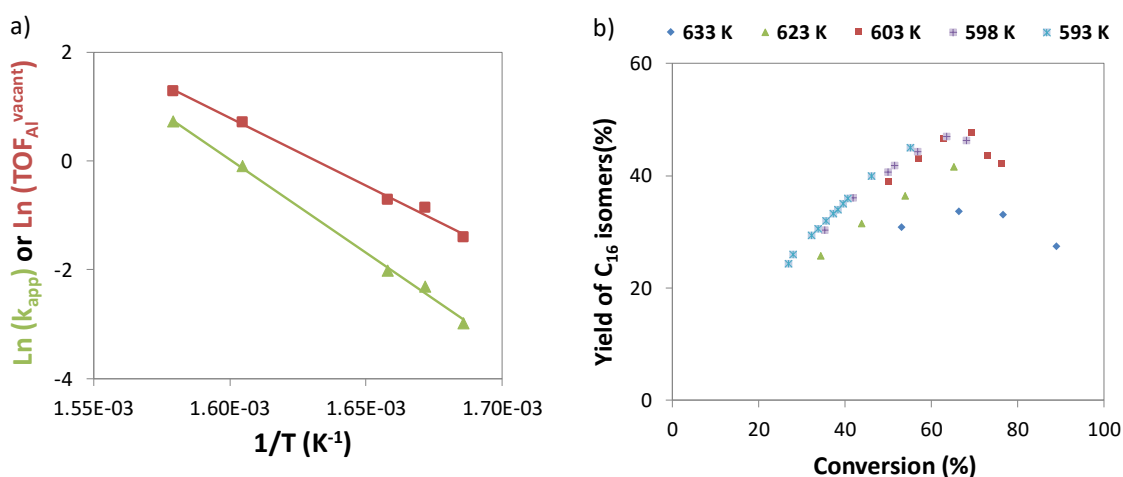
19 The Ni-decorated MoS₂ slabs had average length of 3.2 nm (Table SI5). The corresponding
20 Ni/Mo molar ratio was about 0.28 (Table SI5) and so somewhat lower than the global Ni/Mo
21 molar ratio of 0.40 (Table SI3). To quantify the active sites for HDH reactions, i.e. the MoS₂ sites
22 promoted with Ni (so-called mixed sites)^{37, 38}, a methodology based on X-ray photoelectron
23 spectroscopy²⁹ was employed. Accordingly, the concentration of NiMoS sites in the catalyst
24 (n_{NiMoS}) was about 0.28 mmol g⁻¹. The corresponding NiMoS sites to Al^{IV} sites molar ratio was
25 2.9 (Table 3). Based on NiMoS sites concentration, the amount of Mo incorporated in the
26 NiMoS site could be assessed. It corresponded to ca. 15% of the overall Mo deposited on the
27 catalyst. Such low values were somewhat expected as only the Mo-edge sites of the slabs are
28 available to be promoted by nickel.

29 An uniform distribution of the metals along the extrudate diameter both before, as previously
30 reported as well²⁹, and after sulfidation (Figure SI1a) was observed, validating the protocol
31 employed to generate the NiMoS active phase. Concerning the location of the metals in the

1 support, both nickel and molybdenum were preferentially located in the alumina (Figure SI1b).
 2 Nevertheless, whereas only 1% of all Mo was deposited in the zeolite, that was 7% in the case of
 3 Ni. Such distribution is in line with the electrostatic interactions between the metal species and
 4 the support components in solution (vide SI.3 for full discussion). As, for slabs of 3.2 nm, Ni/Mo
 5 molar ratios above 0.5 no longer result in the decoration of Mo atoms leading instead to the
 6 decoration of Ni atoms (forming thus NiS_x sites)³⁷, most of the nickel in the zeolite shall be
 7 under rather inactive forms. In practice, virtually all the HDH function was located in the alumina.

8 4.2 Catalytic behaviour

9 The evolution of the apparent kinetic constant for *n*-C₁₆ consumption (which is equivalent to the
 10 catalytic activity per catalyst mass according to our definition of the latter) with temperature was
 11 represented by an Arrhenius' plot in Figure 2a). An excellent fit of experimental data to
 12 Arrhenius' equation was observed. The apparent activation energy estimated amounted to 286
 13 kJ mol⁻¹. This contrasts with the activation energy of 188 kJ mol⁻¹ over Pt/HUSY and the typical
 14 range reported in literature (150-190 kJ mol⁻¹)^{13, 34}. This difference can be tentatively explained
 15 by the presence of NH₃, because ammonia adsorbs on the acid sites of the HUSY zeolite and
 16 its coverage decreases with temperature (chemisorption being exothermic). As a consequence
 17 the amount of protonic sites available for catalysis increases when the temperature is
 18 increased. As a matter of fact, literature on *n*-C₁₆ hydroconversion reported an increase in the
 19 apparent activation energy, in presence of NH₃, of about 100 kJ mol⁻¹³⁴. In cyclohexane
 20 hydroisomerization, we had also observed an increase in activation energy of about 90 kJ mol⁻¹
 21²⁹. In both studies, this increment has been attributed to the contribution of NH₃ desorption
 22 enthalpy to the apparent activation energy.



23 **Figure 2:** a) Arrhenius' plot for catalytic activity (\blacktriangle) and turnover frequency per available
 24 protonic site (\blacksquare). Symbols stand for experimental data and lines for model fitted Arrhenius'
 25 equation. Units of k_{app} : $\mu\text{mol g}_{zeolite}^{-1} \text{s}^{-1}$. Units of $\text{TOF}_{\text{Al}^{\text{vacant}}}$: s^{-1} (molar basis). b) Yield of feed
 26 isomers as function of conversion at different reaction temperatures. Catalyst:
 27 NiMoS/(Al₂O₃+HUSY).

1 Figure 2b) depicts the evolution of C₁₆ isomers yield with conversion obtained at different
2 reaction temperatures for NiMoS/(Al₂O₃+HUSY) catalyst. The yield of feed isomers at a given
3 conversion remarkably increased with decreasing temperatures. From 633 to 623 K, the C₁₆
4 isomer yield at 60% of conversion increased from 34% to 42%. Further increase to 47% was
5 achieved by decreasing the temperature to 603 K. From 603 to 598 K, the differences between
6 C₁₆ isomer yield curves were more subtle. The nearly overlapping curves at 598 and 593 K
7 pointed as well to a greater effect at higher temperatures.

8 A remarkable impact of temperature on C₁₆ isomer yield was hence evident over
9 NiMoS/(Al₂O₃+HUSY) catalyst. This kind of temperature effect has been explained in
10 hydrocracking by the gap in the activation energies of the reactions occurring in the two
11 functions of the catalyst: the apparent activation energy of the HDH reactions is lower than that
12 of the acid-catalysed reactions and then the latter are more temperature-dependent³⁵. Hence,
13 as temperature decreases, the kinetic constant ratio of HDH reactions to acid-catalysed
14 reactions will increase. This will lead to a better balance of the acid function by the HDH
15 function. However, in the case of an excess of HDH to acid activity (i.e. in sufficiently well-
16 balanced catalysts), the kinetic constants ratio will have no practical effect on the metal-acid
17 balance: the catalysts would still be well-balanced, just the excess of HDH activity would be
18 smaller. In fact, no temperature dependence was found over the Pt/HUSY herein (Figure 1b).
19 The occurrence of unique curve for feed isomer yield as function of conversion has been vastly
20 reported in the literature for well-balanced catalysts over relative wide range of operating
21 conditions^{28, 35, 36}. In particular, a similar feed isomers yield curve can be obtained within a
22 temperature span of 65 K³⁶.

23 In summary, the noteworthy decrease in feed isomer yield with increasing temperature might
24 have been due to the change in the kinetic constants ratios with temperature, but other causes
25 cannot be ruled out. Particularly, in presence of analogous partial pressures of ammonia, the
26 number of available Brønsted sites for reaction (i.e. non-inhibited by NH₃) has been determined
27 to be quite small (below 2%) and dependent on the temperature at stake, due to NH₃ desorption
28 with increasing temperatures²⁹. Consequently, the effective HDH to protonic sites ratio is also
29 temperature-dependent in presence of ammonia.

30 **Impact of NH₃ adsorption on NiMoS-to-acid sites ratio**

31 The concentration of vacant protonic sites (n_{Al}^{vacant}) at a given temperature and NH₃ partial
32 pressure corresponds to the fraction of vacant sites at that temperature and pressure times the
33 overall concentration of sites (i.e. n_{Al}) – Eq. (2). The fraction of vacant sites has been previously
34 determined for similar temperatures and analogous ammonia partial pressures²⁹. The data and
35 calculation details can be found in SI.5. It shall be noted that the goal is to get hold of the order
36 of magnitude of the fraction of vacant sites and qualitatively assess the effect of temperature on

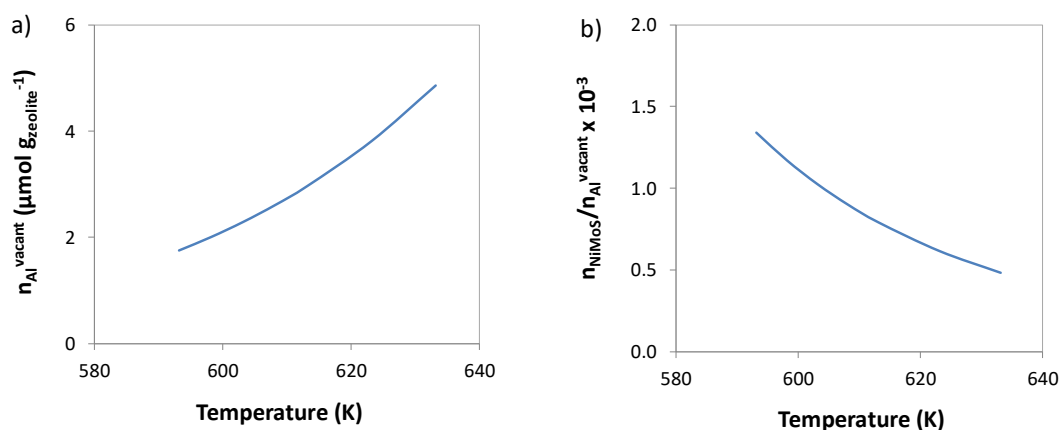
1 it. As the estimations are based on Arrhenius' equation, which contains an exponential term, the
2 values themselves shall be taken cautiously.

$$n_{Al}^{vacant}(T, P_{NH_3}) = \psi_V(T, P_{NH_3}) \times n_{Al} \quad \text{Equation (2)}$$

3 In order to evaluate the effect of temperature in presence of ammonia, the estimated
4 concentration of vacant (i.e. non-inhibited) Brønsted acid sites was plotted as a function of
5 temperature (Figure 3a). The concentration of protonic sites vacant for reaction almost triples
6 within the range of studied temperatures. The concentrations at stake correspond to the
7 inhibition of 99.4 to 99.8 % of the $820 \mu\text{mol g}_{zeolite}^{-1}$ Brønsted sites in the HUSY zeolite. It is
8 worth mentioning that, as this corresponds to nearly saturation of the protonic sites, these
9 results (and the corresponding conclusions) are relative insensitive to ammonia pressure
10 changes within comparable orders of magnitude²⁹.

11 To determine the ratio of the number of available NiMoS sites to the number of available
12 Brønsted sites ($n_{NiMoS}/n_{Al}^{vacant}$) as a function of temperature, the effect of temperature on the
13 available NiMoS sites should also be investigated. Although NiMoS sites are known to be Lewis
14 acid sites, its acidity is much lower than that of a zeolite, particularly the HUSY one employed.
15 Regarding H₂S, it is known to inhibited HDH molybdenum mixed sites³⁸⁻⁴². Nevertheless, H₂S
16 inhibiting effect on these sites is expected to be several orders of magnitude weaker than that of
17 NH₃ on Brønsted sites⁴³. For instance, under comparable operating conditions, a 4-fold
18 reduction in the hydrogenating activity of NiMoS catalysts has been reported, as compared to
19 H₂S-free reaction medium⁴¹. Hence, even though both NH₃ and H₂S will also inhibit the HDH
20 sites in the NiMoS/(HUSY+Al₂O₃) catalyst, the fraction of non-inhibited active sites should be
21 considerably higher than that of protonic ones. Therefore, for the purpose of this study, the
22 concentration of available NiMoS sites was hence assumed to be independent of the operating
23 conditions and similar to the values of n_{NiMoS} previously obtained (Table SI5).

24 Figure 3b depicts the estimated evolution of $n_{NiMoS}/n_{Al}^{vacant}$ with temperature. Decreasing the
25 reaction temperature can substantially augment the $n_{NiMoS}/n_{Al}^{vacant}$, by means of the decreasing
26 number of vacant Brønsted sites due to greater NH₃ adsorption. In particular, from to 633 to
27 593 K, the molar sites ratio almost tripled. It is worth noting that this represents a greater growth
28 in the $n_{NiMoS}/n_{Al}^{vacant}$ than that obtained by increasing the Mo content from 9 to 17%.



1 **Figure 3:** Estimated evolution of a) the concentration of vacant acid sites in the HUSY zeolite
 2 and b) the molar ratio of NiMoS to vacant Brønsted sites with temperature, under reaction
 3 conditions.

4 The estimation of the concentration of vacant sites (n_{Al}^{vacant}) enabled the calculation of the
 5 turnover frequency over the Brønsted acid sites (TOF_{Al}^{vacant}). When compared to the evolution of
 6 catalytic activity with temperature (Figure 2a), a clear lower slope is observed for the turnover
 7 frequency. The apparent activation energy was estimated in 187 kJ mol^{-1} , in the typical range
 8 for the hydroconversion of *n*-hexadecane, whereas that for catalytic activity amounted to 286 kJ
 9 mol^{-1} . The abovementioned hypothesis on the contribution of NH_3 desorption enthalpy to the
 10 apparent activation energy in the NiMoS/(HUSY+ Al_2O_3) catalyst was, hence, further supported,
 11 along with the importance of ammonia inhibition of the Brønsted acid sites.

12 In view of the previous observation, the superior C_{16} isomers yields reached at lower reaction
 13 temperatures (Figure 2b) can be tentatively explained by the larger $n_{NiMoS}/n_{Al}^{vacant}$ ratios. As the
 14 temperature decreased, the gain in isomer yield diminished and overlapped curves were even
 15 observed at the lowest temperatures. This pointed towards an approach to the plateau of
 16 maximal C_{16} isomer yield for temperatures about 593 K, because, even if the metal to acid sites
 17 ratio continuously increased (Figure 3b), the isomer yield became constant. In other words, the
 18 maximal C_{16} isomers yields observed at such temperatures most likely correspond to the
 19 performance of a well-balanced catalyst. In such case, the hydrogenation/dehydrogenation
 20 reactions are much faster than the acid-catalyzed ones, limiting consecutive reactions and,
 21 hence, maximizing the feed isomers yield^{13, 44}.

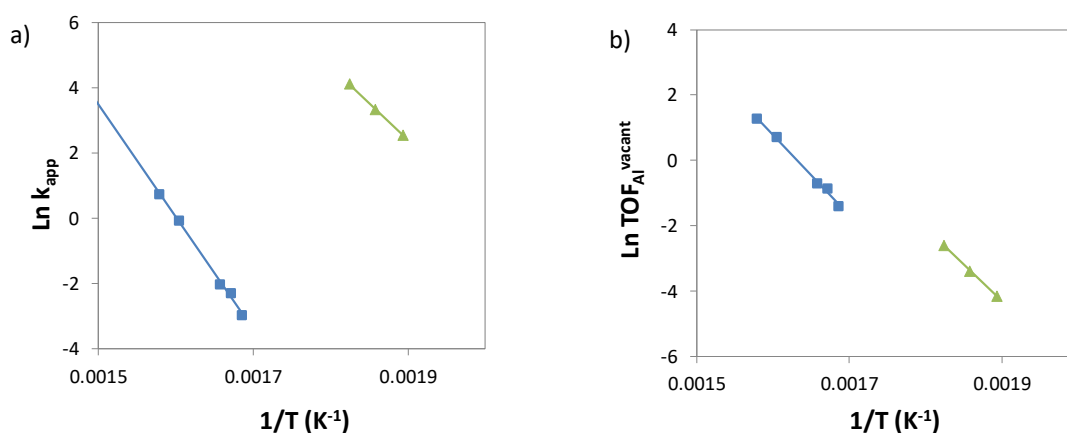
22 In summary, the temperature effect on the catalyst hydroisomerisation selectivity observed can
 23 be explained by both the improvement of balance between the two catalytic functions, due to
 24 either a higher kinetic constants ratio of HDH reactions to acid-catalysed ones or a higher metal
 25 to acid sites ratio. Most likely, the two phenomena took place simultaneously explaining hence
 26 the considerable increase in the maximal C_{16} isomers yield for temperature intervals as narrow
 27 as 5 K. Conversely, the observed improvement in the balance of acid function by HDH function
 28 implied that catalysts were far from the optimal balance at the initial operating conditions (i.e.
 29 high temperatures). Hence, the insensitivity of the catalytic performance to increased

1 molybdenum loadings, observed in the preliminary tests (vide SI.4), implies a relatively constant
2 the metal-acid balance regardless of the Mo loading. In other words, the HDH activity remained
3 similar although the active metal content was doubled (from 9% to 17%).

4 5 Academic vs. industrial case studies

5 5.1 Activity

6 To disregard the simple effect of the lower zeolite content in the NiMoS/(Al₂O₃+HUSY) catalyst,
7 the activity was normalized per zeolite mass. Due different temperature ranges, the catalytic
8 activity was compared through an Arrhenius's plot (Figure 4a). Pt/HUSY exhibited a significantly
9 higher catalytic activity than NiMoS/(Al₂O₃+HUSY) for a much lower reaction temperature. In
10 order to compare directly the catalysts activity, the Arrhenius' equations of NiMoS-catalysts
11 were extrapolated to the same level of activity of Pt-catalysts. In doing so, it became noticeable
12 that the reaction temperature would have to be increased by ca. 125 K in order for NiMoS/(
13 Al₂O₃+HUSY) catalyst to attain the activity level (zeolite mass) of Pt/HUSY one².



14 **Figure 4:** Arrhenius's plot for a) catalytic activity and b) turnover frequency per vacant protonic
15 site over Pt/HUSY (▲) and NiMoS/(Al₂O₃+HUSY) (■) catalysts. Symbols stand for experimental
16 data and lines for regression of the Arrhenius equation. Units of k_{app} : $\mu\text{mol g}_{zeolite}^{-1} \text{s}^{-1}$. Units of
17 TOF_{Al}^{vacant} : s^{-1} (molar basis).

18 5.2 Turnover frequency over available protonic sites

19 The evolution of the turnover frequency over the available protonic sites with temperature is
20 shown in Figure 4b). It should be borne in mind that, for Pt/HUSY, the concentration of
21 Brønsted sites in the catalyst corresponded to that of Al^{IV} species, as no NH₃ was fed in this

² Due to the large difference in the activation energies, the comparison of catalytic activities is very temperature-dependent. For instance, when considering 548 K (i.e. the lowest temperature at which Pt/HUSY was tested), the activity ratio amounts to $3 \cdot 10^6$ in favor of Pt/HUSY catalyst. When considering 633 K (i.e. the highest temperature at which NiMoS/(HUSY+Al₂O₃) was tested), the activity ratio decreases to $7 \cdot 10^3$. Hence, reaction temperature was preferred to compare catalyst activity.

1 case. In the case of NiMoS/(Al₂O₃+HUSY), in order to account for the inhibition of the acid sites
2 by NH₃, the concentration of vacant protonic sites (as estimated in Section 4.2) was employed.
3 NiMoS/(Al₂O₃+HUSY) revealed superior TOF_{Al^{vacant}} than Pt/HUSY but it was also tested at
4 higher temperatures. Comparing both catalysts at similar temperatures, the turnover
5 frequencies over NiMoS/(HUSY+Al₂O₃) were of comparable order of magnitude to those over
6 Pt/HUSY catalysts. Especially, when comparing with very large gap between catalytic activities
7 (Figure 4a). Although precaution is advisable as data was not (and cannot be) obtained at
8 similar temperatures, the results seemingly pointed out a comparable intrinsic activity of
9 available Brønsted site regardless of the differences in catalysts composition.

10 Taking into account the concentration of vacant acid sites, the apparent activation energies
11 were found to be remarkably similar for Pt/HUSY (188 kJ/mol) and NiMoS/(Al₂O₃+HUSY) (187
12 kJ/mol) catalysts. The activation energies seemed hence to be intrinsic to the zeolite function
13 and independent of the HDH function. These results pointed to the rate-determining character of
14 the acid-catalyzed reactions in the conversion of *n*-hexadecane, in agreement with the well-
15 balanced character of Pt/HUSY. For NiMoS/(Al₂O₃+HUSY), the similarities to Pt/HUSY strongly
16 supported that the plateau in terms of activity per protonic site was also reached with the
17 NiMoS-acid sites ratios at stake. As discussed above, for NiMoS/(Al₂O₃+HUSY), the apparent
18 activation energy corresponds to the apparent overall activation energy (as calculated by the
19 Arrhenius' equation for catalytic activity) subtracted by enthalpy of NH₃ desorption from the acid
20 sites.

21 **5.3 Selectivity towards isomerization**

22 Figure 5 provides the evolution of C₁₆ isomers yield as a function of *n*-hexadecane conversion
23 for both catalysts. Over NiMoS/(Al₂O₃+HUSY), the evolution of C₁₆ isomers yield was influenced
24 by the reaction temperature (593-633 K), as previously discussed, whereas a unique tendency
25 for isomer yield was obtained over Pt/HUSY catalyst regardless of the temperature at stake
26 (528-548 K). The comparison between catalysts performance is thus temperature-dependent.
27 At 633 K, the maximal C₁₆ isomers yield reachable over NiMoS/(Al₂O₃+HUSY) catalyst was
28 considerably lower than the one obtained over Pt/HUSY catalyst. However, as temperature was
29 decreased over the NiMoS-catalyst, the isomers yield increased for a given conversion (see
30 Figure 2b). In fact, the evolution of C₁₆ isomers yield with conversion at the lowest investigated
31 temperature (593 K) over NiMoS/(Al₂O₃+HUSY) was remarkably similar to that of Pt/HUSY
32 (even though the maximum of C₁₆ isomers yield curve could not be measured over this NiMoS-
33 catalyst as the maximum achievable conversion decreased with decreasing temperature). The
34 catalysts revealed hence the same selectivity in feed isomers for a temperature sufficiently low
35 over the NiMoS/(Al₂O₃+HUSY), notwithstanding the reaction temperature gap of up to 65 K.

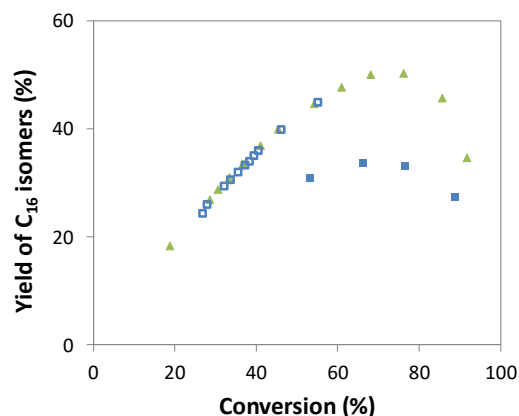


Figure 5: Yield of C₁₆ isomers as a function of conversion over Pt/HUSY at 528-548 K (▲) and NiMoS/(Al₂O₃+HUSY) catalyst at 633 (■) and 593 K (◻).

In Section 4.2, this temperature effect was associated with an improvement of the metal-acid balance. The performance of NiMoS/(HUSY+Al₂O₃) catalyst at the lowest temperatures corresponded hence to that of a well-balanced catalyst. The similar results to Pt/HUSY, which was known to be well-balanced, reinforced that conclusion. In other words, when well-balanced, Pt/HUSY and NiMoS/(HUSY+Al₂O₃) catalysts apparently hold the same intrinsic performance, in spite of the differences in terms of catalyst composition and operating conditions.

As a final note, the distribution of cracked products was observed to be markedly different (Figures SI4 and SI5). Secondary cracking was more extensive over NiMoS/(HUSY+Al₂O₃) than over Pt/HUSY and the isomerization degree was also lower (vide SI.6 for discussion). Therefore, when it comes to cracking products, regardless of the good balance between catalytic functions, the gap in the reaction temperatures dictated a different product distribution.

6 Discussion

In this section, the differences between the laboratorial and the industrial-like catalytic systems are analysed at the active sites and mechanistic level. For comparison purposes, an in-between catalytic system, representing a step in the scaling-up of a hydrocracking catalyst (hence denominated “pilot”), was also included. In this system, the HDH function was Pt (as in the academic-like one), but it was deposited in the shaped HUSY support and located on the alumina phase (as in the industrial-like one)⁴⁵. In that case, the catalyst was evaluated in pure *n*-hexadecane hydroconversion, i.e. in the absence of any contaminants like H₂S and NH₃ (Table 4). Before the mechanistic considerations, the characteristics of NiMoS-based catalyst at the active site level are compared to those of platinum ones. Finally, the gained knowledge will be applied in view of potential solutions for improvement of hydroconversion catalysts.

6.1 Hydrogenating-dehydrogenating function: Pt vs. NiMoS

As mentioned in the Introduction, the hydrogenating function of a hydroconversion catalyst can be granted either by a promoted transition metal sulphide or by a noble metal. The selection is mainly determined by the low sulphur resistance of noble metals. Notwithstanding the fact that both phases promote hydrogenation/dehydrogenation reactions, their active sites are fundamentally different. For Pt-based catalysts, the active sites are surface Pt⁰ atoms, whilst, in NiMoS-based catalysts, the active sites are Ni-promoted MoS₂ sites. Moreover, the accessible Pt sites can be easily probed through chemisorption methods whereas no routine characterization method enables to probe the accessible NiMoS sites. In a previous work²⁹, a methodology to determine the number of NiMoS sites, based on amount of Ni engaged in NiMoS phase, was developed. While this allows to finally quantify the amount of NiMoS sites, it should be kept in mind that this corresponds to the overall sites and not only to the accessible ones, as when measuring the chemisorption of probe molecule, e.g. CO on Pt⁰ sites.

The results revealed that Mo promoted sites (NiMoS) corresponded to only 15% of the overall Mo in the catalyst. Concomitantly, the deposition of Pt in either the HUSY zeolite or in the shaped extrudates leads to metal dispersions between 40²⁷ and 90%^{28, 45, 46}, with shaped supports typically featuring higher dispersions. Potentially, because ionic exchange is preferred in the latter whereas incipient wetness impregnation is commonly employed in powder zeolites. Therefore, in NiMoS-based catalysts, the metal loading should be superior to that in Pt-based catalyst in order to achieve the same amount of HDH sites.

In this study, due to the documented low HDH activity of NiMoS sites compared to Pt sites, the difference was even made larger: Pt loading was 0.7% whereas Mo loading was 12%. This gave rise to HDH sites concentrations in the catalysts of 20 μmol g⁻¹ and 280 μmol g⁻¹, respectively. Despite the potential overestimation of the concentration of accessible NiMoS sites might be overestimated to some extent by this technique, the NiMoS-based catalyst had more than 10-fold the HDH sites existing in Pt-based catalyst. More importantly, the preliminary tests (vide SI.4) indicated that apparently the Mo content cannot be used to tailor the activity of the HDH function as easily as in Pt catalysts.

6.2 Metal-acid balance and intimacy

Both hydrogenating functions were able to properly balance the zeolitic acid functions, according to the comparable turnover frequency over Brønsted sites and C₁₆ isomers yields, but the metal to acid sites ratios differed enormously. Particularly, the plateau of maximal feed isomers yield was reached in Pt/zeolite catalysts for a metal to acid sites molar ratio of ca. 0.02 (at 543 K) whilst, over NiMoS-catalysts, a metal to acid sites ratio of 1.3·10³ was required (at 593 K). Although the number of accessible NiMoS sites may have been somehow overestimated and tests occurred at higher reaction temperatures, the metal-to-acid sites ratio

1 required was five orders of magnitude greater than the n_{Pt}/n_{Al} , clearly corroborating the
2 substantially greater hydrogenating activity of Pt^0 sites compared to NiMoS sites, previously
3 referred in literature ^{47, 48}.

4 In the investigated conversion range, similar yield in C_{16} isomers was reached for catalysts
5 working at a temperature differing 65 K. Although the low activity of the NiMoS catalyst did not
6 allow for high conversion experiments, all evidence indicate that the reaction temperature does
7 not affect the isomerisation selectivity when catalysts are well-balanced. This agrees with the
8 unique feed isomer yield curve typically reported in literature over a relatively wide range of
9 operating conditions for HUSY zeolite ^{35, 36}. Furthermore, it points out a comparable activation
10 energy of isomerization and cracking pathways, as already suggested in literature for particular
11 cases ³⁵. Otherwise, the isomerization selectivity at a given conversion of well-balanced
12 catalysts would be influenced by the reaction temperature.

13 The disclosure of a similar intrinsic behaviour of well-balanced zeolites in such dissimilar
14 catalytic systems allowed several insights. First of all, despite the numerous steps during
15 catalyst preparation (comprising shaping, metal deposition, and thermal treatments), the zeolite
16 performance can be preserved by judiciously designing the experimental protocols.
17 Furthermore, inhibition by ammonia of most of the acid sites only reduced the number of
18 available sites, while their intrinsic performance was not affected. Conversely, reports in
19 literature showing a beneficial effect of NH_3 inhibition to selectivity have commonly attributed
20 this to the suppression of the strongest acid sites. Herein, this effect was proposed to be mainly
21 a consequence of the different degree of metal-acid balance, while the HUSY zeolite selectivity
22 remained unchanged. This also supports the argument that selectivity of well-balanced HUSY
23 zeolite is mainly determined by confinement effects ⁴⁹.

24 The number of acid-catalysed steps that may occur before hydrogenation depends also on the
25 distance between HDH and acid sites ^{15, 16, 45}. For instance, in the pilot-like catalyst, with
26 platinum nanoparticles located in the alumina binder, a lower maximal C_{16} isomers yield was
27 observed. This was attributed to the longer distance between HDH and acid sites, as compared
28 to Pt/HUSY ⁴⁵. The critical Pt-Brønsted acid site distance was disclosed to be determined by the
29 size of zeolite clusters which was in the 200-20000 nm range, being hence much larger than the
30 typical distance between Pt particles. Herein, for NiMoS-based catalysts, the case is similar:
31 NiMoS sites were located on the alumina and so the critical distance between NiMoS and
32 Brønsted acid sites should still be half the size of zeolite clusters, so 100-10000 nm. Even
33 though a similar longer distance would be expected for both shaped catalysts, either Pt or
34 NiMoS based, no impact of the longer HDH-acid sites distance was noticeable in the latter case.
35 As a matter of fact, the requirement for the critical distance (R) intimacy between HDH and acid
36 sites, for a given support (i.e. fixed diffusivity, D) at fixed operating conditions (i.e. fixed olefins
37 concentration, $[O]_{eq}$), is determined by the rate of the overall reaction rate, in this case the rate
38 of consumption of n -hexadecane (Eq. 3) ¹⁵. In well-balanced catalysts, as the HDH reactions

1 are at equilibrium, the overall reaction rate is determined by the acid-catalysed reactions. In that
 2 way, the much lower activity of NiMoS-based catalysts, due to the significant inhibition of the
 3 HUSY zeolite by NH₃, compared to Pt-based ones could be at the origin of the irrelevance of
 4 HDH-acid sites distance in industrial-like system.

$$R^2 < \frac{[O]_{eq}D}{r_{overall}} \quad \text{Equation (3)}$$

5

6 **Table 4:** Parameters controlling the performances (activity and isomerisation selectivity) of the
 7 three catalytic systems in the hydroconversion of *n*-hexadecane.

Catalytic system	Academic ¹³ & this study	Pilot ⁴⁵	Industrial This study
Support	Powder HUSY	Shaped Al ₂ O ₃ +HUSY	Shaped Al ₂ O ₃ +HUSY
HDH function	Pt	Pt	NiMoS
Location of HDH function	Zeolite	Alumina	Alumina
Metal-acid intimacy	+++	-	++
Metal-acid balance	++	+++	+
Limiting factor of catalytic performance	Balance	Intimacy	Balance
Solutions	Increasing Pt loadings	Deposition of Pt on zeolite Reducing the size of zeolite clusters	Increasing NH ₃ inhibition (e.g. lower reaction temperature) Reduction of zeolite content Increase hydrogenating capacity of HDH function

8 6.3 Industrial applications

9 From a performance point of view, the sole substantial modification from the academic to
 10 industrial system was the catalytic activity: NiMoS-catalyst would require an increase of 125 K in
 11 the reaction temperature to reach the same catalytic activity of Pt/zeolite catalyst. This much
 12 lower activity in industrial-like catalytic system can be mostly attributed to the inhibition of more
 13 than 99.4% of the Brønsted sites by ammonia. The concentrations of available acid sites at
 14 stake (in the order of a few micromoles per zeolite gram) were about two orders of magnitude
 15 lower than the 820 μmol g_{zeolite}⁻¹ in the HUSY zeolite and the typical concentration of Brønsted
 16 sites over non-inhibited zeolites⁵⁰. In addition, the dilution of the zeolite, in the “industrial”
 17 catalyst, by both the binder and the metals introduced, also accounted for the reduction in the
 18 activity (9-fold loss). The product of these two effects leads to an estimated loss in activity
 19 between 1500 and 4500 times. From the catalyst point of view, enhanced activities in
 20 hydroconversion industrial catalysts can be hence achieved by mitigating the acid sites

1 inhibition, but the gain in activity would be followed by a loss in feed isomers yield. In other
 2 words, the small number of available Brønsted sites in industrial catalytic systems (as compared
 3 to bench) results in a far lower hydroconversion activity but enables, at the same time, an
 4 acceptable metal-acid balance in presence of a substantially weaker HDH function.

5 The presence of basic molecules in the feedstock ensures hence the proper balance between
 6 the catalytic functions of industrial catalysts. Furthermore, one may hint that, as long as the
 7 HDH function is granted by a promoted transition metal sulphide (i.e. a function with relatively
 8 weak hydrogenating activity), industrial catalysts will be restrained to a much lower activity than
 9 bench catalysts if an acceptable isomerization selectivity has to be granted. In fact, when
 10 necessary, more extended inhibition of the Brønsted sites by NH₃ or lower zeolite contents can
 11 be used to improve the metal-acid balance and hence the isomerization selectivity.

12 In spite of the approach to the industrial operation (for VGO hydrocracking) in terms of operating
 13 conditions, some discrepancies still persisted (Table 1). These were tentatively filled in by taking
 14 into account the literature consensus on the role of each operating variable, as summarized in
 15 Table 5. It is worth mentioning that some relevant topics to industrial operation are still barely
 16 studied in open literature. Particularly, most accounts are on short and long chain paraffins and
 17 only a few studies have been devoted to the hydrocracking mechanism of complex naphthenic
 18 and aromatic molecules, and their mixtures^{12, 46, 51-54}. Hence, the focus was herein put on the
 19 well-known effects of operating conditions on long chain *n*-paraffins hydrocracking.

20 The foreseeable effects point to opposite directions precluding a straightforward answer.
 21 However, the significant impact of temperature in metal-acid balance together with the expected
 22 higher reactivity of the industrial feedstock hint at a lower degree of metal-acid balance in
 23 industrial operation, as to compared to the tests performed in this study. Therefore,
 24 improvements on industrial catalyst performance might be achieved by more hydrogenating
 25 HDH functions, including improvements on molybdenum sulphides⁵⁵, and/or by an optimal
 26 control of the metal-acid balance via a careful manipulation of the temperature. Playing with the
 27 acid site inhibitor partial pressure is also a possibility, but, due to the very high coverage of the
 28 acid sites, this is expected to have limited impact²⁹.

29 **Table 5:** Differences and expected outcomes of modification of the operating conditions from
 30 the ones used in this work to VGO hydrocracking ones, for a given bifunctional catalyst.

Parameter	Feedstock	NH ₃ partial pressure	Total pressure	Temperature
Change	Longer paraffins	Increase	Increase	Increase
Effects	Greater reactivity ^{35, 56}	Less available Brønsted sites (this study)	Lower reactivity gap between molecules ^{57, 58} Superior metal-acid balance ³⁵	More available Brønsted sites (this study) Lower kinetic constant ratio of HDH to acid

	reactions ³⁵			
Expected outcome	Superior activity Inferior isomerization selectivity , due to lower metal-acid balance and, potentially, low intimacy	Minor if Brønsted sites are nearly saturated	Unpredictable as effects are contradictory	Superior activity Inferior selectivity , due to lower metal-acid balance

1 Finally, the key role of temperature in the catalyst performance shall be taken into account when
 2 designing and selecting catalysts for scale-up and industrialization. As the effect of temperature
 3 depends on how balanced the catalyst is, it is strongly dependant on both the catalyst and the
 4 operating conditions. Hence, in a first, more fundamental stage, the temperature effect shall be
 5 suppressed in order to evaluate the catalyst intrinsic performance. During scale-up, the effect of
 6 temperature shall be then thoroughly evaluated. Due to the numerous differences imposed by
 7 the scale, extrapolation of relative performances from bench to industrial scale should be done
 8 extremely cautiously.

9 7 Conclusion

10 The main goal of this work was to shed light at the parameters controlling the performance of
 11 industrial-like NiMoS/(Al₂O₃+HUSY) hydrocracking catalysts, bridging the gap to the most
 12 academically studied Pt/HUSY ones. For the same reactants partial pressures but mimicking
 13 the presence of contaminants in the industrial case study (by co-feeding H₂S and NH₃), well-
 14 balanced industrial-like catalyst revealed to be as selective as the laboratorial one, but much
 15 less active (temperature gap at iso-conversion of 125 K) due to the inhibition of most part of
 16 Brønsted sites by NH₃. The turnover frequency per non-inhibited protonic site was however
 17 comparable to that of Pt/HUSY catalysts. Hence, as long as the zeolite is not modified during
 18 catalyst preparation (comprising shaping and metal deposition), the zeolite intrinsic performance
 19 is not affected by the choice or location of HDH function, nor by the presence of NH₃.

20 Concerning the HDH-acid sites molar ratio, under the chosen operating conditions, both
 21 industrial-like and laboratorial-like catalysts could achieve the optimal balance between
 22 functions but the ratio of HDH to acid sites required was some orders of magnitude greater in
 23 NiMoS-catalysts the ones reached in Pt-based catalysts. Such extraordinarily high HDH to acid
 24 sites ratios over NiMoS-catalysts is mostly ensured by the partial inhibition of Brønsted sites. In
 25 that way, one may hint that, as long as the HDH function is granted by a promoted transition
 26 metal sulphide (i.e. a function with relatively weak hydrogenating activity), industrial catalysts
 27 will be restrained to a much lower activity than bench catalysts in order to keep an acceptable
 28 selectivity.

1 In summary, ensuring optimal metal-acid balance and intimacy is critical to assess the intrinsic
2 behaviour of bifunctional catalysts and achieve optimal performance in industrial applications.
3 Given the predominant role of the metal-acid balance in industrial-like conditions, the key
4 parameters playing a role in the latter are temperature, via the inhibition of acid sites, and the
5 hydrogenating function. Therefore, in addition to careful design of the operating conditions, the
6 development of more efficient industrial catalysts shall focus not only on less diffusion-
7 restrained acidic materials, but also on the engineering of more active hydrogenating functions,
8 ensuring thereby the most out of the cutting-edge materials developed in the last decades.

9

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16

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