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**Full Length Article**

**Selective synthesis of acetone from isopropyl alcohol over active and stable CuO–NiO nanocomposites at relatively low-temperature**



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CuO–NiO nanocomposite catalysts were synthesized by an oxalate precipitation route. The basicity of the catalysts was measured using adsorption–desorption of acetic acid. The cata- lytic conversion of isopropanol to acetone was carried out in a conventional fixed bed flow type reactor at 200 °C using N2 as a carrier gas. The results indicated that, the addition of NiO to CuO greatly enhanced the electrical conductivity via the creation of new charge car-

riers (Cu+–Ni3+). In addition, the results revealed that the catalyst containing 30 wt.% NiO

calcined at 400 °C possesses the highest catalytic activity with 98% conversion and 100% selectivity to acetone. The remarkable catalytic performance of the synthesized catalysts is attributed to the decrease of the energy gap, the increase in the concentration of charge

carries (Cu+–Ni3+), and to the creation of more weak and intermediate basic sites.

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# Introduction

Acetone is an important material in many laboratory and in- dustrial applications such as, a chemical intermediate in pharmaceuticals and as a solvent for vinyl and acrylic resins, lacquers, alkyd paints, inks, cosmetics and varnishes. It is also used in the preparation of paper coatings, adhesives, and heat- seals coatings and is also employed as starting material in the synthesis of many compounds [[1–3]](#_bookmark5). Acetone can be produced

industrially via two routes, the first is the cumene hydroper- oxide oxidation and the second is the dehydrogenation of isopropyl alcohol (IPA). It is estimated that, at present, nearly 85% of acetone world capacity is based on cumene hydroper- oxide route for the co-production of acetone and phenol. About

0.62 tons of acetone is produced per ton of phenol obtained. The production of acetone from only cumene would require a balancing of the market with the phenol product from this process. Therefore, there is a need of alternative routes for acetone production other than the cumene process.

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Dehydrogenation of IPA over a metal, metal oxide or salt cata- lysts is the second route for the manufacture of acetone. The preparation of nanostructured materials has received consid- erable attention owing to their unique properties and interesting potential applications [[4–10]](#_bookmark6). Metal oxide nanomaterials are in- teresting solids due to their surface acid–base properties and oxidation–reduction potentials and hence they constitute the largest family of heterogeneous catalysts [[11–16]](#_bookmark7). Several tran- sition metal oxides exhibit the oxidation–reduction behavior due to the ease with which they can be used as an oxidizing agent and then readily be generated [[17–21]](#_bookmark9). In this context, Turton et al. [[22]](#_bookmark10) mentioned that a single pass conversion of 85–92% with respect to IPA with reactor conditions of 2 bars and 350 °C is generally achieved. El-Shobaky et al. [[23]](#_bookmark11) have studied the catalytic conversion of IPA over CuO/MgO treated with K2O. They stated that the investigated system behaved as selective catalyst for dehydrogenation of isopropyl alcohol. In addition, the surface properties, catalytic activities, and cata- lytic selectivities toward IPA conversion of NiO/Al2O3 and CuO/ Al2O3 catalysts were investigated by Ashour [[24]](#_bookmark12). The author concluded that the conversion of isopropanol at a reaction tem- perature of 260–350 °C proceeded to propene and to acetone with different selectivities. Moreover, the influence of precur- sor of MgO and preparation conditions on the catalytic dehydrogenation of IPA over CuO/MgO catalysts has been studied by El-Molla et al. [[25]](#_bookmark13). Furthermore, the physicochemi- cal properties of individual and binary Ni and Ce oxide systems have been studied by Deraz and Al-Arifi [[26]](#_bookmark14). They concluded that at a reaction temperature of 250–450 °C, NiO and also CeO2 acted as dehydrogenation catalysts whereas their mixture containing 12 wt.% NiO acted as dehydrogenation and a de- hydration catalyst for isopropanol conversion. The maximum conversion obtained was 47% at a reaction temperature of 450 °C with selectivity to acetone of 62.2%. The surface and catalytic properties of NiO/MgO system doped with Fe2O3 have been studied by Shaheen et al. [[27]](#_bookmark15) and they concluded that pure and variously doped solids behave as dehydrogenation and de- hydration catalysts leading to the formation of acetone and propene.

In our previous work [[28]](#_bookmark16), CuO–NiO mixed nanocomposites

were synthesized by oxalate route and the catalysts were char- acterized by TG, DTA, XRD, TEM, nitrogen sorption and surface excess oxygen.

However, the highly catalytic conversion of IPA selectively to acetone over nano CuO–NiO mixed oxide systems, to our knowledge, has not yet been reported. Therefore, the chal- lenge of this work was devoted to synthesize acetone from IPA over CuO–NiO nanocomposites as competitive catalysts with those previously reported.

# Experimental

## *Materials*

Acetone, ethanol and oxalic acid dihydrate were of analyti- cally pure reagent and were used without further purification. Copper nitrate Cu(NO3)2.3H2O and nickel nitrate Ni(NO3)2.6H2O were supplied by WINLAB company for laboratory chemicals (>99%).

## *Preparation of pure and mixed oxide nanocomposites*

Pure Nano CuO and NiO were prepared by precipitation method as described previously [[12,28–30]](#_bookmark8). The precipitates were dried in the air overnight at room temperature to get the precur- sors. Pure CuO and NiO were obtained by calcination of their precursors from 400 up to 700 °C in a muffle furnace under air atmosphere for 3 h. CuO–NiO mixed oxide systems were pre- pared by co-precipitation method as described previously. The CuO–NiO nanocomposites with different ratios of NiO (1–50 wt.%) were obtained by annealing their precursors from 400 up to 700 °C in a muffle furnace under air atmosphere for 3 h.

## *Apparatus and techniques*

### *Electrical conductivity*

The electrical conductivity measurements were carried out with the method described by Said et al. [[31]](#_bookmark17). The powder sample is compressed between two platinum discs whose ends are con- nected to the electrometer. The whole sample is then placed inside an electric oven which opens along its generatrix. The oven is of a non-inductive type and has no temperature gra- dient in its central part. A temperature controller is adjusted within ±1 °C in the range of 100–300 °C. A Keithley Electrom- eter model 610 C instrument is used for measuring the resistivity.

### *Determination of basic sites*

The basicity of pure CuO, pure NiO and CuO mixed with 30 wt.% NiO catalysts calcined at 400 °C was investigated by the satu- ration of the basic sites with acetic acid according the following procedure: After measuring the conversion activity of IPA at steady state conditions over the investigated catalysts, at a re- action temperature of 200 °C, the catalyst was injected with different volumes of glacial acetic acid in the stream of reac- tants using N2 as a carrier gas and following up the activity variations. Moreover, the strength of the basic sites was studied by following up the change in the catalytic activity of IPA with the desorption of acetic acid from the presaturated samples with time at 200 °C.

### *Catalytic activity*

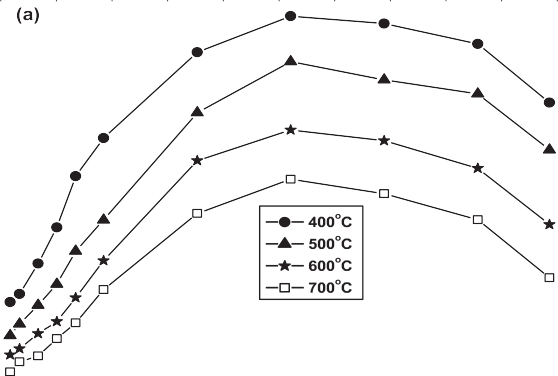
The activity of the catalysts toward dehydrogenation of iso- propyl alcohol (IPA) in the gas phase was carried in a conventional fixed bed flow type reactor. The gases after re- action were chromatographically analyzed by an FID on a Unicam ProGc using 10% PEG 400 glass column (2 m). The re- action conditions were: 0.5 g catalyst weight, 1.6% IPA reactant was fed into the reactor after nitrogen had been bubbled through thermostated IPA at a total flow rate of 110 mL min−1 and the reaction temperature was 200 °C. The measurements of conversion, yield and selectivity (in %) were made after 1.5 h to achieve steady state conditions.

# Results and discussion

## *Electrical conductivity measurements*

The electrical conductivity measurements of the pure CuO, pure NiO and CuO–NiO mixed nanocomposite catalysts calcined at

2CuO  2NiO  Cu2O  Ni2O3



(1)

or

Cu2  Ni2  Cu  Ni3 (2)

On the other hand, the decrease in the electrical conduc- tivity of the CuO–NiO mixed oxides upon increasing the wt.% NiO above 30 may correspond to a possible decrease in the mo- bility of charge carriers [[37]](#_bookmark22). The observed decrease in the electrical conductivity values of the pure and mixed oxide nanocomposites upon increasing the calcination tempera- ture from 400 to 700 °C may be attributed to the decrease in the values of SBET [[28,39,40]](#_bookmark16).

The effect of IPA vapor admission on the electrical con- ductivity of the pure oxides and CuO–NiO mixed catalysts calcined at different temperatures has been studied and the results are shown in [Fig. 1(b)](#_bookmark1). The experimental conditions used are similar to that used in catalytic activity runs. The results indicated that a similar trend (as in [Fig. 1(a)](#_bookmark1)) was observed, but



with lower log σ values. This behavior entirely reflects that IPA reactant decreases the charge carriers (Cu+ and Ni3+) created

in these catalysts via the electrons injected from IPA [[40]](#_bookmark23).

On the other hand, it was established that the value of ac- tivation energy provides a better information about the conductivity changes. Therefore, log σ values were plotted as a function of the reciprocal of absolute temperature and the

activation energies ΔEσ were calculated from the slope of the straight lines of the following Arrhenius relationship [[41]](#_bookmark24).

  \*eEKT

(3)

#### Fig. 1 – Variation of log σ with the % mixing of NiO for CuO– NiO nanocomposites calcined at 400–700 °C (a) without isopropyl alcohol, and (b) with isopropyl alcohol.

400–700 °C are carried out and graphically presented in [Fig. 1(a)](#_bookmark1). The results show that the electrical conductivity of the pure NiO is higher than that of CuO whereas both conductivities are lower than that of the mixed oxide nanocomposites for all calcination temperatures. In addition, the electrical conduc- tivity of the CuO–NiO mixed oxide systems increases with increasing the wt.% NiO reaching the maxima at 30%. Above this maxima, there is a continuous decrease in the conduc- tance values with further increase of NiO up to 50 wt.%. Moreover, upon increasing the calcination temperature from 400 up to 700 °C, log σ values decrease for the pure and mixed catalysts.

However, it is well known that CuO is a p-type semicon- ductor [[32]](#_bookmark18) and its electrical conductivity is due to the presence of Cu+ ions in its non-stoichiometric structure. Also, NiO behaves

also as a p-type semiconductor and its p-typeness exists from

a hopping mechanism [[33]](#_bookmark19) via an electron transfer between Ni3+ and Ni2+. So, the observed increase in the electrical con- ductivity of the CuO–NiO mixed oxide systems may be attributed to the increase in the concentration of charge car- riers (Ni3+ and Cu+) [[34,35]](#_bookmark20). The creation of these charge carriers due to a synergistic effect [[17,36–38]](#_bookmark9) can be explained accord- ing to the following proposed mechanistic synergism:

where σ is the electrical conductivity, σ\* is the pre-exponential factor, ΔEσ is the activation energy by conduction, K is the Boltzmann constant and T is the absolute temperature.

The graphical representations of the linear form of the Arrhenius plot for each sample are given in [Fig. 2(a–c)](#_bookmark2). It can be seen from these figures that log σ varies linearly against 1/T for the pure and mixed oxide systems, indicating that one and the same conduction mechanism is established for all the in- vestigated samples in the temperature range of 100–300 °C.

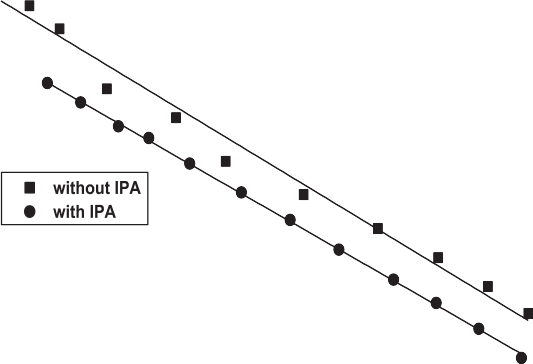
Values of ΔEσ are obtained by least square fitting of the ob- tained lines.

The activation energy values of the electrical conduction of the catalysts under investigation (with and without IPA) are calculated and cited in [Table 1](#_bookmark1). The results reveal that the values

of ΔEσ of pure oxides in the absence of IPA possess the highest values. Upon mixing CuO with 30 wt.% NiO, the activation

|  |  |  |
| --- | --- | --- |
| **Table 1 – Activation energy values (with and without IPA) for pure CuO, pure NiO and CuO mixed with 30 wt.% NiO nanocomposites calcined at 400 °C.** | | |
| Catalyst | ΔEσ (eV)  Without IPA | With IPA |
| Pure CuO | 0.83 | 0.79 |
| CuO mixed with 30 wt.% NiO | 0.12 | 0.03 |
| Pure NiO | 0.50 | 0.40 |
| IPA: Isopropyl alcohol. | | |







#### Fig. 2 – Variation of log σ versus 1/T for (a) pure CuO, (b) pure NiO, and (c) CuO mixed with 30 wt.% NiO, nanocomposites calcined at 400 °C.

energy progressively decreases, reaching to (0.12 eV). This de- crease in ΔEσ value of CuO mixed with 30 wt.% NiO catalyst reflects again the pronounced increase in the creation of the charge carriers (Cu+ and Ni3+). On the other hand, the activa- tion energy values measured in the presence of IPA indicated that the ΔEσ variation with NiO loading exhibits similar be- havior, but with lower values than that measured in the absence

of IPA. The reason for such behavior is due to the injected electrons by IPA into the catalyst surface during the dehydro- genation process [[35]](#_bookmark21). These electrons are trapped in or on the catalyst surface, causing a downward bending of the conduc- tion band toward the Fermi level and consequently decrease

the activation energy ΔEσ. Therefore, the decrease of ΔEσ should facilitate the electron exchange between the conduction and

valence bands [[42]](#_bookmark25). Moreover, its width is an important factor in controlling the redox mechanism, the number of chemisorbed molecules in the course of a catalytic reaction and the nature

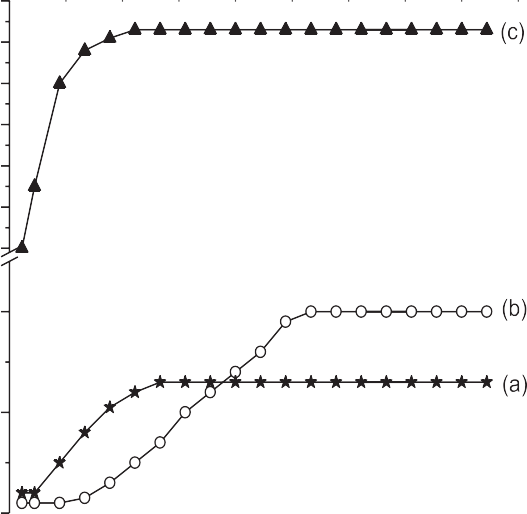
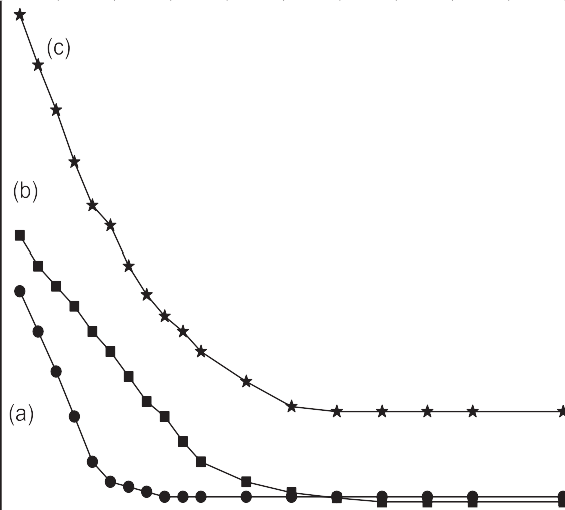
of the chemical bonds formed between the molecules and the catalyst surface. These factors control at the same time the ac- tivity and selectivity of the catalyst as well as the mechanism of the catalytic reaction.

## *Basicity of catalysts*

Saturation of the active basic sites by acetic acid over pure oxides and CuO mixed with 30 wt.% NiO catalysts calcined at 400 °C during the IPA conversion was studied and the results obtained are represented in [Fig. 3(a–c)](#_bookmark3). Curve (a) shows the in- fluence of acetic acid additions on the catalytic dehydrogenation of IPA over NiO catalyst. It appears that the saturation of the basic sites on NiO by acetic acid was obtained after injection of 1.8 μL. In addition, the IPA conversion decreases from 44% to 1% with increasing the volume of acetic acid up to 3 μL. Curve

(b) shows the influence of acetic acid injection on the IPA









#### Fig. 3 – Effect of acetic acid injections on the catalytic conversion of IPA to acetone over (a) pure NiO, (b) pure CuO and (c) CuO mixed with 30 wt.% NiO nanocomposites calcined at 400 °C.

#### Fig. 4 – Activity variation of IPA with reaction time over presaturated catalysts with acetic acid, (a) pure NiO, (b) pure CuO and (c) CuO mixed with 30 wt.% NiO nanocomposites calcined at 400 °C.

activity over pure CuO. It illustrates that the IPA conversion de- creases from 54% to 1% with increasing the volume of acetic acid reaching its steady state after addition of 4 μL. On the other hand curve (c) represents the effect of poisoning the active basic sites CuO mixed with 30 wt.% NiO. It shows that upon addi- tion of acetic acid by 3.5 μL, the conversion activity decreases from 98% to 19%. These results revealed that the acetic acid approximately saturated the available active basic sites re- sponsible for dehydrogenation of IPA over pure oxides whereas 19% is still working in case of the mixed oxide (30 wt.% NiO). In addition, the distribution of the basic sites on the mixed oxide is quite improved.

In an attempt to shed light on the strength of the basic sites over the catalyst presaturated with acetic acid, the activity varia- tion of IPA at the reaction temperature of 200 °C was measured with time and the results are presented in [Fig. 4(a–c)](#_bookmark3). Curve

(a) shows that desorption of acetic acid from the surface of pure NiO has been almost rapid where the conversion of the IPA and the yield of acetone attained a steady state after 12 min. The value of % conversion and yield of acetone at the steady state was 13% only. This means that about 30% of acetic acid was retained on the catalyst surface as indicated in [Fig. 3(a)](#_bookmark3) for the unsaturated NiO. This behavior reflects that the strength

of ≈ 71% of the available basic sites on NiO is strong.

Curve 4(b) shows that the desorption of acetic acid from CuO catalyst occurs slowly where the steady state value of conver- sion was obtained after 25 min. Furthermore, the conversion at the steady state reaches 20% compared to 54% for the un- saturated catalyst. These results reflect that the basicity of CuO catalyst is reduced by 34%. This behavior suggests that the re-

tained acetic acid (34%) on the surface of pure CuO may be due to the presence of ≈ 63% strong basic sites which need higher temperature to overcome the binding energy between the acetic

acid and the strong basic sites. According to the results ob- tained from the desorption of acetic acid from the surfaces of both pure oxides, it is interesting to mention here that the amount of acetic acid retained is approximately equal. This may support that the distribution of the basic sites over both oxide catalysts surfaces almost the same. In addition, the major strength of these sites is strong and uncomfortable for con- version of IPA to acetone.

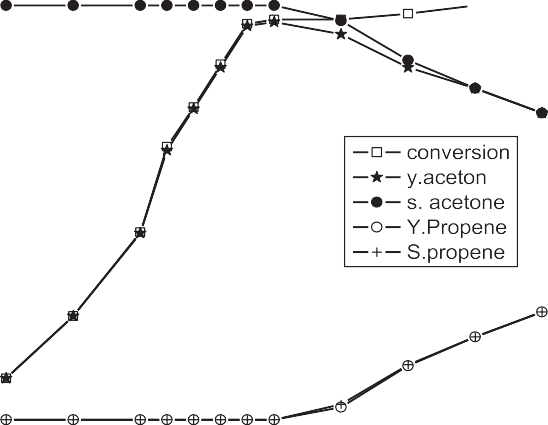
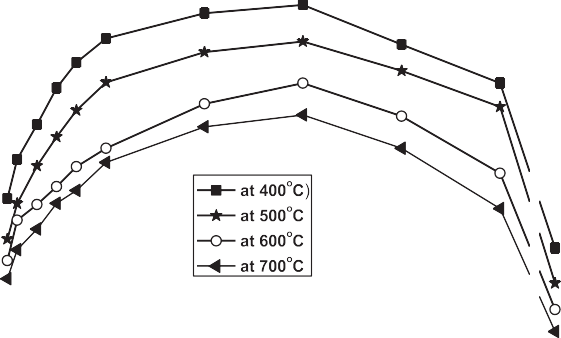
Curve 4(c) represents the effect of time on the desorption of acetic acid from CuO mixed with 30 wt.% NiO catalyst. It shows that desorption of acetic acid occurs rapidly and hence the conversion of the IPA and the yield of acetone increase rapidly with time. Also, the sample nearly restores its activ- ity ~95% after about 10 minutes. The remained value of ~3% may be attributed to the adsorption of acetic acid on strong basic sites on the catalyst surface.

The comparison between the desorption studies of acetic acid over the catalysts under investigation reveals that CuO mixed with 30 wt.% NiO catalyst restored almost its original activity rapidly compared with CuO and NiO catalysts. This con- firms that the creation of weak and intermediate basic sites on the surface of the catalyst containing 30 wt.% NiO en- hances the reaction of IPA to acetone and consequently the addition of NiO into CuO promotes its basicity to be comfort- able for dehydrogenation of IPA [[43–45]](#_bookmark26).

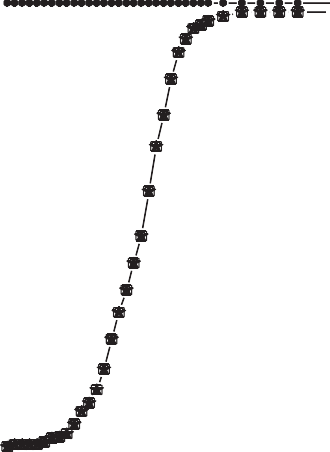
## *Catalytic activity*

The dehydrogenation of IPA over CuO–NiO nanocomposite systems calcined at 400, 500, 600 and 700 °C was carried out and the results are presented in [Fig. 5(a)](#_bookmark4). It shows that IPA con- verts to acetone (major) and propene (minor) over pure CuO and CuO mixed with 1 and 3 wt.% NiO whereas upon









#### Fig. 5 – (a) Variation of IPA conversion with the % mixing of NiO for CuO–NiO nanocomposite catalysts calcined at

**400–700 °C, (b) effect of reaction temperature, and (c) effect of reaction time on the catalytic dehydrogenation of IPA over CuO mixed with 30 wt.% NiO nanocomposites calcined at 400 °C.**

increasing the wt.% NiO up to 50%, IPA conversion proceeds to acetone only with selectivity 100%. From these results, one can observe that pure CuO possesses catalytic activity toward IPA conversion to acetone higher than that of pure NiO [[46]](#_bookmark27). Such results may be correlated with the higher values of the surface area and the amount of surface excess oxygen as dis- cussed earlier [[28]](#_bookmark16). On the other hand, it is well known that NiO precedes the dehydrogenation reaction for hydrocarbons and alcohols, thus the formation of acetone as a major product is quite expected. Hence, the observed increase in the selec- tivity toward acetone formation due to the increase of NiO ratio suggests an increase in the number of active sites respon- sible for dehydrogenation of IPA. So, the addition of NiO (1–30 wt.%) into CuO as presented in [Fig. 5(a)](#_bookmark4) leads to a con- tinuous increase in the conversion of IPA and selectivity toward acetone reaching a maximum yield of (98%) over the catalyst

containing 30 wt.% NiO. Above this maximum, further addi- tions of NiO lead to a decrease of both conversion and yield of acetone. The observed remarkable catalytic activity of the mixed oxide catalysts may be not only due to the presence of the one component sites [[47,48]](#_bookmark28) Cu2+–Cu+, Ni3+–Ni2+, but also due to the creation of new active ion pairs sites (Cu+–Ni3+). Such active sites which existed via the synergism [[17,36–38]](#_bookmark9) are ex- hibiting weak and intermediate strength as discussed above in the basicity section. The improvement in their strength is playing a major role for the dehydrogenation of IPA. In this context, El-Molla et al. [[25]](#_bookmark13) have reported that the presence of medium-strength basic sites on the surface of catalyst cata- lyzes the rate determining step of the energetically easier Hα abstraction from adsorbed 2-propoxide forming carbanion in- termediate that finally leads to acetone. In addition, it was reported that Cu2+, Cu+ and Cu0 are the active and selective sites

responsible for IPA dehydrogenation into acetone over CuO/MgO catalysts [[25,49]](#_bookmark13). They concluded that, at a reaction tempera- ture of 175–275 °C, the different investigated solids behave mainly as a dehydrogenation catalyst yielding acetone beside MIBK (mesityl isobutyl keton) formation via aldol condensa- tion mechanism. At lower temperature, 175 °C, the reaction is completely selective to MIBK, and on increasing the reaction temperature up to and above 200 °C, selectivity to acetone starts to begin reaching to its high value (maximum 92%) at a reac- tion temperature of 275 °C. Moreover, the catalytic performance of CuO/MgO treated with K2O catalysts toward IPA conver- sion, at a reaction temperature of 150–400 °C, was reported [[23]](#_bookmark11). They stated that the system behaved as selective catalyst for dehydrogenation of isopropyl alcohol to acetone (major) with

selectivity >80% besides propene as a minor product.

Therefore, from the comparison between our CuO–NiO mixed oxide nanocomposite systems with that previously re- ported systems [[22–27]](#_bookmark10), we can conclude that our system has two advantages: (1) the reaction temperature for complete IPA conversion to acetone, 200 °C, is relatively low compared with the previous reported systems, and (2) the selectivity to acetone as the desired product, reached 100%. These two advantages make our system, which was prepared by a facile route (nanocomposite), a competitive one than those reported previously.

In fact a good contribution of our system for proceeding IPA to acetone may be supported with the results of electrical con- ductivity as illustrated from the behaviors of both [Fig. 5(a)](#_bookmark4) and

[Fig. 1(a and b)](#_bookmark1) with the wt.% NiO. So, the decrease in ΔEσ as shown in [Table 1](#_bookmark1) should enhance the electron exchange

between IPA and the catalyst surface during the reaction and consequently increases the conversion and selectivity toward acetone formation. Moreover, the decrease in the catalytic ac- tivity of the mixed oxides above 30 wt.% may be due to the decrease in their conductivity and the mobility of charge car- riers [[37]](#_bookmark22). On increasing the calcination temperature above 400 °C, the results of catalytic activity of the pure and mixed oxides exhibit similar behavior as that observed for the cata- lysts calcined at 400 °C but with lower values. These results can be discussed in terms of the possible change in the con- centration of cation pairs acting as active sites for the catalyzed reaction [[50]](#_bookmark29) and a possible decrease in the SBET of the cata- lysts [[28,39,51]](#_bookmark16).

To achieve the best conditions that will give the maximum yield of acetone over the catalyst containing 30 wt.% NiO cal- cined at 400 °C, the factors affecting the catalytic conversion of IPA into acetone were studied in the following.

### *Effect of catalytic reaction temperature*

The effect of reaction temperature on the catalytic conver- sion of IPA in the range 100–300 °C was studied. The percentages of conversion, yield and selectivity are represented in [Fig. 5(b)](#_bookmark4). It shows that the IPA conversion and acetone yield increase monotonically with increasing the reaction temperature reach- ing the value of 98% at the reaction temperature of 200 °C. Above 200 °C, propene started to appear and it increases with in- creasing the reaction temperature up to 300 °C. These findings indicate that the active basic sites responsible for the dehy- drogenation of IPA are working well at the reaction temperature of 200 °C.

### *Effect of catalyst weight*

The effect of the weight of catalyst containing 30 wt.% NiO on the percentages conversion, yield and selectivity of IPA dehy- drogenation was studied at 200 °C and the results are shown in Fig. S1, Supplementary material. It shows that the % con- version of IPA increases with increasing the catalyst weigh from (0.1–0.8) reaching its steady state value of 98% with selectiv- ity of 100% toward acetone at a catalyst weight of 0.5 g. These data reflect that as the catalyst weight increases, the number of active basic sites responsible for the dehydrogenation process is normally expected to increase. On further increase in the catalyst weight up to 0.8 g a little amount of propene not ex- ceeding 2% is observed.

### *Catalyst stability*

To evaluate the stability of the catalyst containing 30 wt.% NiO, the dehydrogenation of IPA to acetone was investigated and the results are presented in [Fig. 5(c)](#_bookmark4). It shows that the maximum conversion of IPA (98%) with selectivity 100% to acetone was obtained after 60 min from the introduction of reactant fol- lowed by a steady state till 60 h, which means that the catalyst under test has a good stability toward production of acetone.

# Conclusions

The main conclusions derived from the results obtained can be summarized as follows:

* Pure and mixed solids of CuO and NiO nanocomposites have been synthesized by direct, facile co-precipitation method (oxalic acid rout).
* The addition of NiO to CuO resulted in increasing the elec- trical conductivity, whereas the activation energy of conduction decreases.
* Incorporation of NiO with CuO creates more weak and in- termediate basic sites which play the main role for the selective dehydrogenation of IPA into acetone.
* The catalyst containing 30 wt.% NiO calcined at 400 °C pos- sesses the highest catalytic activity with 98% conversion and 100% selectivity to acetone at the reaction temperature of 200 °C, and it exhibits good stability with duration time up to 60 h.
* The remarkable activity and selectivity of the catalysts under investigation are correlated well with both electrical con- ductivity and basicity.

# Appendix: Supplementary material

Supplementary data to this article can be found online at [doi:10.1016/j.ejbas.2016.08.004](http://dx.doi.org/10.1016/j.ejbas.2016.08.004).

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