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SOLUTION COMBUSTION SYNTHESIS OF (Co₄Ni)Cr₂O₄ PIGMENTS: INFLUENCE OF INITIAL SOLUTION CONCENTRATION

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Abstract

Initial solution concentration effect was studied on the synthesis of mixed spinels Co₁₋₄NiₓCr₂O₄ (0≤ₓ≤1) obtained by Solution Combustion Synthesis. Fd-3m spinel structure was developed in all range of compositions analysed, regardless of the concentration. However, structural characteristics such as ion rearrangement and crystal size showed a noticeable dependence on the initial concentration, being the spinel network more ordered and with higher crystallite size as the concentration increased. Cell parameter, however, presented dependence on composition but not on initial solution concentration.

All as-synthesized pigments showed a significant colouring power in ceramic glazes without any significant influence of initial solution concentration. Therefore, a second thermal treatment was not needed. The coloured glazes covered a broad range of tones in the green section of colour space, which evolved as a function of composition.

Keywords: Powders: chemical preparation (A); colour (C); spinels (D); solution concentration
1 Introduction

Submicronic materials with characteristic structural dimensions have been of interest for more than 20 years and continue to attract attention because of their unique properties [1]. Such materials have found a variety of applications in different branches of industry including microtechnology, biotechnology and surface coatings, as well as energy storage and conversion devices, such as fuel cells.

In the case of the traditional ceramic sector, submicronic powders are greatly welcomed when inkjet technology is wanted to be applied. Nanosized ceramic pigments are needed in order to develop a good quality inks, which are able to pass through the printer nozzles without damaging the mechanical system. At the same time, pigments must be able to develop intense colours. These inks are currently manufactured by micronizing conventional ceramic pigments down to average diameters of 0.2-0.6 µm [2]. To prevent print head nozzles from clogging, the most important requirement is to ensure that 99% of the pigment particles are less that 1µm in diameter [3,4]. This fact demands a very different approach to the comminution stage: from controlling average particle size during jet milling (conventional pigments) to ensure that practically all particles are submicronic.

A wide variety of techniques are used to obtain materials at microscale in industry. The most basic one used in ceramic sector is high-energy milling. However, this process is energy intensive and frequently consumes spare parts of the mill, resulting in high costs. Furthermore, this milling process presents the drawback of damaging ceramic pigment’s crystalline structure and, as a consequence, their tinting strength is reduced [5]. All these inconvenient have encouraged companies and research centres to foster in the development of innovative synthetic routes with a lower cost of production of nanopigments with the optimum structure.

**An alternative methodology that has stood out because of its capability for obtaining fine particles** is the so-called solution combustion synthesis (SCS). This method is a combination of combustion and a reactive solution approach. SCS involves a self-sustained reaction in a
solution of metal nitrates and different fuels as urea or glycine. The reaction between fuel and oxygen-containing species formed during decomposition of nitrates provides conditions for rapid high-temperature interaction [6]. The initial liquid solution of nitrate precursors, after fast preheating to moderate temperatures (500°C) self-ignites over the whole volume (volume solution combustion mode (VSC)) leading to the formation of a very spongy mass with tailored composition.

**SCS was reported to be a technology able to** synthesize simple and complex oxides used as pigments or advanced ceramics [7-9]. It contributes to nanopigment development with special features [10-13] because of its characteristic combustion process. On one hand, the reactant mixing is really effective since starting media is an aqueous solution, which favours cations to be mixed on a molecular scale. On the other hand, under the appropriate conditions the high temperatures (~1500°C) reached in the solution combustion synthesis method allow obtaining oxide products, as ceramic pigments are, with a high purity and good crystallinity [6]. Finally, as the whole combustion process is carried out in a few minutes (short reaction time), with the release of secondary products as a high volume of gases, there is an inhibition of particle size growth and a spongy microstructure development. These facts contribute to obtain a solid product with a very low bulk density, easy to mill.

Despite all advantages that SCS technology provides, the published studies are mainly focused on characterizing synthesized products and practically no information is available about the effect of modifying the process variables like selected fuel, initial solution concentration or maximum temperature. **Therefore,** the aim of this study is centred on evaluating the effect of the initial solution concentration over the synthesis of pigments Co$_{1-x}$Ni$_x$Cr$_2$O$_4$ carried out by means of solution combustion synthesis. The solid solutions between spinels CoCr$_2$O$_4$ and NiCr$_2$O$_4$ were selected as a result of being two of the most frequently pigments used in the ceramic industry, according to the Color Pigments Manufacturer Association (CPMA) [14]. The study is also intended to evaluate whether the final pigments are chemically and thermally stable against the ceramic glaze once applied.
2 Material and methods

\[\text{Co}_{1-\Psi}\text{Ni}_\Psi\text{Cr}_2\text{O}_4 (0 \leq \Psi \leq 1, \text{ in steps of 0.2})\] pigments were synthesized from their corresponding \(\text{Co, Ni and Cr nitrates mixed with urea as fuel and water as a solvent}\) (All reactants used were from Panreac Química, S.A.U. Spain). Urea proportion was calculated following the reaction stoichiometry in order to reach the combustion of the whole mixture. Table I shows the initial solutions’ compositions. The proportion of water was changed in order to study the initial solution concentration effect in the synthesis of the pigments. Consequently, three complete series with different initial solution concentration of spinel precursors were prepared: 2.4 M, 1.2 M and 0.6 M, which from now on will be referenced as high, medium and low solution concentration, respectively.

The first aim was to maintain the volume of solution in each experiment constant, in order to avoid increasing the number of variables to be taken into account. However, due to the high volume of the combustion product obtained in some experiments, it was necessary to readjust the original solution volume so as to avoid any contamination of the kiln chamber and their refractories.

All solutions were poured in a 700-mL pyrex container of 14 cm in diameter which was inserted in a preheated kiln at 500 °C (BLF 1800, Carbolite Furnaces Ltd, UK) during 20 min of soaking time. Then, the kiln was turned off and the as-synthesized pigment was progressively cooled down to ambient temperature.

Every pigment was wet milled to break the agglomerates in a ball mill using water as a fluid and agate jars (Pulverisette 5, Fritsch GmbH, Germany). Afterwards, the pigments were dried under infrared lights and sieved with a 200-µm mesh.

Pigments were characterized from a chemical, microstructural, mineralogical, physical and colourimetric point of view. Chemical composition was determined by an energy-dispersive X-ray microanalysis instrument (EDX) (Genesis 7000 SUTW, EDAX, USA) coupled to a FEG-SEM (Field Emission Guns-Scanning electron microscope) (QUANTA 200F, FEI Co, USA)
with which the microstructural characterisation was carried out. Adsorption/desorption
isotherms and specific surface area values were determined according to the BET method
(Brunauer-Emmet-Teller) using nitrogen gas as adsorbate (Tristar 3000, Micromeritics, USA).
Additionally, pore size distribution was obtained from the adsorption isotherms using the
Barrett–Joyner–Halenda method (BJH).

Identification of crystalline phases and the posterior measurement of crystallite size and cell
parameters evaluation were carried out by XRD (Theta-Theta D8 Advance, Bruker, Germany),
with CuK radiation (λ = 1.54183 Å). The generator applied an intensity light source of 45 kV
and 40 mA. XRD data were collected in a 2θ from 5 to 90° with a step width of 0.015° and a
counting time of 1.2 s/step by means of a VÂNTEC-1 detector. Raw data were refined by
Rietveld method using 4.2 version of the Rietveld analysis program DIFFRACplus TOPAS. A
pseudo-Voight function to describe peak shapes was assumed. The refinement protocol included
the background, the scale factors and the global-instrument, lattice, profile and texture
parameters.

To evaluate colour development, pigments were mixed in a proportion 2/98 wt% into a
transparent single-fired porous tile glaze (chemical composition: 0.5% Na₂O 4.0 % K₂O, 15.3%
CaO, 0.9 MgO, 9.0% ZnO, 7.4% Al₂O₃, 3.0% B₂O₃, 59.5% SiO₂) and fired in an electric
laboratory kiln according to a thermal cycle of single-fired floor tiles (maximum temperature
1100 ºC and 6 min of soaking time). Spectrophotometric curves of the glazed tiles were
obtained (Color Eye 7000A, X-Rite Inc, USA), and CIELab* chromatic coordinates were
calculated using CIE Illuminant D65 and CIE 10° standard observer.

3 Results and discussion

As-obtained pigments presented a highly fluffy texture, filling the whole synthesis glass
container. Bulk density was quite low which made them easy to disaggregate with a low effort.
Pigment colours presented an evolution around the green tones, from a bluish green to a greyish
green.
3.1 Chemical composition

EDX analysis of some of the synthesized pigments showed a good correlation with theoretical values according to the initial solution compositions (Fig. 1). When pure spinels were synthesized ($\Psi=0.0$ and $\Psi=1.0$) the differences among values were smaller than in solid solution spinels. However, it has to be taken into account that EDX method is semiquantitative and little oscillations around the exact value can be produced in some samples due to the lack of planarity of the spongy samples. This latter phenomenon is responsible for modifying the signal received by EDX detectors in order to determine the molar content. The lack of planarity of the samples affects particularly the oxygen percentage determination. In previous reports published about solid solutions of spinels, a good correlation between experimental and theoretical composition was demonstrated in all $\Psi$ range [15]. For that reason, not all pigments developed in this study have been analysed by EDX, only the most representative pigments in this case.

According to obtained results, from a chemical point of view, SCS method allows obtaining homogeneous solid solutions of spinels with molar compositions practically adjusted to the theoretical ones.

3.2 Crystalline structures

Diffraction patterns showed that the main crystalline phase obtained in all the range of compositions was of spinel type, concretely the Fd-3m class (face-centred structure) [16]. However, as Ni$^{2+}$ content increased in the composition ($\Psi>0.0$), a rise in the proportion of a secondary phase with eskolaite structure (Cr$_2$O$_3$) [17] was also observed in the samples. Fig. 2 shows the complete diffraction patterns for all composition range of (Co,Ni)Cr$_2$O$_4$ samples with both crystalline structures (spinel and eskolaite) identified by XRD at 1.2M concentration. According to Rietveld data, the eskolaite increasing effect was more pronounced when the solution concentration decreased (Fig. 3). This phenomenon pointed out that not all the trivalent ions were integrated into the spinel structure, although the percentage of eskolaite was not higher than a 5% in any case. Despite this circumstance, it was possible to consider that SCS
technique is suitable for synthesizing spinel pigments because spinel-type industrial ceramic pigments frequently contain low proportions of unreacted oxides.

On the other side, all synthesized pigments presented a high crystallinity, higher than 98 ± 4% according to Rietveld analysis. Practically no amorphous phase was observed in the samples. Nevertheless, spinel structure evolution in the solid solutions indicated the presence of an improvement in ion rearrangement to obtain a more defined structure, reaching a maximum for the spinel with $\Psi=0.8$ (Fig. 4). This phenomenon was deduced after observing that spinel main peak evolved towards higher heights and narrower and more defined shapes. Such behaviour suggested that the progressive enrichment with Ni$^{2+}$ ion improved the capacity of obtaining a more ordered spinel structure. On the other hand, the absence of Co$^{2+}$ provokes an obvious negative effect on spinel structure development because, as $\Psi$ increases, a trend to increase eskolaite phase percentage at the cost of spinel formation was also observed.

Regarding the initial solution concentration, an important effect was detected based on the progression of the main peak intensity of the diffraction patterns because, as the initial solution concentration increases, the intensity peak becomes higher and more defined, indicating the rearrangement of a more ordered spinel structure. In Fig. 4 diffraction patterns of the pigments synthesized with the most concentrated solutions showed the highest peaks, which meant a better capability to obtain a more ordered structure. Such behaviour can be interpreted considering that as the solution concentrates, the volume of liquid in the synthesizing container is reduced. This phenomenon yields a faster heat transfer, which evaporates water in shorter times, limiting the possibility of ion segregation in the drying mixture and improving reactant contact during the combustion process. It can be supposed that as the concentration increases, the bulk density of the dry mixture of reactants is higher, so the energy generated by the reaction allows process to reach higher temperatures, which facilitates ions mobility and their rearrangement as spinel network.

Taking into account Fig. 3 and 4, it is possible that a relation between obtaining a more ordered structure and the eskolaite proportion exists. It could be related to different effects taking place
during the synthesis. On one hand, it would be possible that some chromium could not be able
to join the spinel structure. Therefore, the chromium positions would either be filled by other
ions, such as Co$^{3+}$ or be left as a vacant. On the other hand, due to the important effect of the
fluorescence in the diffraction pattern of the pigments, it could be possible that some Co and Ni
compounds (oxides or hydroxides) were also present in the samples, but in proportions under the
detection limit of XRD [18-21]. Consequently, the molar relation Co/Ni versus Cr was
maintained in the spinel, although the network order was not favoured. To distinguish between
both effects, it would be necessary the use of analytical techniques, capable of detecting
crystalline phases in very low proportions.

The crystallite size was calculated in each one of the pigments (Fig. 5). The data showed a
progressive decrease of crystallite size as the cobalt content was reduced (from $\Psi=0$ to $\Psi=0.8$).
However, the absence of cobalt ($\Psi=1$) changed the global trend and crystallite size sharply
plummeted, up to practically a half the previous data points. The cobalt enrichment in the
structure seemed to favour the growth of the formed crystallite, whereas the Ni$^{2+}$ ion exerted the
opposite effect.

Molar concentration also influences crystallite size evolution since the most concentrated
solution generates pigments with bigger crystallites than the diluted ones. This result agrees
with the proposed hypothesis about the effects of thermal treatment based on X-ray diffraction
patterns. According to which, it was interpreted that a higher solution concentration limits the
possibility of ion segregation and, at the same time, improves reactant contact during
combustion process, favouring ion rearrangement as spinel network.

No clear trend was observed in lattice parameter data against initial solution concentration (Fig.
6). As expected, the lattice parameter showed a clear relation with composition, as predicted by
Vegard’s law. Theoretical lattice parameters evolved from 8.340 Å for CoCr$_2$O$_4$ spinel [22] to
8.315 Å for NiCr$_2$O$_4$ spinel [23]. However, a positive deviation from the theoretical line was
observed. In fact, experimental values presented a curved shape in which a positive deviation
from theoretical values is observed in $\Psi<0.8$, but in $\Psi=1$, the trend inverted, being the
theoretical values higher than the obtained ones. Deviations from theoretical path were previously described in some works by Mestre et al. [9], Shou-Yong et al. [24] and Nlebedim et al. [25], but no explanation has been proposed.

The general trend of lattice parameters of the cell was consistent with the lower ionic radius of the Ni$^{2+}$ (0.49 Å) with respect to the Co$^{2+}$ (0.56 Å) in tetrahedral coordination [26]. The lower the ion size is, the shorter the distance between ions becomes and, therefore, the lattice parameter decreases since dimensions of the spinel cell are reduced [15].

3.3 Morphological characterisation

The study of morphology and grain size was carried out over the as-synthesized samples, prior to the wet-milling process, since in previous studies some modifications in the composition and structure were observed due to the hydration of certain non-perfectly reacted phases [15].

Obtained micrographies demonstrated the development of round-shaped grains in all samples (Fig. 7), regardless of the composition and solution concentration. In NiCr$_2$O$_4$ spinels case ($\Psi=1$) grains were near spherical, whereas a cobalt enrichment promoted more angular grains with a geometry more similar to theoretical crystalline habit of spinel [27].

Molar concentration exerted a significant influence in grain development since in all the compositions tested an increase in concentration favoured a grain growth in different degrees. The effect observed was similar to the effect over the crystallite size, but not equal. It can be assumed that an improvement in ion homogeneity, thanks to higher solution concentration, agrees with the fact that both the crystallite and, consequently, the grain experienced a growth [28]. Besides the general growth observed, it has to be highlighted the specific case of $\Psi=0.6$, where a singular phenomenon is produced. The grain size difference between the product of the diluted solution and the product of the concentrated one is the highest in comparison to rest of the samples. Grains in the pigment obtained from the concentrated solution with $\Psi=0.6$ grew in a considerable way, acquiring a shape that practically reflects the theoretical crystalline habit of spinel. These grains are pretty angular and their faces fitted together perfectly, reducing the
intragrain porosity of the system. By contrast, the grain size of the sample obtained with $\Psi=0.8$ and 2.4 M (the highest crystallite size), only shows a grain size slightly higher than the sample with $\Psi=1.0$. This fact demonstrates that grain and crystallite sizes are not influenced in the same way by composition and concentration.

Specific surface data of the pigments showed lower values when initial solutions were more concentrated, which totally corroborates the behaviour observed in scanning electron microscopy analysis (SEM). The samples with most ordered structure presented less intergranular porosity and, consequently, lower specific surface area. At the light of the results (Table 2), it can be certainly said that all analysed samples exhibited well-sintered grains since all specific surface values evolved in a narrow range unlike the observed values in previous works in which they can reach values up to 200 m$^2$/g in CoAl$_2$O$_4$ spinels [9, 15]. In addition, although in a subtle way, the absence of Ni$^{2+}$ in the composition improved sintering because specific surface area reduced slightly.

To study the porosity of pigments, BJH curves were calculated from the analysis carried out on isothermal curves (Fig. 8). No significant differences were observed among obtained curves. Nevertheless, mention can be made of the fact that the presence of certain small-size porosity (< 10 nm) in the pigments obtained from 0.6M solutions could be responsible for the subtle increase observed in specific surface area.

### 3.4 Colouring power

All synthesized pigments presented a high colouring power in the selected glaze. No defects were observed on the surfaces like pinholes or colour heterogeneities. These facts are indicative of a good integration of the pigments into the ceramic glaze, without any unfavourable interaction. This behaviour is of relevant importance in order to determine the stability degree of ceramic pigments synthesized by SCS.

Final glazes covered a broad colour palette in the green sector of colour space evolving with the composition, from a blue-green tone to a greyish-green. The spectrophotometric curves (Fig. 9)
showed a progressive variation as the pigments enriched in Ni$^{2+}$, which can be related with the absorption bands corresponding to electron transitions of Co$^{2+}$, Ni$^{2+}$ and Cr$^{3+}$ ions located in spinel crystals [29, 30]. In fact, as Ψ increased, the reflectance band characteristic for blue colour was reduced (350<\lambda<450 \text{ nm}), which was promoted by the gradual elimination of Co$^{2+}$ [31]. Red reflectance band (\lambda>700 \text{ nm}) also evolved negatively as Ψ increased. On the other hand, the yellow reflectance band (550<\lambda<680 \text{ nm}) was strengthened because of the Ni$^{2+}$ content increase. All these changes generated less bluish colours with higher green components.

The CIELab* coordinates, increased progressively with Ψ but with different trends. L* coordinate experienced a 10 units’ increase, indicative of the development of brighter colours as the proportion of Ni$^{2+}$ was higher (Fig. 10 a). b* coordinate showed the gradual elimination of blue component and, as a consequence, the evolution towards colours with yellow component (Fig. 10 b). The range of variation in this case was the highest, practically 20 points, which was a significant change in this coordinate. Nevertheless, the coordinate that seemed to experience the smallest change regarding its evolution with Ni$^{2+}$ was a* coordinate (Fig. 10 c). Its evolution showed a low-defined minimum for Ψ=0.2, and the green component reduces progressively as the proportion of Ni$^{2+}$ increases. When Ψ=1.0, a significant change is produced, due to the complete elimination of cobalt, which resulted in a sharp decrease in green component. Such variation was produced in an interval of 8 points, a minor change in comparison to the rest of the cases, but visually appreciable in the resulting colour. In addition, this coordinate showed an effect on initial solution concentration, since the green component was higher as the initial solutions were more concentrated.

According to colorimetric results, it was possible to obtain a practically pure green shade when spinel composition moved around Ψ=0.5 since no traces of neither blue nor yellow tones appeared in the colour.
4 Conclusions

Molar concentration of the initial solution effect on the evolution of the solution combustion synthesis has been evaluated in the present study. In this case, SCS method allows synthesizing the Fd-3m spinel structures, characteristic of ceramic pigments with the solid solution composition (Co,Ni)Cr$_2$O$_4$. The presence of Ni$^{2+}$ in the composition favours the appearance of a secondary phase with eskolaite structure, in a percentage lower than 5% wt. Certain influence of initial solution concentration was observed related to crystallinity parameters of the pigments. The products of the most concentrated solutions showed a more ordered spinel network and a higher crystallite size than the diluted ones. Spinel lattice parameter was not influenced by the initial solution concentration. Nevertheless, it was directly affected by composition attending to Vegard’s law, regardless the positive deviation observed from theoretical trend. Pigment morphology was modified depending on both composition and mixture concentration. The most Ni-rich spinel presented spherical grain shapes; however, samples with cobalt showed more angled grains, which were more similar to theoretical crystalline habit of a perfect spinel. For composition $\Psi=0.6$ and the highest concentration, grain sizes experienced a noticeable growth with well-defined grain borders and a compact structure.

Colouring power was intense in all cases and it was not highly affected by the concentration of initial solution. The colours generated in the selected glaze covered a broad palette of green shades. A progressive Co$^{2+}$ content reduction provokes an increase in luminosity, a progressive reduction in green component and a change from a bluish hue towards a greenish one. In this study it has been possible to develop a green tone practically pure with the composition $\Psi=0.5$.

SCS method has been validated as an effective technique for spinel pigments synthesis since the products have high colouring power, and they are stable against variations in the molar concentration of the precursor’s solution.
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Figure captions

Figure 1 EDX chemical analysis of SCS spinels comparing experimental and theoretical values

Figure 2 XRD phase identification for all composition range of (Co,Ni)Cr₂O₄ samples at 1.2M concentration (○ spinel, ● eskolaite)

Figure 3 Mass percentage of secondary eskolaite phase in pigments versus composition and initial solution concentration

Figure 4 Evolution of the XRD main peak of the pigments versus Ψ and concentration of initial solution

Figure 5 Evolution of crystallite size depending on parameter Ψ

Figure 6 Comparison of cell parameters of the spinel with the prediction of Vegard’s law, based on the ICCD data from the spinels CoCr₂O₄ (Ψ=0.0) and NiCr₂O₄ (Ψ=1.0)

Figure 7 Micrographies obtained by SEM of selected pigments

Figure 8 BJH curves showing pore size distribution of selected samples

Figure 9 Reflectance curves of the glazes that contain the synthesized pigments (2.4M samples), indicating the bands related to electron transitions of Ni²⁺, Co²⁺ and Cr³⁺ ions located in spinel crystals

Figure 10 Evolution of the glaze’s chromatic coordinates versus studied parameters of incorporated pigments (Ψ and initial solution concentration): a) L*, b) b* and c) a* coordinates
### Table 1 Composition of initial solutions

| Ref. | $\Psi$ | Urea (g) | Co(NO$_3$)$_2$·6H$_2$O (g) | Ni(NO$_3$)$_2$·6H$_2$O (g) | Cr(NO$_3$)$_3$·9H$_2$O (g) | H$_2$O (mL) |
|------|--------|----------|---------------------------|---------------------------|---------------------------|-------------|
| S1   | 0.0    | 24.0     | 17.46                     | 0.00                      | 48.0                      | 25 / 50 /100 |
| S2   | 0.2    | 24.0     | 13.96                     | 3.49                      | 48.0                      | 25 / 50 /100 |
| S3   | 0.4    | 24.0     | 10.47                     | 6.98                      | 48.0                      | 25 / 50 /100 |
| S4   | 0.6    | 20.0     | 5.82                      | 8.72                      | 40.0                      | 21 / 42 /84  |
| S5   | 0.8    | 16.8     | 2.44                      | 9.77                      | 33.6                      | 18 / 35 /70  |
| S6   | 1.0    | 16.8     | 0.00                      | 12.21                     | 33.6                      | 18 / 35 /70  |

### Table 2 Specific surface area values (m$^2$/g) for selected samples

| Composition | Initial solution concentration |
|-------------|--------------------------------|
|             | Low                            | High                           |
| $\Psi = 0.0$| 11.5                           | 10.6                           |
| $\Psi = 0.6$| 15.4                           | 12.3                           |
| $\Psi = 1.0$| 16.2                           | 11.3                           |
Figure 2
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Figure 4
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Figure 6

Concentration

- ○ 0.6M
- + 1.2M
- × 2.4M

Vegard's law
Figure 9

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