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¹ Chemistry Synthesis, Characterization and Photocatalysis of KSr ² 2 Nb 5 O 15 Doped with Nickel

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7 Abstract

17

Potassium and strontium niobate (KSr2Nb5O15) of tungsten structure (TB) bronze was 8 doped with nickel (Ni2+) with stoichiometry type KSr2NixNb5-xO15-? onde x = 0.25; 0.50 9 and 0,75. The powders were prepared by chemical synthesis using the modified Polyol 10 method, and characterized by infrared spectroscopy with Fourier transform (FT-IR), by X-ray 11 diffraction (XRD), including the network parameters (relative intensity - IR(KSNNi0,50) ? 12 IR(KSNNI0,25) ? IR(KSNNI0,75)) and crystallite size. The effects of doping (Ni2+) in the 13 process of photo degradation were studied, the absorbance as a function of time for the Basic 14 Blue dye 41 result from the photo degradation performed with the photo catalyst KSNNi0,50 15 presented a lower absorbance value, and the spectrophotometer was used in the UV-vis region. 16

Index terms— heterogeneous catalysis; oxidative catalysts; photo catalysts, niobate; tetragonal tungsten
 bronze structure; basic blue 41.

20 1 Introduction

he chemical contamination of the water covers a large spectrum of pollutants. The textile and leather industries 21 are mainly responsible for the discharge of these large quantities of dyes. The highly colored effluents from these 22 industries are affecting the nature of water, inhibiting the penetration of sunlight and reducing photosynthetic 23 24 reactions, ATAR et al. (2008) [1]. Approximately 10-15% of these dyes are not fixed to the substrates during the 25 dyeing process, as some dyes used in the textile industry and their biodegradation byproducts may have a high degree of toxicity, mutageni city and carcinogenicity to humans; FRAGAA, ZANONIA (2009) [2]. These Dye 26 effluents are difficult to treat by conventional methods and cannot be completely degraded. Heterogeneous photo 27 catalysis has aroused great interest, with the aim of efficiently purifying waste water containing dyes; ZHANG 28 et al (2009) [3]. 29

Basic Blue 41 dye boasts an azo chromophore group and is used in the dyeing of synthetic fibers such as polyamide, polyester and viscose. Basic (cationic) dyes, which mostly have hydrolysis stability, are pH sensitive and are soluble in aqueous medium. Therefore, they require more efficient treatment methods for their complete removal in surface waters and effluents; FRAGAA, ZANONIA (2009) [2].

The photo catalytic reactions on semiconductor surfaces are processed according to the basic steps of: 34 35 excitation with light of energy greater than the band gap (Eg), of the semiconductor, generation of electron/gap 36 pairs (e-/h+); imprisoning electrons and gaps by adsorbed species. The mechanism of redox processes mediated 37 by semiconductors in aqueous media promotes the formation of the hydroxyl radical (OH?), powerful oxidizing agent, generated by promoting the oxidation of the water / hydroxyl adsorbed by the lacuna, making the photo 38 catalytic process highly efficient to oxidize most of the organic compounds. The efficiency of this process is 39 related to the lifetime of the gaps (h+bv) and retardation of the recombination velocity of the generated loads 40 (e-/h+) in the semiconductor; FRAGAA, ZANONIA (2009) [2]. 41

Although titanium dioxide is currently the most widely used, niobate based photo catalysts have been extensively studied because of their excellent photo catalytic properties, such as KNb 3 O 8 , K 6 Nb 10.8

O 30, K 4 Ce 2 M 10 O 30 (M?Ta, Nb), NiM 2 O 6 (M?Nb, Ta), K 4 Nb 6 O 17, BiNbO 4, NiO-KTiNbO 44 5, etc. Among these niobate photo catalysts, K 4 Ce 2 M 10 O 30 (M?Ta, Nb), NiM 2 O 6 (M?Nb, Ta), 45 NiO-KTiNbO 5 e K 4 Nb 6 O 17 high photo catalytic activity was found in the field of water decomposition. The 46 KNb 3 O 8, K 6 Nb 10.8 O 30 and BiNbO 4 were studied for the degradation of dyes. However, the potassium 47 and strontium type niobates (TTB) KSr 2 Nb 5 O 15 showed high photo catalytic activity in the degradation of 48 red acid G under irradiation UV degrading and breaking the nitrogen double bond (-N?N-), and also causes the 49 double bonds of the benzene and naphthalene rings to split up, the degradation rate exceeds 85% and its kinetics 50 remains the first order; ZHANG et al. (2009) [3]. 51 Recent studies of the KSr 2 Nb 5 O 15 attests its photo catalytic activity resulting in methy lene blue photo 52

oxidation as a model reaction. These niobates KSr 2 Nb 5 O 15 present higher catalytic activity with lower
 calcination time, and under very low concentration conditions under irradiation UV; MATOS et al. (2017) [4].

The present work aims to analyze the photo catalytic potential of materials with stoichiometry KSr 2 (The infrared spectral absorption spectroscopy was performed in a spectrophotometer of the brand SHIMADZU model

⁵⁷ IRAffinity-1. The spectral range used was in the medium infrared region (3500 -450 cm -1), with resolution ⁵⁸ of 8 cm -1 and 120 scans. The samples were dispersed in KBr in the proportion of 1:100. Potassium Bromide

tablets (KBr) were prepared using a tablet of the mark PIKE de 13 mm diameters in hydraulic press of 10 ton.

Each tablet was produced by macerating 1,5 mg of the sample, ceramic powders, with 0,30 g de KBr in agate mortar. The mixture with KBr was pressed in the pellets by 10 min. The tablet thus obtained was analyzed in

62 a spectrophotometer.

⁶³ 2 iii. X-ray diffractometry (XRD)

The precursor post-ceramics, the solid solutions of KSr 2 (Ni x Nb 5-x)O 15-?, where the x = 0.250; 0.500 e 0.750 producing stoichio metrically the following materials: KSr 2 Ni 0.25 Nb 4.75 O 15-?, KSr 2 Ni 0.50 Nb 4.5 O 15-? and KSr 2 Ni 0.75 Nb 4.25 O 15-? and treated at temperatures of 1250°C, were characterized by X-ray diffraction using a diffracto meter SHIMADZU (model XRD-6000), with radiation Cu K ? (??=1.54060

⁶⁸ Å), operating at 40kV and 30mA, in the range of 5? 2?? ? 80, with scan time of 1,00°/min, steps of 0,02° and

time per step equal to 1,20s. The slits of divergence and scattering used were $1,00^{\circ}$ and the receiving slot of 0,30 mm.

71 **3 a. Network Parameters**

The network parameters are calculated using the "Least Squares". The positions 2?? 0, the plans hkl, the type of the crystalline structure, the number of interactions to be executed, are the program data.

⁷⁴ 4 b. Average crystallite size

The values of the mean crystallite size for the solid solutions of KSr 2 (Ni x Nb 5-x)O 15-?, where a) x = 0.25; b) x = 0.50 and c) x = 0.75 were calculated by the Scherrer equation, equation (1). Year 2019© 2019 Global Journals (D D D D) D = K ?? cos ? (1)

⁷⁸ being ? the width of the peak where the intensity is half of its maximum value denominated (peak width at ⁷⁹ half height), ? is the angle corresponding to the diffraction, ? is the wavelength of the Cu (1,5406 Å), k is the ⁸⁰ constant of proportionality, called the particle shape factor (TB = 0,89) NUFFIELD (1986) [21].

⁸¹ 5 c) Evaluation of photo catalytic potential

For the study of photo catalytic activities, we used the degradation material Basic Blue 41 in aqueous solutions. The photo catalytic tests will be performed in order to maximize the activation of the post-ceramics as a catalyst.

⁸⁴ 6 d) Photo catalytic procedure of Basic Blue 41 dye

To investigate the photo catalytic activity of the ceramic powders, a photo catalytic reactor. This low power photo catalytic reactor has a source of irradiation, a light bulb with dimensions (length 438,0 mm, diameter 26,0 mm) HSN ® 15 W G13 which emits radiation at the wavelengths UVC between 200 to 280 nm, that is, specifically 250 nm, which corresponds to the range of the electromagnetic spectrum of ultraviolet C (rays UV-C). In this way the photo catalytic activity of the material was studied using as comparative parameter tests with UV-C. For the light bulb the measured irradiance was 0,3 mW/com 2 during the photo catalytic test stage and the measured irradiance of 1,8 mW/cm 2.

Basic blue 41 solutions were prepared by means of a dilution of the stock solution, thus obtaining a concentration of 40 ppm. The pH of the solution was then measured through a pH meter (GEHAKA) adjusting the pH to 8 in some tests with the aid of a few drops of sodium hydroxide 0,01mol.L -1 and in other tests the natural pH of the solution was maintained for the purpose of comparison with the original methodology using the ceramic powders. Was heavy 0,125g of catalyst, in order to obtain optimum concentration of catalyst (0,55 g.L -1) in a volume of 1000 mL of solution (12,5 mg/L).

The first step consists of the photo catalytic tests of a period of 1 hour, where at the beginning of the first 60 minutes the post-ceramics were added in solution of Basic Blue 41, with the flow of water circulating through the reactor, with the help of a small compressor, in order to reach the equilibrium of adsorption-desorption in the dark, this is the step in the dark, process of accommodation of the material to solution. The last 30 minutes

- with the reactor lamp on and continuous flow. In the last minutes (photolysis) at 59 minutes, a 5.0 mL aliquot of the sample was collected by means of a volumetric pipette, starting the removal of the first sample, identified
- 104 (t = 0).

The second step of the photo catalytic tests were the collections every 15 minutes totaling 1 h, and identified = $1, 2, 3 \in 4$.

The third step of the photo catalytic tests, the collections were performed every 30 minutes, totaling 2 h and identified by $t = 5, 6, 7, 8, 9 \in 10$, totaling 11 samples collected.

In the fourth step, the collected samples were accommodated in test tubes enclosed in boxes avoiding exposure to light. These test tubes were centrifuged, 3000 rpm by 20 minutes, in order to separate the catalyst from the solution, 0,5 mL of the sample (supernatant) with the aid of a Pasteur pipette, and diluted 5 mL of distilled and deionized water, measured with a volumetric.

In the fifth step, the supernatants were analyzed in a UV-vis spectrophotometer and programmed for wavelength reading (?) 611 nm of the solution, recording the absorbance of the samples.

¹¹⁵ 7 e) Determinação da Eficiência e Parâmetro Cinético de ¹¹⁶ Degradação

The photo catalytic activity of post-ceramics KSr 2 (Ni x Nb 5-x)O 15-? , where x = 0.25; 0.50 and 0.75 the kinetics of the disappearance of Basic Blue 41 as test molecule.

In order to establish the photo catalytic behavior of niobate based materials, both degradation kinetics and direct photolysis in the absence of solids were followed. The disappearance of the Basic Blue dye 41 was reported in terms of the conversion (X) obtained by the following expression of the efficiency represented by the concentration

122 equation (2): $X = ?? C \circ ? C t C \circ ?? . 100 ? ? (A o ? A t) A o ? . 100 (2)$

Where Co is the initial concentration of Basic Blue dye 41, Ct is the concentration in reaction time t, and ao and at are the initial absorbance and the absorbance at the reaction time t. In this way he analyzed and studied the efficiency and kinetics of degradation of the dye Basic Blue 41.

126 **8 III.**

127 9 Results and Discussions a) Infrared with Fourier Transform (FT-IR)

Figure 1 [22]; YEBIN, GUOHUA, HUA (2003) [23] and BERGAMASCHI (2000) [24][22-24], this system has 129 characteristic bands that identify the presence of an "envelope" and with the addition of nickel (Ni-O) to KSr 2 130 Nb 5 O 15; YEBIN, GUOHUA, HUA (2003) [23] tends to displace the system bands. Symmetrical stretching 131 is attributed (? s) and antisym metric (? as) DENIO et al. (2010) [22] and BERGAMASCHI (2000) [24]. 132 Figure 1 (a) -(KSNNi 0.25) shows a portion of the spectrum in the infrared region where it is characterized by 133 an envelope in the region between 450 -1100 cm -1 for the system KSr 2 (Ni x Nb 5-x)O 15-? where a) x =134 0.25. In this figure, the spectra show bands of strong intensity below 1000 cm -1, characteristic of niobatos. The 135 bands identified have wavelengths in 571, 590, 672, 781, 844 and 924 cm -1 such band refer to the oxide-metal 136 bonds. 137

Figure 1 (b) -(KSNNi 0,50) shows parts of the spectrum in the infrared region where it is characterized by an envelope in the region between 450 -1100 cm -1 for the system KSr 2 (Ni x Nb 5-x)O 15-? where b) x = 0.50. In this figure, the spectra show bands of moderate intensity below 1000 cm -1, characteristic of niobatos.

The bands identified have wavelengths in 548, 587, 660, 795, 852 and 918 cm -1 such band are attributed to the oxide-metal bonds.

Figure 1 (c) -(KSNNi 0,75) shows parts of the spectrum in the infrared region where it is characterized by an envelope in the region between 450 -1100 cm -1 for the system KSr 2 (Ni x Nb 5-x)O 15-? where c) x = 0,75. In this figure, the spectra show bands of moderate intensity below 1000 cm -1, characteristic of niobatos.

The bands identified have wavelengths in 560, 583, 653, 791, 864 and 923 cm -1 such band are attributed to 146 the oxide-metal bonds. Table 1 ??O). The medium and wide asymmetric bands in the regions of 672 cm -1 to 147 781 cm -1, observed in the spectrum of KSr 2 (Ni 0,25 Nb 4,75)O 15-? they move to regions between 660 cm -1 148 to 795 cm -1 spectra of the solid solutions of KSr 2 (Ni 0.50 Nb 4.50)O 15-? and this system for 653 cm -1 to 149 791 cm -1 for the system KSr 2 (Ni 0,75 Nb 4,25)O 15-? . The bands in the region between 844 cm -1 to 924 cm 150 -1 can be attributed to the symmetrical (Nb-O-Nb) LANFREDI, FOLGUERAS-DOMÍNGUES, RODRIGUES 151 (1995) [25]. The displacement of these bands to the region of smaller wave number in the spectra of the solid 152 solutions of the KSr 153

¹⁵⁴ 10 b) X-ray diffractometry (XRD)

Figure 2 shows the X-ray diffracto grams obtained for the precursor powders of the stoichio metric system based on niobate KSr2(NixNb5-x)O15-? where x = 0.25; 0.5 and 0.75 heat treated at 1250 ° C for 1 hour, in

an oxygen atmosphere. X-ray diffracto grams show an increase in the definition of diffraction peaks to 1250°C, 157 associated with the decrease of the microde formation of the net and increase of the structural stability. According, 158 LANFREDI et al. (2005) [26], in solid solutions of KSr 2 (Ni x Nb 5-x)O 15-? is adopted the valence of the 159 Ni 2+, since the oxidation state +3 of nickel cation (Ni) has been rarely detected. The substitution of radium 160 cations (r) such as the Sr 2+ (rSr 2+ = 1,18 Å) by cations of lightning (r) small, as the Ni 2+ (rNi 2+ = 0,69 161 Å) not favorable. In addition, the cations of Ni 2+ show strong preference for octahedral coordination, the same 162 coordination of niobium (Nb). In this sense, the ionic radius of Nb cations in a high oxidation state (rNb 5+=163 0,64 Å), similarity with the ionic radius of the Ni 2+. However, the valence difference is equal to three units, 164 which is not favorable. However, the best similarity of the ionic Ni 2+ occurs for the Nb 5+ partially reduced 165 to Nb 4+ (rNb 4+ = 0.68 Å). Here it is important to comment that the Nb 4+ represents a partial reduction of 166 the niobium cation, whereas a completely reduced state is given by the niobium with valence 3+, Nb 3+, (rNb 167 3 + = 0.72 Å). 168

The structural characterization of the postceramics constituted by the KSr 2 (Ni x Nb 5-x)O 15-? where x=169 0,25; 0,50 e 0,75 obtained by the modified Polyol chemical method was investigated and analyzed by means of 170 the X-ray diffraction technique. The standard XRD for the system KSr 2 (Ni x Nb 5-x)O 15-? with a range of 171 2? of 5° -80° is shown in Fig. 1 (0,25; 0,50 \pm 0,75). According to the crystal data file JCPDS 34-0108 (2000) [27], 172 173 the system is of tetragonal type and presents spatial group P4bm (100). The diffracto grams of the samples that were prepared and calcined at 1250 ? / 1h fit the pattern, and in some angles (??) present significantly lower 174 175 displacements, which may consider that these samples consist of single phase for the compounds with variation of doping. The Diffracto gram shown in figure 2 (0,25) referring to the system KSr 2 (Nb 5x)O 15-? where x 176 = 0.25 presents formation of a single crystalline phase based on the tetragonal symmetry of the crystallographic 177 sheet JCPDS: 34-0108 (2000) [27] related to the niobium oxide strontium and potassium (KSr 2 Nb 5 O 15). as 178 for the values of the müeller index (hkl), of 2??, of ??, of the interplanar distances (nm) and the intensities can 179 be seen in table 1 below. 180

Figure 2 (0.50) shows the X-ray diffracto gram obtained by the system precursor powder KSr 2 (Ni x Nb 5x)O 15-? where x=0,50 a 1250 °C/1h calcined in an oxygen atmosphere. The diffracto gram, similar to figure 1 (0.25), shows crystalline phase formation indicated in the indexing of the crystallographic data sheet JCPDS: 34-0108 (2000) [27] tetragonal symmetry for this system KSr 2 Nb 5 O 15 the narrow peaks indicating an increase in the crystallinity of the calcined material are observed in this diffracto gram (Fig. 2 (0.25, 0.50 and 0.75)) the 1250 °C / 1h.

Figure 2 (0.75) shows the X-ray diffracto gram obtained by the KSr 2 (Ni x Nb 5-x)O 15-? where x=0,75 calcined 1250 $^{\circ}$ C / 1h and obtained in an oxygen atmosphere. This characterization showed the formation of the monophasic and crystalline powder (KSNNi 0,75). According to the data of this record, the phases found have a tetragonal structure compatible with the spatial group P4bm (100). i.

¹⁹² 11 Network Parameters

The values given in tables 2 and 3 refer to the crystallographic sheet JCPDS: 34-0108 (2000) [27] shows the network parameters used to obtain the results of the interplanar distances and the ??. The relative intensity (IR) for stoichio metric systems ii.

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¹⁹⁷ 13 Crystallite size

198 The crystallite size of the solid solutions KSr 2 (Ni x Nb 5-x)O 15-? where x = 0.25; 0,5 and 0,75 was determined using the program Jade 8 Plus. The widening of the mean width at half height (FWHM) of the diffraction peaks 199 of the experimental diffracto gram was considered. The Jade 8 Plus program calculates the mean crystallite size 200 by applying the Scherrer equation; AZAROFF, BUERGUER (1958) [28]. Where D is the average crystallite size, 201 k is the proportionality constant, which depends on the shape of the particles (TB = 0.89) 2,? is the wavelength 202 of the Cu (1,5406 Å), ? is the width at half height of the corrected peak and ? the angle corresponding to the 203 diffraction. The instrumental factors were corrected using the Silicon (Si) standard. The crystallite size KSN 204 pure and doped with Nickel Method Modified polyol calcined for 1 hour, follows table ??. The largest crystallite 205 size was observed for solid solution KSr 2 (Ni 0,75 Nb 0,25)O 15-? . The increase in the mean crystallite size with 206 increasing the value of x to 0.75 is related to a greater distortion of the unit cell, increasing the diffusion process 207 208 and nucleation of the crystals; MELO (2007) [29]. These values show that the increased dopant concentration 209 promotes an increase in structural anisotropy in the material; DANTAS et al.

(2009) [30]. According to the values in Table ??, the value of the average size of network crystallite increases as the x value increases from 0.25 to 0.75. This effect is a consequence of the high degree of doping of the host structure, where an excess of nickel cations causes a disorder in the crystalline lattice to form defects, caused by non-stoichio metry of the structure; WANG et al. (2012) [31]. Year 2019 In this diffracto gram between 20 and 35 displayed on the 2?? scale there is a close junction between peaks in relation to figure 2, the crystal plug, JCPDS: 34-0108 (2000) [27] follows partially offset from the peaks. The values of the interplanar distances, the relative intensity and the 2?? of the diffracto gram coincide with the values listed on this sheet. The lines of this 217 chart coincide with the diffracto gram peaks, however, it is possible to notice a singular difference between these

218 peaks, as regards the relative intensity with the lowest intensity (Figure 2 -c) KSNNi 0,75) in relation to the

relative intensity of b) KSNNi 0,50 in Figure 2, where in this diffracto gram the intensity of the peaks is much

220 more expressive and defined, but of a considerable narrow width.

221 14 c) Avaliação do potencial fotocatalítico: Determinação

da eficiência e Cinética de degradação Figure ?? -a) KSNNi 0,25 ; b) KSNNi 0,50 and c) KSNNi 0,75 shows the 222 absorbance as a function of time, the adsorption in the dark for 1h and the degradation rate with the reactor 223 connected for 3h together with the photo degradation kinetics with irradiation of UV light. Figure 4 shows the 224 rate of discoloration over time for a) KSNNi 0,25 ; b) KSNNi 0,50 and c) KSNNi0 ,75 and Fig. 5 shows the 225 Ln (C 0/ C t) depending on the time for the systems (a) KSNNi 0,25; b) KSNNi 0,50 and c) KSNNi 0,75 to 226 investigate the degradation of Basic Blue 41. The performance of the materials synthesized here can be observed 227 that the rate of adsorption is very fast for all catalysts, reaching the equilibrium of the dye after 30-60 min. 228 Thus, photocatalytic tests with UV light irradiation were performed after an initial 60 min adsorption period. It 229

 $_{\rm 230}$ $\,$ is interesting to note that the amount of Basic Blue 41 adsorbed has decreased over time.

²³¹ 15 d) Efficiency and rate of degradation of KSNNi 0,25 ;

 $_{\rm 232}$ $\,$ KSNNi 0,50 and KSNNi 0,75 $\,$

The first test was performed with the stoichio metry material KSr 2 (Ni x Nb 5-x)O 15-? where x = 0.25233 irradiated with UV light and catalyst 0,10g and concentration of 12.5 mg.L -1 of the type dye (Basic Blue 41). 234 Figure ?? shows the absorbance versus time of Basic Blue 41 using the material KSNNi 0.25. The moment the 235 material is added KSNNi 0.25 the elapsed time solution of 1 h without UV irradiation at a drop in absorbance of 236 2.35%, This percentage is due to the fact of an accommodation of the solution to the surface of the catalyst, being 237 that at the moment of the adsorption there was no degradation of the solution of Basic Blue 41. However, for 238 239 the same material KSNNi 0,25 considering the last three hours the rate of discoloration was 91,35 % considering 240 its absorbance, which leads us to believe that the degradation of Basic Blue 41 can be attributed to the photo catalytic effect of KSr 2 (Ni x Nb 5x)O 15-? where x = 0.25 under irradiation UV. 241

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243 17 C

The second test performed with stoichiometry material KSr 2 (Ni x Nb 5-x)O 15-? where x = 0.50 irradiated 244 245 with UV light and catalyst 0,10g and concentration of 12,5 mg.L -1 of the dye (Basic Blue 41). Figure ?? shows 246 the absorbance versus time of Basic Blue 41 using the material Figure ?? shows the absorbance versus time of 247 Basic Blue 41 using the material KSNNi 0,50. The moment the material is added KSNNi 0,50 the elapsed time of 1 h, without UV irradiation at a drop in absorbance of 4,00 %. This percentage is due to the fact that a solution 248 249 solution to the surface of the catalyst, indicating a larger surface, and that at the moment of adsorption there was no degradation of the solution of Basic Blue 41. However, for the same material KSNNi 0,50 considering the 250 last three hours the rate of discoloration was 97,51 % considering its absorbance, which leads us to believe that 251 the degradation of Basic Blue 41 can be attributed to the photo catalytic effect of KSr 2 (Ni x Nb 5-x)O 15-? 252 where x = 0.50 under irradiation UV. 253

The third test performed with the stoichiometry material KSr 2 (Ni x Nb 5-x)O 15-? where x = 0.75 irradiated 254 255 with UV light and catalyst 0,10g and concentration of 12,5 mg.L -1 of the dye (Basic Blue 41). Figure ?? shows 256 the absorbance versus time of Basic Blue 41 using the material KSNNi 0,75. The moment the material is added KSNNi 0,75 the elapsed time of 1 h, without UV irradiation at a drop in absorbance of 3,00 %, This percentage is 257 due to the fact that a solution to the surface of the catalyst, indicating a larger surface, and that at the moment 258 of adsorption there was no degradation of the solution of Basic Blue 41. However, for the same material KSNNi 259 0.75 considering the last three hours the rate of discoloration was 97.51 % considering its absorbance, which leads 260 us to believe that the degradation of Basic Blue 41 can be attributed to the photo catalytic KSr 2 (Ni x Nb 5-x 261)O 15-? where x = 0.75 under irradiation UV. 262

It is important to emphasize that the adsorbed samples presented expected results due to the chemical nature 263 of Basic Blue 41, this dye being a strong Lewis base, and its adsorption is thermodynamically favored by Lewis 264 acidic solids, such as the materials synthesized here. According to surveys; MATOS et al. (2017) [4], in fact, 265 these materials that are Lewis acids constituted of niobatos present high electronic affinity of their ions Nb (86,1 266 267 kJ / mol). This condition induces the agreement that niobate based catalysts have a more acidic surface pH and 268 therefore have a high affinity for basic amines such as Basic Blue 41 with a high dissociation constant (pKb) 269 and a high neutralization potential. Absorbance readings at the maximum absorption wavelength of the "basic blue 41" dye were performed for the photo catalytic tests with the photo catalysts KSNNi 0.75; KSNNi 0.50270 and KSNNi 0,25 . According to figure $\ref{solution}$ -a) KSNNi 0,25 ; b) KSNNi 0,50 and c) KSNNi0 ,75 a decrease in the 271 absorbance values versus time for all tests was observed indicating a decrease in the concentration of the basic 272 blue 41 dye in solution resulting from the photo degradation. Among the photo catalytic tests performed, the 273 photo catalyst test KSNNi 0,50 presented lower absorbance value. 274

From Figure 4-a) KSNNi 0,25; b) KSNNi 0,50 and c) KSNNi0,75 an increase in the rate of decolorization as a 275 function of time is observed for all the photo catalytic tests performed, indicating a decrease in the concentration 276 of the dye "basic blue 41". The photo catalyst that presented the highest discoloration rate was the KSNNi 0,50 277 so it was the most efficient. On the other hand, the Figure ?? -a) KSNNi 0,25; b) KSNNi 0,50 and c) KSNNi 0,75 278 279 show very similar kinetic trends in the photo catalytic degradation of Basic Blue 41 for all photo catalysts KSr 2 (Ni x Nb 5-x)O 15-? where x = 0.25; 0.5 and 0.75. To compare the photo activity, the first order apparent rate 280 constants (k app) were estimated from linear regression of kinetic data, and assuming that Basic Blue 41 photo 281 degradation follows a first order reaction rate mechanism; FRAGAA, ZANONIA (2009) [2] e ATAR, OLGUN, 282 ÇOLLAK (2008)[1] [1][2]. We attribute this photo activity due to the presence of apical oxygen atoms that are 283 284 very reactive and are attached to the niobium in the octahedron [NbO 6] MATOS et al. (2017) [4]. Analysis of the kinetic curves in all the photo catalytic tests performed shows a first order kinetics, the data shown in Table 1 285 confirm such observations. The photo catalytic test with the photo catalyst KSNNi 0,5 presented a higher speed 286 constant and a shorter half-life, being the most efficient. According to the results, the nickel doping in the host 287 structure of the KSr 2 Nb 5 O 15 promotes an increase of the photo catalytic activity until a substitution of 10% 288 in mol of Ni 2+, and a decrease in photo catalytic activity can be observed for the KSN doped with 15 mol%. 289

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²⁹¹ **19 IV**.

²⁹² 20 General Discussions

Solid state diffusion is a mass transport medium occurring within the solid materials according to an atomic 293 movement occurring in stages. The mechanisms promoting this mass transport are realized by means of an 294 295 exchange of atoms which is located in a normal position of the network with an adjacent gap or by a migration 296 from an interstitial position to an adjacent empty interstitial position in which the host metal, the interstitial atomic species diffuse more rapidly, CALLISTER (2013) [32]. There are some factors that influence diffusion and 297 depends on both the diffusing host component and the temperature, this diffusion coefficient being a function 298 of temperature. In semiconductors, heat treatments promote the diffusion of impurities that are transported 299 into the host cell and it may also occur that such transport carries these impurities further into the host cell, 300 generating a more suitable concentration distribution. 301

302 The diffracto grams show formation of the TTB structure with monophasic and crystalline characteristics even with doping's made with the nickel metal forming the system KSr 2 (Ni x Nb 5-x)O 15-? where x varies 303 304 its doping, x = 0.25; 0.50 and 0.75. The calcination at 1250 ° C, the grains present expansion evidenced in the 305 diffracto graphic peaks exposed in Figure 2, where these peaks are pointed and narrow, what causes the system 306 to present crystallinity and to be monophasic. This process favors the absence or even decrease (elimination) of defects found in the crystalline lattice due solely to structural homogeneity. By means of this phase expansion, 307 308 we have an increase in crystallinity, which can be evidenced by decreasing the width of the diffraction peaks, increasing in intensity and in numbers. The diffracto gram (Figure 2) shows the formation of the phase with 309 structure of tetragonal symmetry (TB). With the thermal treatment of the precursor powder performed at 1250 310 C / 1h the values of the interplanar distances, relative intensity and 2?? of the experimental diffracto gram 311 coincide with the crystallographic JCPDS_34-0108 (2000) [27], this sheet refers to the phase of KSr 2 Nb 5 O 15 312 No other secondary phase has been identified, showing that the compound obtained is monophasic. For solid 313 solutions with x ? 0.75 diffracto grams showed the formation of a single phase associated with KSr 2 Nb 5 O 15 . 314 315 The valence of the Ni 2+ in solid solutions of KSr 2 (Ni x Nb 5-x)O 15-?, since the oxidation state 3+ of the nickel (Ni) cation has been rarely detected. The substitution of cations of large radius (r) such as, for example, Sr 316 2+ (r Sr 2+=1,18 Å) by cations of small radius (r), such as Ni 2+ (r Ni 2+=0,69 Å) not favorable. In addition, 317 the cations of 2+ show strong preference for octahedral coordination, the same coordination of niobium (Nb). In 318 this sense, the ionic radius of Nb cations in a high oxidation state (r Nb 5+=0.64 Å), similarity with the ionic 319 radius of the Ni 2+ . However, the valence difference is equal to three units, which is not favorable. However, 320 the best similarity of the ionic Ni 2+ occurs for the 5+ partially reduced to Nb 4+ (r Nb 4+ = 0.68 Å). Here it 321 is important to note that the Nb4 + represents a partial reduction of the niobium cation, whereas a completely 322 reduced state is given by the niobium with valence 3+, Nb 3+, (r Nb 3+=0.72 Å); LANFREDI et al. (2005) 323 [26]. The considerable increase in the network parameters may be associated with the partial substitution of 324 325 ions Nb 5+ by the ions Ni 2+ in the crystalline lattice, where there is a decrease in the covalent character of the 326 bond Nb-O with the increase in the ionic character of the bond Ni-O; LANFREDI et al. (2015) [33]. In fact, the 327 results of the FT-IR (Fig. 1) showed a correlation between the wave number of the bands and the displacement 328 magnitude of the decentralized Nb location as a function of the increased doping of the powders KSr 2 Nb 5 O 15; LANFREDI, FOLGUERAS-DOMÍNGUES, RODRIGUES (1995) [25]. The partial replacement of niobium 329 cations by nickel cations in the host structure can be accompanied by the formation of oxygen vacancies from 330 the charge compensation mechanism due to the partial reduction of niobium; LANFREDI et al. (2015) [33]. 331 Thus, the formation of oxygen vacancies can be accompanied by disproportionation of cations Nb 5+ for Nb 3+332 , in which it presents a larger ionic value due to its punctual loading, where r Nb 3+? r Ni 333

334 **21 2**+

?r Nb 5+, thus justifying the decrease in the volume value of the unit cell. Furthermore, nickel cations specifically
occupy the position of the Nb(1). This may be due to the ion ray value and the preferential octahedron occupation.
This occupation results in a degree of distortion of the octahedron of the [Nb(2)O 6], where this distortion is
necessary for an accommodation of the nickel cations to occur in the formation of the structure.

In the host structure KSr 2 Nb 5 O 15, the niobium has preference in occupying the octahedral sites, NbO 6 Cation substitution Nb 5+ by cations Ni 2+ can cause distortions in the octahedra as well as the creation of a sub level of energy resulting from the formation of gaps due to the difference of electrons between the cation Nb 5+ and doping cation, resulting in a decrease in the band gap. A higher degree of distortion and structural defects can be expected as a result of increased nickel doping. However, doping with 0.75 mol of Nickel can cause such a disorder, in such a way that the electronic mobility of the electrons is hampered, requiring a greater energy for electronic transitions of the conduction band to the valence band.

In terms of the first-order apparent rate, it should be noted that all niobate-based materials presented less 346 photo activity when compared to TiO 2. The conversion of Basic Blue 41 is probably attributed to a series of 347 chemical reactions occurring on the outer surface of these doped Nb-based materials Ni 2+. Thus, a deeper 348 analysis of the photo catalytic activity can be performed to obtain the overall reaction rate considering the 349 surface concentration of Basic Blue 41, since this primary fraction of Basic Blue 41 molecules will undergo photo 350 degradation. The photo catalytic degradation of Basic Blue 41 can be considered as a unimolecular catalytic 351 surface reaction, where the adsorption of Basic Blue 41 followed by photo degradation under irradiation UV; 352 ATAR, OLGUN, QOLAK (2008) [1]. The reaction rate for the degradation of Basic Blue 41 was faster for photo 353 catalysts with KSr 2 (Ni x Nb 5-x)O 15-? where x = 0.50 (KSNNi 0.50). These results suggest that changes 354 in the crystalline structure of the niobate-based material, especially when increased binding occurs Nb-O(6) can 355 notably influence photo catalytic activity. Probably, it is the most active oxygen in the sense of probability of 356 357 the transfer of charge to the molecules of O 2 and therefore the longer the length of such a connection Nb-O 358 the more this distortion occurs, and thus the greater the efficiency of electron transfer and the greater the photo 359 catalytic activity. Otherwise, it can serve as a more active center, a trap; JUAN, JORGE, JEAN-MARIE (2001) [34] and JUAN, JORGE, JEAN-MARIE (1998) [35] [34][35] on the nature of the dominant species of reactive 360 361 oxygen.

362 V.

363 22 Conclusões

The chemical synthesis based on the modified Polyol method proved to be adequate for the preparation of monophasic and crystalline post- The lines of the crystal, JCPDS: 34-0108 (2000) [27], coincides with the peaks of the diffracto gram, however it is noticed a singular difference of these peaks, relative intensity has lower intensity (figure 2 -c) KSNNI 0,75) in relation to the relative intensity of KSNNI 0,50 in Figure 2, where in this diffracto gram the intensity of the peaks is much more expressive and defined, but narrower and sharper, which demonstrated in descending order of relative intensities I R(KSNNi0,50) ? I R(KSNNI0,25) ? I R(KSNNI0,75) ;

1

D D D D) (© 2019 Global Journals

[Note: C]

Figure 1: Table 1 :

370

 $^{^1\}odot$ 2019 Global Journals Chemistry Synthesis, Characterization and Photocatalysis of KSr 2 Nb 5 O 15 doped with Nickel

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Hkl	2??	??	d interplana	r Intensidade
			(nm)	
(400)	28.626	14.313	3.1	12
(311)	32.076	16.038	2.8	100
(002)	46.013	23.01	1.9	36
(422)	57.205	28.6025	1.6	31
(731)	61.297	30.65	1.5	6
(622)	67.112	33.56	1.4	7

Figure 2: Table 2 :

3

 $\mathbf{2}$

Hkl	2??	??	d interpla-	Intensidade
			nar (nm)	
	27,9	$13,\!95$	3,2	1.003,5
KSr 2 (Ni 0,25 Nb 4,75	$29.7 \ 32,3$	$14,85\ 16,15$	$3,0\ 2,8$	$1.300,3\ 2.308,9$
)0 15-?				
	27,9	$13,\!95$	3,2	2.449,4
KSr 2 (Ni 0,50 Nb 4,50	29,7 $32,2$	$14,85\ 16,1$	$3,0\ 2,7$	3.047,3 5.104,6
)0 15-?				
	27,9	$13,\!95$	3,2	1.631,8
KSr 2 (Ni 0,75 Nb 4,25	$29,8\ 32,2$	$14,9\ 16,1$	$2,9\ 2,7$	$1.865, 1\ 3.455, 5$
)0 15-?				, ,

Figure 3: Table 3 :

1

Catalyzer	KSNNi0,25	KSNNi0,50	KSNNi0,75
K (min)	0,01125	0,01571	0,00663
t 1/2 (min)	61,61	44,12	104,55
R 2	0,95017	0,97412	0,88381
Kinetic	$Ln(C \ 0 \ /C \ t \) = -$	$Ln(C \ 0 \ /C \ t \) = -$	$Ln(C \ 0 \ /C \ t) =$
	0,20445 + 0,01125	0,32642 +	0,03394 +
Equation	x	$0,0151 \ {\rm x}$	$0,00663 \ {\rm x}$

Figure 4: Table 1 :

KSr 2 (Ni x Nb 5-x)O 15-? , where x = 0.25; 0.5 and 0.75 with stoichiometry can be controlled. In addition, the production of the post-ceramics with a shorter calcination time than those prepared by conventional mixing of oxides. Due to the high polarity of the polyols, the inorganic salts (precursors) are easily solubilized; The nucleation and growth steps occur at the boiling point of the polyalcohol (eg, ethylene glycol at 197 ° C, diethyleneglycol at 246 ° C and tetraethyleneglycol at 314 ° C) and the use of high temperatures (at the boiling point of the polyalcohol) produces materials with high crystallinity;

Absorption spectroscopy in the infrared region showed a displacement of the bands characteristic of the Nb-O for regions of low frequencies with increasing concentration of nickel cations. This displacement shows that the bonds between niobium and oxygen cations, for the nickel doped solid solutions, are shorter than KSr 2 Nb 5 O 15:

- A decrease of the absorbance values as a function of time was observed for the concentration of the basic blue 41 dye in solution, resulting from the photo degradation of the photo catalytic tests performed for the photo catalyst KSNNi 0,50 which presented lower absorbance value.
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