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Analysis of 100gsm A4 Copy Paper Sample and the X-Ray Diffraction Pattern of its Ash Content on Ignition at 900 °C and at 525 °C and Some of its Mechanical, Structural and Optical Properties

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Abstract

The residue on ignition at 900 °C of the copy paper sample Multipaper Fabriano 100g/m² was determined to be 12.96%w/w and its XRD pattern showed the presence of lime, syn CaO, peaks placed at 20 values of 32°, 37°, 37°, 38°, 48°, 54°, 64°, 67°, 67°, 80°, 89°, 92°, 103°, 107° which corresponded to crystal planes of 111, 002, 211, 220, 221, 311, 311, 400, 411, 420, 422 and 500. The high values of tensile force, made of 100% chemical pulp of 4.91KN/m in the machine direction and of 3.28KN/m in the cross direction of the paper sample correlated with an alkaline pH value of 8.8 and a low value of residue on ignition. Also, the average low roughness Bendtsen of the paper of 64ml/min related with the low pick resistance measured to be 1.18m/s in the machine direction and 0.19m/s in the cross direction. The residue on ignition, of the same copy paper sample, at 525 °C was also determined to be 22.8%w/w and calcite, magnesian, gallium iron sulfide was identified from their XRD pattern. The band gap energy E_{g} from the diffuse reflection spectroscopy study of the 100gsm A4 copy paper sample was calculated to be 1.1654879939663700eV at room temperature. The crystallographic phase of the residue on ignition at 900 °C was determined by X-Ray diffraction (XRD) and the powder diffraction pattern of CaO was indexed by trial and error and all of the peaks were assigned their hkl values. The Miller indices of the reflections were determined and also the separation of planes. The average crystallite size τ of the pattern of CaO was determined to be on average below 60nm, by using the Scherrer equation. A graphical representation of the line broadening of the diffraction peaks as a function of the crystallite size was also presented. The peak broadening factor β which was equal to FWHM of the reflections of the sample was related to the micro-strain ε and a graph was plotted and from its intercept the lattice strain was calculated to be 4.719727188044x10⁻⁴rad.

Keywords: Residue on ignition at 900 °C; Residue on ignition at 525 °C; XRD pattern; Lime; Magnesian; Full width at half maximum (FWHM); Reflectance spectrum; Tauc plot; Band gap energy

Introduction

Paper contains, apart from cellulose, in small proportions resins, gum, proteins and minerals from wood. The minerals are mainly silicon, chlorides, sulphur, phosphates, carbonates, iron and magnesium oxides, calcium, potassium and sodium salts. There are several minerals used in paper filling and coating such as kaolin clay, calcium carbonate, titanium dioxide and talc, for improving paper properties [1,2]. In order to produce good quality paper wood adhesives have to be utilized also [3]. Starting in the 1960s in Europe there has been a strong trend in the direction of calcium carbonate fillers [4]. Because the presence of calcium carbonate generally causes the pH of a papermaking system to remain above a value of 7, the term "alkaline papermaking" has been used. The higher extract pH of paper made under alkaline papermaking conditions has had a favorable side benefit, a much

slower deterioration of the strength properties of paper upon storage, compared to paper made under acidic conditions with aluminum sulfate [4].

The identification of the species present in the ash on ignition of a A4 100gsm copy paper sample from its powder diffraction pattern is based upon the position of the lines in terms of 2θ and their relative intensities. The diffraction angle 2θ is determined by the spacing between a particular set of planes, with the aid of the Bragg equation [5]. This distance d is calculated from the known wavelength of the source and the measured angle. Line intensities depend upon the number and kind of atomic reflection centers that exist in each set of planes. Powder x-ray diffraction is one of the primary techniques used to examine the physic-chemical composition of unknown materials [6]. Materials are generally made up of atoms arranged in crystal microstructures. Minerals are crystalline inorganic solids possessing well-defined chemical composition [7].

The present work aims at presenting technical information about the crystal structure of the minerals present in a paper sample and its connection with its properties both mechanical and optical, as well structural. Paper is a complex composite material. Its structure characteristics as well as its surface influence its printability. The roughness among others is important for the absorption and the spreading of inks, and also determines the amount of coating of the paper sample [8]. The characterization of the paper surface is crucial for the understanding of light reflection and scattering that controls the level of gloss of the paper. The paper's optical properties have a large impact on the print quality because the paper is visible between the printed area and defines also the background reflectance for the toner layer [9].

Materials and Methods

Instruments and materials

The pH value of the paper sample was determined using a Metrohm 716 DMS Titrino, Swiss made, pH-meter and corresponded to the equilibrium pH for 2g of the paper sample dispersed in 100ml of water at 25 °C (298K). The measurement was done in duplicate and the average value was reported. For the residue on ignition method the principle required weighing the paper sample pieces in two heat-resistant crucibles of 50ml capacity and igniting both of them at 900 °C in an oven. Also, in two other crucibles of 50ml capacity the paper sample pieces were ignited at 525 °C. The ash mass was determined, in both cases, by weighing the crucible after the paper sample had been ignited for 1 hour in the oven. The European reference test methods used to determine the ash content were ISO 2144 [10] and ISO 1762 [11] respectively. An oven Carbolite England, OAF 11/1 maximum temperature 1100 °C, was used to incinerate the paper sample test pieces inside the crusibles for both temperatures.

The A4 paper sample tested was conditioned in a standard atmosphere according to standard ISO 187 [12], at the temperature of 23 °C and relative humidity of 50%. The basic paper properties i.e.,

basis weight (ISO 536), thickness and density (ISO 534), moisture content (ISO 287) were determined according to standardized methods. A spectrophotometer CM-3630 was used for measuring the optical properties of the A4 100gsm copy paper sample, ISO brightness (ISO 2470), opacity and transparency (ISO 2471). It was a model BCMTS M Type 40605 with Touchscreen M software of the company Frank-PTI Germany. The reflectometer had a reference standard for its calibration. It had also a non-fluorescent calibration plate of ceramic, a non-fluorescent working standard, Konica Minolta, made in Japan, no20286101. It had a black cavity, made in Japan, for calibration of the low end of the photometric scale.

A digital precision micrometer TMI Model No 49-61-01-0002, S.N. 33421-01 with range 0-1.270mm, was used for measuring the thickness of the A4 copy paper sample. The tensile testing machine Zwick Roel was used for measuring the tensile strength of the paper sample. The machine extended the paper test pieces of dimensions 15mmx210mm at 20mm/min constant rate of elongation and measured the maximum tensile force. It had a strength force of 2.5KN. The machine had also a long-stroke extensometer placed directly on the paper test pieces for the measurement of their elongation. The machine was connected with a computer LG. Two clamps for holding the paper test pieces of 15mm width were used. The clamps grab the test pieces firmly along a straight line across the full width of the test pieces and adjusted the clamping force pneumatically. A guillotine IDEAL 1043 GS made in Germany was used for cutting the paper test pieces to dimensions of 15mmx210mm.

A conditioning chamber that was capable of providing and maintaining standard conditions of temperature and humidity, where the paper test pieces were pre-conditioned at 23 °C \pm 2 °C and 30%r.h. \pm 5%r.h. for 24hours and were conditioned at 23 °C \pm 1 °C and 50%r.h. \pm 2%r.h for 16hours. The paper samples were conditioned for 16 hours at 23 \pm 1 °C temperature and 50% \pm 2% relative humidity in accordance with the ISO 187 standard [12]. Weighing's were carried out on an electronic balance Sartorius Basic plus, AG Göttingen Germany BP 221S, with maximum capacity 220g and accuracy to 0.1mg.

Pick velocity was determined according to ISO 3783 [13]. For measuring the picking or the pick velocity an IGT Global Standard Tester 1, with printing force 350N and printing velocity 4.00m/s, and a high-speed inking unit 4, software Version 3.01, consisting of 5 inking drums having contact with the top-roller, was used. Thus, the determination of the pick resistance IGT of the paper sample was calculated. The IGT pick test oil was of middle viscosity with number 404.004.020 and consisted of poly-isobutene's. The tester had an aluminum disc 10mm wide with smooth edges, and a diameter of 65.0mm. A paper pack consisting of 6 layers of paper with a total thickness of 1.5mm was used. An ink pipette for applying an accurate quantity of pick-test oil to the inking device, having a volume of 2ml, was used. The distributing time of the ink was 15s and the inking time of the aluminum disc was 10s. The speed during inking time was 0.4m/s and the temperature was kept at 23.2 °C. The ink layer thus formed was 8µm thick. Also, a

pick-start viewer was used to determine the starting point of pick. It had a light source providing grazing incident light to the paper test piece in order to observe pick velocity. Petroleum ether 60-80 °C of analytical reagent grade solvent was used to wet the cleaning rag, lint-free, which served as cleaning aid. A tank filled with water with a built-in thermometer which kept the ambient temperature near the printing device at 23 °C with an accuracy of 0.1 °C was used. Test prints were made in accordance with the ISO 3783 [13] standard.

The specular gloss was determined according to TAPPI T480 om-09, on a Glossmeter T480A Model Technidyne Corporation, New Albany, Indiana USA, S.N. 481313. Water absorptiveness was assessed as Cobb value 60s as per ISO 535. The surface roughness and porosity were determined using Bendtsen roughness and air permeability tester ([14], ISO 5636-2). More specifically the Bendtsen roughness [14] and air permeability were measured by a Messmer Buchel roughness and air permeance tester Bendtsen method model No.K513, TMI INC, TMI K513, S.N. 8008-01 with accessories. The crystalline structures and phase identifications, of the CaO present and also of calcite, magnesian, gallium iron sulfides, in the ash contents of the paper sample, were investigated by a Rigaku Miniflex 300/600 Benchtop X-ray diffractometer using a CuKa radiation (λ =1.5418Å) with a current of 15mA and an anode voltage of 40kV.

Results and Discussion

Tensile Properties and other physical and optical properties of the A4 copy paper sample

A copy A4 paper Multi-paper Fabriano $100g/m^2$ was tested in this work which was not a common A4 copy paper 80gsm. The paper properties such as grammage, thickness, Bendtsen roughness, tensile strength, pick velocity IGT, were measured. The A4 copy paper sample 100gsm Multi-paper Fabriano had a tensile index 47.90 $\frac{KNm}{Kg}$ lengthwise (Machine Direction or MD) and 32.0 $\frac{KNm}{Kg}$ crosswise (Counter machine Direction or CD) (Table 1,2) [15]. The grammage of the A4 copy paper sample was 102.5g/sq.m [16]. The elongation at break of the dry A4 copy paper sample was 3.01% lengthwise (MD) and 6.29% crosswise (CD) (Figure 1 & 2). The copy paper sample had thus more paper fibers oriented in the machine direction (MD) than in the cross direction (CD) and was stronger in the machine direction in tension. The density of the 100gsm paper sample was calculated to be 0.9234g/cm³ and its specific volume was calculated to be 1.0829cm³/g [17].

Table 1: Tensile strength of the copy paper sample A4 in KN/m in the two directions of the paper MD and CD.

Dry conditioned A4 copy paper sample

Tensile strength lengthwise MD=4.91KN/m±0.9KN/m

Tensile strength crosswise CD=3.28KN/m±0.2KN/m

Table 2: Tensile strength of the copy paper sample A4 in MPa in the two directions of the paper MD and CD.

Dry conditioned A4 copy paper sample

Tensile strength lengthwise MD=0.3273MPa±0.060MPa

Tensile strength crosswise CD=0.2187MPa±0.013MPa







Figure 2: Tensile force of the dry A4 multi-paper fabriano 100gsm, copy paper sample strips along the counter machine direction (CD) as a function of the strain of its paper specimens.

The thickness of the 100gsm copy paper sample was measured to be 0.1111mm or 111µm ± 5.5µm. The pick velocity IGT was determined to be 1.2m/s lengthwise in MD, and 0.2m/s crosswise in CD at 23 °C according to ISO 3783 [13]. The pick velocity was defined as the velocity at which picking started. The picking of paper was defined as the damage of the paper surface caused by the tensile force during the printing operation. Lower was the surface strength of the paper sample the lower was its pick velocity, as in this sample, whereas a pick velocity of around 2m/s would be more adequate. The pick resistance at 23 °C was then calculated to be P(23)=1.2*51.2/52.0=1.1815m/s in the MD direction and P(23)=0.2*51.2/ 52.0m/s in the CD direction, whereas the general relation applied P(23)= $v_{p}*\frac{VT}{V23}$ and P(23) was the pick resistance at 23 °C in m/s, v_n was the mean velocity at the point where pick began in m/s, V_{T} was the viscosity of the pick test oil used at average temperature T=23.1 °C during the test in Pa.s, being 51.2Pa.s, and $V_{_{23}}$ was the viscosity of the pick-test oil at 23.0 °C in Pa.s, being 52.0Pa.s. The Bendtsen roughness of the paper was measured to be 67ml/min for side A upwards and 61ml/min for side A downwards. The pH value of the paper [18] was determined to be 8.8 at 20.2 °C (293.35K) and its ash content on ignition at 900 °C was found to be 12.96%w/w. The low pick resistance was related to this scale of roughness Bendtsen for this value of pH 8.8, below pH 9.0 and ash content on ignition below 13%w/w. The high value of lightness L* and ISO brightness, beside a positive value of colour coordinate a* and a negative value of colour coordinate b* revealed that the copy paper sample had a natural blue-red look (Table 3). The outdoor daylight CIE Whiteness D65/10°, UV adjusted was measured to be 165 whiteness units. The copy paper sample had thus a high whiteness and also had a high ISO brightness. The ISO brightness

was defined as the diffuse reflectance of a thick stack of paper sheets i.e., 10 paper sheets, at the mean wavelength of about 457nm. The paper whiteness correlated with the lightness (L* value) and color (b* value). The measurement of each side of Bendtsen roughness revealed no difference in surface roughness between the two sides. The small Bendtsen roughness of 64ml/min on average, meant that the surface of the paper sample was very smooth.

Table 3: Surface, mechanical, and optical properties of A4 copy paper Multi-paper Fabriano 100gsm are given as mean value.

L*/a*/b*: 93.03/4.05/-12.63
Specular gloss (%) : Side A: 7.3, Side B: 5.3
Basis-weight (g/m ²):102.5g/m ²
Thickness (μm):111μm ± 5.5
Water absorptiveness Cobb 60s (g/m ²):Side A:20.8, Side B: 19.8
Bendtsen roughness (ml/min):Side A:67, Side B:61
Air permeability (ml/min): 458
ISO brightness (%): 100.95
Opacity (%):95.87
Transparency (%): 11.82
Tensile strength (KN/m): MD:4.9KN/m±0.9KN/m, CD:3.3KN/ m±0.2KN/m
Moisture content (%w/w):4.9 ± 0.6
Density (Kg/m ³):923.4

The high density of the paper sample may have resulted from the presence of fillers present in it and from the brought distribution of the fiber dimensions resulting in a good alignment of fibers and the web connectivity [18]. Despite high air permeability and high density, water absorptiveness, Cobb value, was average to high, meaning that less sizing agents were added in the paper's production process. A low surface roughness coincided with a low value of specular gloss [19]. The ash content on ignition of the copy paper sample at 900 °C was determined to be 12.96%w/w, and the ash content on ignition of the copy paper sample at 525 °C was determined separately to be 22.8%w/w.

X-ray diffraction pattern of the ash content on ignition at 900 °C, of the copy paper Multi-paper Fabriano sample

The ash on ignition, of the paper sample Multi-paper Fabriano 100gsm, at 900 °C was determined to be 12.9%w/w (Figure 3 & 4). For diffraction studies the sample ash on ignition of 0.2779g was ground to a fine homogeneous powder and the phase of the species syn CaO, lime, from its powder diffraction pattern was based upon the

position of the lines in terms of 2θ and their relative intensities. The wavelengths were: Target: Cu, the X-ray type: Ka, Ka1:1.54056(ang.), Ka2:1.54440(ang.), Kb:1.39220(ang.) and the Kalpha12 ratio: 0.4970 and the horizontal polarization was 0.500. The optical device had an incident mirror, the incident monochromator: Use, the receiving monochromator: fixed monochromator, the optics: slit conditions: Incident slit, receiving slit#1, Receiving#2. The Measurement had Measurement axis: 2Theta/Theta, Target: Cu and wavelength type: KAlpha(KAlpha1+KAlpha2). The X-ray diffraction pattern of the ash on ignition at 900 °C was lime syn CaO. The resulting X-ray powder diffraction data for syn CaO together with the observed 2θ , the d-spacings as well as the relative intensities of the reflections are given in Figure 7. The results of the XRD analysis confirmed the formation of CaO species of diameter less than 60nm in the ash on ignition at 900 °C of the copy paper sample tested.



Figure 3: XRD pattern of the ash on ignition at 900 °C of a 100g/m² A4 copy paper sample Multi-paper Fabriano.





The ash on ignition, of the paper sample Multi-paper Fabriano 100gsm, at 525 °C was determined to be 22.78%w/w. For the diffraction studies the sample ash on ignition of 0.4860g was ground to a fine homogeneous powder. The identification of the species $CaCO_3$ (calcite) in the presence of MgCO₃ (magnesian) and GaS, FeS (gallium iron sulfide) also present (Figure 4) from its powder diffraction pattern was based upon the position of the lines in terms of 2 θ and their relative intensities. Apart from calcite and

magnesian it was possible to recognize a third phase gallium iron sulfide in the ash at 525 °C of the paper sample tested (Figure 5 & 6). XRD diagrams were taken for each ignition temperature separately [20]. The X-ray source was CuKa radiation with wavelength of λ =1.54Å. The sample i.e., ash on ignition at 900 °C, was scanned in step-scan mode in area between 3° and 110°20 with step size of 0.02°20 (Figure 6 & 7).



Figure 5: More detailed XRD	pattern syn CaO, o	of the ash on ignition at 900	°C of a 100g/m ² copy paper sample.
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X-Ray	40 kV, 15 mA	Scan speed / Duration time	10.0000 deg/min
Goniometer	Mini Flex 300/600	Step width	0.0200 deg
Attachment	Specimen Rotation Attachment	Scan axis	Theta/2-Theta
Filter K-beta(x1)		Scan range	3.0000 - 110.0000 deg
CBO selection slit -		Incident slit	1.250deg
Diffracted beam mono. None		Length limiting slit	10.0mm
Detector D/teX Ultra		Receiving slit #1	13.0mm (Open)
Scan mode CONTINUOUS		Receiving slit #2	13.0mm

Figure 6: Measurement conditions of the XRD pattern of the syn CaO (lime).

No.	2-theta(deg)	d(ang.)	Height(counts)	FWHM (deg)	Int. I (counts deg)	Int. W(deg)	Asym. factor
1	32.1464(17))	2.78221(14)	1032(32)	0.286(4)	344(4)	0.333(14)	0.56(4)
2	37.261(3)	2.4112(2)	1745(42)	0.110(11)	226(100)	0.13(6)	1.5(3)
3	37.343(14)	2.4061(9)	1335(37)	0.14(4)	219(93)	0.16(7)	1.5(3)
4	37.486(11)	2.3973(7)	1546(39)	0.26(3)	474(55)	0.31(4)	1.5(3)
5	48.269(6)	1.8839(2)	12(3)	0.127(12)	2.12(12)	0.18(6)	0.93(14)
6	53.8231(18)	1.70188(5)	1374(37)	0.260(5)	412(5)	0.300(12)	0.57(5)
7	64.130(3)	1.45098(6)	438(21)	0.239(8)	122(2)	0.278(19)	0.76(10)
8	67.308(3)	1.38999(6)	296(17)	0.124(7)	43(7)	0.14(3)	0.90(10)
9	67.451(7)	1.38739(13)	311(18)	0.221(16)	80(7)	0.26(4)	0.90(10)
10	79.636(3)	1.20293(3)	174(13)	0.210(7)	47.7(10)	0.27(3)	0.60(9)
11	88.530(3)	1.10361(3)	180(13)	0.226(8)	54.3(12)	0.30(3)	0.83(15)
12	91.491(2)	1.07546(2)	454(21)	0.218(5)	126(2)	0.278(17)	1.05(13)
13	103.353(4)	0.98187(3)	364(19)	0.235(4)	107.2(11)	0.295(19)	0.64(5)
14	107.37(16)	0.9560(10)	3.3(18)	0.16(15)	0.6(6)	0.2(3)	1(5)

Figure 7: Peak list of the XRD pattern of syn CaO. The x-ray diffractogram of the ash on ignition of the paper sample at 900oC showed a single-phase lime CaO.

 $CaCO_3$ + heat \rightarrow CaO + CO₂ (900°C)

Figure 8: Scheme of chemical reaction of CaCO3 present in the paper sample to becoming CaO during ignition of the copy paper sample in the oven at 900 °C.

Crystallite size basic theory for the XRD pattern of syn CaO

The full width at half maximum (FWHM) of the reflection depended both on the diffractometer geometry and the setups such as the slits, the optics, the 2θ angle and the detector, also on the sample itself. For ideal conditions the standard powder diffractometer may have produced diffractograms with peak widths

(FWHM) between 0.03° and $0.1^{\circ} 2\theta$ over the whole angular range. Here the peak width (FWHM) was between 0.110° and $0.286^{\circ} 2\theta$. The peak broadening caused by the sample, i.e., ash on ignition at 900 °C, could have been explained by the presence of very small crystallites and micro-strain.

Particle size calculation of the XRD phase CaO (lime)

For Rietveld refinement (WPPF) the background was calculated with a β -spline function. B-spline was selected as the function expressing the background. The function β -spline used a smooth curve passing through the calculated points at the background level as the function expressing the background (Table 4).

Table 4: The grain size of CaO nanoparticles present in the ash on ignition at 900 °C of the copy paper sample.

2θ of the intense peak (deg)	θ of the intense peak (deg)	FWHM of intense peak (β) radians	Size of the particle (D) in nm
32.1464	16.0732	0.004992289	28.91101145
37.261	18.6305	0.001920111	76.22447283
37.343	18.6715	0.002443778	59.90512436
37.486	18.743	0.004538444	32.27023901
48.269	24.1345	0.002216856	68.55402066
53.8231	26.91155	0.004538444	34.27017215
64.13	32.065	0.004171878	39.22846771
67.308	33.654	0.002164489	76.97648485
67.451	33.7255	0.003857678	43.22637227
79.636	39.818	0.003665667	49.25886152
88.53	44.265	0.003944956	49.09278755
91.491	45.7455	0.003805311	52.22698315
103.353	51.6765	0.004102056	54.52319264
107.37	53.685	0.002792889	83.85044595
	53.46561686		

The Crystallite size calculation, and micro-strain ϵ , related to peak broadening, for the XRD pattern of syn CaO (lime)

The Scherrer equation [21] described the relation between the broadening of the reflection and the average crystallite size τ , by the following relation:

$$\tau = \frac{K * \lambda}{\beta * \cos \theta} \quad (1)$$

whereas $\boldsymbol{\beta}$ was the broadening of the reflection due to small

crystallite sizes and was equal to FWHM of the reflection of the real sample, K was approximately 0.9 and λ was the wavelength and θ the diffraction angle.

Figure 9 described the line broadening effect given by decreasing crystallite sizes. Its influence was stronger than instrumental broadening for crystallite sizes which were much smaller than 1 μ m, as could be seen in Table 4 & 5. Figure 9 described the line broadening effect given by the crystallite sizes. The diffraction peak width may have been containing microstructural information.

Table 5: In Table 5 we represented the line width $(o2\theta)$ i.e., Int. W(deg) (Figure 7) along with the particle dimension as calculated in Table 4.

Line Width (o20)	Particle Dimension (Å)
0,333	289,1101145
0,13	762,2447283
0,16	599,0512436
0,31	322,7023901
0,18	685,5402066
0,3	342,7017215
0,278	392,2846771
0,14	769,7648485
0,26	432,2637227
0,27	492,5886152

0,3	490,9278755
0,278	522,2698315
0,295	545,2319264
0,2	838,5044595



Figure 9: Line broadening as a function of the crystallite size.

In order to distinguish between broadening by strain and broadening by crystallite size the reflection profile was analyzed. It was calculated the product of β by $\cos\theta$, whereas β was the broadening of the reflection due to small crystallite sizes and

was equal to FWHM of the reflection of the real sample and θ was the diffraction angle. Also, sin θ was calculated, and a graphical representation was constructed of sin θ versus β cos θ (Table 6).

Table 6: In table 6 we represented $\beta \cos\theta$ whereas β was FWHM in radians (Table 4) along $\sin\theta$, whereas θ was the diffraction angle.

βcosθ	Sinθ
0,004737134	0,27686522
0,001819494	0,31946379
0,002315161	0,32014179
0,004297768	0,32132377
0,002023076	0,40888004
0,00404696	0,45261447
0,003535443	0,530881
0,001801719	0,55417631
0,003208458	0,55521464
0,002815534	0,64035103
0,002825059	0,69797796
0,002655524	0,71624714
0,002543688	0,7845221
0,001654016	0,80577327

From the plot of structure $\beta^*\cos\theta$ on y axis against sin θ on x-axis (Figure 10 &11) we calculated the strain component from the slope which was -0.00213252rad. The negative strain component indicated that the micro strain was compressive. Also, from the plot

of structure $\beta^*\cos\theta$ on y-axis against sin θ on x-axis we calculated the particle size component from the y intercept which was 0.004006175rad (Table 7).

Table 7: In Table 7 we represented $\frac{\beta}{4\tan\theta} = \varepsilon$ i.e., the micro-strain, along β which was the FWHM of the reflection of the sample in radians.

β/4tanθ	β
0,004331651	0,004992289
0,001423866	0,00190111

0,001807918	0,002443778
0,003343799	0,004538444
0,001236962	0,002216856
0,002235324	0,004538444
0,001664894	0,004171878
0,000812791	0,002164489
0,001444693	0,003857678
0,001099215	0,003665667
0,001011872	0,003744956
0,000926888	0,003805311
0,000810585	0,004102056
0,000513177	0,002792889







Figure 11: The peak broadening β was related to micro-strain $\varepsilon = \frac{\beta}{4 \tan \theta}$ and the lattice strain was calculated to be 0.0004719727188044rad or 4.719727188044x10⁻⁴ rad from the intercept of the graph.

Estimation of the band gap energy Eg from the DRS study of the A4 copy paper sample

A graph was plotted of $(K/S hv)^2$ versus hv. The extrapolation of the straight line to $(K/S hv)^2=0$ axis (Tauc Plot) gave the value

of the band gap energy E_g . The ISO brightness was measured to be 100.95%. Its opacity was 95.87% and transparency 11.82%. The K/S value according to equation (2)

$$\frac{K}{S} = \frac{(1-R)^{2}}{2R}$$
 (2)

was calculated to be 49.47995295. Equation (2) was the Kubelka-Munk equation at any wavelength. It was applicable in the limiting case of an infinitely thick sample like a paper sheet, where thickness and sample holder have no influence on the value of reflectance (R) [22].

The band gap energy of the chromophores presents in the cellulose of the $100g/m^2$ A4 copy paper sample was estimated to be 1.1654879939663700eV for permitted indirect transitions, or 1.867111765E-19Joules, at room temperature (Figure 12). The band gap was estimated from the function trend in the excel spreadsheet with the data calculated for the whole range of

wavelengths 400-705nm. A pH value of 8.80 led to an optical band gap value of 1.1654879939663700eV which was a pH lower than 9, or 10, but still resulting in the paper's nanoparticle aggregation. The trend in the band gap was a decreasing energy band gap because of the presence of calcite CaCO₃ and magnesian MgCO₃ in the paper polymer matrix, compared to a previous band gap of 1.1697157220054000eV at a pH value of 9.6 [23], where only calcite was present. The concentration of ions specifically H⁺ and of OH⁻ also affected the band gap of the paper sample towards a lower band gap. The indirect bandgap thus decreased when the pH lowered from 9.6 to 8.8 [24].





General Indexing Method (trial and error) for the XRD pattern of syn CaO (lime)

The general indexing method was applied in this study whereas certain parameters were known, such as the wavelength of the radiation, the θ and 2θ values of the reflections from the experiment, the crystal system, the constants a, b, c, α , β , γ , of the crystalline material from the experiment or from the literature. The principle of the method was the determination of the values of the hkl indices of a reflection for which the squared difference of the experimental and theoretical value of a quantity became minimal, i.e.,

$$/Q_0 - Q_c / = minimum (3)$$

Where as
$$Q_0 = \left(\frac{1}{d_0}\right)^2 = \left(\frac{2\sin\theta o}{\lambda}\right)^2$$
 (4), and $Q_C = \left(\frac{1}{dc}\right)^2$ (5)

The quantity $\frac{1}{dhkl}$ was a quantity whose experimental value was given by Bragg's equation by solving for $\frac{1}{dhkl}$ and raising it to the square, which was the quadratic form of the Bragg equation.

The procedure was to successively giving h,k,l, simple integer values (positive, or negative or 0), until the best possible agreement between the experimental value and the corresponding theoretical value was obtained. Thus, we determined the hkl indices of the

reflections which corresponded to each specific θ . The crystal system of lime syn, CaO, identified by experiment was cubic. The space group was Fm3m with unit cell with a=b=c=4.81Å and the corners were α = β = γ =90° and the wavelength was λ =1.5406Å.

From the peak list in Figure 7, we read 2θ =32.1464° from the experiment. Then

$$Q_0 = d_0 = \frac{\lambda}{2\sin\theta} = \frac{1.5406}{2\sin16.0732} - \frac{1.5406}{0.55373044} = 2.782220172$$
(6)

The equidistance d_{hkl} with respect to the cell constants and indices was calculated as in equation (7):

$$Q_{C} = Q_{hkl} = \frac{1}{(2.782220172)^{2}} = 0.129187408 \text{Å}^{-2}$$
(7) and
$$Q_{hkl} = \frac{1}{dhkl^{-2}2} = \frac{h^{2} + k^{2} + l^{2}}{a^{2}} = \min (8)$$

From where $\frac{h^2 + k^2 + l^2}{(4.81\text{ Å})^2} - 0.129187408\text{ Å}^2 - 2 = \min$ (9). Then it was revealed that $h^2 + k^2 + l^2 = 2.98889279 \sim 3$ and then hkl=111. The indices of the reflection $\theta = \frac{32.1464}{2}$, were hkl=111.

From the peak list in Figure 7, we read $2\theta{=}37.261^\circ$ from the experiment. Then

$$Q_0 = d_0 = \frac{\lambda}{2\sin\theta} = \frac{1.5406}{2\sin18.6305} = \frac{1.5406}{0.63892757} = 2.411227927$$
(10)

The equidistance d_{hkl} with respect to the cell constants and indices was calculated as in equation (11):

$$Q_{c} = Q_{hkl} = \frac{1}{(2.411227927)^{2}} = \frac{1}{5.814020116} = 0.171998028 \text{Å}^{-2}$$
(11) and
$$Q_{hkl} = \frac{1}{dhkl^{2}} = \frac{h^{2} + k^{2} + l^{2}}{(4.81 \text{\AA})^{2}} = \min (12)$$
$$\frac{h^{2} + k^{2} + l^{2}}{(4.81 \text{\AA})^{2}} = 0.171998028 \text{\AA}^{-2} = \min (13)$$

Then it was revealed that $h^2+k^2+l^2=3.979363576\sim4$, and then hkl=002, or hkl=200, or hkl=020.

From the peak list in Figure 7, we read $2\theta{=}37.343^\circ$ from the experiment. Then

$$Q_0 = d_0 = \frac{\lambda}{2\sin\theta} = \frac{1.5406}{2\sin18.6715} = \frac{1.5406}{0.640283581} = 2.406121359$$
(14)

The equidistance d_{hkl} with respect to the cell constants and indices was calculated as in equation (15):

$$Q_C = Q_{hkl} = \frac{1}{(2.406121359)^2} = \frac{1}{5.789419994} = 0.172728874 \text{ Å}^{-2}$$
 (15) and

$$Q_{hkl} = \frac{1}{dhkl^2} = \frac{h^2 + k^2 + l^2}{(4.81\text{ Å})^2} = \min$$
(16)

From where $\frac{h^2 + k^2 + l^2}{(4.81 \text{ Å})^2} - 0.172728874 \text{ Å}^{\wedge} - l = \min (17)$ Then it was revealed that $h^2+k^2+l^2=3.996272502\sim4$, and then hkl=002, or hkl=200, or hkl=020.

From the peak list in Figure 7, we read $2\theta{=}37.486^\circ$ from the experiment. Then

$$Q_0 = d_0 = \frac{\lambda}{2\sin\theta} = \frac{1.5406}{2\sin18.743} = \frac{1.5406}{0.642647547} = 2.39727049$$
(18)

The equidistance d_{hkl} with respect to the cell constants and indices was calculated as in equation (19):

$$Q_{C} = Q_{hkl} = \frac{1}{(2.39727049)^{2}} = \frac{1}{5.746905802} = 0.17400668 \text{ Å}^{\wedge} -2 \text{ (19) and}$$
$$Q_{hkl} = \frac{1}{dhkl^{2}} = \frac{h^{2} + k^{2} + l^{2}}{(4.81 \text{ Å})^{2}} = \min (20)$$

$$\frac{h^2 + k^2 + l^2}{2} - 0.17400668 \text{ Å}^{\wedge} - 2 = \min$$

From where $(4.81\text{ Å})^2$. Then it was revealed that $h^2 + k^2 + l^2 = 4.025835949 \sim 4$, and then hkl=002, or hkl=200, or hkl=020.

Indexing process of powder diffraction pattern was done and Miller Indices (hkl) to each peak was assigned in first step. Indexing has been done in two different methods and data were in Table 8-10 [25]. In Table 9 we found a dividing constant and the values of the third column became approximately indegers. Here the constant was calculated to be 25.403(=102.057-76.654).

Table 8: X-ray powder diffraction data of the syn lime (CaO) with all the reflexions hkl.

No.	2-theta(deg)	d(ang.)	h	k	l	Separation of plane (nm) in the cubic lattice
1	32.1464(17))	2.78221(14)	1	1	1	0.2782210002
2	37.261(3)	2.4112(2)	0	0	2	0.2411200001
3	37.343(14)	2.4061(9)	0	0	2	0.2406100001
4	37.486(11)	2.3973(7)	0	0	2	0.2397300002
5	48.269(6)	1.8839(2)	2	1	1	0.1883900003
6	53.8231(18)	1.70188(5)	2	2	0	0.1701880001
7	64.130(3)	1.45098(6)	2	2	1	0.145098
8	67.308(3)	1.38999(6)	3	1	1	0.1389990001
9	67.451(7)	1.38739(13)	3	1	1	0.1387390001
10	79.636(3)	1.20293(3)	4	0	0	0.120293
11	88.530(3)	1.10361(3)	4	1	1	0.110361
12	91.491(2)	1.07546(2)	4	2	0	0.107546
13	103.353(4)	0.98187(3)	4	2	2	0.0981869999
14	107.37(16)	0.9560(10)	5	0	0	0.0956

Table 9: Simple peak indexing for the XRD pattern of syn CaO.

Peak position, 20	1000*sin ² θ	1000*sin ² θ / 25	Reflection	Remarks
32.1464	76.654	3	(111)	$1^2 + 1^2 + 1^2 = 3$
37.261	102.057	4	(002)	$0^2 + 0^2 + 2^2 = 4$
37.486	103.249	4	(002)	$0^2 + 0^2 + 2^2 = 4$
53.8231	204.859	8	(220)	$2^2 + 2^2 + 0^2 = 8$

Table 10: Peak indexing from d-spacing for the XRD pattern of syn CaO.

Peak Position, 20	d	1000/d ²	(1000/d²)/42.815	hkl
32.1464	2.78221	129.187	3.02	111
37.261	2.4112	172.0020132	4.02	002

37.486	2.3973	174.0023963	4.06	002
53.8231	1.70188	345.2567124	8.06	220

Four main peaks at 20 values of 32.1464, 37.261, 37.486 and 53.8231 degrees corresponding to (111), (002), (002) and (220) planes of CaO were observed. In Table 10 we found a dividing constant and the values of the third column became approximately indegers. Here the constant was 42.815 (= 172.0020132-129.187).

The separation of planes in the cubic lattice of syn CaO.

The Miller indices were useful for expressing the separation of planes (Table 8). The separation of the 111 plane in the cubic lattice was calculated to be 0.2782210002nm.

Thus,
$$Q_{hkl} = Q_{111} = \frac{1}{d111^2} = 0.129187408 \text{ Å}^{-2}$$
 and $d_{111} = 0.2782210002 \text{ nm}.$

The separation of the 002 plane in the cubic lattice was calculated to be 0.2411200001 nm.

Thus,
$$Q_{hkl} = Q_{002} = \frac{1}{d002^2} = 0.172002013 \text{ Å}^{-2}$$
 and $d_{002} = 0.2411200001 \text{ nm}.$

The separation of the 002 plane in the cubic lattice was calculated to be 0.2406100001nm.

Thus,
$$Q_{hkl} = Q_{002} = \frac{1}{d002^2} = 0.172731941 \text{ Å}^{-2}$$
 and $d_{002} = 0.2406100001 \text{ nm}.$

XRD was applied to identify qualitatively the powder's ash on ignition crystalline phases. A diffractogram was constructed by plotting the produced pattern to link the diffraction angle to the diffracted intensity. The CaCO₃ present along with MgCO₃, GaS and FeS in the copy paper sample and identified by XRD, improved the mechanical and the optical properties of the final product.

For the A4 copy paper sample 100gsm studied in this work it was of interest to carry out a study of the powder residue on ignition x-ray powder diffraction structure, among its other properties. The small bandgap of 1.1654879939663700eV of the copy paper sample could classify the material as semiconductor which contained four types of cations of Ca, Mg, Ga, Fe and anion S [26].

Conclusion

This study reported the presence of CaO present in the residue on ignition of a copy paper sample at a temperature of 900 °C for one hour time. The presence also of CaCO₃, MgCO₃, GaS and FeS was also revealed in the residue on ignition of the same copy paper sample, but at a temperature of 525 °C for also a one-hour time. The mineralogical changes that had occurred in the ashes on ignition, of the paper sample, as a function of temperature, were identified by X-Ray Diffraction (XRD). Calcite (CaCO₃) was decomposed at 900 °C to CaO. A higher particle size of 53.5nm in the phase of CaO coincided with a lower band gap value of the paper material studied. Finally, XRD data could provide useful information and also helped correlate microscopic observations with the bulk sample such as copy paper.

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