New Ultramarine Generations from Egyptian Raw Materials

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Abstract

New ultramarine generation have been prepared by the solid state mixing and firing technique from Egyptian Raw Materials for the first time. The raw materials used includes kaoline, Na₂CO₃, charcoal and sulphur precursors. The action of NH₄Cl on the obtained ultramarine has been studied using X-ray, UV absorption and DC-electrical conductivity, whereas a mid et esr appears. Results of TGA, DTA and electrical conductivity have been undertaken and interpreted for the first time.

I. Introduction

Kowalak et al.¹ found that alternative methods of ultramarine synthesis involving zeolites as principal substances were developed in their lab contrary to the conventional technology. Introduction of sulfur radical precursors into the zeolites allows one to obtain ultramarine with almost negligible emission of SO₂ upon calcination. The polysulfides S₃²⁻ and S₆²⁻ seem to be the most effective precursors. Modification of hydrothermal synthesis of sodalite in the presence of sulfur compounds (including sodium thiosulfate) don’t result in products of deep blue colouration. Thermal crystallization of the mixture containing kaolin and sodium polysulfides, which were carried out under the industrial conditions, resulted in the product similar to the common ultramarine.

Gobeltz et al.² investigated the encapsulation of the chromophores into the sodalite structure during the synthesis of the blue ultramarine pigment from metakaolin, sulfur, sodium carbonate and a reducing agent. The insertion of the chromophores occurs during the formation of the sodalite structure and also after its formation by diffusion into the sodalite cages. The possibility of diffusion of the chromophores into the sodalite structure was proved independently by synthesis of blue ultramarine pigments from sodalite, previously synthesized and mixed with a source of polysulfindes.

Kowalak³, investigated that ultramarine analogs are manufactured by roasting zeolites A, X, Y or sodalite impregnated with S radical (Chromophore precursors and optionally reducing agents, e.g. pitch. The precursors are S or alkali metal (preferably Na) polysulfides and the calcination is carried out preferable at 973 – 1173 K in the presence of partial absence of the air followed by reheating at 723 – 1023 K in the air for example, a little pigment was manufactured by roasting comminuted zeolite A mixture with Na₂S. 9H₂O Combined with pitch.

The loss of weight of the reaction mixture during calcination is < 5% and SO₂ emission is significantly lower than in the traditional procedures employing kaolin instead of zeolites.

Tae et al.⁴ studied the reaction mechanism in synthesis of ultramarine blue from kaolin, S, Na₂CO₃, SiO₂ and rosin by thermal analysis, a powder X-ray diffraction analysis and Ramon scattering. The mixture of the raw materials was calcined at 820°C for 5 hrs. (heating rate; 0.1°C/min) to form the ultramarine green, which was the intermediate products of the ultramarine blue. The ultramarine blue was finally prepared with the oxidation of ultramarine green at 500°C.
In the calcination process, the Na sulfides were generated by the reductive reaction of Na$_2$CO$_3$, S and rosin at 400 – 500°C, and the NaAlSiO$_4$ was formed by the synthesis of anhydrous amorphous kaolin and the Na sulfides at ~ 600°C, and the ultramarine green begun to form at ~ 700°C with the reaction of the NaAlSiO$_4$ and the Na sulfides. In the oxidation process, the S atoms transformed the $(S^2_-)$ ions of the ultramarine into $(S^3_-)$ ions, and these reactions caused the ultramarine green to convert the ultramarine blue. The lattice parameter of ultramarine green was increased during oxidation.

Kowalak$^5$, showed that a reaction mixture containing kaolin, soda or other alkaline compound, an agent facilitating incorporation of anion radicals $(S^3_-)$ into sodalite structure, and reducing agent is roasted at 1023 K and then at ~ 829 K to give ultramarine. The agent facilitating incorporation of anion radicals $(S^3_-)$ into sodium structure is selected from alkali metal sulfides, preferably sodium sulfide.

Leschewski and Moller$^6$, deduced that when blue ultramarine is fused with HCOONa a white product results without loss of S. This white substance is leuco-ultramarine and is reconnected into coloured product by atmo, by removal of alkali by means of S, by treatment with Cl water of CI gas or fusion with alkali chlorates or nitrates, blue ultramarine boiled with ethylene chlorohydrine loses most of its Na content by removal of alkali and becomes white. Fusion with NaCl does not restore the colour but fusion with Na$_2$S produces a delicate blue and boiling Na$_2$S solution gives a delicate green.

Jaeger$^7$, showed that the ultramarines are stable toward alkalies but unstable toward acids. Na can be replaces by Li (bluish violet), Rb, Cs, K, Tl (violet), Ag (light yellow), Mg (light blue) and other bivalent metals (colourless) by treatment in aqueous solution at 100 - 120°C. Replacement of S by Se or Te yields blood-red or yellow ultramarine resp. They posses hydraulic properties and resemble permutities in many respects. The substitution of Na by Ag leads to an equal dependent on concentration and temperature and that of Ag by alkali goes more readily with iodides than with chlorides. Replaceability and reaction velocity increase with decrease in at. wt. of the alkali metal. All these ultramarines yield spectrograms identical with those of nosean and haunynte. The centered cubic lattice has a 9.13 A.u. The space groupings of the atoms, certain of which appear to be “wandered” are discussed.

Hoffmann$^8$, showed that the sodalite complex possesses an inert silicate component, Na$_4$Al$_4$Si$_6$O$_{24}$ which in the presence of Na$_2$SO$_3$ takes part in reaction as Na$_6$Al$_4$Si$_6$O$_{24}$. Ultramarine substrates exhibits characteristic irradiation effects corresponding to their chemical compound and these present a series of colour changes opposite in direction to those obtained with polysulfide S. Irradiation of white-green and violet ultramarine produces colour alterations which no longer occur if the return of the dissociation. S in blue ultramarine into the original complex is prevented. The behaviour of blue ultramarine is not contrary to the assumption that ultramarine is definite S atoms of the various substrates.

Kook et al.$^9$ showed that ultramarine blue was synthesized with Kaolin, sulfur, sodium carbonate, silica and rosin, and its reaction mechanism was investigated. The mixture of the raw materials was calcined at 820°C for 4 hrs (heating rate: 2°C/min) to form the ultramarine green, which was the intermediate of ultramarine blue. The ultramarine blue was finally prepared with the oxidation of the ultramarine green at 500°C. The reaction products and the structural changes of ultramarines appeared in this synthesis were established by power X-ray diffraction analysis and Raman Scattering. The sodium sulfide and the NaAlSiO$_4$ were generated at 500°C and 620°C respectively in calcination process and the ultramarine green was appeared at near 740°C with the reaction of sodium sulfates and NaAlSiO$_4$. Also, the formation of ultramarine blue was caused by the sulfur atoms generated from the oxidation of sodium sulfide which existed in ultramarine green.

Ultramarine blue pigments have been investigated by EPR and Raman spectroscopies, as well as by
colorimetry-for the first time, correlations between EPR, Raman, and colourometric parameters of these pigments have been established. It’s now established that there isn’t only one blue ultramarine but different shades of blue depending on the relative concentrations of blue and the yellow chromophores. The greenish and reddish shades of blue are characterized by EPR. Raman and colourometric parameters as suggested by N- Gobeltz et al.\textsuperscript{10}.

Balkus and Kowalak\textsuperscript{11} worked on the crystallization zeolites (A, X, hydroxysodalite) in the presence of sulfur compounds in order to incorporate them into the intracrystalline cavities already on nucleation of the zeolite structure. This method of encapsulation has been proved successful for various metal complex compounds.

The major goal of the present investigation is to discover new ultramarine generations from Egyptian raw materials for inorganic pigments industries.

Materials

These above samples (kaolin, anhydrous sodium carbonate, charcoal and sulfur precursor are finely ground together and tightly packed into a small porcelain crucible which is then covered; the crucible is heated to at least a bright red for 1.5 hrs. Then the cover is removed and after firing for an additional 2 hrs and cooling, the ultramarine easily separated forms the uncoloured materials. Composition and condition of preparation of the ultramarine samples are shown in table (1).

X-ray Diffraction analysis: The prepared samples were subjected to Cu-K\(\alpha\) X-ray analysis using Schimadzu (Japan) X-ray diffractometer, using Ni-filter at room temperature in the range \(2\theta = 10-70^0\)C.

Infrared Absorption Spectral Measurements: The room temperature IR absorption spectra were recorded using the solid KBr disk technique and a-Perkin Elmer infrared spectrophotometer in the range 4000 - 200 cm\(^{-1}\).

Electronic Absorption Spectral (UV/VIS) Measurements: The ultramarine (blue and violet) samples were subjected to UV/Visible absorption spectral measurements using the Nujol mull (glycerine suspension) solid technique and the results were compared to those of commercial ultramarine (produced by Reckit’s Colours of Polifarb). A PYEUNICAM SP 800 spectrophotometer was used in the range 900 to 200 at room temperature.

Thermogravimetric analysis (TGA): TGA is measure of the amount and rate weight change of material, as a function of increasing temperature (weight change in temperature). Our measurements were undertaken in the temperature range 0 – 800\(^{\circ}\)C in air using Schimadzu (Japan) Thermal Analysis.

Differential thermal analysis (DTA): DTA is a technique in which a record is made of the temperature difference between the sample and the reference material against time or temperature. DTA records every enthalpy, exothermic and endothermic as caused by structural or chemical changes. In the temperature range (10 mv/cm) from 20 up to 600 with pure Al\(_2\)O\(_3\) as a reference material using DT-30 Schimadzu thermal analyzer in air.

Electronic Spin Resonance Measurements: The Electron Spin Resonance spectra were recorded at X-band frequencies on JES-FE 2 XG, ESR spectrometer at room temperature at the Central Research Lab, Tanta University, Egypt. The magnetic field was swept from 0 to over 8 KG and the calibrated energy was 1 KG with a digital gaussmeter. Several selected samples were investigated and showed identical features. DPPH was used as internal reference.

DC-Electrical Conductivity Measurements: The DC-electrical conductivity was measured using the two terminals DC-method. The pellet was inserted between spring loaded copper electrodes. A Kiethley 175-Multimeter (USA) was employed from room temperatures up to 573 K. The temperature was measured by a calibrated chromel-alumel thermocouple placed firmly at the samples. Measurements were conducted in such a way that at each temperature a sufficient time was allowed for thermal equilibrium.

Results and Discussion
Fig. 1 display the Cu-Kα X-ray diffraction pattern of ultramarine samples. It can be easily seen that, the kaolinite phase disappeared completely from X-ray pattern of the ultramarine. This suggests the complete formation of ultramarine instead of kaolinite phase. As deduced by Stoky et al. for both natural (Lazurite, Laps lazuh) and synthetic ultramarine show the structure of sodalite. From (ASTM card No. 3 - 0334) ultramarine sample have the formula (Na₄Al₃Si₃O₁₂Cl) sodium aluminium silicate. As illustrated in table 1, the sulfur source affects the structure and the colour of the resulting ultramarine samples, Thus XRD pattern given in fig. 1 show that calcination with sodium polysulfides leads to sodalite structure.

Since the sodalite structure with formula Na₄Al₃Si₃O₁₂Cl with cages of 0.6 nm in diameter are stable and they can host variety of molecules (e.g. NaCl in Natural sodalite). Even large molecules containing as many as seventeen atoms can be encapsulated. So encapsulation of elemental sulfur and sulfur precursor is available. The calcination at high temperature (~ 800°C) was usually conducted with reductive agent (Pitchtar). The resulting samples show intense blue colouration similar as natural ultramarine. The interesting structure transformation towards sodalite was noticed for impregnated zeolites A, X, Y upon heating at temperatures higher than 600°C. X-ray patterns also display that most of peaks are characteristic to synthetic ultramarine nearly at the same positions as those of sodalite structure but my be shifted to lower 2θ i.e. increase d-spacings. This is ascribed to the encapsulation of either sulfur element or sulfur precursor in the sodalite cages. These are of larger diameter than that of chloride in natural sodalite. In natural sodalite the maximum intensity is that of peak at 2θ - 24.52° but in our synthetic ultramarine samples, the maximum intensity is that of peak at 2θ ≅ 24° for samples (3) and (4) that containing Na₂S₂O₃ and Na₂SO₃ which is similar to that of natural sodalite on the other hand we found that the maximum intensity is that of peak at 2θ ≅ 32° for sample (5), (6) (These samples are of blue and intense blue colour. This is interpreted on the basis of formation of other ultramarine phase.

Fig. 2 shows the effect of adding NH₄Cl on the values of 2θ and (d Å) of ultramarine blue (containing S and Na₂SO₃). We found that, the maximum intense peak is that appear at 2θ = 28.33° shifted to lower 2θ value in a comparison to ultramarine blue containing (sulfur element and Na₂SO₃). Nearly all characteristic 2θ values of our synthetic ultramarine shifted to lower 2θ value (higher d Å) values mean that Cl⁻ ions replaces S₃⁻ radical and thus causing fading of blue colour. It has been found that upon the addition of NH₄Cl to ultramarine sample prepared by using elemental sulfur on the colour and structure of the produced ultramarines.
sulfur and Na$_2$SO$_3$ as sulfur precursor we found that the blue colour undergo faddening and become violet. The maximum intense peak at $2\theta \cong 32^\circ$ that characteristic for sodalite, shifts to lower 2$\theta$ value (increased d-spacing) whereas $S_3^-$ radicals are responsible for blue colour.

![Figure-2: The X-ray diffraction patterns of the prepared ultramarine blue and violet containing:](image)

- a) Sulfur as sulfur precursor.
- b) Sulfur as sulfur precursor and NH$_4$Cl

The IR absorption spectral bands of ultramarine samples appear in the medium 535 cm$^{-1}$ and 1123 - 1129 cm$^{-1}$ region$^{24}$ (Fig. 3. and Table 3). It can easily be seen that, the IR spectra modes in this region are believed to originate from $S^{3-}$ & $S^{2-}$ in a sodium alumino-silicate matrix Na$_8$[Al$_6$Si$_6$O$_{24}$]Sn$^{18}$. It has been found that the IR absorption bands at 1123 - 1125 cm$^{-1}$ suffers shift to shorter wave number (red shift) in ultramarine samples containing S element to ultramarine samples containing S and Na$_2$SO$_3$ as sulfur precursor associated with increase in the intensity of the blue colour of the synthetic ultramarine$^{19}$. The IR absorption spectral band at 624 cm$^{-1}$ is due to Si-O stretching vibration$^{20}$. The IR absorption bands appearing at $\cong 3428$ cm$^{-1}$ and 1627 cm$^{-1}$ are characteristic for water of crystallinity.

The ultramarine samples were subjected to UV/visible absorption spectral measurements using glycerin suspension$^{15}$ and they were comparable to those of commercial ultramarine (produced by Reckit’s colours of polifarb). The estimation was verified by electronic spectral assignments. The correlation between the absorbance measured for the samples and the wave number. Fig. 4 indicates the maximum intensity of the band at $\sim 610$ nm for the sample prepared with poly sulfide (5,6). The highest intensity of blue colour of the above two samples could result from feasible formation of $S_3^-$ radicals from $S_3$ anion$^{21}$. The maximum of this absorption peak are slightly shifted towards higher frequency blue shift (604 nm - 606 nm) compared to that of commercial ultramarine 610 nm. There is other band notable in the spectral at $\sim 370$ nm. This band is assigned to S$_2$ species$^{21}$. The absorbance of this band is considerably higher for polysulfides than for sulfide or elemental sulfur. The intensity of this peak does not depend considerably on sulfur number$^{21}$. The band appears at $\sim 610$ nm was shifted to lower frequency for nearly prepared samples (1), (2), (3), (4), (5), (6), (7). We deduced that the maximum intensity of this peak depends considerably on the calcination time as well as on the sulfur number (or sulfur precursor). It was
noted that high firing temperature during ultramarine synthesis led to deep blue colouration (see fig. 4). There is also another electronic band appearing at \( \sim 238 \text{ nm} \) the absorbance of this band increase with colour intensity and sulfur precursor increases. It has been found that the absorbance of the band at 604 nm decreases as sulfur precursor decrease and also by the addition of ammonium chloride to either prepared samples or to its starting mixtures (fig. 4).

![Absorbance spectra](image)

**Wavelength (nm)**

**Figure-4:** Electronic absorption spectra of the prepared ultramarine samples containing:

- a) Sulfur element
- b) Sulfur element and NH\(_4\)Cl
- c) Na\(_2\)S
- d) Na\(_2\)S\(_2\)O\(_3\)
- e) Contain Na\(_2\)SO\(_3\)
- f) Sulfur and Na\(_2\)SO\(_3\)
- g) Sulfur and Na\(_2\)SO\(_3\) (fired for 10 hr)

Figs. 5 and 6 represent the TGA and DTA curves of ultramarine samples containing sulfur as sulfur precursor, the two endothermic peaks at 95.13\(^\circ\)C and 772.93\(^\circ\)C and three exothermic peaks with maximum at 170.23\(^\circ\)C, 245.42\(^\circ\)C and 353.95\(^\circ\)C. The first endothermic peak at 95.13\(^\circ\)C which is accompanied by 1.84% weight loss (see TGA curve fig. 5) is associated to dehydration of humidity water content. The second endothermic peak at 772.93\(^\circ\)C which is accompanied by 1.10% weight loss (see TGA curve fig. 6) is due to CO\(_2\) evolution and oxygen uptake (oxidation) during the formation of ultramarine blue. The first exothermic peak at 170.23\(^\circ\)C is accompanied by 5.44% weight gain (see TGA curve) is assigned to the first step of partial oxidation of charcoal and evolving CO/CO\(_2\) during the formation reaction. The two exothermic peaks at 245.42\(^\circ\)C and 353.95\(^\circ\)C which are accompanied by 1.95% and 1.68% weight gain (see TGA curve fig. 5) are assigned to the second step in the partial oxidation of charcoal and evolving CO/CO\(_2\) during the formation reaction. The weight loss caused by evolving SO\(_2\) is almost negligible. Thus, result of TGA and DTA for these faint blue ultramarine samples are in full agreement and support one another.

**Figure-5:** The thermogravimetric analysis (TGA) of ultramarine sample containing sulfur element

For the sample with sodium sulfite (Na\(_2\)SO\(_3\)) as sulfur precursor fig. 7 and 8 represents the TGA and DTA curves of the sample. Two endothermic peaks are observed with maxima at 102.65\(^\circ\)C and 442\(^\circ\)C. Other three exothermic peaks are observed with maximum at 197.02\(^\circ\)C, 242.36\(^\circ\)C and 360\(^\circ\)C. The first endothermic peak at 102.62\(^\circ\)C which is accompanied by -5.08% weight losses (see TGA curve fig. 9) is associated to the dehydration of humidity H\(_2\)O content. The second endothermic peak at 440\(^\circ\)C (see DTA curve fig. 8) is assigned to oxygen uptake (oxidation) and CO\(_2\) evolution during the formation of ultramarine.

The first exothermic peak at 197.02\(^\circ\)C which is accompanied by -5.08% weight loss (see DTA curve fig.8) is assigned to the first step in the
partial oxidation of charcoal and evolving CO/CO₂. The two stepwise exothermic peaks at 242.38°C and ≈ 360°C which is accompanied by 1.75% weight gain (see DTA curve Fig. 8) are assigned to the second step in the partial oxidation of charcoal and evolving of CO/CO₂. The weight loss caused by evolving of SO₂ is negligible. Thus, results of TGA and DTA are in full agreements for these blue ultramarine samples and support one another.

Figures: 9 and 10 display the ESR spectra of blue ultramarine (containing S and Na₂SO₃ fired at 750°C for 3 h) and deep blue ultramarine (containing S and Na₂SO₃ fired at 750°C for 10 h), respectively. Their calculated g values are found to be (g = 1.93) for the blue ultramarine and (g = 1.94) for the deep blue ultramarine as shown in Figs. (9,10). It was established that the presence of S₃⁻ radicals were confirmed by ESR and their ESR signal anisotropy can result from the extra cages S₃⁻ radicals. In our investigated ultramarine sample, ESR data indicated further anisotropy comparing with earlier investigators. This explains the more colour deepening in our deep blue ultramarine sample regarding our mode of synthesis and raw material included to modify the product for industry.
Applying the Arrhenius plot equation [23]

\[ \sigma = \sigma_0 e^{-\Delta E/KT} \]

Where \( \sigma \) is the electrical conductivity at temperature \( T \), \( \sigma_0 \) is the pre-exponential factor, \( K \) is the Boltzmann's constant and \( T \) is the absolute temperature. The intrinsic activation energy for electric conduction is calculated to be 0.041 eV in the temperature range 498 – 673 K. Thus the corresponding energy gap is evaluated to be 0.082 eV. The ultramarine sample (3) behaves as a semiconductor in the range 293 – 688 K, on the same proceeding rule, at the temperature range 298 – 473 K investigated; this ultramarine sample (3) behaves as extrinsic semiconductor. The 473 K is the transition temperature from extrinsic semiconductor to intrinsic semiconductor. In the range 473 – 673 K, there is intrinsic semiconduction. After applying Arrhenius plot, the activation energy for electric conduction is calculated to be 0.816 eV in the temperature range investigated. It is generally accepted that the presence of \( S_3^- \) is a prerequisite step for semiconduction in prepared samples and this formation of radical cause current flow. For ultramarine sample (5), the material is semiconductor in the temperature range 398 – 838 K. In the range 398 – 563 K this ultramarine sample (4) behaves as intrinsic semiconductor while in the temperature range 563 – 838 K behaves as extrinsic semiconductor. The 563 K being the transition temperature from extrinsic to intrinsic semiconductor. Applying the Arrhenius plot, the activation energy for electric conduction is calculated to be 0.044 eV in the temperature range 563 – 838 K. The induced metallic conduction mechanism in the extrinsic range (low temperature) is due to thermal decomposition. Our prepared ultramarine sample (6) is semiconductor in the range 313 – 753 K. In the range 313 – 463 K ultramarine sample (6) behaves as extrinsic semiconductor with a transition temperature of 463 K from extrinsic to intrinsic semiconductor. Applying the Arrhenius plot, the activation energy for electric conduction is calculated to be 0.816 eV in the temperature range 313 – 753 K. In the range 313 – 463 K ultramarine sample (6) behaves as intrinsic semiconductor while in the temperature range 463 – 753 K behaves as extrinsic semiconductor. The 463 K being the transition temperature from extrinsic to intrinsic semiconductor. Applying the Arrhenius plot, the activation energy for electric conduction is calculated to be 0.044 eV in the temperature range 463 – 753 K. For our prepared ultramarine violet sample (7), there are two conduction mechanism appear to operate as follows: The first being the metallic conduction mechanism in the extrinsic range (318 – 403 K) may due to the presence of \( \text{NH}_4 \) at low temperature and this greatly enhance some rearrangements of the electronic band structure of our prepared ultramarine violet sample. This inflection in the conductivity temperature (fig. 11) is interpreted on the basis of the occurrence of phase changes as detected from the DTA curve indicating its endothermisty accompanied with colour faddening. The second being the semiconducting mechanism in the temperature range 408 – 623 K, After applying Arrhenius plot, the activation energy for electric conduction is calculated to be 0.073 eV in the temperature range 453 – 623 K, thus the corresponding energy gap is calculated to be 0.146 eV.

From Table (3) it can easily be seen that the activation energy for intrinsic semiconduction and hence the corresponding \( E_g \) decrease in the direction from sodium thiosulfate \( \text{Na}_2\text{S}_2\text{O}_3 \)
containing ultramarine (0.041 eV). Thus the semiconductivity is modified by sulfur addition, then Na$_2$SO$_3$, NH$_4$Cl + S, Na$_2$SO$_3$ + S and further by Na$_2$S$_2$O$_3$.

![Figure-12: The variation of DC-electrical conductivity (log$\sigma$) versus reciprocal of absolute temperature (1000/T) K$^{-1}$ for ultramarine samples](image)

**Table-1: The effect of the sulfur precursor on the structure and the colour of the prepared ultramarines**

<table>
<thead>
<tr>
<th>Sample</th>
<th>$S^{3-}$ precursor</th>
<th>Colour</th>
<th>Calcination temperature</th>
<th>Time (hrs)</th>
<th>Structure after calcination</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ultramarine blue</td>
<td>S</td>
<td>Light blue</td>
<td>400°C</td>
<td>2</td>
<td>Unknown</td>
</tr>
<tr>
<td></td>
<td>S$_2$O$_3$</td>
<td>Light blue</td>
<td>750°C</td>
<td>4</td>
<td>Sodalite</td>
</tr>
<tr>
<td></td>
<td>S$_2$O$_3$</td>
<td>Blue</td>
<td>750°C</td>
<td>6</td>
<td>Sodalite</td>
</tr>
<tr>
<td></td>
<td>SO$_3$</td>
<td>Blue</td>
<td>750°C</td>
<td>4</td>
<td>Sodalite</td>
</tr>
<tr>
<td></td>
<td>SO$_3$</td>
<td>Blue</td>
<td>750°C</td>
<td>6</td>
<td>Sodalite</td>
</tr>
<tr>
<td></td>
<td>Na$_2$SO$_3$ + S</td>
<td>Blue</td>
<td>750°C</td>
<td>4</td>
<td>Sodalite</td>
</tr>
<tr>
<td></td>
<td>Na$_2$SO$_3$ + S</td>
<td>Deep blue</td>
<td>750°C</td>
<td>4 then 6</td>
<td>Sodalite</td>
</tr>
<tr>
<td></td>
<td>Na$_2$S</td>
<td>Gray</td>
<td>750°C</td>
<td>6</td>
<td>Sodalite</td>
</tr>
<tr>
<td>Ultramarine violet.</td>
<td>Na$_2$SO$_3$ + S + NH$_4$Cl</td>
<td>Violet</td>
<td>750°C</td>
<td>4</td>
<td>Sodalite</td>
</tr>
</tbody>
</table>
Table-2: Characterization and band assignment of the infrared absorption bands (cm⁻¹) of ultramarine samples (1,2,4,5,6)

<table>
<thead>
<tr>
<th>Ultramarine samples</th>
<th>Band assignment of the most characteristic absorption bands</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1) 3427 b</td>
<td>Asymmetric and symmetric stretching vibration of H₂O.</td>
</tr>
<tr>
<td>(2) 342 m</td>
<td>Bending vibration mode of H₂O.</td>
</tr>
<tr>
<td>(4) 3419 b</td>
<td>S₃⁻ &amp; S₂⁻ in a sodium alumi-n-silicate matrix.</td>
</tr>
<tr>
<td>(5) 3428 m</td>
<td>Associated with the Si-O stretching vibration.</td>
</tr>
<tr>
<td>(6) 3429 m</td>
<td>S₃⁻ &amp; S₂⁻ in a sodium alumi-n-silicate matrix.</td>
</tr>
<tr>
<td>(1) 1623 s</td>
<td>Silica spectrum Na-O stretching on vibration and also due to SiO stretching vibration.</td>
</tr>
<tr>
<td>(2) 1628 w</td>
<td></td>
</tr>
<tr>
<td>(4) 1127 s</td>
<td></td>
</tr>
<tr>
<td>(5) 1123 m</td>
<td></td>
</tr>
<tr>
<td>(6) 1123 m</td>
<td></td>
</tr>
<tr>
<td>(1) 1001 v,s</td>
<td></td>
</tr>
<tr>
<td>(2) 1025 w</td>
<td></td>
</tr>
<tr>
<td>(4) 1002 s</td>
<td></td>
</tr>
<tr>
<td>(5) 1026 w</td>
<td></td>
</tr>
<tr>
<td>(6) 623 m</td>
<td></td>
</tr>
<tr>
<td>(1) 534 w</td>
<td></td>
</tr>
<tr>
<td>(2) 624 m</td>
<td></td>
</tr>
<tr>
<td>(4) 624 v,b</td>
<td></td>
</tr>
<tr>
<td>(6) 624 v,s</td>
<td></td>
</tr>
</tbody>
</table>

Table-3: The transport data for the prepared ultramarine blue sample

<table>
<thead>
<tr>
<th>Ultramarine No.</th>
<th>Composition</th>
<th>Intrinsic Activation energy (ΔEo eV)</th>
<th>Energy gap (Eg eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1)</td>
<td>Kaolin + Na₂CO₃ + S + Charcoal</td>
<td>0.041</td>
<td>0.082</td>
</tr>
<tr>
<td>(2)</td>
<td>Kaolin + Na₂CO₃ + Na₂S₂O₃ + Charcoal</td>
<td>0.816</td>
<td>1.632</td>
</tr>
<tr>
<td>(4)</td>
<td>Kaolin + Na₂CO₃ + Na₂SO₃ + Charcoal</td>
<td>0.0435</td>
<td>0.087</td>
</tr>
<tr>
<td>(6)</td>
<td>Kaolin + Na₂CO₃ + S + Na₂SO₃ + Charcoal</td>
<td>0.099</td>
<td>0.198</td>
</tr>
<tr>
<td>(7)</td>
<td>Kaolin + Na₂CO₃ + Na₂SO₃ + S + NH₄Cl</td>
<td>0.073</td>
<td>0.146</td>
</tr>
</tbody>
</table>

References