Investigations of Feldspar-Quartz Raw Materials After Firing: Effect of Various Na$_2$O/K$_2$O Ratio and Synthetic Pigments Addition

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Abstract: Different techniques (chemical analysis, X-ray diffraction, Mössbauer spectroscopy, UV-Vis-NIR spectrophotometry and chromaticity measurements) were used to study the color variation of feldspathic raw materials after firing at 1200 °C. Two varieties of feldspar-quartz grits (with Na$_2$O or K$_2$O prevalence) doped with synthetic pigments (hematite and rutile) were heat-treated in an oxidizing atmosphere. The results obtained indicate that the color of the ceramic body after firing may not only depend on the presence and analytical content of coloring elements (such as iron and titanium), but on their valence state and the chemical characteristics of the melt itself. The prevalence of larger alkali K$^+$ ions over Na$^+$ ions may cause the reduction of the coordination number of iron. This has a significant effect on the length of the Fe–O bond, which in turn influences the capacity for light absorption of a specific wavelength. The results of spectrophotometric measurements showed pronounced disparity between the raw materials and samples doped with coloring oxides. The introduction of synthetic pigments caused clear changes in the spectral dependence of the total absorbance. The absorbance increased in the whole measuring range, in particular in the visible part of the spectrum. Furthermore, a red-shift of the absorption bands towards longer wavelengths attributed to oxygen-metal charge transfer (OMCT) interactions between Fe$^{3+}$ and O$^{2-}$ ions as well as to d-d transitions within Fe$^{3+}$ ions was observed.

Keywords: potassium- and sodium-rich feldspar-quartz; coloring oxides; Mössbauer spectroscopy; UV-Vis-NIR spectrophotometry; color after firing

1. Introduction

Feldspars are the most common raw materials used as fluxes in the ceramics industry. Compared to other components of the ceramic body (e.g., ceramic clays and quartz), the low melting point of feldspars means that they form a vitreous phase contributing to the sintering of a given mixture of constituents in the course of firing. In Poland, the basic source of feldspathic raw materials (feldspar-quartz) are deposits located in Lower Silesia (SW Poland), in the marginal zone of the Strzęgom-Sobótka leucogranite massif. These rocks developed as a result of metasomatic alteration of various types of granite in the contact zone of magma intrusion with surrounding rocks (gabbro, serpentine and amphibolite) of the Carboniferous age [1]. Metasomatic transformation of the rocks was followed by enrichment of their mineral composition in feldspar minerals (plagioclases and microcline-type perthite).

The major domestic manufacturer of feldspar-quartz commodities is the Strzeblowskie Mineral Mines (SKSM) of Sobótka, operating four deposits of granitoids located in the NW part of the outer
zone of the massif. The excavated raw material is characterized by considerable mineral and chemical differentiation and quite a high content of coloring oxides (0.2–0.7% Fe₂O₃ + TiO₂ [2]). To obtain stable and repeatable parameters of commercial product with fixed chemical composition tailored to customer requirements, the raw material is selectively extracted from each deposit. Afterwards, it is examined for chemical composition at consecutive stages of processing (crushing, sieving, and blending). Another crucial parameter that needs to be controlled is the color after firing, as the main customer of these commodities is the domestic ceramic tile industry [3].

The aim of the present study was to assess the influence of chemical composition of the raw material (especially alkanes and coloring oxides) on the color of the sintered body. In addition, this study sought to identify mineral phases that occur as a result of heat treatment of two commercial grades of feldspar-quartz grits from SKSM, with special attention to iron- and titanium-bearing ones, as they are the most common chromophores. The research was also focused on interaction between iron and titanium ions in the course of firing. Appropriate control of the chemical composition of the raw materials used in the ceramics industry has fundamental significance for properties and aesthetic qualities of the final products.

2. Materials and Methods

The raw material from deposits located in the Sobótka region consists of well-characterized mineral components [4,5]. According to previous studies, the main minerals containing coloring elements are: chlorites, micas (biotite, muscovite and sericite), rutile, garnets, feldspars, monazite, zircon, xenotime, epidote (Fe-rich pistacite), magnetite, pyrite and Fe-Ti spinel (titanomagnetite). The basic constituent of the raw materials after firing is the amorphous phase, resulting from thermal decomposition of feldspars and partly quartz, accompanied by single quartz grains and other minerals characterized by high melting temperatures, such as zircon, Fe-Ti spinel, xenotime and monazite [5,6].

The subject of the current research was feldspar-quartz commodities provided by the Strzeblowskie Mineral Mines of Sobótka. The samples represented two commercial grades of feldspathic grits of different moduli K₂O:Na₂O, i.e., with the prevalence of potassium or sodium particles in the chemical composition (K₂O:Na₂O = 1.8 or 0.6, respectively, Table 2).

From each type of the raw material there were four samples prepared: the starting/basic ones (marked 0, Figure 1), and three others doped with controlled quantities of Fe₂O₃—hematite—and TiO₂—rutile (designated as I, II and III), as indicated in Table 1. Thorough homogenization of the basic material with coloring oxides was obtained by dry milling in a planetary ball mill for around 40 min. The pigments used in the experiment were synthetic ferric red TP-303 (Nofar) of 99.2% Fe₂O₃ purity and titanium white Tytanpol ®R-001 of 99.1% TiO₂ purity.

Table 1. The percentages of synthetic pigments added to basic samples of feldspar-quartz raw materials (wt. %).

| Sample | Fe₂O₃ | TiO₂ |
|--------|-------|------|
| I      | 1.5   | -    |
| II     | -     | 0.5  |
| III    | 1.5   | 0.5  |

The chemical compositions of the studied materials were quantified by the wavelength dispersive X-ray fluorescence method (WDXRF) on a Bruker AXS (Karlsruhe, Germany) S8 TIGER spectrometer with a 3 kW Rh anode X-ray tube. The spectrometer is equipped with a Bruker’s Quant-Express software allowing for standardless analysis of elemental composition of the samples. Quant-Express comprises a customized calibration prepared by Bruker using the feldspar-quartz rock of interest and selected certified reference materials. The accuracy of the measurements was determined by the instrument supplier (Table 2). After grinding (d₉₀ = 53.3 μm) and mixing with a few drops of distilled...
water, the 20 g portions of the material were pressed into pellets of ca. 40.4 mm diameter and maximum 10 mm thickness in a stainless-steel matrix of a Herzog TP 20 P hydraulic press with a pressure of 100 kN. The pellets were fired for 51 min in an oxidizing atmosphere in an electric furnace (Nabertherm LS 25/13). The maximum temperature 1200 °C was reached in 22 min (starting from room temperature) with a ramp rate of 54.5 °C/min. After 6 min holding time at 1200 °C the samples were quickly cooled down to 95 °C at a rate of 48.04 °C/min (for 23 min) in a furnace with integrated fans, then removed to air-cool to room temperature. These conditions (temperature and time) are usually adopted in the current fast-firing technology of ceramic tile production. After cooling, the samples were examined by X-ray diffraction, Mössbauer spectroscopy, UV-Vis-NIR spectrophotometry and chromaticity analysis.

![Figure 1. Color after firing of the studied samples: potassium (K) and sodium (Na) types doped with synthetic pigments (marked I-III) plus that of the basic samples (marked 0).](image)

| Table 2. Analyzed weight % composition of the studied feldspar-quartz raw materials. |
|---------------------------------------------------------------|
| Sample | SiO₂ | Al₂O₃ | Fe₂O₃ | TiO₂ | Na₂O | K₂O | CaO | MgO | LoI |
| Max. deviation | ±2.0 | ±0.150 | ±0.010 | ±0.005 | ±1.0 | ±0.30 | ±0.30 | ±0.05 | n.d. |
| Potassium-rich samples | | | | | | | | | |
| 0 K | 75.90 | 13.93 | 0.30 | 0.053 | 2.87 | 5.14 | 0.20 | 0.16 | 1.19 |
| I K | 73.18 | 14.46 | 1.77 | 0.055 | 2.96 | 5.23 | 0.22 | 0.16 | 1.62 |
| II K | 73.56 | 14.79 | 0.34 | 0.566 | 3.05 | 5.33 | 0.22 | 0.17 | 1.59 |
| III K | 72.46 | 14.51 | 1.85 | 0.545 | 3.00 | 5.27 | 0.21 | 0.16 | 1.69 |
| Sodium-rich samples | | | | | | | | | |
| 0 Na | 73.78 | 15.20 | 0.36 | 0.096 | 5.48 | 3.49 | 0.38 | 0.13 | 0.98 |
| I Na | 71.90 | 15.23 | 1.77 | 0.042 | 5.38 | 3.56 | 0.38 | 0.14 | 1.24 |
| II Na | 72.57 | 15.40 | 0.29 | 0.537 | 5.48 | 3.58 | 0.38 | 0.13 | 1.23 |
| III Na | 71.80 | 14.97 | 1.75 | 0.536 | 5.31 | 3.50 | 0.38 | 0.13 | 1.27 |
| n.d.—not determined. | | | | | | | | | |

The mineral phases were identified by X-ray powder diffraction (XRD). The data were collected by a Rigaku MiniFlex 600 X-ray diffractometer (Neu-Isenburg, Tokyo, Japan) with the curved crystal graphite-monochromatized CuKα radiation, operating with a voltage of 40 kV and an emission current of 15 mA. XRD patterns were recorded in the angular range 3–75° 2Θ with a step size of 0.05°, counting time
1 s per step. The examinations were performed for feldspar-quartz samples of both types, before and after firing.

The Mössbauer spectra were collected using the RENON MsAa-3 spectrometer equipped with LND detector with Kr filter and laser interferometer He-Ne at room temperature [7]. A commercial $^{57}$Co (RH) isotope was used as a source of radiation. All spectra shifts were reported versus the total shift in $\alpha$-Fe. The obtained spectra were processed by the MOSGRAF-2009 software. There were four samples analyzed: two samples of sodium feldspar-quartz doped with one and two coloring oxides (I Na and III Na) and two corresponding samples of potassium feldspar-quartz (I K and III K).

The optical properties of selected samples after firing were analyzed based on the spectral dependence of the absorbance ($A$). The spectra were recorded in a broad wavelength range ($\lambda$) of 220 to 2200 nm with a Jasco V-670 UV-VIS-NIR double-beam spectrophotometer equipped with a dedicated 150-mm integrating sphere to perform the measurements of the total absorbance. The measured parameter was the reflectance (scale 0–100%), which was converted into the absorbance and analyzed. Such a calculation was feasible because the samples thickness was incomparably greater than the wavelength of the light used (10 mm vs. 220–2200 nm). The integrating sphere was coated with barium sulfate and the spectra were referenced against white standard—Spectralon. The applied speed was 200 nm/min with a step of 0.5 nm.

A portable Konica Minolta CM-2300d spectrophotometer was used to determine the chromatic coordinates of fired samples. The Konica Minolta CM-2300d instrument of spherical type with horizontal alignment, is suitable for complex color analysis in the visible wavelength range (400–700 nm). Color coordinates $L^*$, $a^*$ and $b^*$ values of the specimens were obtained in CIE 1976 (Commission Internationale de l’Eclairage) space, where $L^*$ refers to darkness-brightness (0–100), $a^*$ to the red ($a^* > 0$) or green ($a^* < 0$), and $b^*$ to yellow ($b^* > 0$) or blue ($b^* < 0$).

3. Results

3.1. Chemical Composition

Table 2 lists the chemical composition of the studied samples. The difference in $K_2O$, $Na_2O$ and $Al_2O_3$ content between samples of sodium and potassium influences the amount of amorphous phase during firing. Furthermore, the basic potassium sample (0 K) contains slightly less $Fe_2O_3$ and $TiO_2$, which are the main color contributors, as compared to its sodium counterpart (0 Na).

3.2. The XRD Analyses

The X-ray diffraction analyses of the studied samples only provided information on the main mineral phases because many of the peaks corresponding to minor phases coincided with major ones. The predominant crystalline phases of the raw (unfired) sample of sodium feldspar-quartz (marked Na raw) were: albite ($d$—4.02, 3.78, 3.67, 3.19 and 2.93 Å) and $\beta$-quartz ($d$—4.29, 3.35, 1.82 and 1.54 Å), as well as—albeit less numerous—microcline ($d$—4.26, 3.84, 3.48, 3.28 and 3.24 Å) (Figure 2).

In addition, some faint reflections of illite ($d$—10.0, 4.46, 3.36 and 2.86 Å) and kaolinite ($d$—7.05, 4.48, 3.58, 3.85 and 2.55 Å) were also registered. The diffraction pattern of the sample after firing (0 Na) evidenced the presence of a characteristic amorphous hump within the $2\Theta$ 15–40$^\circ$ range associated with the existence of the glassy phase, as well as remnants of quartz and albite that were not completely melted. Some small peaks indicating the secondary mullite (probably) precipitation were also registered ($d$—5.37, 3.43, 3.37 and 1.46 Å). Sodium-rich variety samples doped with synthetic pigments (I Na and III Na) gave weak hematite peaks ($d$—2.69, 2.51, 1.68 and 1.46 Å) and a much larger hump of an amorphous phase, which arose mainly at the expense of albite. The thermal decomposition of albite may be attributed to a high iron oxide content. It is well known that in silicate melts iron may occur in both oxidation states, as $Fe^{2+}$ and $Fe^{3+}$ ions [8,9], of which $Fe^{2+}$ acts as a network modifier (flux) and thereby lowers the melting point and viscosity [10,11].
According to the XRD analysis of the unfired sample of the potassium variety raw material (K raw) its main mineral phases are: β-quartz (d—4.26, 3.34, 2.46, 1.82 and 1.54 Å) and microcline (d—4.20, 3.83, 3.30, 3.24, 3.02 and 2.96 Å) (Figure 3). Some smaller reflections of albite (d—4.03, 3.83, 3.77, 3.66, 3.19, 2.95 and 2.56 Å), accompanied by illite (d—10.05, 4.48, 3.35, 2.86 and 2.57 Å) and kaolinite (d—7.20, 4.48, 3.85, 3.58 and 2.56 Å) were also registered. The X-ray pattern of the sample after firing (0 K) shows some amorphous phase formation (but in much smaller amounts than with the sodium variety sample) and the disappearance of the majority of peaks (especially of illite, kaolinite and albite), except for reflections of quartz and some microcline, the intensity of which weakened markedly. The relatively smaller amorphous phase hump may be a result of the higher melting temperature of potassium feldspar and the smaller alkali content in comparison with sodium samples (8.01% and 8.97% K₂O + Na₂O, respectively). Pure albite Na[AlSi₃O₈] melts congruently at 1118 ± 3 °C, while potassium feldspar K[AlSi₃O₈] melts incongruently at 1150 ± 20 °C, forming the liquid phase containing leucite and silica [12,13]. The latter is characterized by much higher viscosity than the liquid
phase arising in the course of heat treatment of sodium feldspars [14]. Thus, at high temperatures, the glassy phase forms slower and in smaller quantities.

Figure 3. XRD patterns of potassium feldspar-quartz samples in a raw state/unfired (K raw) and the samples after firing: without synthetic pigments addition (0 K), as well as doped with 1.5% Fe$_2$O$_3$ (I K), and with 1.5% Fe$_2$O$_3$ + 0.5% TiO$_2$ (III K).

The addition of coloring oxides (I K and III K samples) resulted in the decay of microcline reflections and the appearance of small peaks of mullite (d—5.37, 3.40, 3.39 and 2.29 Å) and of hematite (d—2.69, 2.51, 2.20 and 1.68 Å, albeit of lower intensity than in the X-ray pattern of corresponding sodium samples), as well as raised hump of amorphous phase. The reasons for these differences were similar to the sodium-type material (divalent iron acting as a flux); however, the amount of amorphous phase was notably lower. The peaks of mullite are weaker in the X-ray diffraction pattern of the potassium samples after firing due to the fact that their thermal decomposition takes place at higher temperatures than alkaline feldspars rich in Na$_2$O. When it comes to sodium feldspars, they turn into the liquid phase faster and secondary mullite crystalizes earlier and more easily. Melting at lower temperature resulted in numerous bubbles that appeared on the surface of the I Na sample (Figure 4), while the surface of the I K pellet was almost free of such irregularities. The peaks of rutile or other titanium phases were not identified. This was probably due to their concentration below the detection limit of the XRD method.
Figure 4. Sodium feldspar-quartz sample doped with 1.5% Fe$_2$O$_3$ (I Na) after firing, whose surface is covered with numerous bubbles.

3.3. Mössbauer Spectroscopy

The Mössbauer spectra of both feldspar-quartz varieties doped with synthetic pigments were fitted with three components: one sextet of Fe$_2$O$_3$ and two doublets of Fe$^{3+}$ and of Fe$^{3+/2+}$, respectively. The sextet of well crystalized hematite with characteristic hyperfine parameters, i.e., isomer shift IS (0.37 mm/s), quadrupole splitting QS (−0.21 and −0.23 mm/s) and magnetic field B (around 51 T), accounts for around 2/3 (60–68%) of the spectra area (Table 3, Figures 5 and 6). Undeniably, this should be linked to the added synthetic hematite, and possibly to iron ions released from the lattice of other minerals at elevated temperatures that form fine iron oxide particles dispersed in the aluminosilicate matrix [15–18].

Table 3. The results of Mössbauer spectroscopy studies.

| Sample | Phase | A (%) | IS (mm/s) | QS (mm/s) | B (T) | Г (mm/s) |
|--------|-------|-------|-----------|-----------|-------|----------|
| I Na   | Fe$_2$O$_3$ | 65 ± 2 | 0.37 | −0.21 | 51.1 | 0.28 |
|        | Fe$^{3+/2+}$ | 14 ± 1 | 0.54 | 2.57 | — | 0.90 |
|        | Fe$^{3+}$ | 21 ± 1 | 0.34 | 0.98 | — | 0.90 |
| III Na | Fe$_2$O$_3$ | 68 ± 2 | 0.37 | −0.23 | 50.7 | 0.33 |
|        | Fe$^{3+/2+}$ | 13 ± 1 | 0.58 | 2.53 | — | 0.90 |
|        | Fe$^{3+}$ | 19 ± 1 | 0.34 | 0.91 | — | 0.90 |
| I K    | Fe$_2$O$_3$ | 66 ± 2 | 0.37 | −0.21 | 51.1 | 0.26 |
|        | Fe$^{3+/2+}$ | 13 ± 2 | 0.63 | 2.43 | — | 0.93 |
|        | Fe$^{3+}$ | 21 ± 2 | 0.34 | 0.82 | — | 0.82 |
| III K  | Fe$_2$O$_3$ | 60 ± 2 | 0.37 | −0.23 | 50.7 | 0.33 |
|        | Fe$^{3+/2+}$ | 16 ± 2 | 0.58 | 2.39 | — | 0.74 |
|        | Fe$^{3+}$ | 24 ± 2 | 0.33 | 0.81 | — | 0.90 |

A—relative contribution of the given iron site (sub-spectrum), IS—isomer (total) shift versus α-Fe, QS—quadrupole splitting, B—magnetic hyperfine field, Г—absorber line-width.

The isomer shift of doublets of Fe$^{3+}$ ions, i.e., IS = 0.33–0.34 (Table 3), is typical for iron oxide compounds [17,18]. However, according to some references, the iron phases present in the raw material can decompose in the course of firing and then iron can be partly or completely diluted in the liquid aluminosilicate phase (at the atomic level) with no separated iron oxide phase [15,18,19] or may be incorporated into the lattice of mullite, substituting for Al$^{3+}$ [20].
Figure 5. Room temperature Mössbauer spectra of the sodium-rich samples heat-treated at 1200 °C. (a) I Na doped with Fe$_2$O$_3$, (b) III Na doped with Fe$_2$O$_3$ + TiO$_2$.

Other components of the analyzed Mössbauer spectra, i.e., doublets of Fe$^{3+/2+}$ are characterized by a quadrupole splitting of about 2.39–2.57 mm/s typical for Fe$^{2+}$ (around 2.5 mm/s) and by an isomer shift of about 0.54–0.63 mm/s assigned to Fe$^{3+}$ in octahedral site (CN = 6). The values of IS are generally in accordance with data in the literature, i.e., 0.35–0.55 mm/s [21] or 0.28–0.50 mm/s [17]. Their relative content ranged from 13% to 16% (Table 3). The presence of Fe$^{3+/2+}$ may be a result of the reduction of some Fe$^{3+}$ ions to Fe$^{2+}$ due to their substitution by Ti$^{4+}$, following the charge compensation rule [22], as well as of the equilibrium of the Fe$^{3+}$/Fe$^{2+}$ with the atmospheric partial pressure of oxygen ($p$O$_2$) [23]. The quadrupole splitting value, which gives the information about local symmetry disorders in the crystal lattice, is much higher for Fe$^{3+/2+}$ (2.39–2.57 mm/s) than that for Fe$^{3+}$ (0.81–0.98 mm/s). This can be linked to possible interactions between Ti and Fe cations (the higher QS, the lower symmetry of the electric charge distribution). Another possible explanation of Fe$^{3+/2+}$ presence is in line with the observations of Bingham et al. [24], according to which the clusters of Fe$^{2+}$–Fe$^{3+}$ cations can occur in the melts rich in Fe$_2$O$_3$. Clustering may include Fe$^{2+}$–O–Fe$^{3+}$ and Fe$^{3+}$–O–Fe$^{3+}$ interactions.

The contributions (percentages) of the $^{57}$Fe Mössbauer spectral components described above were designated as the area contributions of the corresponding sub-spectra to the whole cross-section of the spectrum. It should be kept in mind that area contributions of the Mössbauer signals discussed here do not necessarily reflect the actual concentrations of the different iron states. Different bond properties of
the iron atoms in their lattice positions lead to different Mössbauer-Lamb factors (MLF), which give rise to different intensities of the Mössbauer signals. In this study, the area contributions of the Mössbauer spectral components are shown and discussed without correction for the possible different values of the MLF, but in fact the MLF for considered materials should have quite similar values and the effects of differential MLF could lead to errors of about 15% for the Fe$^{3+}$/Fe$^{2+}$ ratio [25,26].

Figure 6. Room temperature Mössbauer spectra of the potassium-rich samples heat-treated at 1200 °C. (a) I K doped with Fe$\text{O}_3$, (b) III K doped with Fe$\text{O}_3$ + TiO$_2$.

The relative shares of various forms of iron in the spectra of I K as compared to III K sample, differ from each other: hematite in I K sample is 66% while in III K it is 60% (Table 3). This appears to be related to the presence of TiO$_2$ and possibly to the potassium nature of the aluminosilicate melt (it is worth noting that the spectra of I Na and III Na samples are almost identical). However, based on the results of Mössbauer spectroscopy measurements, neither the titanium role nor titanium and iron interaction in the case of samples marked III of both varieties of raw materials, can be elucidated.

3.4. UV-Vis-NIR Absorbance Spectra

Spectral dependence of the total absorbance in the wavelength range from 220 to 2200 nm measured for selected feldspar-quartz samples after firing is presented in Figure 7. The spectral line shapes of both examined raw material varieties (0 Na, 0 K) are typical for these types of feldspar [27,28]. The presence of a small amount of Fe$\text{O}_3$ in the raw materials (0.30–0.36%) gives a clear manifestation in their optical
spectra. In the UV region, a distinct absorption band centered at 262 nm (marked #1) can be observed. It may be assigned to ligand-to-metal charge transfer transitions taking place between $O^{2-}$ and $Fe^{3+}$ ions (OMCT) probably in an octahedral FeO$_6$ cluster [20,29]. The visible region of the absorbance spectra shows a series of small bands that can be ascribed to d-d transitions within $Fe^{3+}$ ions. They appear at: 384 (#2), 413 (#3), 448 (#4), 487 (#5), 545 (#6), and 642 nm (#7). These bands may be attributable to $Fe^{3+}$ ions in tetrahedral (#2–#4) and octahedral (#5, #6) coordination [30,31]. Some authors [20] ascribe the absorption band at around 448 nm (#4) to d-d transitions of octahedrally coordinated $Fe^{3+}$. It is worth noting that the aforementioned bands are present in both the potassium and sodium feldspar variety spectrum; however, they are more pronounced in the latter one (0 Na). A weak absorption band at around 860 nm (#5) can be assigned to hematite $\alpha$-Fe$_2$O$_3$ [20,29]. Additionally, between 1000 and 2150 nm a broad band (#9) of weak absorption is observed. Such a band in the near IR may be assigned to the presence of $Fe^{2+}$ ions [20]. The bandwidth can be, according to the literature [32–34], reasonably explained. First and foremost, two bands may overlap, one at around 1100 nm indicating the presence of $Fe^{2+}$ ions in an octahedral sites and another at around 2000 nm corresponding to tetrahedrally coordinated $Fe^{2+}$ ions. The reason for overlapping is probably Jahn-Teller distortion of the octahedrally coordinated iron ions. An additional two bands can be distinguished in the absorption spectra of the examined samples, the first centered at 1420 nm (#10) and the second at 1915 nm (#11). They are much more pronounced in the spectrum of K-type feldspar. According to the literature, these bands may originate from the combination of vibration modes of molecular water occurring in alkali feldspars and quartz in the form of microscopic or sub-microscopic fluid inclusions [34,35]. However, to verify this assumption, the additional investigations would be required. Disparity in absorbance spectra of sodium and potassium feldspar may be attributed to the difference in their chemical properties. The chemical surrounding is affected by atomic properties, especially by ionic radius ($K^+$—1.38 Å, Na$^+$—1.02 Å) and electro-negativity (Pauling scale: $K^+$—0.82, Na$^+$—0.93).

The shape of the absorbance spectra registered for feldspar-quartz samples doped with Fe$_2$O$_3$ synthetic pigment (I Na, I K) is generally similar to these of the basic samples (0 Na, 0 K). Some differences, however, are manifested. First of all, strong absorption band #1 in the UV region is broadened and shifted toward longer wavelengths, up to 390 nm in the visible region. An observed feature can be directly correlated with increased content of $Fe^{3+}$ ions [20]. The red-shift of bands may indicate ligand-to-metal charge transfer transitions occurring between $O^{2-}$ and $Fe^{3+}$ ions as well as the successful incorporation of some $Fe^{3+}$ ions into the lattice of other phases, e.g., mullite as isomorphic substitution for Al$^{3+}$ [20]. The latter assignment seems plausible because the presence of crystalline mullite was evidenced by XRD analysis. As a result, the observed strong absorption may, in fact, be a composite of the mentioned phenomena. Moreover, the intense absorption appeared at around 550 nm. This may be attributed to the simultaneous excitation (magnetic interaction) of adjacent $Fe^{3+}$ cations in the hematite lattice [29]. Above 570 nm the absorbance is much higher than that of undoped materials. The band at 860 nm (#8), assigned to $\alpha$-Fe$_2$O$_3$ (hematite), is significantly stronger than that registered for the raw materials (0 Na, 0 K). Furthermore, a slight shift of the absorption bands arising from $Fe^{3+}$ d-d transitions towards longer wavelengths is also observed. It can be noted that the bands of I K and I Na samples are quite similar and, except for obvious differences in absorbance intensity, resemble the spectrum of undoped sodium sample. Changes in the electronic structure due to increased amount of iron (Pauling electro-negativity of iron is 1.83) eliminate differences caused by distinct chemical properties of sodium and potassium [27,36].

Simultaneous addition of Fe$_2$O$_3$ and TiO$_2$ (III Na, III K) results in further broadening of the strong absorption range (with intensity exceeding 82%) between 220 and 2200 nm. Even though $Fe^{3+}$ bands are less pronounced, their further red-shift in the visible region may be observed. The absorption band at 860 nm (well-defined in the spectra of I K and I Na samples) is much weaker and hardly detectable. Such an effect in combination with the presence of $Fe^{3+}$-related bands could result from incorporation of $Fe^{3+}$ ions into the lattice of other phases, including TiO$_2$ and mullite. It should also be noted that
the broad spectral band in the near IR assigned to Fe$^{2+}$ ions is more distinct than that observed in other spectra.

Figure 7. Optical absorbance spectra of selected samples after firing: undoped (0 K, 0 Na), doped with Fe$_2$O$_3$ (I K, I Na), and doped with both Fe$_2$O$_3$ and TiO$_2$ (III K, III Na).

3.5. Chromaticity Analysis

To quantify the color of the studied samples and to examine its variation with the iron and titanium content, the CIE (1976) colorimetric system was used (Table 4 and Figure 8). After firing of the potassium-type raw material, the color is brighter compared to that of the sodium type (which is confirmed by the L* coordinate values of 87.71% and 75.86%, respectively). This results from the difference in light absorption in the visible region between both varieties of the studied material, especially in the range of ca. 400–550 nm (Figure 7). The absorption registered for the sodium sample (0 Na) is more intense than that of its potassium counterpart (0 K). The lower the absorption, the more light is reflected from the surface of the sample and the brighter color is perceived. As a consequence, the observed color of 0 Na sample is pale pink, with the corresponding a* value (6.05) much higher than that of 0 K sample (1.67), which is light grey in color (Figure 1, Table 4).
Table 4. Color parameters of the studied samples of different modulus $K_2O:Na_2O$ after firing.

| Sample  | Color Parameters of the Samples After Firing |
|---------|---------------------------------------------|
| L*(%)   | a*  | b*  | C*    | h*  | ΔE  |
|---------|-----|-----|-------|-----|-----|
| Potassium-rich variety |
| 0 K     | 87.71 | 1.67  | 9.25  | 9.40 | 79.43 |
| I K     | 37.76 | 16.63 | 10.01 | 19.41 | 47.13 |
| II K    | 84.56 | 2.01  | 15.04 | 15.17 | 82.39 | 5.66 |
| III K   | 37.61 | 5.25  | 4.92  | 7.19  | 43.16 | 44.96 |
| Sodium-rich variety |
| 0 Na    | 75.86 | 6.05  | 9.68  | 11.42 | 57.99 |
| I Na    | 36.59 | 15.91 | 9.34  | 18.45 | 30.41 | 46.14 |
| II Na   | 79.45 | 2.80  | 11.78 | 12.11 | 76.64 | 3.88 |
| III Na  | 35.74 | 3.71  | 2.56  | 4.51  | 34.63 | 44.97 |

L*—brightness; a*—red ($a > 0$); b*—yellow ($b > 0$); $C^* = (a^2 + b^2)^{1/2}$—chroma/saturation of color; $h^* = \arctan(b/a)$—hue; $\Delta E = \sqrt{\Delta L^2 + \Delta a^2 + \Delta b^2}$—total difference of color in relation to basic sample.

Figure 8. Color parameters of the studied samples in CIE space: K—potassium feldspar-quartz samples, Na—sodium feldspar-quartz samples, 0—basic samples, I—samples doped with 1.5% Fe$_2$O$_3$, II—samples doped with 0.5% TiO$_2$, III—samples doped with 1.5% Fe$_2$O$_3$ + 0.5% TiO$_2$.

Mixing with Fe$_2$O$_3$ resulted in decreased brightness L* (to 37.76% and 36.59%, respectively) and hue (h*) of both raw material varieties (Table 4), the change in their coloration was influenced by the hematite added as well as by the breakdown of iron-containing minerals at elevated temperatures with the formation of particles of hematite and Fe$^{3+}$ ions dispersed in the aluminosilicate phase [37–40]. This resulted in strong absorption in the visible region up to 570 nm accompanied by a tint change to dark brown (probably due to d-d electronic transitions in Fe$^{3+}$ ions coupled with Fe$^{3+}$–Fe$^{3+}$ pair excitation in hematite).

Mixing the studied materials with two synthetic oxides (Fe$_2$O$_3$ and TiO$_2$) caused further significant darkening of their color in relation to samples marked I (doped with Fe$_2$O$_3$ only). This was manifested by reduction of brightness L* (to 37.61% and 35.74%, respectively) that resulted from further broadening of the strong absorption range from the visible into the IR part of the spectrum. The addition of TiO$_2$...
caused a distinct change in color of the samples to almost black probably owing to the incorporation of iron ions to the structure of the synthetic rutile and the following charge transfer between Fe and Ti.

4. Discussion

The colors of the fired samples of both raw material varieties doped with two synthetic pigments turned to darker hues than those with the addition of Fe₂O₃ only. This points to an influence of TiO₂. Commercial titanium white is considered to be one of the best white pigments, since pure TiO₂ does not absorb visible light. However, it has been reported that when some iron is present in the structure of TiO₂, a charge transfer between Fe²⁺ and Ti⁴⁺ ions may occur [41–43]. This can contribute to the optical absorption in the UV and visible range resulting in the change in the coloration of examined samples [31,42–44]. Moreover, there are many colors that are caused by the much weaker electronic transitions between d (or f) orbital energy levels of single transition metal ions, such as Fe [43,45]. These transitions may be strongly intensified by magnetic coupling of adjacent Fe³⁺ cations in the crystal structure of iron phases [29].

Additionally, the presence of iron ions as impurity in raw materials can also contribute to the optical absorption in the UV and visible range due to O–Fe charge transfer (OMCT) [31,42,44]. There is evidence indicating that increasing the iron content causes an increase in the intensity of OMCT bands in the ultraviolet, hence shifting the UV edge towards smaller energies in the visible region [23,24]. This feature was clearly visible especially in the optical spectra of the examined samples doped with synthetic hematite.

On the other hand, the investigations performed by Shi et al. [46] demonstrated that a high content of Fe₂O₃ and TiO₂ in the raw material encourages the formation of a metastable pseudobrookite phase (Fe³⁺,Fe²⁺)₂(Ti,Fe³⁺)O₅ coexisting with rutile in the course of the firing. This is in accordance with a ternary phase diagram TiO₂, FeO, Fe₂O₃ [47]. In theory, pseudobrookite should disintegrate during cooling, but in the course of short thermal treatment with relatively low kinetics of this transformation, some remnants of the mentioned phase may remain in the fired material [46,48]. Pseudobrookite can form a solid-state solution with ferropseudobrookite (Mg²⁺,Fe²⁺,Al³⁺,Fe³⁺)Ti⁴⁺₂O₅ [49,50]. The resulting coexistence of Fe²⁺ and Fe³⁺ in the same structure, may presumably lead to the increase in optical absorption and thus to the change in color. However, these phases have not been detected in the examined samples by XRD analysis, and therefore this hypothesis cannot be confirmed.

Numerous researchers have found that Fe can substitute Ti in the high-temperature melt due to the close ionic radii and valence states of these ions [46,51,52]. According to Kim et al. [53], Ti⁴⁺ enhances the stability of Fe³⁺ relative to Fe²⁺ in the glassy phase. This is in accordance with the previous work on the redox equilibria of Fe- and Ti-bearing silica glasses, showing the structural instability of the Fe²⁺–O–Ti⁴⁺ network [54].

Other reasons for color differences between raw materials of the same or similar coloring oxide contents but of distinct chemical character (sodium or potassium), may be their different behavior during the firing process. The raw material with a higher sodium content forms the liquid phase at lower temperatures than its potassium counterpart. This favors faster phase-to-phase interactions and transition from the crystalline to the liquid state. This may be coupled with releasing iron ions from the lattice of other minerals, which form hematite particles that cause permanent coloring of the material after firing [18]. At high temperatures, some Fe³⁺ ions may be accepted by structures of newly formed mullite at the isomorphic position. Fe³⁺ ions can substitute Al³⁺ ions due to the similar ionic radii of these elements, i.e., Fe³⁺ = 0.55 or 0.59 Å depending on CN, and Al³⁺ = 0.54 Å [18,55].

Furthermore, many investigations have demonstrated a different influence of alkalis, such as K and Na, on iron coordination in various types of silica glasses [23,41–44,46–58]. It has been stated that the presence of larger alkali metal cations (e.g., K⁺) facilitates lower metal-oxygen coordination numbers for different transition metal ions, such as Fe³⁺ (ionic radius of K⁺ is 1.38 Å, while the ionic radius of Na⁺ is 1.02 Å). The valence and coordination of the iron ion have a major effect on the Fe–O bond strength (interionic distance). It was found that replacing K₂O by Na₂O in the silica glasses
containing Fe$^{3+}$ ions resulted in the change of average iron coordination from 4 to 5, while the reversed action caused a marked reduction of CN and the accompanying Fe–O distance. Li et al. [59] revealed that alterations in bond strength between the coloring ions (Fe$^{3+}$) and the surrounding ligands (O$^{2-}$) can cause various ligand field splitting energies of Fe-3d levels (for iron present as Fe$^{3+}$ with distorted octahedral geometry) and therefore contribute to different light absorption in the visible light spectrum. Color difference between the fired bodies containing various alkalis (with K$_2$O or Na$_2$O prevalence) is clearly visible in the absorption spectra of the studied samples, especially 0 Na and 0 K (undoped with synthetic pigments).

5. Conclusions

The investigations presented here showed that the color of the ceramic body after firing may depend not only on the presence and analytical content of coloring elements (such as iron and titanium), but on their valence state and coordination that induce specific optical transitions as well as on the chemical characteristics of the melt itself (e.g., the prevalence of potassium or sodium ions). This was confirmed using spectroscopic methods and chromaticity measurements.

According to Mössbauer and UV-Vis-NIR analyses, iron is present in the studied samples as hematite α-Fe$_2$O$_3$, Fe$^{3+}$, Fe$^{2+}$ and Fe$^{3+}$/Fe$^{2+}$ ions (detected by Mössbauer method as Fe cations in oxidation states intermediate between Fe$^{2+}$ and Fe$^{3+}$). Analysis of optical properties of the raw materials reveal absorption bands resulting from ligand-to-metal charge transfer transitions between oxygen (O$^{2-}$) and iron (Fe$^{3+}$) (OMCT) which occur in the UV and stretch into the visible region, especially after doping the examined samples with synthetic pigments. In the visible domain the spectra show a series of absorption bands resulted from d-d electronic transitions within Fe$^{3+}$ ions occupying tetrahedral and/or octahedral coordination. The absorption bands in the near IR region confirm the presence of Fe$_2$O$_3$ (hematite) as well as octahedrally coordinated Fe$^{2+}$ ions. Doping with Fe$_2$O$_3$ pigment resulted in widening of absorption range in the visible region associated with increased content of Fe$^{3+}$ ions as well as in appearance of the strong and well-defined bands assigned to hematite. These optical features indicate that the majority of synthetic pigment remained unaffected in the course of heat treatment. Simultaneous incorporation of Fe$_2$O$_3$ and TiO$_2$ leads to subsequent broadening of the strong absorption range in the visible and near-IR regions. Weak absorption band attributed to α-Fe$_2$O$_3$ in combination with confirmed presence of Fe$^{3+}$ ions may suggest successful doping of iron into TiO$_2$.

The prevalence of larger alkali cations in the melt (K exceeding Na), is likely to cause a reduction of the coordination number of iron, which has a fundamental effect on the Fe–O bond length. This in turn influences the capacity for light absorption in the visible region. In the case of the sodium-rich variety (0 Na), the absorption (resulting from d-d transitions within Fe$^{3+}$ ions) is much stronger than that recorded for its potassium counterpart (0 K). The addition of synthetic hematite caused an increase in the intensity of OMCT bands registered in the ultraviolet (due to O–Fe charge transfer transitions), hence shifting the UV edge towards the visible region. Furthermore, with increasing Fe concentration, more of the iron ions may occur in adjacent sites, and this probably manifested itself as strong absorption in the visible domain, owing to possible transitions resulting from simultaneous excitation of magnetically coupled Fe$^{3+}$ cations in the hematite lattice. Furthermore, a slight shift of the absorption bands arising from Fe$^{3+}$ d-d transitions towards longer wavelengths was observed. As a consequence, the perceived color of both feldspar-quartz varieties changed into dark brown. Doping with both synthetic pigments (Fe$_2$O$_3$ and TiO$_2$) caused intense absorption (absorbance > 82%) in the whole range of visible light resulting in further darkening of the examined materials which color turned almost black. This could result from charge transfer between Fe and Ti, which has an absorption centered in the near-UV region tailing into the visible and the blue end of the spectrum. However, more complex relationships between Fe and other structural components are possible.

It can be concluded that skillful control of the chemical composition of feldspar-quartz raw materials and precise determination of the redox state of iron can help to obtain ceramic bodies of a required color, notwithstanding the content of coloring oxides in the raw material used.
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