

Ceramics International 32 (2006) 393-405



www.elsevier.com/locate/ceramint

The role of counterions (Mo, Nb, Sb, W) in Cr-, Mn-, Ni- and V-doped rutile ceramic pigments Part 2. Colour and technological properties

M. Dondi a, G. Cruciani b, G. Guarini A, F. Matteucci a,*, M. Raimondo a

^a Institute of Science and Technology for Ceramics (ISTEC), Via Granarolo 64, 48018 Faenza, Italy ^b Earth Science Department, University of Ferrara, Corso Ercole 1 d'Este 32, 44100 Ferrara, Italy

Received 17 January 2005; received in revised form 18 February 2005; accepted 20 March 2005 Available online 18 August 2005

Abstract

Industrial rutile pigments are manufactured using several chromophores: Cr (giving an orange hue), Mr (tan), Ni (yellow) and V (gray); a second element, the so-called *counterion* (i.e. Mo, Sb, Nb or W) is always added in order to achieve the desired coloration and/or improve the technological properties (e.g. chemico-physical resistance in ceramic bodies and glazes). The colour of these pigments is determined by both metal-ligand charge transfer ($Ti^{4+} \leftrightarrow O^{2-}$) and crystal field effects (transition metals substituting Ti^{4+} in octahedral coordination). Though the absorbance bands are broad and frequently overlapped, the UV-vis-NIR spectra suggest the occurrence of Cr^{3+} , Mn^{2+} , Mn^{3+} , Ni^{2+} , V^{3+} , and V^{4+} as chromophores. Rutile pigments are suitable for through-body (up to $1250~^{\circ}C$) and glaze applications (up to $1100~^{\circ}C$). The best coloration of porcelain stoneware bodies is achieved with Sb or W as counterions, though the higher stability is ensured by Sb, but in the Ti-Ni-W system. The best glaze colours are accomplished by W-bearing pigments, which however are less stable than Nb- or Sb-containing ones, except than for the V+W coupling. This latter represents a new and very interesting Co-free and Cr-free black pigment for low temperature applications.

© 2005 Elsevier Ltd and Techna Group S.r.l. All rights reserved.

Keywords: B. Spectroscopy; C. Colour; C. Optical properties; D. TiO₂; D. Traditional ceramics; Ceramic pigments

1. Introduction

Several rutile-based ceramic pigments are industrially manufactured using different chromophore elements: chromium (to get orange hues), manganese (brown), nickel (yellow), and vanadium (gray). A second element with high field strength (the so-called *counterion*, i.e. Sb, Nb or W) is always added in order to fulfil the electroneutrality of the structure, so achieving the desired coloration and/or improving the technological properties of pigments (e.g. chemico-physical resistance in contact at high temperature with the liquid phase present in ceramic bodies and glazes) [1,2]. In some cases, even molybdenum is utilised as counterion, though not considered in the DCMA list [2–4].

Despite their intense and brilliant colours, rutile pigments have been progressively confined to low temperature ceramic applications, because of their partial solubility into the glazes over 1000 °C. This drawback can be minimised using special, Ti-rich glaze formulations, that unfortunately exhibit both technological (very high thermal expansion coefficient) and aesthetic limitations (high refractive index, strong brilliance) [5,6]. However, a renewed industrial interest has been growing in the last decade, due to the large request of pigments for through-body colouring of porcelain stoneware tiles, whose requirements are fulfilled by rutile pigments [2,7].

The colouring mechanisms allowing to get the rather wide chromatic palette offered by rutile pigments were not thoroughly investigated, being generally attributed to crystal field transitions [8–14] with some role played by the Ti–O metal–ligand charge transfer (MLCT) [4,10] or the band gap [4].

^{*} Corresponding author. Tel.: +39 0546 699738; fax: +39 0546 46381. E-mail address: matteucci@istec.cnr.it (F. Matteucci).

The main goal of this study is to understand in depth the mechanism responsible for pigment coloration, trying particularly to highlight the relative influence of different counterions, along with the valence of chromophore elements, the change in rutile crystal structure and the occurrence of accessory phases. Secondarily, this paper is aimed at assessing the ability of counterions to modify the technological behaviour of rutile pigments in both ceramic glazes and through-body applications.

This second part follows the former study of the effect of counterions on the rutile crystal lattice and the phase transformations occurring during the pigment synthesis [15].

2. Materials and methods

Sixteen compositions of ceramic pigments were prepared at the laboratory scale with stoichiometry $Ti_{1-2x}A_xB_xO_2$, where A = Cr, Mn, Ni or V; B = Mo, Nb, Sb or W; x = 0.03, plus a reference sample with x = 0. The samples are named by two letters: the first for the chromophore (C, M, N and V for Cr, Mn, Ni and V, respectively) and the second for the counterion (M, N, S and W for Mo, Nb, Sb and W, respectively); the undoped titania is referred to as AA.

Details on raw materials and synthesis conditions are provided in the first part of this study [15]. The pigments were ground in an agate mortar and sieved below 50 μm, then characterised with the UV-vis-NIR spectrophotometer (Perkin-Elmer, λ_{35} , 300–1100 nm range, step 0.3 nm) equipped with an integrating sphere made of BaSO₄. Measurements were performed with diffuse reflectance method using BaSO₄ as a reference, and CIELab coordinates were calculated by the spectra with standard illuminant D_{65} and observer 10° through the software UV WINLAB (Perkin Elmer). CIELab parameters express brightness $(L^* =$ white +, black -) and chroma $(a^* = \text{red} +, \text{green} -;$ b^* = yellow +, blue –). Optical spectra were deconvolved using a PFM software (Origin Lab). As far as colourimetric parameters are concerned, the colour purity is a measure of the saturation calculated using the lever rule from the Yxy diagram (shaped like a horse-shoe); in details the colour purity is calculated as a % ratio of the segment traced out from the white center to the sample point divided for the segment traced from the white center to the border of the Yxy diagram passing from the sample point.

The technological behaviour was assessed by adding (5 wt.%) of the pigment into a porcelain stoneware body and a ceramic glaze applied on a wall tile, then testing its chromatic appearance and stability for firing temperatures in the 1150–1250 °C range (porcelain stoneware) or 950–1150 °C range (ceramic glaze). The colour stability was evaluated by measuring the CIELab parameter $\Delta E^* = (\Delta L^* + \Delta a^* + \Delta b^*)^{-0.5}$ (HunterLab Miniscan MSXP4000) which expresses the total chromatic change with respect to a reference: the lower ΔE^* , the more stable the pigment. The

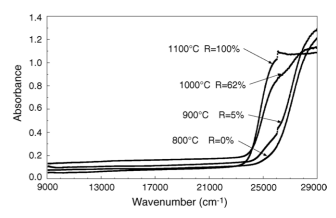


Fig. 1. Absorbance spectra of titania calcined at various temperatures with the amount of rutile formed (R%).

lowest temperature (950 °C for the glaze; 1150 °C for the stoneware) was chosen as reference.

3. Results and discussion

3.1. UV-vis-NIR spectroscopy

The main feature of the UV-vis-NIR spectra of rutile pigments is a characteristic band attributable to the ${\rm Ti}^{4+}\leftrightarrow {\rm O}^{2-}$ charge transfer, which moves from the near UV ($\sim 30,000~{\rm cm}^{-1}$) for anatase to the violet region of visible light ($\sim 25,000~{\rm cm}^{-1}$), so resulting in the pale yellow colour of rutile [16]. A gradual shift of this band can be actually appreciated along with the anatase-to-rutile transformation in the undoped titania (Fig. 1).

This circumstance fosters the overlapping of charge transfer (CT) and crystal field (CF) effects, making very difficult any quantitative interpretation from a rigorous spectroscopic approach, i.e. measuring precisely the CF strength Dq and the interelectronic repulsion Racah's *B* parameter. It was tried to overcome this drawback by eliminating the MLCT band, simply by subtracking the contribution of the undoped rutile from the visible spectrum [10] and deconvolving the expected CF peaks (Fig. 3). The

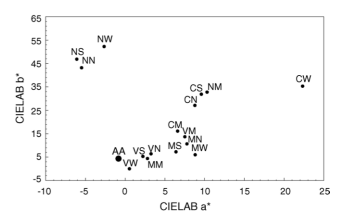


Fig. 2. Colour plot (CIELab a^* vs. b^*) of rutile pigments.

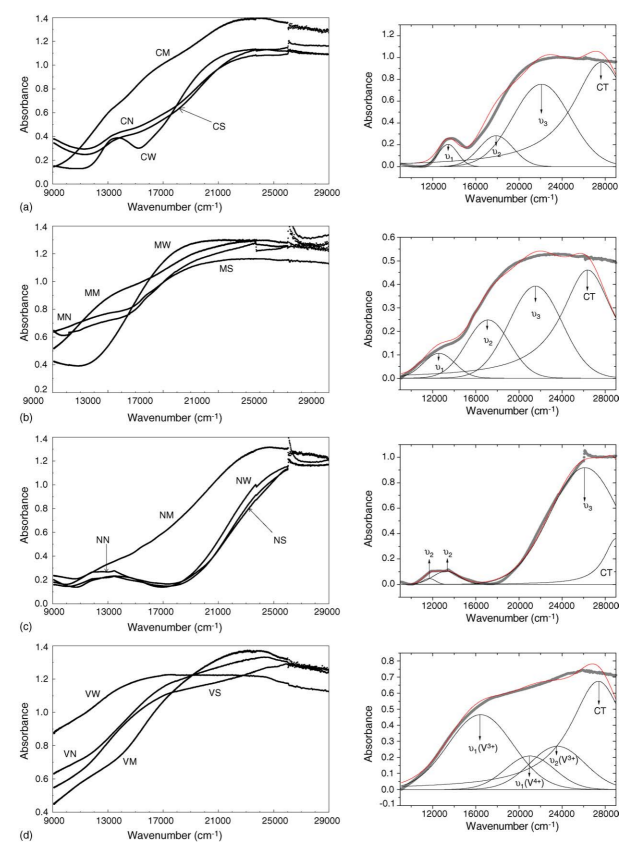


Fig. 3. Optical spectra of rutile pigments obtained by doping with chromium (a), manganese (b), nickel (c) and vanadium (d). In the right column: examples of peak deconvolution for samples CW, MS, NN and VS.

results are broad bands, which are somehow attributable to the main CF peaks of chromophore ions. $Dq_{(cub)}$ and B_{35} values were estimated by Tanabe-Sugano diagrams and spin-allowed transitions, respectively [16–18]; the crystal field stabilization energy (CFSE) was calculated by the Dq values [19]. The nephelauxenic ratio β_{35} , expressing the covalent character of the metal–oxygen bond, was calculated as B_{35}/B_0 , where the B_0 is the free ion value: the lower the β_{35} , the stronger the covalency [17].

A colour plot of the pigments synthesised at $1100\,^{\circ}\text{C}$ is presented in Fig. 2. The samples co-doped with Nb, Sb or W exhibits colours expected for Cr (orange), Mn (brown), Ni (yellow) and V (gray). In contrast, all Mo-bearing pigments present a brownish shade that alters considerably the desired hue.

3.1.1. Chromium-doped pigments

The spectra of Cr-bearing samples are characterised by (Fig. 3a):

- three main parity-forbidden transitions of Cr^{3+} in octahedral coordination, i.e. $\upsilon_1 = {}^4A_2({}^4F) \rightarrow {}^4T_2({}^4F)$, $\upsilon_2 = {}^4A_2({}^4F) \rightarrow {}^4T_1({}^4F)$, and $\upsilon_3 = {}^4A_2({}^4F) \rightarrow {}^4T_1({}^4P)$, which are not resolved, but whose positions can be inferred at $\sim 17,500, \sim 22,500$, and $\sim 42,000 \text{ cm}^{-1}$, respectively;

- two weak Cr^{3+} spin-forbidden transitions, i.e. ${}^4A_2({}^4F) \rightarrow {}^2T_1({}^2G)$ and ${}^4A_2({}^4F) \rightarrow {}^2E({}^2G)$, that are merged in the 13,500–13,600 cm $^{-1}$ range;
- a wide Ti-O CT band, that seems to extend its effect towards the lower energies, so overlapping the CF peaks.

The deconvolution of visible spectra, once the Ti–O CT band was subtracted, allows to estimate a CF strength Dq ranging from 1737 cm⁻¹ (CS) to 1757 cm⁻¹ (CW), that is consistent with the value expected for the Cr–O bond length in the rutile structure (196 pm) [19,20]. Racah's B_{35} values are in the 430–540 cm⁻¹ range (Table 1).

As far as the effect of counterions is concerned, Nb- and Sb-doped pigments give practically the same spectra, with a gently dipping slope to a minimum of absorbance at $\sim\!11500~{\rm cm}^{-1}$, resulting in an orange hue, slightly darker for CN. The presence of tungsten determines both a shift of the band edge to $\sim\!22,\!000~{\rm cm}^{-1}$ and a steeper slope; this circumstance brings about a lower absorbance of CW in the $14,\!000-18,\!000~{\rm cm}^{-1}$ region, with respect to CN and CS, that produces a purer orange colour. Furthermore, the band at $\sim\!13,\!600~{\rm cm}^{-1}$ is clearly more intense in the W-bearing pigment; this feature might be attributed to the $^2E \rightarrow ^2B_2$ transition of W^{5+} [18]. This inference should be in

Table 1 Optical spectroscopic data, crystal field strength (Dq), crystal field stabilization energy (CFSE), Racah's B_{35} parameter and β_{35} (= B_{35}/B_0) of the rutile pigments synthesised at 1100 °C

		Crystal field transitions			Dq (cm ⁻¹)	CFSE (kJ/g ion)	$B_{35} (\text{cm}^{-1})$	β_{35} (1)
		$\overline{v_1}$	v_2	v_3				
Cr ³⁺	CM	17,840	23,520	42,000	1784	-256.2	536	0.57
	CN	17,430	22,860	41,840	1743	-250.3	510	0.55
	CS	17,370	22,390	42,160	1737	-249.4	466	0.50
	CW	17,570	22,380	42,210	1757	-252.3	433	0.46
Mn ²⁺	MM	12,160	16,620	24,200	1337	0	807	0.85
	MN	11,500	16,730	23,610	1364	0	787	0.83
	MS	11,000	16,490	23,620	1407	0	787	0.83
	MW	12,300	17,100	23,490	1296	0	783	0.82
Mn ³⁺	MM	16,180			1618	-116.1	n.d.	
	MN	16,230			1623	-116.5	n.d.	
	MS	15,900			1590	-114.2	n.d.	
	MW	16,610			1661	-119.2	n.d.	
Ni ²⁺	NM	n.d.	12,210	25,870	843	-121.1	911	0.87
	NN	n.d.	12,180	26,660	841	-120.8	965	0.93
	NS	n.d.	12,470	26,270	861	-123.6	920	0.88
	NW	n.d.	12,650	26,580	874	-125.4	929	0.89
V ³⁺	VM	16,400	23,890	41,950	1740	-244.6	790	0.89
	VN	16,260	23,390	42,050	1703	-246.9	738	0.83
	VS	16,610	23,530	42,160	1730	-244.6	702	0.79
	VW	15,730	23,460	42,270	1679	-254.1	849	0.96
V ⁴⁺	VM	20,510			2051	-147.3		
	VN	20,810			2081	-149.4		
	VS	21,260			2126	-152.6		
	VW	20,940			2094	-150.3		

Crystal field transitions: Cr³⁺: $\upsilon_1 = {}^4A_2({}^4F) \rightarrow {}^4T_1({}^4F)$, $\upsilon_2 = {}^4T_2({}^4F)$, $\upsilon_3 = {}^4T_1({}^4P)$; Mn^{2+} : $\upsilon_1 = {}^6A_1({}^6S) \rightarrow {}^4T_1({}^4G)$, $\upsilon_2 = {}^4T_2({}^4G)$, $\upsilon_3 = {}^4E, {}^4A({}^4G)$; Mn^{3+} : $\upsilon_1 = {}^5E({}^5D) \rightarrow {}^5T_2({}^5D)$; Ni^{2+} : $\upsilon_2 = {}^3A_2({}^3F) \rightarrow {}^3T_1({}^3F)$, $\upsilon_3 = {}^3T_1({}^3P)$; V^{3+} : $\upsilon_1 = {}^3T_1({}^3F) \rightarrow {}^3T_2({}^3F)$, $\upsilon_2 = {}^3T_1({}^3P)$, $\upsilon_3 = {}^3A_2({}^3F)$; V^{4+} : $\upsilon_1 = {}^2T_2({}^2D) \rightarrow {}^2E({}^2D)$. Racah's B_0 (cm⁻¹): Cr³⁺: 933; Mn^{2+} : 960; Ni^{2+} : 1042; V^{3+} : 886 [17,18].

agreement with the structural data, which suggest the occurrence of W^{5+} [15].

The addition of molybdenum, in contrast, increases the absorbance all over the spectrum, presumably including Mo⁵⁺ absorbance bands at 13,000–14,000 cm⁻¹ and 19–23,000 cm⁻¹ [18], so producing a dark colour, far from the orange hue of the other Cr-bearing rutiles.

3.1.2. Manganese-doped pigments

The manganese-bearing samples exhibit a very wide band that practically absorbs all the light over \sim 19,000 cm⁻¹, leaving a moderate transmission in the red–orange region, which results in dark brown shades (Fig. 2). This wide band is probably due to the coalescence of CF peaks referable to (Fig. 3b):

- the $^5E(^5D)$ → $^5T_2(^5D)$ transition of Mn $^{3+}$, widely splitted in two bands at $\sim 11,500$ and $\sim 21,000$ cm $^{-1}$, respectively, that result centred at $\sim 16,000$ cm $^{-1}$ [17];
 the prevailing Mn $^{2+}$ transition $^6A_1(^6S)$ → $^4E,^4A(^4G)$ at
- − the prevailing Mn^{2+} transition $^6A_1(^6S) \rightarrow ^4E, ^4A(^4G)$ at $\sim 23,500 \text{ cm}^{-1}$, that is associated to the spin-forbidden bands: $^6A_1(^6S) \rightarrow ^4T_1(^4G)$ at $11,000-12,000 \text{ cm}^{-1}$ and $^6A_1(^6S) \rightarrow ^4T_2(^4G)$ at $\sim 16,500 \text{ cm}^{-1}$.

Among the counterions, Nb and Sb seem to play more or less the same role, being the UV-vis-NIR spectra very

similar, but a slightly higher absorbance for MN. As in the case of CW, the occurrence of tungsten made the band slope steeper, with a higher transmission in the $14,000-15,000\,\mathrm{cm}^{-1}$ range, which explains the redder hue of MW. However, this latter pigment contains surely a significant contribution of Mn^{2+} due to the presence of $\mathrm{MnWO_4}$ [15].

The Mo-bearing rutile is distinguished by an increased absorbance especially in the 14,000–17,000 cm⁻¹ range, that causes a darker shade of brown, maybe connected with the occurrence of multiple valences of molybdenum [18].

3.1.3. Nickel-doped pigments

The optical spectra of these samples are characterised by a strong absorbance band with the edge at $\sim\!\!26,\!000~{\rm cm}^{-1},$ as in undoped rutile, but with a gentler slope, whose onset is at $\sim\!19,\!000~{\rm cm}^{-1}$ (Fig. 3c). Less intense peaks, centred at the red-NIR border (12,000–14,000 cm $^{-1}$) leave the maximum light transmission be around 17,000 cm $^{-1}$, that justifies the deep and brilliant yellow colour of these pigments [12,13].

Two bands at $\sim 12,500$ and $\sim 26,000$ cm⁻¹ are usually attributed to the Ni²⁺ transitions in the octahedral field: from the ground term ${}^3A_2({}^3F)$ to $\upsilon_2 = {}^3T_2({}^3F)$, ${}^3T_1({}^3F)$ and $\upsilon_3 = {}^3T_1({}^3P)$, respectively [16,17]. In the red-NIR region there is another peak at $\sim 13,500$ cm⁻¹, that could be due either to the unusual co-presence of Ni³⁺, e.g.

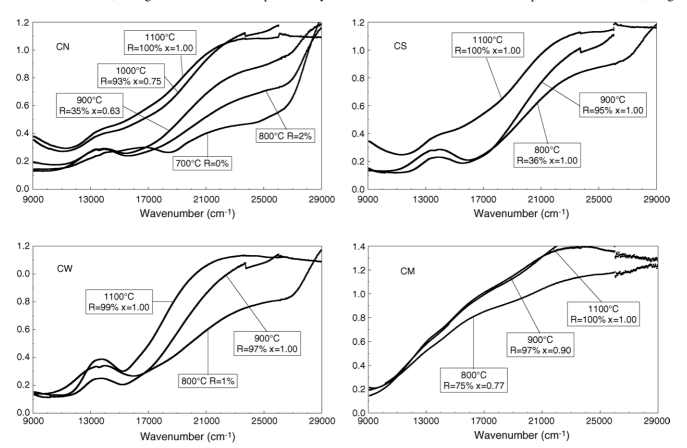


Fig. 4. Absorbance spectra of Cr-doped rutile pigments for increasing temperatures (700–1100 $^{\circ}$ C) with amount of rutile (R%) and Cr into the rutile lattice (x, where the molar fraction 0.03 is normalised to 1).

 $^4\text{T}_1(^4\text{F}) \rightarrow ^4\text{T}_2(^4\text{F})$, or to a considerable splitting of the Ni²⁺ υ_2 transition, as wide as 1190–1410 cm⁻¹, related to the TiO₆ octahedron distortion.

The optical spectra of pigments co-doped with Nb, Sb and W are very similar among each other; NW appears to be yellower since its absorbance band is slightly shifted to lower wavenumbers (i.e. more blue light absorbed), and hence it exhibits the minimum absorbance at $\sim 17,000~\rm cm^{-1}$ (i.e. more yellow light transmitted). The spectrum of NM pigment is completely different from the others Ni-doped samples, showing a much higher absorbance in the entire visible range and particularly a maximum value at $\sim 12,000~\rm cm^{-1}$, as a result of CF and/or CT effects linked to molybdenum multiple valences [18].

3.1.4. Vanadium-doped pigments

The co-doping with vanadium plus niobium, antimony or molybdenum gave similar results in terms of optical spectra, which are characterised by a quite regular slope connecting the NIR to the MLCT band with increasing absorbance up to 24,000 cm⁻¹. The transmitted light is scarce and limited to the red–orange region, so leading to the brownish-gray hue of these materials. In contrast, the Ti–V–W system exhibits a nearly constant absorbance along the entire visible range, that justifies its black colour (Fig. 3d). The different behaviour of this sample has been also confirmed by structural data [15].

The slope of these spectra can be explained by the overlapping of different CF transitions referable to:

- V³⁺ transitions $\upsilon_1 = {}^3T_1({}^3F) \rightarrow {}^3T_2({}^3F)$ at $\sim 16,500 \text{ cm}^{-1}$, $\upsilon_2 = {}^3T_1({}^3F) \rightarrow {}^3T_1({}^3P)$ at $\sim 23,500 \text{ cm}^{-1}$, and $\upsilon_3 = {}^3T_1({}^3F) \rightarrow {}^3A_2({}^3F)$ at $\sim 42,000 \text{ cm}^{-1}$;
- V^{4+} transition ${}^{2}T_{2}({}^{2}D) \rightarrow {}^{2}E({}^{2}D)$ at $\sim 21,000 \text{ cm}^{-1}$.

In reality, this latter band is usually splitted, when V^{4+} is in a distorted octahedral environment, as it occurs for the other d^1 ion Ti^{3+} [19]. For instance, in the turquoise pigment based on V^{4+} -doped zircon, the $^2B_2(^2D) \rightarrow ^2E(^2D)$ transition is splitted at $\sim 12,500$ and $\sim 15,700$ cm⁻¹ [21].

3.2. Colour development

The colour of titania pigments appears to develop contemporarily to the anatase-to-rutile transformation and the solid solution of chromophores and counterions into the rutile structure, though some sequential order is claimed in the literature [4]. These reactions are probably associated with valence changes of some cations in several of the systems under study [4,9,10].

The evolution of optical spectra and colourimetric parameters for increasing calcination temperatures will be discussed for every pigment, trying to attribute the spectral changes to CF or CT transitions as well as to the increasing amount of chromophore ion (as the molar fraction x) dissolved into the rutile lattice. A common feature for every system is the shift of the MLCT band along with the anatase-to-rutile transition in the 25,000–29,000 cm⁻¹ range.

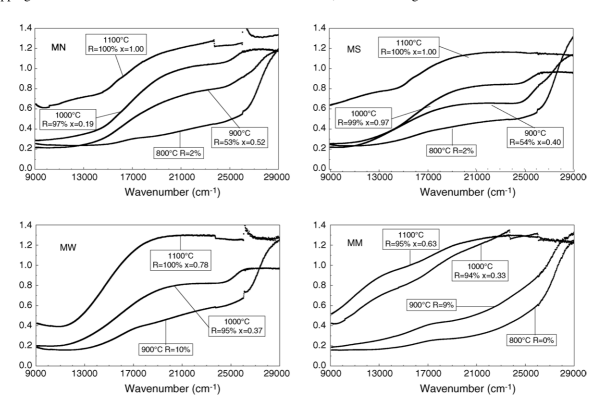


Fig. 5. Absorbance spectra of Mn-doped rutile pigments for increasing temperatures (700–1100 $^{\circ}$ C) with amount of rutile (R%) and Mn into the rutile lattice (x, where the molar fraction 0.03 is normalised to 1).

The *chromium-doped pigments* show a similar evolution that is best appreciable in the system CN (Fig. 4): a clear increase of absorbance occurs for increasing calcination temperature. The peak of Cr^{3+} υ_1 transition in Cr_2O_3 (centred at $\sim 16,000~cm^{-1}$) is present just after firing at 700 °C, then it shifts to a higher wavenumber as chromium is gradually diffused into the rutile lattice, though always with scarce intensity. Stronger effects concern both the υ_2 peak ($\sim 23,000~cm^{-1}$), which appears to be well correlated with the Cr amount in rutile, and the spin-forbidden transitions.

The spectra of manganese-doped pigments evolve with increasing temperature showing two distinct trends, apart from the above-mentioned band shift due to the titania polymorphic reaction (Fig. 5). The first one is the gradual growth of absorbance in the green-violet spectrum from 800 to 1000 °C, which might be tentatively explained by supposing the occurrence of Mn²⁺ in accessory phases [15], so justifying the increased absorbance around 24,000 cm⁻¹ at lower temperatures; a diffusion of Mn³⁺ into rutile could mainly occur at higher temperature, causing the absorbance increment especially at $\sim 16,000 \text{ cm}^{-1}$. However, it is well known that the Mn³⁺ peak undergoes a wide splitting in a distorted octahedral environment, such as that of rutile; this splitting could account for the absorbance in such a wide interval. The second trend is the outstanding increase of absorbance in the red-NIR region passing from 1000 to 1100 °C, which is more evident in MS and MN than in MW that is not easily explainable by CF attributions to manganese transitions.

The *nickel-bearing pigments* exhibit a fully comparable evolution of optical spectra for increasing temperature. Along with the shift of the MLCT band, there is a regular increment of the ${}^3A_2({}^3F) \rightarrow {}^3T_1({}^3P)$ transition at $\sim 23,000~{\rm cm}^{-1}$, with the progressive dissolution of Ni²⁺ in the rutile structure (Fig. 6). The intensity growth of the other peaks is much less evident, even if an appreciably increment of the peak at $\sim 13,500~{\rm cm}^{-1}$ occurred passing from 1000 to $1100~{}^{\circ}C$. This circumstance might be consequent to a partial oxidation of Ni²⁺, as this peak could be attributed to Ni³⁺, or more likely to a progressive splitting of the Ni²⁺ υ_2 transition.

The fundamental change in the *pigments containing* vanadium is the strong absorbance increase in almost the entire visible spectrum, since the shift of the Ti–O CT band is not well appreciable (Fig. 7) as even samples calcined at $800\,^{\circ}\text{C}$ are to a large extent transformed in rutile [15]. This evolution might imply a faster growth of peaks attributable to V^{3+} in respect of those related to V^{4+} . The main difference among the counterions is the marked increment of absorbance in the NIR-red field occurring with the tungsten addition, once the calcination temperature grew from 900 to $1100\,^{\circ}\text{C}$. Despite the small excess of WO₃ in the sample

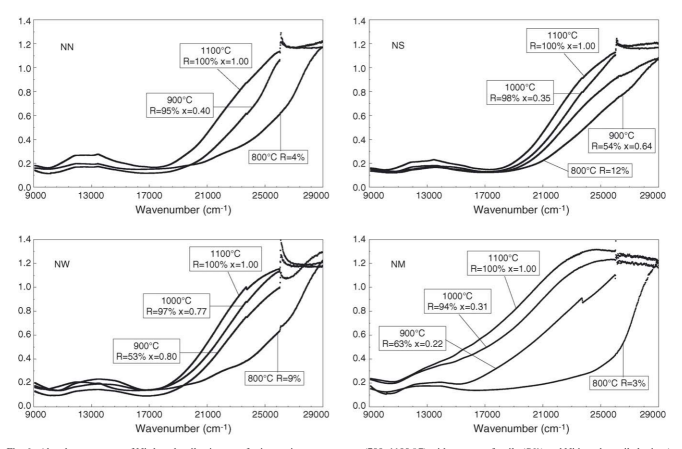


Fig. 6. Absorbance spectra of Ni-doped rutile pigments for increasing temperatures (700–1100 $^{\circ}$ C) with amount of rutile (R%) and Ni into the rutile lattice (x, where the molar fraction 0.03 is normalised to 1).

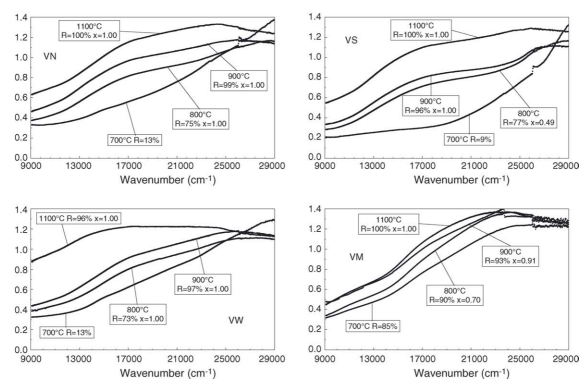


Fig. 7. Absorbance spectra of V-doped rutile pigments for increasing temperatures (700–1100 $^{\circ}$ C) with amount of rutile (R%) and V into the rutile lattice (x, where the molar fraction 0.03 is normalised to 1).

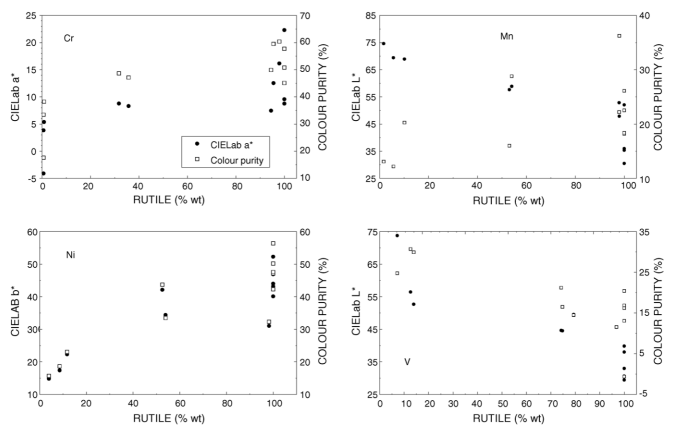
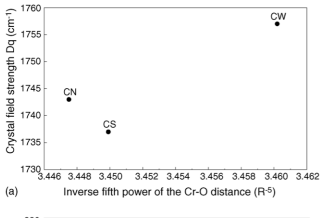


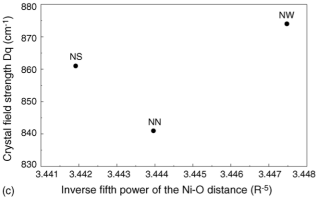
Fig. 8. Colour (CIELab a* or b*) and saturation (colour purity) of pigments fired at various temperatures (700-1100 °C) vs. the rutile amount.

VW, probably a specific contribution by W⁵⁺ should be claimed to account for this increased light absorbtion or, in alternative, the splitted Ti^{3+} υ_1 transition, due to the occurrence of Ti_5O_9 as accessory phase [15].

As a matter of fact, the colour evolves along with the amount of rutile in all systems; nevertheless, this change in hue and saturation goes beyond the total transformation of anatase in rutile, confirming analogous experimental observations [8], though with different effects for the various chromophores. In particular, for increasing firing temperature:

- a more saturated and purer colour is gradually developed in Cr-bearing pigments; however, all samples exhibit a counteracting trend from 1000 to 1100 °C, which increased the colour purity but reduced the red hue (Fig. 8a).
- Mn-doped pigments become regularly darker for increasing firing temperature and rutile amount, with a contemporary growth of purity and brown colour (Fig. 8b).
- Ni-containing pigments show a constant increase of both yellow hue and saturation with the rutile amount, which carries on for the highest temperature too (Fig. 8c).
- The continuous diminuition of L* found in V-bearing pigments indicates a progressive darkening with the increase of rutile content, that brings about a reduction of purity, as the colour becomes gray to black.





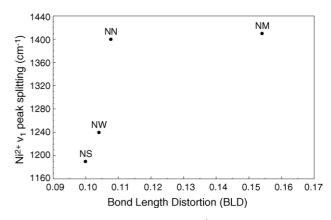
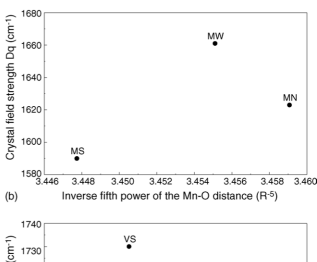


Fig. 9. Splitting of the v_1 transition of Ni²⁺ vs. the distortion of the octahedral site (BLD).

3.3. Colour and structure

The pigment colour depends on the surrounding environment of chromophore element, which can be appraised through the estimate of crystal field strength (Dq) or crystal field stabilization energy (CFSE) [19]. Moreover, the covalent character of metal–ligand bonding may be deduced on the basis of β_{35} parameter, which is the ratio between Racah's B_{35} determined by DRS and the value B_0 measured on the free ion: the lower β_{35} , the higher the covalency [18].



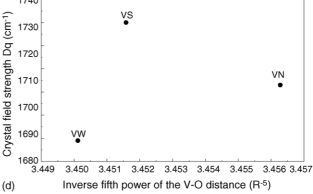


Fig. 10. Crystal field strength Dq vs. the inverse fifth power of the metal-oxygen distance R^{-5} for Cr-doped (a), Mn-doped (b), Ni-doped (c) and V-doped pigments (d).

The main absorbance peaks in the optical spectra of rutile pigments were inferred through a deconvolution procedure, that allowed to get averaged values of Dq, CFSE and β_{35} (Table 1). However, a reliable evaluation of peak splitting δ – attributable to TiO₆ octahedron distortion – is lacking, but for the Ni²⁺ transition ν_2 , whose splitting is in good agreement with distortion values: the wider δ , the larger the bond length distortion (BLD) parameter [15] (Fig. 9).

As expected by the crystal field theory, there is a general dependence of Dq on the metal–ligand distance (R), in particular the inverse of its fifth power (R^{-5}) [17,19]. As a matter of fact, the Dq values measured in rutile pigments respect the overall trend with R^{-5} defined by literature data [19]. However, this relationship is not strictly fulfilled, once appraised in detail, though it can somehow explain the changes in optical spectra of the series co-doped with Nb, Sb or W (Fig. 10). The Mo-bearing samples are omitted, as they are always clearly out of the general trend, likely because the multiple valences of molybdenum make not fully reliable the CF data obtained by peak deconvolution.

More covalent appears to be the Cr–O bonding $(\beta_{35} \sim 0.5)$ while Mn–O, Ni–O and V–O are all in the 0.8–0.9 β_{35} range. There is a general trend of increasing covalent character by changing the counterion: Mo < Nb \leq Sb < W, that is valid just in the Cr-, Mn- and

V-doped series, which in any case seems to be in contrast with the covalence expected on the basis of electronegativity values of these ions [18].

In the V-bearing pigments, there is the exception of VW, that lies completely far away the alignment VN–VS in Fig. 10d. Effectively, VW has structural features remarkably different from those of VN and VS, especially Ti–O distances and bond length distortion [15] which correspond to a peculiar optical spectrum and an almost purely ionic bond ($\beta_{35} = 0.96$).

Generally speaking, counterions seem to play an important role, contributing to modify the crystalline environment of chromophore and therefore the pigment colour. As a matter of fact, Nb- and Sb-doped rutiles exhibit similar colours, while W-doped pigments present always some distinctive features. Analogously, Nb- and Sb-bearing samples have similar values of CFSE, metal–ligand distance, bond length distortion and covalent bond character in each series, generally well distinct from those of the pigments containing tungsten.

3.4. Technological behaviour

The rutile pigments exhibit a different technological behaviour depending on the coupling of chromophore + counterion, that will be discussed separately for the two

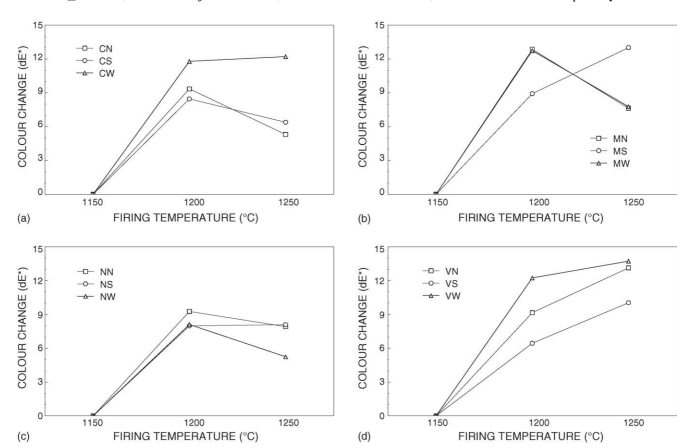


Fig. 11. Technological behaviour of rutile pigments applied into a porcelain stoneware tile body: (a) Cr-doped, (b) Mn-doped, (c) Ni-doped, and (d) V-doped. The lower the colour change (ΔE^*) , the higher the chemico-physical stability of the pigment.

ceramic applications here experimented: porcelain stoneware body and wall tile glaze.

3.4.1. Porcelain stoneware bodies

Contrasting the colour change with increasing firing temperature, it can be seen that, among the chromium-bearing pigments, the sample CW is less stable than CN and CS (Fig. 11a). As far as manganese-doped pigments are concerned, antimony ensures a better colour stability up to $1200\,^{\circ}$ C, but at higher temperatures niobium and tungsten seem to be more performant (Fig. 11b). Just slightly differences arose about pigments containing nickel, being tungsten the best counterion (Fig. 11c). The ranking of increasing stability in the vanadium rutiles is: W < Nb < Sb (Fig. 11d).

All these considerations are referred to colour changes, while the absolute chromatic performance is summarised in the CIELab a^* versus b^* plot (Fig. 12).

Some pigments present an unsuitable technological behaviour, as the colour of tiles is either not significantly different than that of undoped porcelain stoneware PS (e.g. MN, MS and VW) or similar to that achieved adding undoped rutile AA (e.g. MW and VN). The best results in through-body application are obtained with the pigments NW (yellow), CS (yellowish orange) and CW (orange-buff).

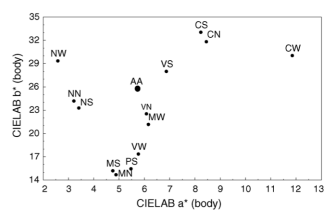


Fig. 12. Colour plot (CIELab a^* vs. b^*) of rutile pigments applied into a porcelain stoneware tile body.

3.4.2. Wall tile glaze

Counterions play a different role also in stabilizing the pigment in ceramic glaze applications. All counterions have more or less the same effect in Cr-doped pigments, with just small differences that suggest an order of increasing stability: W < Sb < Nb, with Mo exhibiting an anomalous behaviour, with the best performance at 1150 °C (Fig. 13a).

The pigments co-doped with Mn and Nb or Sb are very stable, since they do not vary their colour along the 1000–

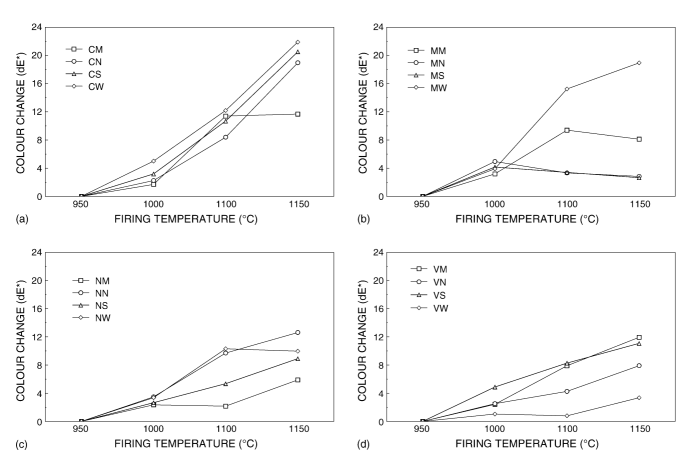


Fig. 13. Technological behaviour of rutile pigments applied into a wall tile glaze: (a) Cr-doped, (b) Mn-doped, (c) Ni-doped, and (d) V-doped. The lower the colour change (ΔE^*), the higher the chemico-physical stability of the pigment.

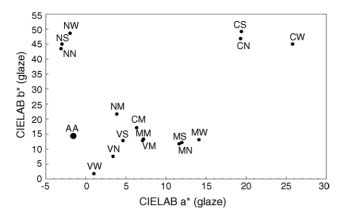


Fig. 14. Colour plot (CIELab a^* vs. b^*) of rutile pigments applied into a wall tile glaze.

1150 °C range. Less stable are the samples MM and particularly MW, which contain MnMo₄ and MnWO₄ respectively [15], that are probably dissolved in the molten glaze, so changing their colour (Fig. 13b).

Within the Ni-bearing pigments, the smaller chromatic variations are shown by the Mo-doped sample, which is however poorly significant since its colour is far from the desired yellow hue. Among the other counterions, there is an increasing stability order: $W \sim Nb < Sb$ (Fig. 13c).

The best properties among the V-containing pigments are ensured by tungsten, since no colour change occurs up to 1100 °C. Antimony and molybdenum counterions are scarcely stable, while VN presents an intermediate behaviour (Fig. 13d).

According to the absolute chromatic performance displayed in the CIELab a^* versus b^* plot (Fig. 14), the most suitable counterion is tungsten, being the best pigments for glaze applications: CW (orange), MW (tan), NW (yellow) and VW (black).

However, the colour performance could be also influenced by a different solubility of rutile in glazes depending on its particle size that, though not systematically analysed, was controlled by sieving during sample preparation. Nevertheless, the technological tests indicated W as the best counterions for all the chromophores, hence the influence of particle size appear not to be pivotal.

4. Conclusions

The colour of rutile pigments is determined by both metal-ligand charge transfer (${\rm Ti}^{4+} \leftrightarrow {\rm O}^{2-}$) and crystal field effects (chromophore elements substituting ${\rm Ti}^{4+}$ in octahedral coordination). The absorbance bands in the UV-vis-NIR spectra are broad and frequently overlapped, so resulting poorly resolved and making difficult any accurate quantitative spectroscopic interpretation.

Optical spectroscopic data suggest the occurrence of the following ions: Cr^{3+} , Mn^{2+} , Mn^{3+} , Ni^{2+} , V^{3+} , and V^{4+} .

These results confirm to a large extent the hypotheses advanced during the structural study of the same pigments [15].

The colour takes place during the pigment calcination in a multistage process, involving:

- shift of the Ti–O charge transfer band towards the visible range, consequent to the anatase-to-rutile transformation;
- development of crystal field peaks of chromophores due to progressive incorporation into the rutile lattice after decomposition of accessory phases;
- overlapping of absorbance peaks, due to valence change of chromophore ions (particularly manganese and vanadium, but probably molybdenum, antimony and tungsten too).

The counterions contribute to change the rutile colour by affecting somehow the crystalline environment of chromophore. In fact, the Nb- and Sb-doped pigments have close values of CFSE, metal-ligand distance, bond length distortion and bond covalency, resulting in very similar colours, while W-doped pigments present always some structural and colourimetric differences.

Rutile pigments proved to be suitable for through-body (up to $1250\,^{\circ}\text{C}$) and glaze applications (approximately up to $1100\,^{\circ}\text{C}$). The best glaze coloration is exhibited by W-bearing pigments, which however are less stable than Nbor Sb-containing ones, except than for the V + W coupling, which represents a new, interesting Co-free and Cr-free black pigment for low temperature applications. Into the porcelain stoneware bodies, the best coloration is achieved with Sb or W as counterions, though the best stability is ensured by Sb, but in the Ti–Ni–W system.

References

- P. Escribano Lòpez, J.B. Carda Castellò, C.E. Cordoncillo, Esmaltes y pigmentos ceràmicos, Faenza Editrice Iberica (2001) 1–300.
- [2] Italian Ceramic Society, Colour Pigments and Colouring in Ceramics, SALA, Modena, 2003, pp. 1–295.
- [3] DCMA, Classification and Chemical Description of the Complex Inorganic Color Pigments, 3rd ed., Dry Color Manufacturers Association, Alexandria, VA, 1991.
- [4] J. Maloney, Titanate pigments: colored rutile, priderite, and pseudobrookite structured pigments, in: H.M. Smith (Ed.), High Performance Pigments, Wiley-VCH, 2002, pp. 53–73.
- [5] R.A. Eppler, Reflectance of titania opacified porcelain enamels, J. Am. Ceram. Soc. 48 (5) (1969) 549–554.
- [6] R.A. Eppler, Crystallization and phase transformation in TiO₂-opacified porcelain enamels, I: Theory. II: Comparison of theory with experiment, J. Am. Ceram. Soc. 52 (2) (1969) 89–99.
- [7] F. Bondioli, T. Manfredini, G.C. Pellacani, Inorganic pigments for ceramic tiles: characteristics and industrial applications, Interceram 48 (6) (1999) 414–422.
- [8] N. Tozzi, R. Bindi, G. Ionescu, Ciclo produttivo e metodiche di controllo dei principali pigmenti ceramici in funzione del loro meccanismo di formazione, Ceramurgia 11 (5) (1981) 192– 199.

- [9] R.A. Eppler, Effect of antimony oxide on the anatase-rutile transformation in titanium dioxide, J. Am. Ceram. Soc. 70 (4) (1987) C64– C66.
- [10] S. Ishida, M. Hayashi, Y. Fujimura, K. Fujiyoshi, Spectroscopic study of the chemical state and coloration of chromium in rutile, J. Am. Ceram. Soc. 73 (11) (1990) 3351–3355.
- [11] M.A. Tena, M. Llusar, J.A. Badenes, J.B. Vicent, M.C. Granana, G. Monrós, Synthesis and structural characterisation of solid solutions $Cr_xTi_{1-2x-y}V_{x+y}O_2$ at atmospheric pressure, Br. Ceram. Trans. 98 (5) (1999) 230–233.
- [12] M.A. Tena, M. Llusar, J.A. Badenes, M. Vicente, G. Monrós, Influence of precursors on formation of TiO₂–CrTaO₄ rutile solid solutions, Br. Ceram. Trans. 99 (5) (2000) 219–224.
- [13] M.A. Tena, A. Mestre, A. García, S. Sorlí, G. Monrós, Synthesis of gray ceramic pigments with rutile structure from alkoxides, J. Sol–Gel Sci. Technol. 26 (2003) 813–816.
- [14] S. Sorlí, M.A. Tena, J.A. Badenes, J. Calbo, M. Llusar, G. Monrós, Structure and color of $Ni_xA_{1-3x}B_{2x}O_2$ (A = Ti, Sn; B = Sb, Nb) solid solutions, J. Eur. Ceram. Soc. 24 (8) (2004) 2425–2432.

- [15] F. Matteucci, G. Cruciani, M. Dondi, M. Raimondo, The role of counterions (Mo, Nb, Sb, W) in Cr-, Mn-, Ni- and V-doped rutile ceramic pigments. Part 1. Crystal structure and phase transformation, Ceram. Int. (2005)10.1016/j.ceramint.2005.03.014.
- [16] A.S. Marfunin, Physics of Minerals and Inorganic Materials. An Introduction, Springer-Verlag, 1979, pp. 1–340.
- [17] M. Wildner, M. Andrut, C.Z. Rudowicz, Optical absorption spectroscopy in geosciences. Part I: Basic concepts of crystal field theory, EMU Notes Mineral. 6 (2004) 93–143.
- [18] A.B.P. Lever, Inorganic Electronic Spectroscopy, 2nd ed., Elsevier, 1984, pp. 1–863.
- [19] R.G. Burns, Mineralogical Applications of Crystal Field Theory, 2nd ed., Cambridge University Press, 1993, pp. 1–551.
- [20] M. Andrut, M. Wildner, C.Z. Rudowicz, Optical absorption spectroscopy in geosciences. Part II: Quantitative aspects of crystal fields, EMU Notes Mineral. 6 (2004) 145–188.
- [21] S. Di Gregorio, M. Greenblatt, J.H. Pifer, M.D. Sturge, ESR and optical study of V⁴⁺ in zircon-type crystals, J. Chem. Phys. 76 (6) (1982) 2931–2937.