

Journal of Silicate Based and Composite Materials

A TARTALOMBÓL:

- Synthesis of hydrogarnets in the system Al₂O₃·2SiO₂-SiO₂-CaO-H₂O under hydrothermal conditions
- Rheological behaviour modelling of cement paste with nanotubes and plasticizer
- Sol-gel synthesis and structural characterization of Fe doped barium titanate nanoceramics
- Effect of H₂ on SiO and SiC formation
- Thermal stability and flame retardant properties of plasticized poly(vinyl chloride) hybrid composite for construction application
- Rheology of natural hydraulic lime pastes modified by non-traditional biopolymeric admixtures



2019/6



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Journal of Silicate Based and Composite Materials

TARTALOM

180 Az Al₂O₃·2SiO₂-SiO₂-CaO-H₂O rendszerben hidrogarnetek szintézise hidrotermikus körülmények között

184 Nanocsövek és lágyító hozzáadásával készült cementpaszták reológiai viselkedésének modellezése

Gintautas SKRIPKIUNAS • Ekaterina KARPOVA

■ Mindaugas DAUKSYS

Zdzisław PYTEL

190 Fe-adalékolt bárium-titanát nanokerámia szol-gél szintézise és szerkezeti jellemzése

Mohammed TIHTIH ■ Karoum LIMAME ■ Yahya ABABOU

■ Salaheddine SAYOURI ■ Jamal-Eldin F. M. IBRAHIM

194 A H₂ hatása a SiO és SiC képződésre

Trygve Storm AARNÆS ■ Merete TANGSTAD

198 Építőipari célú lágy PVC hibrid kompozitok hőstabilitása és lángállósága

Ali I. Al-MOSAWI • MAROSSY Kálmán

204 Nem-hagyományos biopolimer adalékszerekkel módosított természetes hidraulikus mészpépek reológiája

Tomáš ŽIŽLAVSKÝ ■ Martin VYŠVAŘIL ■ Pavla ROVNANÍKOVÁ

CONTENT

180 Synthesis of hydrogarnets in the system Al₂O₃·2SiO₂-SiO₂-CaO-H₂O under hydrothermal conditions Zdzisław PYTEL

184 Rheological behaviour modelling of cement paste with nanotubes and plasticizer

Gintautas SKRIPKIUNAS • Ekaterina KARPOVA

■ Mindaugas DAUKSYS

190 Sol-gel synthesis and structural characterization of Fe doped barium titanate nanoceramics

Mohammed TIHTIH ■ Karoum LIMAME ■ Yahya ABABOU

■ Salaheddine SAYOURI ■ Jamal-Eldin F. M. IBRAHIM

194 Effect of H₂ on SiO and SiC formation

Trygve Storm AARNÆS ■ Merete TANGSTAD

198 Thermal stability and flame retardant properties of plasticized poly(vinyl chloride) hybrid composite for construction application

A. I. AI-MOSAWI • Kálmán MAROSSY

204 Rheology of natural hydraulic lime pastes modified by non-traditional biopolymeric admixtures

Tomáš ŽIŽLAVSKÝ ■ Martin VYŠVAŘIL ■ Pavla ROVNANÍKOVÁ

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Synthesis of hydrogarnets in the system Al₂O₃ • 2SiO₂ -SiO₂ -CaO-H₂O under hydrothermal conditions

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Abstract

The paper summarises the experimental data on the products of synthesis of mixtures containing metakaolinite (AS $_2$), silica (SiO $_2$) and lime (CaO) in various proportions, under hydrothermal conditions at 180 °C. The actual compositions of reacting mixtures were determined by the molar ratios Ca/(Al+Si), Al/(Si+Al) and C/S, in the respective ranges (0.75÷1.5), (0.2÷0.5) and (1.0÷3.0). Compositions of the reacting mixtures correspond to stoichiometric ratios of particular oxides, being the components of hydrogarnets from the series $C_3AS_3 - C_3AH_6$, whose composition varies in accordance with the formula $Ca_3Al_2(SiO_4)_{3x}(OH)_{4x}$, for x in the range (0.0-2.0). The mineralogical characteristic of thus obtained products of synthesis are evaluated basing on XRD, DTA, TG, IR and SEM+EDAX test data. Results obtained to date reveal that the hydrogarnet C_3AH_6 is the most frequent product obtained in the analysed systems. It coexists with other products of the synthesis, belonging to the group of hydrated calcium silicates, chiefly revealed as an amorphous phase C-S-H and tobermorite ($C_5S_6H_6$).

Keywords: metakaolinite, hydrogarnets, hydrated calcium silicates, hydrothermal conditions Kulcsszavak: metakaolinit, hidrogénhidrát, hidratált kalcium-szilikátok, hidrotermikus körülmények

1. Introduction Hydrogarnets of

Hydrogarnets of the series $A_3B_2\{[TO_4]_{3-x}(OH)_{4x}\}$, x ranging from 1 to 3, have the structure of garnets in which the process of partial or complete isomorphic substitution occurs, referred to as 'hydrogarnet substitution', in accordance with the formula 4(OH) \leftrightarrow $(SiO_a)^4$. Cations of metals A and B occur in respective configurations 8 and 6 and the tetrahedron, cation T is mostly silicon [1]. Because of the possibility of isomorphic substitution as a continuous process, a series of solid solutions will be formed, their names depending on extreme terms in the given series [2]. Grossular ($Ca_3Al_2[SiO_4]_3$) - hydrogarnet ($Ca_3Al_2(OH)_{12}$) is a well known hydrogarnet series whose elemental composition varies in accordance with the general formula $Ca_3Al_2[SiO_4]_{3-x}(OH)_{4x}[3]$. It is assumed that those structures in the series are most stable in which x = 1, for example hydrogrossular (Ca₃Al₂[SiO₄]₂(OH)₄). This series is of great importance in the sector of manufacturing of construction materials based on mineral binders in the form of cement or lime [4-6]. This series is of particular importance for ceramic engineers since its components determine the functional features of those construction materials, particularly their durability and strength, in the context of their susceptibility to carbonisation. These impacts, however, are not clearly defined, as several aspects are involved, yet it is a well-established fact that the higher the silica contents in hydrogarnets, the better the chemical resistance of construction materials containing those mineral binders.

Depending on the extent of 'hydrogarnet substitutions', physical properties of individual components of the series of solid solutions will vary [7, 8]. In the case of the series $C_3AS_3 - C_3AH_6$, the increase in the number of substitutions of the tetrahedrons $[SiO_4]^{4-}$ by the complex $[(OH)_4]^{4-}$ is accompanied by the continuous change (increase) of the elementary cell parameters,

from $a_0 = 1.185$ for grossular to $a_0 = 1.256$ for hydrogarnete. Furthermore, as the number of complexes $[(OH)_4]^4$ increases, apart from the lattice parameters, the position and intensity of reflexes will change, too. Besides, some diffraction lines will disappear whilst new ones will appear instead. At the same time, the growth of the elementary cell is accompanied by the decrease of the refractive index 'n' from 1.743 for C_3AS_3 to 1.605 for C_3AH_6 and by the change of density. With the increase of the

amount of water present in the hydrogarnete structure in the

form of hydroxyl groups, the dehydroxylation effect becomes

more intense and the temperature of its maximum on the DTA

curve goes down from about 750-850°C for C₂AS₂ to (350-

2. Materials

500)°C for C₃AH₆ [9-11].

Reaction mixtures for the synthesis of hydrogarnets contain the following materials:

- metakaolinite METASTAR 501 (symbol ME) whose composition is shown in *Table 1*.
- burnt lime obtained by calcination analytically pure CaCO₃ at the temperature 1050°C for 3 hours (symbol C)
- silica powder (manufacturer: Quartzwerke GmbH), commercially available as SIKRON-Feinstmehl SGL-300 (symbol S)
- analytically pure aluminium oxide γ-Al₂O₃ (symbol A)
- distilled water (symbol H)

SiO₂ Al₂O₃ CaO MgO Fe₂O₃ Na₂O K₂O TiO₂ MnO P₂O₅ L.I.O 52.22 41.41 0.08 0.26 0.49 0.01 1.73 0.01 0.01 0.13 1.66

Table 1 Chemical analysis of metakaolinite METASTAR 501 1. táblázat METASTAR 501 metakaolinit kémiai elemzése

3. Experimental

The synthesis of hydrogarnets of the series $C_3AS_3 - C_3AH_6$ was conducted under hydrothermal conditions. Chemical composition of the reaction mixtures was varied, determined by the relevant molar ratios: C/(S+A), A/(S+A) and C/S. Those molar rations were determined by the complex value x present in the general formula defining the composition hydrogarnets of the given series. For each considered value of x a reaction mixture was prepared in the form of dry components, whose chemical composition would represent the theoretical hydrogarnate composition for the given value of x. Synthesis of hydrogarnets was conducted for the following values of x: 0; 1.0; 1.5; 2.0. Accordingly, the formulas expressing the composition of particular hydrogarnets and the corresponding molar ratios agree well with the data in *Table 2*.

Sample	Value	Hydrogarnet formula		Molar ra	tios
designation	of "x"			C/(S+A)	A/(S+A)
ME - 0	0	$\begin{array}{l} {\rm C_3AI_2[SiO_4]_3} \\ ({\rm 3CaO\cdot AI_2O_3\cdot 3SiO_2}, {\rm C_3AS_3}) \end{array}$	1.0	0.75 (3/4)	0.25 (1/4)
ME - 1	1.0	$C_3AI_2[SiO_4]_2(OH)_4$ (3CaO·AI ₃ O ₃ ·2SiO ₂ ·2H ₂ O, C ₃ AS ₂ H ₂)	1.5	1.0	0.33 (1/3)
ME - 1.5	1.5	$\begin{array}{l} {\rm C_{3}AI_{2}[SiO_{4}]}_{1.5}{\rm (OH)_{6}} \\ {\rm (3CaO\cdot AI_{3}O_{3}\cdot 1.5SiO_{2}\cdot 3H_{2}O, C_{3}AS_{1.5}H_{3})} \end{array}$	2.0	1.2 (6/5)	0.4 (2/5)
ME - 2	2.0	C ₃ Al ₂ [SiO ₄](OH) ₈ (3CaO·Al ₃ O ₃ ·SiO ₂ ·4H ₂ O, C ₃ ASH ₄)	3.0	1.5	0.5

Table 2 Chemical composition of synthesised hydrogarnets 2. táblázat A szintetizált hidrogarnetek kémiai összetétele

Reaction mixtures are prepared in stabilised and reproducible conditions. The procedure involved weighing the required amounts of basic components: metakaolinite (ME) and lime (C). Furthermore, certain amounts of silica powder (S) or aluminium oxide (A) were added to make up for the deficiency of silica or aluminium in metakaolinite and to obtain the desired chemical composition of reaction mixtures, determined by the molar ratios given in Table 2. Afterwards the mixtures are homogenised in an agate mortar, in the first stage with no presence of water, and then with water added. The amount of distilled water added to the mixture is determined by the need to ensure the complete hydration of calcium oxide and to achieve such consistency of the slurry that the ratio of water 'w' to dry substances 's' should equal 0.8. Thus prepared slurry is then transferred to Teflon crucibles which are then placed inside pressure cylinders acting as laboratory autoclaves. The slurry is then subjected to hydrothermal treatment. The samples are autoclaved at the temperature 180°C, which corresponds to the saturated vapour pressure of 1.002 MPa, for a variable period of time: 8, 24 and 72 hours. The time required to reach the specified conditions is about 1.5 hours and the autoclaves are cooled down to the ambient temperature in an unconstrained manner.

The phase composition of thus obtained materials is analysed using the XRD, DTA, TG, IR and SEM+EDAX methods.

4. Results

The analysis of the phase composition of thus obtained products uses mostly the XRD data. X-ray tests were performed using an X-ray diffractometer (Philips PW 1040). X-ray patterns were registered in the angle range CuK_{α} 5 - 60° 20 and the presence of mineral phases was established on the basis of ICPDS-ICDD file Copyright 2005. Test results in the form of XRD data are compiled in *Fig. 1*.

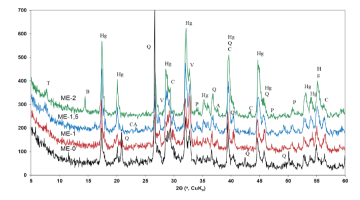


Fig. 1 XRD data of samples synthesised for 24 hours 1. ábra A 24 órán át szintetizált minták XRD adatai

Designations: Hg-hydrogarnate; Q-quartz; P-portlandite; C- calcite; V-vaterite; B-boehmite; A- aluminium oxide; CA-carboaluminate; T- tobermorite;

IR spectra (400 – 4000 cm⁻¹) are registered using the standard technique involving the KBr tablet transmission. Measurements are taken with a Fourrier spectometer Bio-Rad FTS 60MV with the resolution capacity 4 cm⁻¹ for 256 repetitions. Test results in the form of IR spectra of the investigated samples are shown in *Fig. 2*.

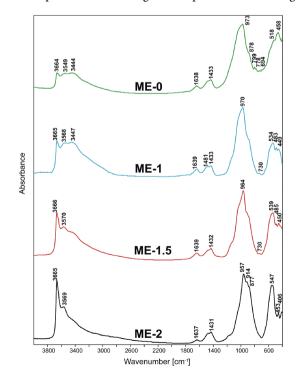


Fig. 2 IR spectra of the investigated materials 2. ábra A vizsgált anyagok IR-spektrumai

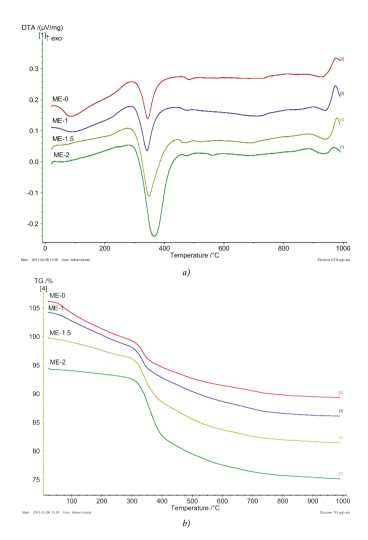
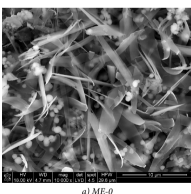
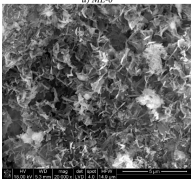
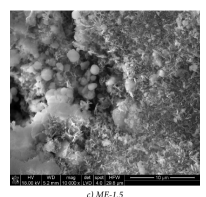


Fig. 3 Thermal treatment data: a) DTA curves; b) TG curves 3. ábra Hőkezelési adatok: a) DTA görbék; b) TG görbék







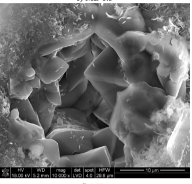


Fig. 4 Microstructure of investigated materials 4. ábra A vizsgált anvagok mikroszerkezete

Investigations by the thermal methods DTA and TG are conducted using the device NETZSCH STA 449 F30 *Jupiter*. Measuring is done on the carrier DTA-TG, in the temperature range 20-1000°C, the heating rate 10° C/min, in the presence of air with Al_2O_3 in the crucibles. Test results in the form of DTA and TG curves are shown in *Fig. 3*.

The microstructure of autoclaved materials is analysed using a scanning microscope NOVA $_{\rm NANO}$ SEM 200 (FEI COMPANY) equipped with a micro-analyser EDAX. Samples to be tested are first sprinkled with coal powder. The most distinctive images of microstructure of investigated sample profiles are shown in *Fig. 4*.

5. Discussion

The analysis of X-ray test data (*Fig. 1*) reveals that for each investigated sample the basic product of synthesis will be hydrogarnete, accompanied by tobermorite. Furthermore, for hydrogarnets we observe most characteristic line shifting for the specified value of 'd', which evidences the formation of solid solutions. The locations of reflexes from the planes (2,1,1) and (6,4,0) and their half-width indicate that in each case we obtain a hydrogarnete, whose composition varies in accordance with the Vegard's law (correlation factor 0.999), for *x* ranging from 2.38 to 2.65. It is reasonable to suppose that the composition of this product falls between the compositions of hydrogarnets given by the formulas: $C_3Al_2(SiO_4)_{0.62}(OH)_{9.5}$ and $C_3Al_2(SiO_4)_{0.35}(OH)_{10.6}$. Actually, that means that in each case the recorded extent of silica substitutions ('hydrogarnet substitution') is less than the predicted (theoretical) value [8, 12].

The analysis of IR spectra in Fig. 2 reveals that the band responsible for OH⁻ group substitutions is that registered for the wave number 3666 cm⁻¹. Its intensity grows with the increasing number of substitutions, and its small half-width is indicative of the fact that OH groups are distributed in an ordered manner within the structure of the material. The spectral line 3570 cm⁻¹ displays similar though less pronounced behaviour. The spectral line corresponding to the wave number 3400 cm-1 displays an entirely different behaviour and the large band half-width indicates that it can be associated with tensile vibrations of OHgroups, distributed in the statistical manner in the structure of the material, or, more probably, it can be associated with water molecules physically connected with the material's surface. This conclusion can be drawn from the changes of the band intensity correlated with that of the spectral line 1630 cm⁻¹, associated with deformative vibrations of H-O-H within the water molecule [13].

The IR spectrum for the sample ME - 0 reveals a spectral line 973 cm⁻¹, indicative of the occurrence of asymmetric tensile vibrations in Si-O and the spectral line 458 cm⁻¹ associated with bending vibration in SiO₄ tetrahedrons, which is a distinctive feature of ortho-silicates, including hydrogarnets. The spectrum also reveals the presence of a doublet of spectral lines 780-800 cm⁻¹, indicative of vibrations symmetrically stretching the Si-O-Si bridges, which is a distinctive feature of β -quartz. This doublet is absent in subsequent spectra, which suggests that the samples do not contain free quartz. Furthermore, a spectral line is revealed corresponding to the wave number 914 cm⁻¹, characteristic of Al-OH vibrations ($LK_{Al} = 6$, octahedron). The band intensity tends to increase in subsequent spectra, which can indicate that a part of OH groups are associated with aluminium, specifically with AlO, tetrahedrons. Of particular importance is the fact that position of a band with the highest intensity tends to shift from the wave number 973 cm⁻¹ (sample ME - 0) to 957 cm⁻¹ (sample ME - 2), which can indicate that substitutions of OH groups in a large extent depolymerize the structure, which tends towards the ideally mono-silicate structure. Furthermore, the spectra reveal the lines for the wave number 1432 cm⁻¹, associated with CO₃²⁻ groups, which is confirmed by the presence of carbonates.

DTA and TG curves for the investigated materials, shown in Fig. 3, are very similar. That can indicate that the analysed samples have a similar composition in qualitative terms. Quantitative differences involve the intensity and temperature of the predominant endothermic peak associated with dehydroxylation of hydragarnets. The enhanced peak intensity and the increase of its maximum temperature from 344.4°C (sample ME - 0, x = 0) to 366.4°C (sample ME - 2, x = 2) evidence the formation of the series of hydrogarnets with the decreasing number of silicate substitutions. However, the scale of the temperature increase suggests that the number of 'hydrogarnate substitutions' $4(OH) \rightarrow [SiO_A]^4$ inadequate in relation to the specified value of x, determining the composition of reaction mixtures. That is why the chemical composition of thus formed hydrogarnets does not represent their predicted (theoretical) composition [14].

The microstructural analysis of investigated samples (*Fig. 4*) reveals the co-existence of two basic products synthesised under hydrothermal conditions: tobermorite formed in the shape

of elongated crystals, and hydrogarnets with a typical crystal morphology. In a majority of investigated samples these products form a heterogeneous mixture of statistically distributed crystals.

One can conclude, therefore, that regardless of the initial composition of reaction mixtures, in each case the synthesis conducted under hydrothermal conditions will yield two products with crystalline structure: tobermorite and hydrogarnate, containing silicon. However, the extent of silicate substitutions in the structure of hydrogarnets is lower than expected and fails to reflect the chemical composition of mixtures used in the process of synthesis, no matter what the time of duration.

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Rheological behaviour modelling of cement paste with nanotubes and plasticizer

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Abstract

Rheological properties of cement pastes determine the technological properties of concrete mixtures and have a significant influence on the concrete placement, including pumping, self-compacting, 3D printing and others. The different types of additives and admixtures allow managing by the technological and rheological properties of cement systems. The lignosulphonate (LS), sulfonated naphthalene and melamine formaldehyde (NF), polycarboxylate ether (PCE) and acrylic polymer plasticizers, which change the consistency of concrete mixtures, are more widely used like plasticizing admixtures in construction practice nowadays. The carbon nanotubes are a perspective tool in managing by the rheological and technological properties of concrete in view of their outstanding mechanical, thermal, optical, electrical and some other properties.

The rheological behaviour of cement pastes with nanotubes and different types of polymer plasticizing admixtures was analyzed in the course of the current research using rotating rheometer with coaxial cylinders.

The cement pastes and concrete mixtures have the non-linear shear rate and shear stress dependence and require the application of non-linear rheological models such as Herschel-Bulkley or modified Bingham model. The approximation of experimental flow curves and evaluation of the rheological parameters such as yield stress, plastic viscosity and pseudoplastic indexes were performed based on the Herschel-Bulkley model in the course of the present research.

The testing of cement pastes in the presence of different plasticizing admixtures with and without MWCNT revealed the differences in rheological parameters depended on applied plasticizer. The molecular structure of plasticizing admixtures, small size, high aspect ratio and surface area of MWCNT, physical and chemical interactions between plasticizers and MWCNT can be a reason of observed changes in the rheological properties of cement pastes. The addition of MWCNT contributes to the obtaining of the cement paste of low dilatancy with definite flowability, yield stress and viscosity.

Keywords: cement paste, plasticizer, nanotubes, yield stress, viscosity, rheological model Kulcsszavak: cementpép, folyósítószer, nanocsövek, folyáshatár, viszkozitás, reológiai modell

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1. Introduction

The exploitation properties and durability of concrete structures depend on the properties of concrete mixtures and concrete placement technologies. Pumping of concrete mixtures is one the general way which is used for concrete transportation at the construction sites nowadays. The main rheological properties of concrete mixtures such as yield stress and plastic viscosity can be used for more accurate prediction of concrete mixtures behaviour in the course of a concreting. However, the evaluation of rheological parameters at the construction site is a challenging task caused by the complexity of rheological testing equipment. Therefore, the researchers are searching for a correlation between the main rheological parameters and technological properties of concrete mixtures [1-6]. For example, E. Secrieru et.al. investigated the effect of pumping on the rheological properties of fresh concrete. They performed a full-scale pumping experiment on a ready-mix concrete with the usage of viscometers for rheological test and revealed that pumping increases yield stress and reduces the viscosity of concrete mixture [1]. The research [2] reported the dependence between rheological parameters of concrete mixture and high pressure caused by concrete pumping.

In addition to pumping, the interest of researchers is devoted to the design of self-compacting concrete (SCC) [7-9] and development of 3D concrete printing technologies [10-12]. Based on the rheological parameters, the authors of the research [7] characterized the surface quality of SCC. They established that yield stress lower than 100 Pa and plastic viscosity lower than 10 Pa·s contribute to the surface defects of concrete caused by segregation. As for 3D printing technology, S.C. Paul et.al. studied 3D concrete printing technology and noticed that better pumpability of 3D printable cementitious materials is reached than thixotropy value is greater than 10000 N mm rpm [10].

The fundamental rheological model, which is used for the description of the flow behaviour of cementitious systems, is the Bingham model. The significant amount of researches demonstrate the usage of this model for the explanation of the rheological behaviour of cement systems [13-15]. The Bingham model characterizes the linear dependence between shear stress (τ) and shear rate (γ) of cement system and is described by the Equation (1):

$$\tau = \tau_0 + \mu_{pl} \cdot \gamma \tag{1}$$

where: τ - shear stress, Pa;

 τ_0 - yield stress of the cement paste, Pa;

 μ_{pl} - plastic viscosity of the cement paste, Pa·s;

 γ - shear rate, s⁻¹.

The rheological behaviour of modern concrete modified by different types of additives and admixtures require an application of the non-linear rheological model. The number of research presents the non-linear character of flow behaviour for modified cement systems [16-23]. The Herschel-Bulkley model and modified Bingham model are used to characterize a non-linear flow behaviour of cement systems. They described by the Equations (2) and (3), respectively:

$$\tau = \tau_0 + K \cdot \gamma^n \tag{2}$$

$$\tau = \tau_0 + \mu \cdot \gamma + c \cdot \gamma^2 \tag{3}$$

where: τ - shear stress, Pa;

 τ_0 - yield stress of the cement paste, Pa;

K - consistency factor, Pa·sⁿ;

μ - plastic viscosity, Pa·s;

γ - shear rate, s⁻¹;

n - shear thinning/ thickening index (n<1; n>1, respectively);

c - second order term, Pa·s² [16].

The usage of non-linear rheological model, such as Herschel-Bulkley rheological mode enable to derivate the pseudoplastic index of cement systems what is impossible in case of approximation according to the linear Bingham model. For example, A. Yahia tested the rheological behaviour of cement mortar with water-reducing admixtures and established the shear-thickening response of polycarboxylate plasticizer based on the Herschel-Bulkley model [21].

The increasing interest in the improvement of cement composites by addition of nano additives require an understanding of their action mechanism in the cement system. The analysis of rheological properties and their characterization by the rheological models is one of a way to describe the behaviour of nano additives in cement systems. As well as for the usage of chemical and mineral additives in cement paste, the rheological behaviour of nanomodified cement systems is described by non-linear rheological models. The research [23] presents the results of rheological testing of cement pastes modified by nano-SiO₂, nano-TiO₂, carbon nanotubes and nanofibers. The approximation of obtained rheological data was performed by a modified Bingham model.

The present research is focused on the analysis of the influence of different types of plasticizing admixtures (PL) and multi-walled carbon nanotubes (MWCNT) on the rheological

behaviour of cement pastes. The rheological parameters were evaluated based on the Herschel-Bulkley model.

2. Materials and testing methods

2.1 Materials

The Portland cement without mineral additives CEM I 42.5 R conforming to EN 197-1 with water consumption of 26.6% and fineness by Blain of 3552 cm²/g was used as a binder.

The MWCNT suspension was prepared from the pellets "Graphistrength CW 2-45" produced by the company "Arkema" (France). The pellet contains 45wt. % of MWCNT and 55 wt. % of carboxyl methyl cellulose (CMC). The MWCNT was characterized by the filament length of 0.1 –10 µm and the diameter of 15–20 nm. Distilled water was used as a dispersion medium for MWCNT. The lignosulphonate (LS), sulphonated naphthalene formaldehyde (NF) and polycarboxylate ether (PCE) plasticizers were applied in the research as a dispersant to distribute the MWCNT in the volume of water suspension and as a dispersing agent for cement pastes.

2.2 Testing methods

The ultrasonication by using a Bandelin Sonopuls HD 3400 ultrasonic homogenizer (400 W, 20 kHz) with probe VS 200 T (Ø25 mm, amplitude –82 µm) for 6 min was applied to distribute the MWCNT in water. The more detailed description of a homogenization process of MWCNT suspension is presented in the research [14]. 4 types of MWCNT suspensions, including MWCNT suspension without PL and with LS, NF and PCE plasticizers, were obtained. The concentration of PL was of 3.8% in all aqueous MWCNT suspensions excluding MWCNT suspension without PL. The dosage of PL was chosen to reach the dosage of 1% bwoc in cement pastes as more widely used plasticizer dosage in cement systems.

The rheological test was carried out by rotational rheometer Rheotest RN 4.1 with coaxial cylinders (the scheme of cylinder measuring system is presented in *Fig. 1*) according to the mode presented in *Fig. 2* at 5, 60, 120 min after cement paste mixing. 200 g of cement was used for preparing of cement paste for each rheological test. Water to cement ratio (W/C) was 0.30 and 0.25 for cement paste modified by MWCNT suspension without and with PL, respectively. The MWCNT suspensions were used as mixing water for the cement paste modification in the course of the rheological test. MWCNT dosage was of 0.06% bwoc in the course of rheological tests.

The main rheological parameters, such as yield stress (τ_0) and plastic viscosity (μ) were determined in the course of approximation of flow curve (dependence between shear stress (τ) and shear rate (γ) in the range from 0.1 to 100 s⁻¹ based on Herschel-Bulkley model.

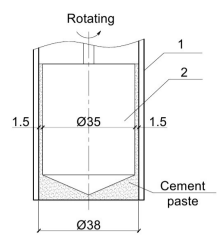


Fig. 1 Scheme of cylinder measuring system Rheotest RN 4.1 1. ábra A Rheotest RN 4.1. mérő rendszer vázlata

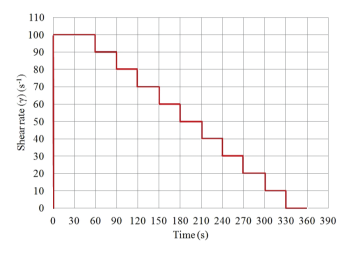


Fig. 2 Shear rate mode of the rheological test 2. ábra A reológiai teszt a nyírási sebesség meghatározásához

3. Results of Investigation

3.1 Influence of chemical admixtures on the flow behaviour of cement paste

The modification of cement paste by PL of different chemical structure leads to the changes in their rheological parameters in different ways. *Fig. 3* represents the flow behaviour of cement pastes without and with the addition of LS, NF and PCE plasticizers in 5 min after cement paste mixing. The approximation of the obtained rheological curves was performed according to the Herschel-Bulkley model. The established rheological parameters, such as plastic viscosity, yield stress and shear thinning/thickening indexes identified the dependence between flow behaviour of cement pastes and applied type of PL. The rheological parameters obtained in the course of the approximation of flow curves by Herschel-Bulkley are listed in *Table 1*.

The significant decrease of yield stress value up to 0.3 Pa was reached in case of application of PCE plasticizer in the cement paste. The yield stress for LS and NF plasticizers were 11.4 and 5.2 Pa, respectively.

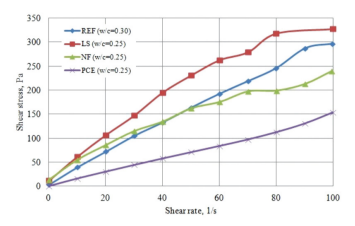


Fig. 3 Flow curves of cement pastes with different PL in 5 min after cement paste mixing

3. ábra Különböző PL tartalmú cementpépek folyási görbéi 5 perc alatt a cementpép keverése után

Specimen	W/C ratio	τ _ο , Pa	μ, Pa·s	n
Reference	0.30	3.6	4.0	0.9
LS	0.25	11.4	13.5	0.7
NF	0.25	5.2	6.0	0.8
PCE	0.25	0.3	0.6	1.2

Table 1 Rheological parameters of cement paste with different plasticizing admixtures obtained based on the Herschel-Bulkley model

1. táblázat A Herschel-Bulkley modell alapján kapott különböző folyósítószert tartalmazó cementpépek reológiai paraméterei

The plastic viscosity gained a smaller value for PCE plasticizer and was of 0.6 Pa·s in comparison with plastic viscosity of 13.5 and 6.0 Pa·s for LS and NF plasticizers, respectively.

Also, the difference in the rheological behaviour of the cement pastes with different PL was observed based on pseudoplastic indexes. The LS and NF plasticizers characterized by the indexes 0.7 and 0.8, respectively, that identify their shear-thinning flow behaviour. The cement paste with PCE plasticizer demonstrated the shear thickening flow behaviour with the index of 1.2.

The w/c ratio for the reference sample was of 0.30 caused by testing procedure. The values of the yield stress, plastic viscosity and pseudoplastic index were reached of 3.6 Pa, 4.0 Pa·s and 0.9, respectively. Despite the differences in w/c ratio for the reference sample and samples with PL, it can be concluded that PCE plasticizer has greater plasticizing efficiency in cement paste in comparison with other PL. Mainly, the observed phenomenon can be explained by the differences in the molecular structure of polymers which causes the differences in their action mechanism in the cement systems.

3.2 Influence of MWCNT with and without chemical admixtures on the flow behaviour of cement paste

The present section shows the effect of MWCNT on the rheological behaviour of cement pastes. The separate and combine usage of MWCNT with PL were tested. The flow curves of cement pastes modified by MWCNT with and without PL are presented in *Fig. 4-7*. The values of yield stress, plastic viscosity and shear thinning/ thickening indexes are shown in

Table 2. The values were obtained based on their approximation according to the Herschel-Bulkley rheological model.

Fig. 4 presents the flow curves for cement paste without and with MWCNT prepared in the same w/c ratio of 0.30. The addition of MWCNT to the cement paste leads to the slight increase of yield stress and plastic viscosity in comparison with the reference sample by 8 and 5 %, respectively. The shear thinning index was changed from 0.94 for reference to 0.98 for nanomodified cement paste.

The addition of various types of PL in combination with MWCNT had a different impact on the rheology of cement pastes.

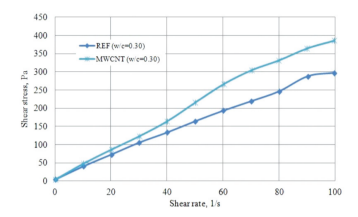


Fig. 4 Flow curves of cement paste with and without MWCNT in 5 min after cement paste mixing

4. ábra A cementpép folyási görbéi MWCNT-vel és anélkül 5 perc alatt a cementpép keverése után

Fig. 5 presents the results of rheological testing of cement pastes modified by MWCNT in the presence of LS plasticizer. The addition of MWCNT to cement paste with LS slightly increases the yield stress by 5% and not significantly decreases the plastic viscosity by 2%. The modification of cement paste by MWCNT changes the shear thinning index from 0.70 to 0.64.

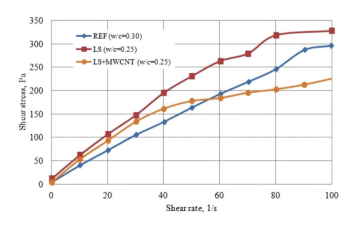


Fig. 5 Flow curves of cement paste with and without MWCNT in the presence of LS in 5 min after cement paste mixing

5. ábra A cementpép folyási görbéi MWCNT-vel és anélkül LS jelenlétében 5 perc alatt a cementpép keverése után The cement paste modified by MWCNT in the presence of NF plasticizer characterized by the increase of yield stress and plastic viscosity by 4% and 20%, respectively. The shear thinning was changed from 0.83 for cement paste modified by NF to 0.74 for cement paste with MWCNT and NF. The flow curves of cement pastes modified by MWCNT in the presence of NF are represented in *Fig.* 6.

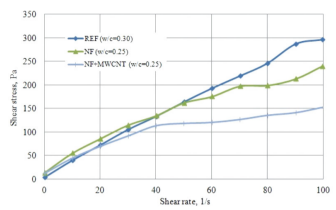


Fig. 6 Flow curves of cement paste with and without MWCNT in the presence of NF in 5 min after cement paste mixing

6. ábra A cementpép folyási görbéi MWCNT-vel és anélkül NF jelenlétében 5 perc alatt a cementpép keverése után

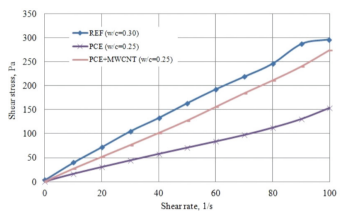


Fig. 7 Flow curves of cement paste with and without MWCNT in the presence of PCE in 5 min after cement paste mixing

7. ábra A cementpép folyási görbéi MWCNT-vel és anélkül PCE jelenlétében 5 perc alatt a cementpép keverése után

The remarkable changes in rheological parameters of cement pastes were observed in case of their modification by MWCNT in the presence of PCE plasticizer. The flow curves of cement paste modified by MWCNT in the presence of PCE plasticizer are shown in *Fig. 7*. The addition of MWCNT to cement paste with PCE plasticizer led to the increase of the yield stress by 33% in comparison with cement paste modified only by PCE plasticizer. The plastic viscosity of nano modified cement paste increased in 2 times in comparison with cement paste with PCE plasticizer. The cement paste with PCE plasticizer characterized by shear thickening flow behaviour. The modification by MWCNT decreased the shear thickening index from 1.20 to 1.18.

The absolute values of tested rheological parameters of cement paste with MWCNT in the presence of LS, NF and PCE plasticizers are given in *Table 2*.

Specimen	W/C ratio	τ ₀ , Pa	K, Pa∙s ⁿ	n
Reference	0.30	3.6	4.0	0.94
MWCNT	0.30	3.9	4.2	0.98
LS	0.25	11.4	13.5	0.70
LS+MWCNT	0.25	12.0	13.3	0.64
NF	0.25	5.2	6.0	0.83
NF+MWCNT	0.25	5.4	4.8	0.74
PCE	0.25	0.3	0.6	1.20
PCE+MWCNT	0.25	0.4	1.2	1.18

Table 2 Rheological parameters of cement pastes with and without MWCNT in presence of plasticizing admixtures obtained based on Herschel-Bulkley model 2. táblázat A cementpépek reológiai paraméterei MWCNT-vel vagy anélkül folyósítószerek jelenlétében a Herschel-Bulkley modell alapján

MWCNT has an impact on the rheological properties of cement pastes. The small size, high aspect ratio, specific surface and porous structure of MWCNT can be a reason for the observed influence on the cement paste. These features of MWCNT contribute them to act physically and chemically in the cement paste. The usage of MWCNT in combination with various polymer PL can increase or decrease their impact on the rheological properties of the cement pastes in dependence on the chemical structure of PL and its ability to interact with MWCNT.

4. Discussions of results

As it was mentioned above, the Herschel-Bulkley and modified Bingham rheological models give more information about the rheological behaviour of the studied cement systems [16-23].

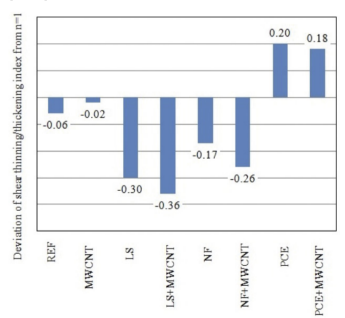


Fig. 8 Deviation of shear thinning/thickening indexes from n=1 in absolute units 8. ábra A nyírási vékonyodási / vastagodási indexek eltérése n = 1-től abszolút egységekben

In the course of the present research, the approximation of the rheological flow curves by Herschel Bulkley model identified the changes of shear thinning/thickening indexes for cement pastes modified by MWCNT in the presence of PL. The shear thinning indexes for cement pastes with MWCNT in the presence of LS and NF were changed from 0.70 to 0.64 and from 0.83 to 0.74, respectively. The cement pastes with MWCNT in the presence of PCE plasticizer were characterized by the changes of shear thickening index from 1.20 to 1.18. The *Fig.* 8 presents the deviation of the pseudoplastic indexes of cement pastes from n=1 in absolute units.

Based on the obtained results, it can be concluded that the addition of MWCNT to cement paste in the presence of various polymer PL leads to the decrease of dilatancy of cement paste. The level of changes in pseudoplastic index is depended on the type of applied PL and can be explained by its molecular structure.

5. Conclusions

- 1. The cement pastes have non-linear rheological behaviour. The processing of rheological data based on non-linear Herschel-Bulkley or Bingham rheological models allows obtaining more rheological parameters with high accuracy.
- 2. The various types of polymer plasticizing admixtures have a different influence on the cement paste flow behaviour. The application of LS, NF increases the thixotropy of cement pastes. The usage of PCE plasticizer revealed the increase of dilatancy of cement paste in comparison with the reference sample without admixtures.
- 3. The MWCNT have an impact on the rheological parameters of cement paste. The complex usage of MWCNT with LS and NF revealed the increase of yield stress by 5 and 4%, respectively. The plastic viscosity decreased by 2% in case of a modification of cement paste by MWCNT and LS. The usage of MWCNT with NF plasticizer led to an increase of the plastic viscosity of the cement paste by 20%. The more significant changes in rheological parameters of cement pastes were obtained for cement pastes modified by MWCNT in the presence of PCE plasticizer. The yield stress increased by 33% and plastic viscosity enhanced in 2 times in comparison with cement paste without MWCNT. The evaluation of shear thinning/ thickening indexes of nanomodified cement pastes identified that MWCNT contribute to the decrease of dilatancy of cement pastes. The observed differences in rheological behaviour of nanomodified cement paste in the presence of different plasticizing admixture can be explained by physical and chemical interactions between polymers and MWCNT.
- 4. Modification by MWCNT allows obtaining the cement paste of low dilatancy with definite flowability, yield stress and viscosity.

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Sol-gel synthesis and structural characterization of Fe doped barium titanate nanoceramics

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Abstract

Fe-doped barium titanate (BFe $_x$ T) nanoceramics were successfully prepared by a simple sol-gel process, these materials are very interesting and could be a candidate for many optoelectronic applications. X-ray diffraction (XRD) patterns of the obtained samples, heat-treated at a quite low temperature (800 °C/3h) revealed that BFe $_x$ T nanoceramics crystallized into a tetragonal phase perovskite structure. The occupation of the Ba and Ti sites by Fe in the BaTiO $_x$ lattice and the evolution of the different parameters (crystallite size, lattice parameters and strain) as functions of the Fe doping have been discussed in details using several characterization techniques including XRD and Fourier Transformation Infrared (FT-IR).

Keywords: Sol-gel, Barium titanate nanoceramics, Fe-doped barium titanate nanoceramics, X-ray Diffraction

Kulcsszavak: Szol-gél, bárium-titanát, nanokerámia, Fe-adalékolt bárium-titanát, röntgendiffrakció

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1. Introduction

The popularity of the research in the area of the high-tech ceramics is largely grown lately [1-31], since the discovery of the ferroelectricity phenomenon, barium titanate BaTiO₂ (BT) has been a fundamental member of the ferroelectric perovskite family. This compound has been used for a long time in many industrial sectors including full swing multilayer ceramic capacitors (MLCCs) [32], the realization of the ferroelectric random access memories (FRAMs) [33], thermistors manufacturing [34] and detection of polluting gases such as CO [35]. However, multilayer capacitors have long been at the forefront of BaTiO₂ applications. The BaTiO₂ nanoceramics can be synthesized by various methods such as solid-state reaction [36], hydrothermal synthesis [37], sol-gel [38] and co-precipitation method [39], compared to other methods, solgel is considered as a simple and easy method which can allows production of a high purity BaTiO₂ with very fine particles of controllable size, moreover, it is possible to easily modify the physical properties of BT since these are very sensitive to doping on the A-site (Ba_DTiO₃), B-site (BaTi_DO₃) and substitution on both sites (co-doping) Ba_DTi_DO₃ which mainly depends on their ionic radii [40]. Recently, the doping of BaTiO₃ nanoceramics by 3d transition metals (Fe, Co, Mn, Ni...) have attracted much attention, 3d transition elements are more likely to be considered as dopants due to their electric and high magnetic properties. To date, a number of research works have

reported the effect of the 3d transition metals on the structural and physicochemical properties of BaTiO₃ nanoceramics. Rani et al [41] have studied the effect of Fe doping on structural, magnetic and magnetoelectric properties of BaTiO, prepared via solid-state reaction route. Khirade et al [42] have reported the effect of Fe doping on the structural, optical and electrical properties of BaTiO₃ nanoceramics synthesis by sol-gel process. Maikhuri et al [43] have demonstrated the influence of A- and B-site substitution on the structural and magnetic properties of BaTiO₃. In the present work, Fe doped BaTiO₃ nanoceramics have been prepared using sol-gel method, the prepared samples were characterized using X-Ray Diffraction and FT-IR. The analysis of X-ray patterns was used to calculate the crystallite size, lattice parameters and lattice strain. The aim of this investigation is to study the effect of Fe dopant on the structural properties of BaTiO₃, In particular, the occupation of Ba and/or Ti sites by Fe in the BaTiO₃ structure.

2. Method of elaboration

Pure and Fe-doped BaTiO₃ were synthesized using sol-gel method, barium acetate trihydrate (Ba(CH₃CO₂)₂.3H₂O), Iron acetate Fe(C₂H₃O₂)₂ and titanium alkoxide Ti[OCH(CH₃)₂]₄ were used as precursors, lactic acid (CH₃CH (OH)COOH) was used as peptizing agent, acetic acid was added to dissolve Iron acetate, and distilled water as solvent. As shown in *Fig. 1*, the first step is to prepare a colloidal solution of TiO₂, to do this; the

titanium alkoxide was added to an aqueous solution of lactic acid and $\rm H_2O$ with continuous stirring at 70 °C. After 24 hours of reaction, the white precipitate obtained is changed into a clear homogeneous solution. In a second step, the colloidal solution produced was added in stoichiometric quantities to iron and barium acetates. The obtained transparent sol was converted to a translucent gel after stirring at 90 °C. Fine powders were prepared from the gel after drying at 90 °C and grinding. Finally, the nanopowders produced were calcined in air at a temperature of 800 °C for 3 h in a programmable oven. Phase identification of the prepared BFexT was performed using X-ray diffraction (Cu Kα radiation, $\lambda = 1.5405980$ Å) and Fourier transform infrared.

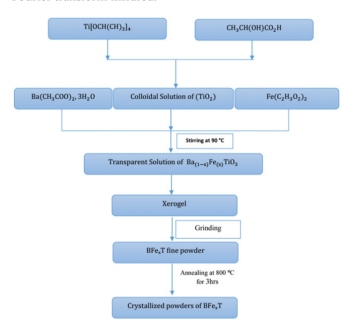


Fig. 1 Flow chart of the preparation of Fe doped BaTiO₃ nanoceramics by sol-gel

1. ábra $\stackrel{\frown}{A}$ Fe-adalékolt BaTiO $_3$ nanokerámia szol-gél eljárással történő előállításának folyamatábrája

3. Results and discussion

3.1 Structural studies

Fig. 2 shows the X-ray diffraction patterns of Ba_{1-x}Fe_xTiO₃ (x = 0, 1, 3%) samples prepared via sol-gel method. XRD patterns of pure and Fe-doped BaTiO3 show a single tetragonal phase without any evidence of secondary phases. The XRD pattern of the samples can be indexed to the tetragonal perovskite structure with P4mm space group, which is in great concurrence with JCPDS No. 05-0626. An important fact revealed in Fig. 2 is that the peak of (101) of Ba, Fe TiO₃ (x=0.01) is shifted to higher 2θ angle compared to the pure BaTiO₂, while the peak of (101) of Ba_{1.x}Fe_xTiO₃ (x=0.03) is shifted to lower 2θ angle. The Peaks shifted to lower and higher angles indicate an increase and decrease in volume (V) of the unit cell respectively as shown in Table 1. Fe²⁺ has a smaller ionic radius (0.78 Å) compared to that of Ba^{2+} (1.35 Å). The slight shift of the peak positions in BFe1T with respect to BaTiO, is may be attributed to the Fe2+ ionic size difference, which could eventually replace the Ba²⁺ ions in the BaTiO₃ lattice [40] and leads to a reduction in

the volume of the unit cell. In addition, the lower angle shift in the peaks of BFe3T can be assigned to the substitution of Fe $^{2+}$ ion by Ba $^{2+}$ and Ti $^{4+}$ ions, but with a predominance of the Ti site occupancy in the BaTiO $_3$ lattice. In contrary, and because of the reason that the Fe $^{2+}$ ionic radii is larger compared to Ti $^{4+}$ (0.605 Å), it can be observed that the substitution of Ti-sites by Fe $^{2+}$ create oxygen vacancies to neutralize the charge. Therefore, an increase in the unit cell volume is predicted as the unit cell parameters a and c have increased.

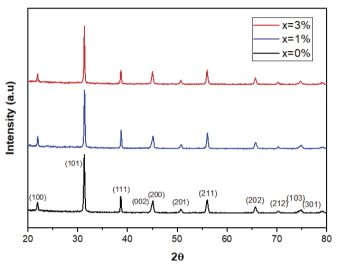


Fig. 2 X-ray diffraction patterns of prepared Fe-doped barium titanate nanoceramics

2. ábra Az elkészített Fe-adalékolt bárium-titanát nanokerámiák röntgendiffraktogramja

The crystallite size of the pure and Fe-doped BaTiO₃ was calculated from the X-ray diffraction patterns according to (101) peak using Debye-Scherrer formula [44]:

$$D = \frac{0.9\lambda}{\beta \cos \theta} \tag{1}$$

Where D is the crystallite size, λ is x-ray wavelength, θ is the diffraction angle and β is full width at half maximum (FWHM) of the (101) peak. The average crystallite size was calculated by settling the highest intensity peak. As listed in *Table 1*.

Moreover, the lattice strain (ϵ) of all samples was estimated using the following equation [45]:

$$\varepsilon = \frac{\beta}{4\tan\theta} \tag{2}$$

The estimated lattice parameters a and c, unit cell volume (V), position of the peak (101), lattice strain and crystallite size are listed in *Table 1*. It was found that the average crystallite size (D) increased with Fe concentration x. However, the lattice constant (c) and lattice strain have been decreased after doping. The increase in crystallite size is due to the big difference between the ionic radius of Fe²⁺, Ba²⁺ and Ti⁴⁺ [31, 42]. The shift in the position of (101) in the XRD patterns and the change in the volume of unit cell after doping reveal that Fe is totally soluble into BaTiO₃ lattice.

3.2 FT-IR analysis

The Fourier transform infrared spectroscopy (FT-IR) spectrums of BaFe_xTiO₃ (x=0, 1, 3%) are shown in *Fig. 3*. The corresponding spectra show absorption bands in the wave number ranging from 4000 – 450 cm⁻¹. The presence of a prominent peak at about 500 cm⁻¹ in the pure BaTiO₃ spectrum corresponds to the vibration of Ti-O bond in the crystal lattice [46]. All the samples show a fingerprint of Ti-O and Ti-O-Ti bonds between 500 cm⁻¹ and 750 cm⁻¹ which is a molecular fingerprint of BaTiO₃. The wavenumber of the absorption peak of Ti-O bond increases after doping, the peak is shifted from 481.12 cm⁻¹ in pure BaTiO₃ to 491.23 cm⁻¹ in 3% Fe doped BaTiO₃ which indicates the change of the unit cell size. Moreover, the bands observed in the region from 1425 cm⁻¹ to 1449 cm⁻¹ could be attributed to BaCO₃ phase present in the prepared samples [47].

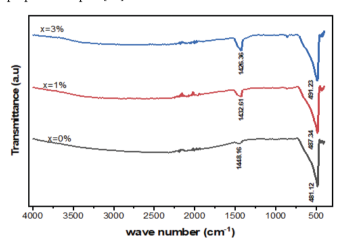


Fig. 3 FT-IR spectrums of pure and Fe-doped barium titanate at different Concentrations (x=0, 0.01 and 0.03)

3. ábra A tiszta és a Fe-adalékolt bárium-titanát FT-IR spektrumai különböző koncentráció esetén (x=0; 0,01 és 0,03)

х	a (Å)	c (Å)	V (ų)		Crystallite size (nm)	
0	3.9943	4.0191	64.12245	31.31533	34.56	0.003883
0.01	3.9949	4.0164	64.01643	31.36792	36.38	0.003584
0.03	4.0013	4.0167	64.30898	31.33112	42.46	0.003159

Table 1 The position of the peak (101), lattice parameters, Volume of the unit cell, lattice strain and crystallite size of BaFe $_x$ TiO $_3$ (x=0, 1 and 3%) nanoceramics 1. táblázat A csúcs helyzete (101), a rácsparaméterek, az egységcellák térfogata, a rácsos törzs és a BaFe $_x$ TiO $_3$ (x=0; 1 és 3%) nanokerámia kristálymérete

4. Conclusion

To summarize, pure and Fe doped barium titanate nanoceramics were synthesized using sol-gel method, the obtained powders were calcined at 800 °C for 3h, the calcined samples were characterized using XRD and FT-IR spectroscopy. The XRD results confirm the presence of single tetragonal phase for both pure and Fe doped materials. This measurement reveals that the Fe²⁺ ion may replace Ba²⁺ for x=1% or the both Ba²⁺ and Ti⁴⁺ but with a predominance for the Ti-site in the case of x=3%. FT-IR measurements showed that the elaborated samples have perovskite structure, which

is in a good agreement with XRD results. The modification in the structural properties after doping indicates that Fe has successfully doped into BaTiO, lattice.

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Effect of H₂ on SiO and SiC formation

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Abstract

Based on previous research, hydrogen has an enhancing effect on the kinetics of the Si-O-C system [1-3]. In this study silicon carbide (SiC) formation, from a reaction between SiO(g) and carbon, were investigated in a hydrogen atmosphere, an argon atmosphere, and an argon atmosphere containing 10% methane, at temperatures between 1495 °C and 1695 °C. The SiO(g) was generated from pellets comprised of a 2:1 ratio of silica (SiO $_2$) and SiC. The SiO(g) generation was monitored through analysis of the CO(g) content of the off-gas, the results indicated that the hydrogen atmosphere had an enhancing effect on SiO(g) generation through this reaction. Samples were retrieved from various locations within the set-up: the crucible, the reaction chamber roof, and from the condensation chamber. The reaction products were imaged in SEM using secondary electrons.

SiC was found growing throughout the reaction chamber, when using an argon atmosphere, a layer of SiC was found covering the graphite parts, whereas utilising a hydrogen atmosphere, or argon with 10% methane, the SiC would instead grow as whiskers. Thermal decomposition of methane is expected to be a challenge in this temperature range [4].

Keywords: silicon, SiC, hydrogen, methane, whiskers Kulcsszavak: szilícium, SiC, hidrogén, metán, whisker

1. Introduction

Silicon is currently produced in a submerged arc furnace from a reaction between SiO₂ and carbon. SiO₂ is supplied from high purity quartz, and commonly used carbon sources include: wood chips, charcoal or coal [5]. One of the most important intermediate species occurring during silicon production is SiO(g). It is formed in the hot lower region of the furnace from a reaction between SiO₂ and silicon carbide (SiC), reaction (1), and is consumed in the cooler upper region where it reacts with the carbon in the charge to form SiC, reaction (2). When SiO(g) is cooled down it will form Si and SiO₂ according to reaction (3), it may also form SiO₂ and SiC according to reaction (4). These reactions are commonly called the condensation reactions [5-6].

$$2 SiO_2 + SiC = 3 SiO(g) + CO(g) SiO(g) + 2 C = SiC + CO(g)$$
(1)

$$2 SiO(g) = 2 SiO_2 + Si 2SiO(g) + 2 CO(g) = SiO_2 + SiC$$
(2)

$$2 SiO(g) = 2 SiO_2 + Si$$
(3)

$$2 SiO(g) + 2 CO(g) = SiO_2 + SiC$$
(4)

Producing silicon consumes large amounts of energy. It is supplied, in part, from electricity, and in part from solid carbon materials. Finding a replacement to the carbon materials could lead to significant reductions in CO_2 emissions. Thus far there has been some research into possible alternatives to carbon for reducing metal oxides [7-9]. Most of the work has been on the use of methane hydrogen mixtures [10-11]. Despite the fact that $\mathrm{CH}_4(\mathbf{g})$ contains carbon, which will lead to $\mathrm{CO}_2(\mathbf{g})$ emissions, it may allow a more efficient process to be designed, or if bio-gas was utilised the net- $\mathrm{CO}_2(\mathbf{g})$ emissions would also be lowered. While some metals, such as iron, may be readily reduced to its

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metallic form by hydrogen, silicon has a much stronger affinity to oxygen making the reduction process more challenging [9]. One of the key challenges that face the use of CH₂(g) in silicon production is that the gas is unstable at high temperatures, and will undergo thermal cracking to H₂(g) and carbon according to reaction [4]. According to its thermodynamic equilibrium, CH₄(g) will be almost completely transformed to carbon and H₂(g). However, literature reports that methane cracking often occurs far away from its thermodynamic equilibrium if it does not happen on a suitable catalyst [12]. Furthermore, the deposited carbon is a poor catalyst for this reaction, so as CH₄(g) cracks it will form a layer which limits further cracking [4], [13]. Similarly, SiO(g) will form a condensate at low temperatures. Thus, there is a large temperature gap between the regions where each of the gasses are stable. Therefore, it is important to research how far away from their respective equilibrium pressures it is possible to go.

$$CH_{4}(g) = C + 2 H_{2}(g)$$
 (5)

This study aims to investigate the reaction between SiO(g) and carbon, and how it is influenced by different hydrogen containing atmospheres. In addition, the extent of thermal cracking of $CH_4(g)$ will be examined, and whether it is possible to have $CH_4(g)$ and SiO(g) react.

2. Experimental

SiO(g) and CO(g) were produced from pellets made from pure quartz and pure SiC which was crushed to a size of 5 mm. The quartz and SiC was pelletized and mixed with a molar ratio of 2:1, which follows the stoichiometry of reaction (1). A schematic drawing of the set-up is shown in Fig. 1. It consists of two main parts: a reaction chamber which is the bottom section, and a condensation chamber which is the top section. The reaction chamber contained a graphite crucible, which was filled with approximately 5 g of pellets before each experiment. An alumina lance was inserted into the reaction chamber from the top of the set-up. The condensation chamber consisted of a graphite tube surrounded by graphite wool insulation. While preparing the experiment, the alumina tube was placed through the middle of the condensation chamber in such a way that it protruded 2.5 cm into the reaction chamber. Additionally, a C-type thermocouple was passed through the lance so that its tip was in the center of the reaction chamber. After inserting the lance, SiC particles were poured into the space between the lance and the condensation chamber wall, which were added to collect reaction products.

After the set-up had been prepared according the description in the previous paragraphs, it was placed into the furnace. The furnace was a resistance heated vertical tube furnace. The off-gas from the furnace went through a gas analyser which detected the CO(g) content. First, the furnace was flushed with argon and evacuated down to 180 mbar, then it was refilled with argon as the temperature program commenced.

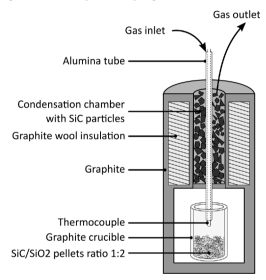


Fig. 1 Set-up used to investigate the reaction between the SiO(g) and CO(g), and carbon in an Ar(g) and an $H_2(g)$ atmosphere

1. ábra A SiO(g) és CO(g), valamint a szén reakciója Ar(g) és $H_2(g)$ atmoszférában

The temperature measured within the reaction chamber was observed to be lower than the furnace set-point, which was 1750°C, 1650°C or 1550°C. Fig. 1 shows the temperature measurements from the reaction chamber. It shows that the real temperature in the reaction chamber is around 55°C lower than the set-point of the furnace, the thermocouple is measuring the temperature in the centre of the reaction chamber. The parameters which were varied between experimental runs were the holding temperature, which was either 1595°C, 1695°C, or 1495°C and the process gas mixture which was a $\rm H_2(g)/\rm CH_4(g)$ gas mix that was varied between 0% $\rm CH_4(g)$ and 10% $\rm CH_4(g)$. The experimental matrix displayed in Table 1 shows the number of experiments performed for each combination of parameters. After each experiment, samples

were taken from various regions of the set-up. They were taken from the crucible rim, the ceiling of the reaction chamber, the graphite ring on the bottom of the condensation chamber, the alumina lance, and SiC chunks in the condensation chamber. The samples were placed into a SEM and imaged with a secondary electron detector.

		Methane content in a CH4(g)/ Ar gas mix		
	Pure H ₂ (g)	0%	10%	
1695°C	2	1	0	
1595°C	2	1	1	
1495°C	0	1	0	

Table 1 Experimental matrix, the number in each cell indicates number of parallels 1. táblázat Kísérleti mátrix, az egyes cellákban szereplő számok a párhuzamok számát jelzik

3. Results/Discussion

3.1 Generation of SiO(g) from the raw material

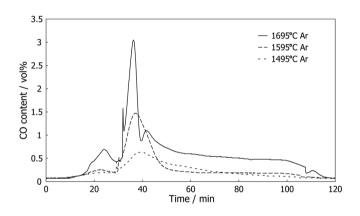


Fig. 2 CO(g) content of the off-gas for experiments in an Ar atmosphere at various temperatures

2. ábra A kiáramló gáz CO(g)-tartalmának vizsgálata Ar atmoszférában, különböző hőmérsékleteken

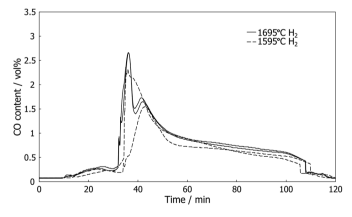


Fig. 3 CO(g) content of the off-gas for experiments in an $H_2(g)$ atmosphere at various temperatures

3. ábra A kiáramló gáz CO(g)-tartalmának vizsgálata H₂(g) atmoszférában, különböző hőmérsékleteken

Isothermal experiments in an argon/CH₄(g)/H₂(g) atmosphere were performed. The reaction was monitored through measurement of the CO content of the off-gas. *Fig.* 2 shows the off-gas measurements for the experiments in an

argon atmosphere at 1495°C, 1595°C and 1695°C. *Fig. 3* shows the off-gas measurements for the experiments in a $\rm H_2(g)$ atmosphere at 1595°C and 1695°C.

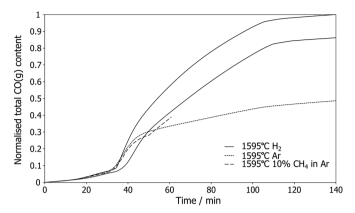


Fig. 4 Normalised integral CO(g) formation during experiments in pure $H_2(g)$, Ar(g) and Ar with 10% $CH_2(g)$, at 1595°C

4. ábra Normál integrált CO(g) képződés vizsgálata tiszta H₂(g), Ar(g) és Ar 10% CH₂(g) összetételű atmoszférában 1595°C-on

Fig. 4 shows a comparison of the normalised integral CO(g) content of the off-gas, with $H_2(g)$, Ar, and Ar with 10% $CH_4(g)$, at 1595°C. The most obvious trend of the off-gas CO(g) content is that it increases with temperature, at least in an Ar(g) atmosphere. Which is an indication of the raw material being converted to SiO(g) and CO(g) at a more rapid pace. However, when the atmosphere is exchanged with pure $H_2(g)$, the effect of temperature is not so obvious. Furthermore, after the initial peak in the CO(g) content, the curves follow each other closely for the rest of the temperature program. Thus, as a result of using an $H_2(g)$ atmosphere, the SiO(g) content in the reaction chamber is increased.

The weight of the pellets which were recorded before and after each experiment are consistent with what was seen in the CO(g) content of the off gas. *Table 2* shows what weight change was seen at different temperatures and atmospheres, evident that much more of the pellets were transformed to SiO(g) in an H₂(g) atmosphere than in Ar(g).

Temperature	Mass change in Ar(g)	Mass change in H ₂ (g)
1695°C		-57.9%
1595°C	-14.4%	-45.7%
1495°C	-12.4%	-

Table 2 Mass change of the raw material for each experiment 2. táblázat A nyersanyag tömegváltozása a kísérleteknél

3.2 Formation of SiC

SiC crystals could be seen growing in the rim of the graphite crucible. At the cooler temperature, the crystals are much smaller and large ones are more sparsely distributed on the surface. In an $H_2(g)$, or Ar(g) with 10% $CH_4(g)$ atmosphere the surface of the crucible is covered in SiC whiskers. Growth of SiC whiskers in $H_2(g)$ containing atmospheres has been seen previously. The most common explanation for growth of SiC in $H_2(g)$ containing atmospheres is that presence of hydrogen allows $CH_4(g)$ to form in small quantities. This would allow for an enhanced gas phase transfer of carbon which is necessary to

grow long SiC whiskers [1], [14]. Improved diffusion in $H_2(g)$ may also contribute. Reaction (6) and (7) show how $CH_4(g)$ might provide a means for gas phase carbon transfer to the whisker tips.

$$C + 2 H2(g) = CH4(g)$$
 (6)

$$SiO(g) + CH_4(g) = SiC(s) + CO + 2H_2(g)$$
 (7)

In Si containing systems the growth mechanisms of whiskers most often cited are vapour-liquid-solid (VLS), where there is a liquid impurity that acts as a catalyst to the whisker growth, vapour-solid (VS) where there is no catalyst, and oxide assisted growth (OAG) where there is layer of molten SiO_2 around the tip [15]. However, regardless of which of these growth mechanisms are present in this work, all of them require gas phase transfer of carbon from the graphite parts to the whisker tip. The two options are through $\mathrm{CO}(g)/\mathrm{CO}_2(g)$ or $\mathrm{H}_2(g)/\mathrm{CH}_4(g)$. Considering that the equilibrium $\mathrm{CH}_4(g)$ pressure is larger by a factor of 1000, compared to $\mathrm{CO}_2(g)$, in this temperature range $\mathrm{H}_2(g)/\mathrm{CH}_4(g)$ is expected to play a bigger role [16-17].

4. Conclusions

The $H_2(g)$ atmosphere had an enhancing effect on the SiO(g) formation in the raw materials. Additionally, the SiO(g) formation in an $H_2(g)$ atmosphere was much less affected by temperature than in a pure Ar(g) atmosphere.

SiC formation in the current experimental set-up was enhanced in an $H_2(g)$ atmosphere compared to an Ar(g) atmosphere. There are a few possible explanations, the presence of $H_2(g)$ allowed for a kinetically more favoured reaction path to SiC formation. The increased SiO(g) formation leads to a higher SiO(g) content throughout the reaction chamber which would increase SiC formation. Faster diffusion in the $H_2(g)$ atmosphere may also play a role in the improved kinetics. Furthermore, the SiC formed in an $H_2(g)$ atmosphere grew as whiskers, while in an Ar(g) atmosphere it grew as a layer covering the graphite. The difference is suspected to be caused by the formation of $CH_4(g)$ from the $H_2(g)$ reacting carbon, and the $CH_4(g)$ allows for fast gas phase carbon transfer to the whisker tip.

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2020 The 3rd International Conference on Composite Materials Science and Technology (ICCMST 2020) will be held in Cavtat, Croatia, during June 10-14, 2020. It is a great pleasure for ICCMST to invite prospective authors initiating the discussion on the challenges that need to be timely overcome and addressing key questions in the field of Composite Materials Science and Technology.

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Thermal stability and flame retardant properties of plasticized poly(vinyl chloride) hybrid composite for construction application

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Abstract

Current research is investigating how to improve constructions efficiency and at the same time increase their fire resistance. Oxydtron is used as admixture for concrete to improve its properties, but Oxydtron has shown unpredictable behavior when used as filler for plasticised PVC where the flame retardation and stabilization at high temperatures of the polymer was improved after Oxydtron additions. Thermal tests were carried out to demonstrate the ability of Oxydtron to improve thermal resistance of the plasticised PVC and these tests included: Limiting oxygen index, Dehydrochlorination, and Dynamic mechanical analysis. The results of these tests proved the efficiency of Oxydtron as effective filler for plasticised PVC by increasing the flame retardation and thermal stability.

Keywords: thermal stabilizing, flammability, oxydtron, plasticized poly(vinyl chloride) Kulcsszavak: hőstabilitás, éghetőség, oxydtron, lágyított PVC

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1. Introduction

When Prometheus stolen the fire from the Hephaestus god of blacksmiths in Mount Olympus according to the Greek mythology and gave it to humans to help them overcome the difficulties of their daily lives related to heating, lighting and cooking; he didn't realize that fire would be useful to humans but also would bring destruction to them if they didn't use it safely. So, they have tried to find the methods to extinguish fire. Then they thought of something entirely new: creating materials that could resist fires reduce their risk and even completely stop it, thus, the concept of flame retreading appeared for the first time; and the subsequent discovery of materials (flame retardants) have the ability to break the cycle of fire [1,2]. As the concept of flame retardation developed, it was necessary to develop the way of thinking in this field; now, the search for new and unconventional materials in industry is imperative to balance environmental considerations for the safe use of non-polluting materials (or at least with limited damage) and to maintain the best possible achievement of new materials compared to traditional materials. Where the purpose is not only to use new engineering materials in the industry, but to be more efficient or at least maintain the same level of performance of the traditional materials currently available which can be termed metaphorically - the sustainability of properties - as with the sustainability of natural materials, otherwise there will be no benefit from Use of such materials [3-6]. Every year, fires cause a significant human and financial losses in all countries, and the proportion of these losses varies from one country to another depending on the equipment and techniques used to reduce the fires risk, as shown in *Table 1* and Fig. 1, which are represented the economic-statistical evaluation of fire costs according to the center of fire statistics report issued in the year 2016 [7]. As shown in *Table 1*. Many countries still don't attach much importance to fire protection, but spend much of their gross domestic product (GDP) to extinguish the fires after ignite, but it is best to be concerned with how to prevent the fire from happening.

In general, the principle of reducing fire rates depends on the provision of conventional fire extinguishing equipment; use of materials with low flammability ranges or improve their thermal resistance if they are highly flammable such as polymers; appropriate design of the buildings to provide outlets to escape when the fire to reduce the loss of human, and all the above points are relevant to the important factor which is the deployment of a "fire prevention culture" that is fully dependent on people's awareness of fire hazards and the extent to which they apply safety arrangements to prevent fires or reduce it [8,9]. Poly(vinyl chloride) is one of the most important types of polymers used today in the plastics industry and has maintained its position in this industry despite its long discovery and the emergence of many new polymers due to its distinctive properties. Poly(vinyl chloride) can be produced in more than one state, which is rigid and flexible. The important thing is that changing poly(vinyl chloride) from rigid to flexible state there's no need to change its chemical formula through polymerization but through additives only, which is not available in the majority of other polymers. These additives improve many of the polymers properties in general, and one of these properties is thermal resistance, where all polymers not just poly(vinyl chloride) have low thermal resistance to high temperatures which considered an obstacle to the use of such materials in applications requiring high thermal resistance. Therefore, there is a high risk of using them in civil and industrial applications without increasing their resistance to combustion [3,10-20].

N	Country			Cost in porti	on of GDP, %		5	Expenditure
		Direct losses	Indirect losses	Cost of fire service	Fire protection in buildings	Fire insurance administrations	$\sum c_i$	losses C3 + C4 + C5
		$\mathbf{C_{_1}}$	C ₂	C ₃	C ₄	C ₅	i=1	C1 + C2
1	Australia	0.07	-	0.17	-	-	-	-
2	Czech Republic	0.07	-	-	0.16	-	-	-
3	Finland	0.17	0.011	0.19	-	0.03	-	-
4	France	0.20	-	=	-	-	-	-
5	Germany	0.12	0.014	-	-	-	-	-
6	Hungary	0.02	-	0.13	-	-	-	-
_ 7	Italy	0.02	-	=	0.35	0.04	-	
8	Japan	0.12	0.060	0.26	0.12	0.09	0.650	2.61
9	Netherlands	0.15	-	0.21	0.31	-	-	-
10	New Zealand	0.12	-	0.16	0.24	-	-	-
11	Poland	0.09	-	0.16	-	-	-	-
12	Singapore	0.04	0.027	0.03	0.40	0.02	0.517	6.72
13	Spain	0.08	-	-	-	-	-	-
14	Sweden	0.18	0.060	0.13	0.20	0.05	0.620	1.58
15	UK	0.13	0.008	0.20	0.29	0.10	0.728	3.84
16	USA	0.10	0.007	0.29	0.29	0.12	0.807	6.54
	Avrg.	0.12	0.027	0.18	0.26	0.05	0.652	3.40

Table 1 Economic-statistical evaluation of fire costs [7] 1. táblázat Tűzesetek költségeinek gazdasági elemzése [7]

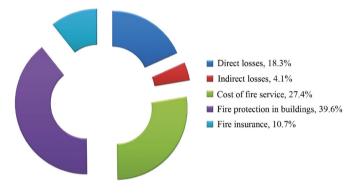


Fig. 1 Economic-statistical evaluation graph of fire costs [7] 1. ábra A tűzesetek költségeinek gazdasági elemzése [7]

Oxydtron one of the non-traditional additive that we used in this study. Those who hear about Oxydtron for the first time are thought to contain nanoparticles as indicated by the name but this is not true where the cement is a micro particles powder (about 2-3 μm) and just the particle's coating layer which consists of modifier materials is nano size (from 10-100 nm); which gives this material special characteristics in addition to its low cost. Oxydtron contains many of the high-temperature oxides and carbonates that make it one of the most important and alternative options for conventional retardants used today [21-23].

2. Methodology

2.1 Materials

The raw materials used for producing tests samples illustrated in *Table 2* which can be described as follows: (1) PVC S-5070 powder (under trademark Ongrovil*) which produced and supplied by BorsodChem Zrt., Hungary; (2) the plasticizer DOP (Di-Octyl Phthalate) supplied by DEZA, a. s. CO.,

Valašské Meziříčí, Czech Republic; (3) Calcium-Zinc based stabilizer (under trademark Newstab-50) which supplied by Betaquímica CO., Barcelona, Spain; (4) the lubricant Wax-E (under trademark Licowax*E) supplied by Clariant International Ltd, Muttenz Switzerland; and (5) Oxydtron (NC) (under trademark Oxydtron-Oxydtron type A) supplied by Bioekotech Hungary Kft.

Component	PVC	basic	PVC modifier		
	PVC S-5070	DOP	Newstab-50	Wax-E	Oxydtron
pphr	100	70	1.5	0.3	0-5 (wt.%)

Table 2 Raw materials for producing test samples and their dosage 2. táblázat Próbatestek készítéséhez használt alapanyagok és adagolásuk

2.2 Materials processing

The main mixing process involved mixing all the raw materials listed in *Table 2*, which accomplished using a mechanical mixer type MTI 10 Mischtechnik at 150 °C; in order to obtain homogeneity in the mixture. After the main mixing process, the Oxydtron is added with the weight fraction (0, 1, 3, and 5) wt.%. The new hybrid blend has been mixed by small electric mixer to provide homogeneous distribution of Oxydtron within the poly(vinyl chloride) powder.

2.3 Samples fabrication

1. Limiting oxygen index samples: L.O.I samples were produced according to ISO 4589-2 standard [24] by using also laboratory two roll mill type Schwabenthan at the same processing conditions. A sheets with 0.4 -0.6 mm thickness made by rolling and then these sheets compressed by a hydraulic press type Bürkle at 300 and 20 bar pressure and 175 °C to the desired L.O.I sample shape.

- 2. Dehydrochlorination test: a particle of PVC and Oxydtron additives with the same dimensions of congo-red test was used in this test at 170 °C and according to ISO 182-3:1993 standard [25].
- 3. Dynamic mechanical analysis test: ISO 6721-11:2012 standard [26] was used to produce the samples. The DMA test samples were a strips fabricated by using extrusion machine type GÖTTFERT at 170 °C temperature and 60 rpm speed at uniform conditions (pressure, temperature, and compression).

2.4 The tests

- 1. Limiting oxygen index test: This test was done at BorsodChem Zrt., Laboratory of Vinyl Technology by using Stanton Redcroft FTA flammability unit.
- 2. Dehydrochlorination test: Dehydrochlorination ranges have been measured by Metrohm 763 Thermomat device found also at BorsodChem Zrt., Hungary.
- 3. Dynamic mechanical analysis test: Dynamic mechanical thermal analyser MK III manufactured by Rheometric Scientific, Inc. and found at BorsodChem Zrt., Hungary, used for testing the samples. The temperature range is -60 °C to 120 °C.
- 4. Scanning electron microscopy (SEM): In order to analyse the elemental composition and structure of Oxydtron, the Carl Zeiss EVO MA10 SEM was used for performed this analysis as shown in *Fig. 2* and *Table 3*. SEM device found at Institute of Physical Metallurgy, Metal Forming and Nanotechnology, University of Miskolc, Hungary.
- Oxides analysis: ICP-AES spectrometer was used for the analysis of Oxydtron components. The results of the analysis are listed in *Table 4*. ICP-AES spectrometer found at Institute of Chemistry, University of Miskolc, Hungary.

Element	Wt.%	At %	Net Inte.	Bkgd Inte.	Inte. Error	P/B
С	7.57	14.33	22.07	1.00	4.06	22.07
0	34.78	49.43	121.40	1.50	1.68	80.93
Na	0.64	0.63	6.53	3.87	10.56	1.69
Mg	0.77	0.72	11.23	5.73	7.74	1.96
Al	2.09	1.76	36.63	7.03	3.55	5.21
Si	10.45	8.46	195.23	9.40	1.37	20.77
S	1.48	1.05	27.37	9.90	4.58	2.76
K	1.08	0.63	18.13	8.00	5.88	2.27
Ca	38.85	22.04	563.83	7.70	0.78	73.23
Ti	0.17	0.08	1.70	6.00	39.75	0.28
Fe	2.12	0.86	12.63	3.67	6.46	3.45

Table 3 EDAX- SEM analyses for structural composition of Oxydtron 3. táblázat Az Oxydtron osszetételének SEM – EDAX analízise

Oxides	Ratio, wt.%
Al_2O_3	4.47
Ca0	58.0
Cr ₂ O ₃	0.006
Fe ₂ O ₃	2.67
K ₂ 0	0.78
MgO	1.20
Mn_2O_3	0.05
Na ₂ O	0.31
SO ₃	2.50
SiO ₂	21.44
TiO ₂	0.274
Zn0	0.138

Table 4 Chemical composition of Oxydtron analyzed by ICP-AES
4. táblázat Az Oxidtron kémiai osszetételének meghatározása ICP-AES módszerrel

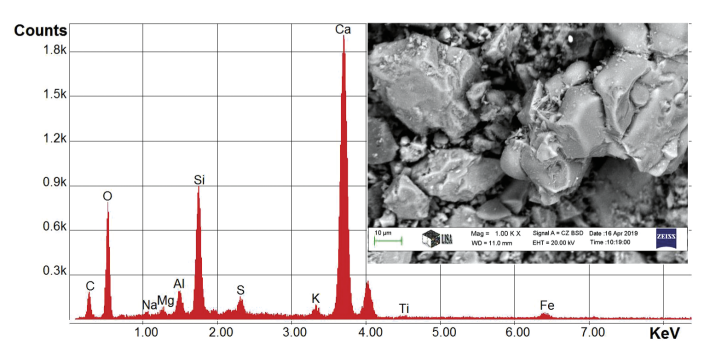


Fig. 2 SEM-energy dispersive X-ray microanalysis for Oxydtron 2. ábra Az Oxydtron SEM energiadiszperzív Röntgen mikroanalízise

4. Results and descussion

Flame retarding results for plasticised PVC containing Oxydtron obtained by limiting oxygen index (L.O.I) test shown in Fig 3. From the first bar of this figure which represents the L.O.I. result of plasticised PVC basic formulation that the value of L.O.I is low (21.6%). The low flame resistance considered a normal state for plastics and this low resistance because of the crystalline structure of these materials (especially organic) therefore it will ignite easily and rapidly when exposed to direct fire [1,3]. And in order to modify this critical behavior of plastics at high temperatures (PVC in this study), the additives are added to increase their thermal resistance. These additives can be used as fillers or coating layers [27,28]. After adding 1% Oxydtron as a new flame retardant additive for plasticised PVC fire resistance of the polymer improves as shown in the second bar of Fig. 3 where the L.O.I value has been increased to (22.3%). The flame resistance of plasticised PVC is increasingly rising with increasing Oxydtron additives quantity which can be observed in the third bar of Fig. 3 which represents the L.O.I value of plasticised PVC containing 3% Oxydtron where the L.O.I value reached to (24.8%), while it will reach a value of (26.4%) as shown in the fourth bar which represents L.O.I value of plasticised PVC containing 5% Oxydtron. This unexpected behavior of Oxydtron stabilizing and flame retarding is due to Oxydtron composition which consists of many compounds such as calcium carbonate (CaCO₂) which considered a universal stabilizer for PVC and also silicon dioxide (SiO₂) in addition to many other compounds, and these two materials (CaCO₃ and SiO₂) will act as a synergist agent with the Newstab-50 (original stabilizer) and creating additional stabilizing synergistic action causing increased thermal resistance and stability of plasticised PVC. Also, the calcium carbonate considered a flame retardant agent and its presence within the Oxydtron composition will help to increase the flame resistance of PVC.

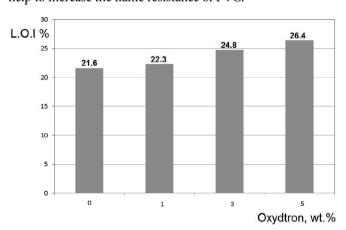


Fig. 3 Limiting oxygen index (L.O.I) values for plasticised PVC-Oxydtron hybrid

3. ábra Az Öxidtron tartalmú lágy PVC keverékek oxygen indexe (LOI)

The crystalline structure of PVC tends to release chlorine containing compounds strongly at high temperatures, as shown in *Fig. 4*, which represents a dehydrochlorination test for plasticised PVC basic formulation at 170 °C; where the chlorine containing compounds loss is very rapid and the slope of the curve is very sharp (high inclination). This is a normal

behavioure for PVC due to the low thermal resistance in high temperatures (for polymers) and this resistance decreases as the release of chlorine containing compounds increases where the crystalline structure of PVC is strongly deformed; and the bonds between atoms begin to break down and huge amounts of chlorine containing compounds are released [29]. This is observed from the high slope of the curve (0.000134) and at this point, the PVC reached the degree of total degradation.

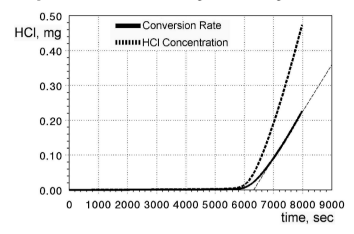


Fig. 4 Dehydrochlorination test of plasticised PVC basic formulation at 170 °C 4. ábra A PVC alapreceptúra sósavlehasadás vizsgálata 170 °C-on

Plasticised PVC becomes more resistant to high temperatures after of Oxydtron addition, as shown in Fig. 5 which represents the dehydrochlorination results of plasticised PVC containing 1% Oxydtron at 170 °C; where the slope of the conversion rate curve decreased from (0.000134) in case of plasticised PVC basic formulation (without additive) to (0.000073) after Oxydtron addition, where the tendency of plasticised PVC to release chlorine containing compounds has been reduced and increased thermal stability and degradation resistance accompanied by extended degradation time range for plasticised PVC basic formulation from (2.21 hours) to (4.75 hours) after 1% Oxydtron addition. When the weight fraction of Oxydtron additives increases the thermal resistance stability of plasticised PVC will rise and this behavior can be distinguished by the Fig. 6 and Fig. 7 which represent the dehydrochlorination results of plasticised PVC containing 3% and 5% Oxydtron at 170 °C respectively. From these figures, we notice that the slope of conversion rate curve will decrease, i.e. the degradation resistance has been increased, where in case of of 3% Oxydtron content the slope of conversion rate curve becomes (0.000062) with an increase in the total degradation time to (4.84 hours), and in case of of 5% Oxydtron content the slope of conversion rate curve will reduce to (0.000053) and increasing the total degradation time to (5.20 hours).

From Fig. 8 which represent the Dynamic Mechanical Analysis (DMA) for plasticised PVC-Oxydtron hybrid composite, we can note that the Oxydtron additives increase the glass transition temperature (Tg), where the glass transition temperature shifts to higher temperatures rising from (-3.26 °C) in case of plasticised PVC basic formulation to (0.30 °C) when 1% of Oxydtron has been added and continues to rise to (3.50 °C) and (4.66 °C) when added 3% and 5% of Oxydtron respectively. Clearly, there is an interaction between

PVC and Oxydtron molecules, which causes the reduction of kinetic energy of plasticised PVC, and reducing the segmental movement, which probably prevents the molecules from assembling in crystalline systems, so the relaxation time of this movement increases. So the polymer will consist of long physically crosslinked liquid-like molecules [30].

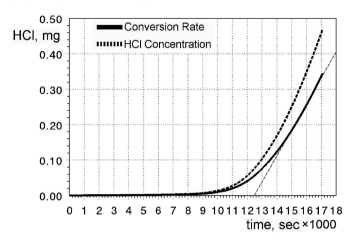


Fig. 5 Dehydrochlorination results of plasticised PVC containing 1% Oxydtron at 170 °C 5. ábra Az 1% Oxidtron tartalmú lágy PVC sósavlehasadás vizsgálata 170 °C-on

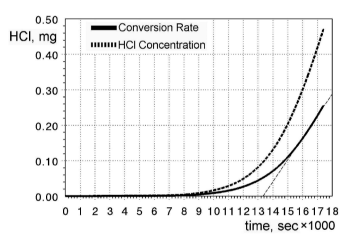


Fig. 6 Dehydrochlorination results of plasticised PVC containing 3% Oxydtron at 170 °C 6. ábra A 3% Oxidtron tartalmú lágy PVC sósavlehasadás vizsgálata 170 °C-on

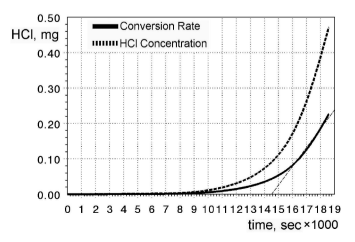


Fig. 7 Dehydrochlorination results of plasticised PVC containing 5% Oxydtron at 170 °C 7. ábra Az 5% Oxidtron tartalmú lágy PVC sósavlehasadás vizsgálata 170 °C-on

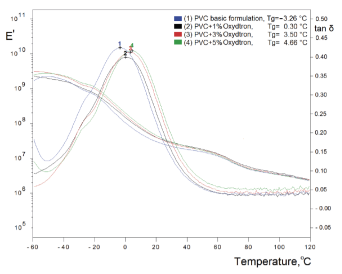


Fig. 8 Dynamic mechanical analysis for plasticised PVC-Oxydtron hybrid composite 8. ábra A lágy PVC - Oxidtron hybrid kompozitok dinamikus mechanikai analízise

5. Conclusions

In general, Oxydtron improves the thermal properties of plasticised PVC, where the enhancement of L.O.I value reached to (22.22%) at 5% Oxydtron content. The degradation rate of plasticised PVC also decreased after adding Oxydtron, where the improvement of time required for total degradation with plasticised PVC containing 5% Oxydtron was (135.30%) longer than plasticised PVC basic formulation. Glass transition temperature was shifted towards higher temperatures after Oxydtron addition, and this indicates that Oxydtron directly affects kinetic energy and segmental movement of plasticised PVC causing to decrease them.

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Rheology of natural hydraulic lime pastes modified by non-traditional biopolymeric admixtures

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Abstract

Viscosity enhancing admixtures, widely used to improve characteristics of concrete and ready-mix mortars, are mainly different derivatives of cellulose. Due to the nature of cellulose processing, the environmental-friendlier alternatives should be studied in order to reduce the impact of the building industry on the environment. The rheological study of natural hydraulic lime (NHL) grouts modified by four different biopolymers is carried out to investigate their behaviour in the NHL-based mortars.

The biopolymers studied are of seaweed (sodium salt of alginic acid (ALGNA) and carrageenan (CG)) and microbial (diutan gum (DG) and xanthan gum (XG)) origin. The effect of addition of these admixtures in the doses of 0.1%, 0.5%, and 1% was studied using hybrid rheometer with DIN concentric cylinders geometry. The flow properties as well as viscoelastic properties were studied. The addition of any of the admixtures led to the increase in yield stress, with DG being the most effective admixture. Desirable increase in consistency coefficient was observed within the pastes with CG and DG addition having growing dosage dependency, the ALGNA addition also increased the coefficient noticeably, but it was furtherly decreased with growing dose of admixture. The fluidity index lower than 1 expressed shear-thinning behaviour of studied pastes, except the pastes with highest dose of admixtures, and all of the XG pastes. The addition of CG and DG supported the stability of the grout expressed as the increase in critical strain, thus prolongation of linear viscoelastic region. The flow strain was increased by all of the studied admixtures promoting the gel-like behaviour of the pastes. Complex modulus and viscosity measured at 1Hz frequency were unaffected by the DG addition while they were increased notably by addition of other admixtures with ALGNA and XG supporting the resistance to deformation of the grouts studied. Correspondingly to complex modulus increase, the loss tangent is diminished, reporting more elastic behaviour of the material.

All of the admixtures studied increased the yield stress, and the influence of most of them had similar trends within other properties. Noticeable differences in efficiency and dosage-dependency were observed. The xanthan gum was overall the worst performing admixture. This was mainly due to higher sensitivity of xanthan to the concentration of bivalent ions in the solution.

Keywords: natural hydraulic lime, rheology, flow properties, viscoelastic properties, biopolymeric admixture

Kulcsszavak: természetes hidraulikus mész, reológia, folyási tulajdonságok, viszkoelasztikus tulajdonságok, biopolimer keverék

1. Introduction

The viscosity enhancing admixtures (VEAs) are used to modify the fresh state properties of building materials, mainly plasters and self-consolidating concrete (SCC) [1, 2]. The VEAs are often of polysaccharidic origin, currently most used are cellulose ethers, but a range of natural gums (e.g. welan gum, guar gum, gum arabic, etc.) is being studied. This study is focused on two biopolymers of microbial origin (xanthan (XG) and diutan gum (DG)), and two obtained from seaweeds (sodium salt of alginic acid (ALGNA) and carrageenan (CG)). These VEAs have been scarcely studied for the use in cementitious material, especially in SCC, and are mainly used in the food industry as gelling and thickening agents and stabilisers [2–4].

The microbial XG and DG are partially in use in building industry, where they reduce segregation and bleeding of SCC mixtures [2, 3, 5]. Xanthan gum, with slump-reducing side effect in utilization in SCC, proposes shear-thinning behaviour [6] in cementitious compositions [2, 7]. While used in the aerial lime-based [8, 9] mortars, the efficiency of XG addition was lower than the expectations based on the results from cement mixtures. This discrepancy is probably caused by a cationic sensitivity of XG, especially in the case of bivalent (e.g. Ca²⁺)

ions [10]. In the other types of materials, xanthan gum based

stabilizer has been successfully used to improve the stability of

fly ash-metakaolin geopolymer binary system [11]. The DG is

of similar use and properties to XG with even less sensitivity to

the conditions such as pH, temperature and salinity [10, 12].

DG is more efficient in the modifying of rheological properties

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of aqueous solution, thus having more dosage-dependent effect [10]. The improved stability of SCC by the DG addition is achieved by the increase in yield stress of the mixture [1]. The ALGNA, acquired from brown seaweed, has properties comparable with the commercially available super-absorbent polymers (SAP) [13]. The impure brown seaweed extract, containing 10-40% of dry weight of alginates was used by León-Martínez et al. as a VEA for SCC with interesting results, the properties of the extract are comparable with the pure sodium alginate, especially in higher concentrations, both, the sodium alginate as well as algae extract express the shearthinning behaviour of aqueous dispersions [14]. The addition of sodium alginate to either aerial or hydraulic lime mortar leads to the increase in yield stress and promotion of shearthickening behaviour of the pastes [8, 15]. Carrageenan (CG) is, as well as more famous gelling agent agar, obtained from red seaweed. There exists a wide range of carrageenan types with different degree of sulfatation, which influences mainly the solubility and gel strength [16]. The CG addition improves the mechanical properties of fly-ash based geopolymers, mainly by creating more condensed structure [17, 18]. The short-term strengths of lime mortars have not been notably improved in the case of hydraulic lime [19] and significantly decreased in the case of aerial lime [9], but with the difference diminishing with further ageing. The aerial lime pastes rheology is improved by increase in yield stress and consistency coefficient, and also by prolongation of period of gel-like behaviour observed by study of viscoelastic properties.

The presented paper studies the influence of the abovementioned biopolymers on the flow and viscoelastic properties of natural hydraulic lime (NHL) based grouts focusing on the dosage-dependency of the biopolymer addition.

2. Materials and methods

2.1 Materials and sample preparation

All the grouts studied were prepared by dry mixing of natural hydraulic lime of NHL 3.5 class according to EN 459-1 (Zement-und Kalkwerke Otterbein GmbH & Co. KG, Germany) and biopolymeric admixture (summary of admixtures used, and their characteristics are stated in *Table 1*) in the dosage of 0.1%, 0.5%, and 1% of the binder weight. Water was then added using constant water:binder ratio of 0.7 and the paste was prepared by mixing for one minute. The grout was then introduced into measuring cup of Discovery HR-1 hybrid rheometer by TA instruments using DIN concentric cylinders measuring geometry tempered to 20°C. The measurement was started by a 60 s preshear at 100 s⁻¹, which begun 5 minutes after water introduction into the dry mixture, followed by 60 s resting time. The curves were analysed using TRIOS 4.0.2.30774 software.

2.2 Methods of measurement

The measuring procedure has been described in detail in previous publication [8], thus only the basic description follows.

Abbrev.	Chemical composition	Commercial name	Viscosity (1% aq. solution, 20°C) [mPa s]	Manufacturer
ALGNA	sodium alginate	-	14.5	Sigma-Aldrich, Co.
CG	carrageenan	Genuvisco CG-131	72.1	CP Kelco
DG	diutan gum	Kelco-Crete DG-F	13 363.8	CP Kelco
XG	xanthan gum	Kelzan AP-AS	7 633.3	CP Kelco

Table 1 Abbreviations, characteristics, and producers of the admixtures used 1. táblázat Az alkalmazott adalékszerek jelölése, jellemzői és gyártói

2.2.1 Flow properties

The results obtained by procedure described in [8] were expressed graphically as flow curves (shear rate vs shear stress) and the downward curves were analysed using Hershel-Bulkley model (Eq. 1) to obtain the variables describing behaviour of the samples.

$$\tau = \tau_0 + k\dot{\gamma}^n \tag{1}$$

where τ - shear stress, $\dot{\gamma}$ - shear rate, τ_0 - yield stress, k consistency coefficient and n - fluidity index which characterizes shear-thinning (n < 1) or shear-thickening (n > 1) behaviour of a material.

2.2.2 Viscoelastic properties

To determine the length of linear viscoelastic region (LVR), characterized by the critical strain γ_C , and flow point γ_F , described as the equilibrium between loss (G') and storage (G') modulus, the small amplitude oscillation test at 1 Hz frequency was carried out. The strain value from the LVR of all samples was chosen to execute the frequency sweep test (0.1 Hz – 10 Hz) in linear conditions. The results were expressed as complex modulus G^* and loss tangent tan (δ) as defined by equations 2 and 3 respectively.

$$G^* = \sqrt{(G')^2 + (G'')^2} \tag{2}$$

$$\tan\left(\delta\right) = G''/G' \tag{3}$$

3. Results and discussion

3.1 Flow properties

The measurement data were expressed as shear rate vs shear stress diagram. For the illustrational purposes, the downward curves of pastes with ALGNA addition are shown in *Fig. 1*. The transition from pseudoplastic (shear-thinning, n < 1) to dilatant (shear-thickening, n > 1) behaviour is observed as a change of shape of curves from convex (REF, ALGNA 0.1% and 0.5%) to concave (ALGNA 1%). The values of yield stress (τ_0), consistency coefficient (k), and fluidity index (n) for all the pastes are compared in *Fig. 2*. The admixture addition increases the yield stress of pastes with further growth with increasing dosage, which is typical for the VEAs in traditional building materials [8, 9, 20–22]. The non-monotonicity of CG and XG addition was observed and described by Cappellari et al. [20] on the cementitious mortars modified by hydroxypropyl

guaran (HPG) and methylhydroxyethyl cellulose, and it is also reported on the NHL pastes modified by HPG with low degree of substitution and hydroxypropyl methyl cellulose [9]. The non-monotonicity is supposed to be caused by the competition between the lubricating and dispersing effect of the admixtures [20]. The most effective admixtures in the lowest dose are CG and XG, while DG is the most efficient admixture in higher doses. The increase in consistency coefficient (k) is connected with improvement of adhesive and anti-sagging properties of grouts. The reversed dosage-dependency of ALGNA is contrary to the aerial lime results [8], and also the inefficiency of XG is much clearer. The extreme drop of k value for CG and DG in highest dosage is probably caused by the high thickness of the tested samples. Lower doses of ALGNA and CG, as well as DG, may be potentially beneficial for the use to modify e.g. tile adhesives. The value of fluidity index, as stated earlier, describes whether the mixture's behaviour is shear-thinning or shear-thickening. The shear-thinning behaviour is more common and mostly desired in the case of building materials [8, 9, 15]. The transition from shear-thinning to shearthickening behaviour is due to water-retentive function of biopolymers [9] that causes a decrease in liquid phase content in the grout, which is basically a suspension of NHL particles in water/water-biopolymer environment, thus the paste draws to the state of "thick suspensions of particles in a liquid", which is the typical example of dilatant fluid [23]. The alginate and DG addition in lower dosage promotes the shear-thinning behaviour of the paste, while high doses of biopolymers and XG in general promote dilatancy of the pastes. A. Azzizi and P. F. G. Banfill [15] observed an opposite trend with the ALGNA addition but using much lower concentrations. Based on the comparison of the two results, it can be assumed that somewhere between the dosage of 0.035% (the highest dose in [15]) and 0.5% of binder weight lies a dose of alginate, at which the thixotropy of the NHL based paste is the most promoted. The studied biopolymers behave differently in comparison to hydroxypropyl derivatives of guar gum, chitosan or cellulose [24] (the same experimental setup on the same measuring device).

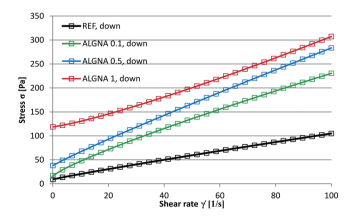
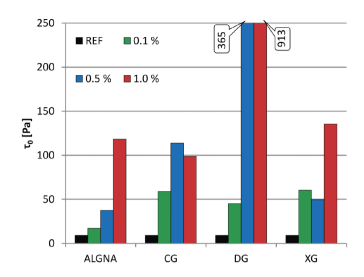
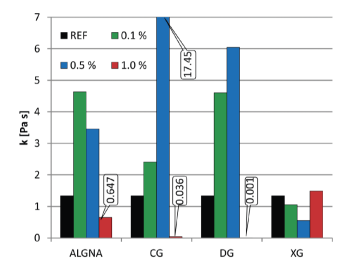


Fig. 1 Downward flow curves of NHL pastes modified by different amount of ALGNA

1. ábra Eltérő mennyiségű ALGNA-val módosított NHL pépek folyási görbéi





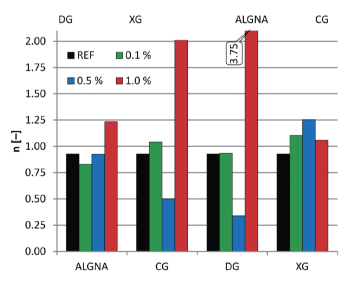


Fig. 2 Comparison of yield stress (τ_0) , consistency coefficient (k), and fluidity index (n) of NHL-based pastes modified by biopolymers

 ábra Biopolimerekkel módosított NHL alapú pépek folyási feszültsége (τ_o), konzisztencia együtthatója (k), and folyékonysági mutatója (n)

3.2 Viscoelastic properties

3.2.1 Amplitude sweep test

The critical strain γ_C determines the length of LVR, thus also the stability of the paste (the longer the LVR, the more stable the paste). The moduli curves in the modulus/oscillation strain system for ALGNA, alongside with the examples of the critical and flow strain values (γ_C and γ_F respectively) are shown in Fig. 3, while in Fig. 4, the strain values for all mixtures are compared. The addition of CG and DG in lowest dose do not affect the value of critical strain, which is furtherly increased with growing dosage of admixture, whereas the CG is more efficient in the dosage of 1%. This behaviour is similar to the one of hydroxypropyl derivatives of guaran, chitosan and cellulose [24], out of which cellulose ether are currently one of the most common VEAs used. The ALGNA decreases the value of γ_c thus the ALGNA-modified pastes behave differently than cementitious pastes modified by marine brown algae extract in which the alginates are present [14]. The XG slightly decreases grouts stability in low dosage and improves it with growing dosage, which is in correspondence with the behaviour of XG aqueous solution in different concentrations [6].

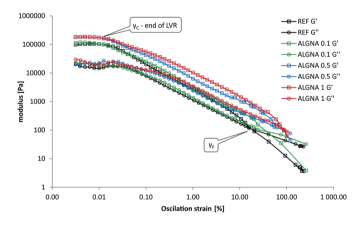
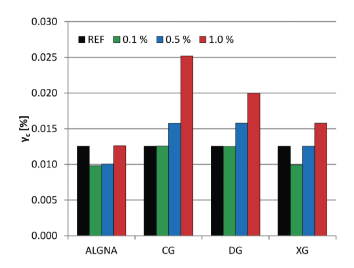


Fig. 3 Storage modulus (G') and Loss modulus (G") of pastes with ALGNA addition 3. ábra ALGNA hozzáadásával készült pépek tárolási (G') és veszteségi modulusa (G")

The second parameter studied, the flow strain γ_{ν} , marks the point of transition from solid viscoelastic behaviour (G' >G") to liquid behaviour (G' < G"). The addition of admixtures markedly increases the flow strain value as seen in Fig. 4. In the case of CG and DG, the highest increase is within the lowest dosage (0.1%) and with growing dosage, the flow strain value decreases. This trend is similar to hydroxypropylmethyl cellulose (HPMC) and hydroxypropyl guaran (HPG) [24] on the same instrumental setup but CG and DG seem to be more efficient. In the case of DG 0.1% the moduli did not intersect in the studied strain range, thus the paste retains viscoelastic properties. The growth of $\gamma_{\scriptscriptstyle F}$ for ALGNA up until dosage of 0.5% is in accordance with results of León-Martínez et al [14], where the increase in flow stress was observed. The XG is in efficiency of affecting the γ_n inferior to the three other admixtures with unclear dosage-dependency.



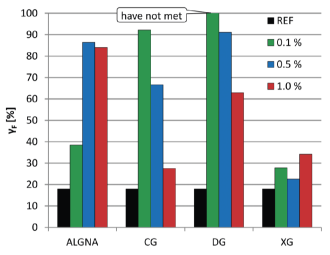


Fig. 4 Comparison of critical strain (γ_c) and flow strain (γ_p) values of tested samples
 4. ábra A vizsgált minták kritikus deformációja (γ_c) és folyási deformációjának (γ_p) összehasonlítása

3.2.2 Flow sweep test

The test was carried out at 0.005% strain, which ensures that all the pastes are in the LVR. The data obtained were visualised using charts similar to Fig. 5 and the values of complex shear modulus (G^*), and loss tangent (tan (δ)) at 1 Hz frequency were recorded for comparison and are presented in Fig. 6. The growth of resistance to deformation of the samples represented by increasing value of G^* in combination with decrease in loss tangent corresponds with the results of amplitude sweep tests. The values of tan (δ) indicate prevailing elastic behaviour of samples with storage modulus being 4-10 times higher than loss modulus. The extremely low tan (δ) value of DG 0.1 % may be interpreted as behaviour of ideally elastic material. The decrease in loss tangent with addition of VEA has been observed also with HPG and chitosan derivatives in lime mortars [25, 26], and hydroxypropyl derivatives of several biopolymers in NHL pastes [24]. On the other hand, HPG and HPMC in cementitious paste did not cause a decrease and starch ether in initial dose caused increase in loss tangent [20]. The gradual increase in tan (δ) with growing dosage, observed especially in the case of CG and DG where the trend corresponds with the growth of flow strain, is also described in aforementioned literature within different admixtures in several binder systems. The studied biopolymers, previously used in aerial-lime pastes [8], exhibited opposite trends in both binder systems, with exception of XG, where the trends are unclear in both cases.

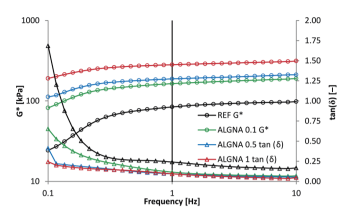
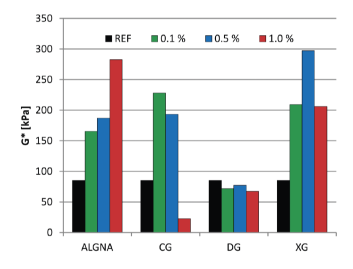


Fig. 5 Results of frequency sweep test for NHL paste with ALGNA
5. ábra ALGNA hozzáadásával készült NHL pépek frekvenciaáram vizsgálatának eredményei



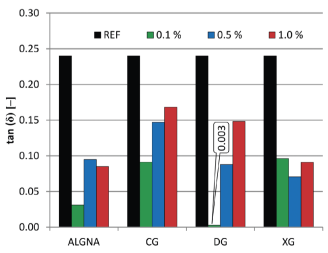


Fig. 6 Complex modulus (G*) and loss factor (tan (δ)) comparison of tested pastes 6. ábra A tesztelt minták komplex modulusának (G*) és veszteség tényezőjének (tan (δ)) összehasonlítása

4. Conclusion

The study informs about influence of addition of biopolymeric admixtures on the rheological properties of natural hydraulic lime pastes.

The addition of biopolymers increased the yield stress of the pastes with growing dosage dependency. The sodium salt of alginic acid, carrageenan and diutan gum increased the consistency coefficient, which is connected with adhesive and anti-sagging properties of mixture, but alginate had dosagedependency trend opposite to the other two admixtures. Addition of these three biopolymers in the dosage of 1% lead to transition of the behaviour of the pastes from pseudoplastic to dilatant, while xanthan gum modified paste showed shear thickening characteristics even in the lowest dose of 0.1%. The study on viscoelastic properties of the modified pastes showed improvement of stability of the grouts modified by carrageenan and diutan gum by prolonging the linear viscoelastic region. The promotion of gel-like behaviour has been observed for all biopolymers but xanthan gum was inferior in efficiency to the other three admixtures. The beneficial effect of the admixtures on the stability of the grouts was also observed by flow sweep tests. The values of complex strain modulus and especially loss tangent support the conclusions.

The use of xanthan gum in natural hydraulic lime pastes is limited due to low efficiency and unclear trends within some of the studied properties. The diutan gum and carrageenan were the admixtures which showed the greatest similarity in behaviour with the hydroxymethylpropyl cellulose, one of the most spread viscosity enhancing admixture in commercial use.

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Libál Lajos György

(1938 - 2019)

Szomorú szívvel tudatjuk mindazokkal, akik ismerték és szerették, hogy életének 82. évében elhunyt Libál Lajos György, az SZTE Üveg szakosztály régi tagja és támogatója.

Középiskolai tanulmányait Budapesten végezte, majd 1965-ben az ausztriai Leobenben szerzett okl. kohómérnöki diplomát.

1981-től Párizsban a RADEX tűzállóanyag-gyártó cég nemzetközi üzletkötője, majd 1988-tól önálló céget alapított. Ezeken keresztül képviselte az európai üvegipar vezető beszállító cégeit, melyek a Tungsram, majd később a GE Hungary üveggyártása egyik legmegbízhatóbb, magas műszaki színvonalat képviselő partnereivé váltak.

Az évtizedek során a segítségével kiemelkedő tudású, nemzetközileg elismert kutatási, fejlesztési szakemberek tartottak színvonalas előadásokat az SZTE Üveg szakosztály konferenciáin, bemutatva a legújabb - elsősorban német – fejlesztéseket, trendeket az üvegipar területén. Az elmúlt években példamutató, jelentős anyagi felajánlást tett az SZTE rendezvények zökkenőmentes lebonyolításához.

80 éves korán túl is aktív "örökifjú" maradt. 1954 óta vívott, és lett korosztályos országos veteránbajnok párbajtőrvívásban.

Bár hősiesen küzdött ellene, élete utolsó nagy csatájában már nem volt esélye a gyilkos kórral szemben.

Nyugodjék békében!

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