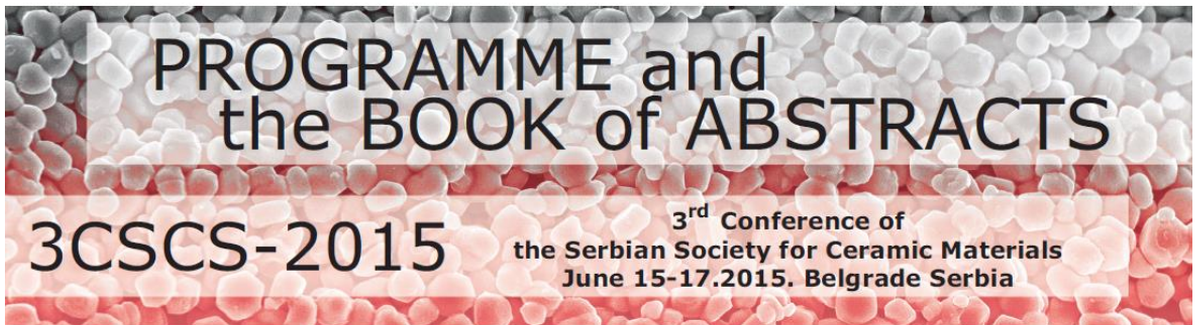


The Serbian Society for Ceramic Materials
The Academy of Engineering Sciences of Serbia
Institute for Multidisciplinary Research - University of Belgrade
Institute of Physics - University of Belgrade
Vinča Institute of Nuclear Sciences - University of Belgrade



Edited by:
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Zorica Branković
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Vladimir V. Srdić

Programme and Book of Abstracts of The Third Conference of The Serbian Society for Ceramic Materilas **publishes abstracts from the field of ceramics, which are presented at international Conference.**

Editors-in-Chief

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Publisher

Institute for Multidisciplinary Research, University of Belgrade
Kneza Višeslava 1, 11000 Belgrade, Serbia

For Publisher

Prof. Dr Sonja Veljović Jovanović

Printing layout

Vladimir V. Srdić

Press

Zonex, Beograd, Serbia

CIP – Каталогизacija у публикацији
Народна библиотека Србије, Београд

666.3/.7(048)

66.017/.018(048)

DRUŠTVO za keramičke materijale Srbije. Konferencija (3 ; 2015 ; Beograd)

Programme ; and the Book of Abstracts / 3rd Conference of the Serbian Society for Ceramic Materials, 3CSCS-2015, June 15-17, 2015, Belgrade, Serbia ; [organizers] The Serbian Society for Ceramic Materials... [et al.] ; edited by Branko Matović ... [et al.]. - Belgrade : Institute for Multidisciplinary Research, University, 2015 (Beograd : Zonex). - 128 str. ; 24 cm

Tiraž 140. - Str. 6: Welcome Message / Branko Matovic. - Registar.

ISBN 978-86-80109-19-0

a) Керамика - Апстракти b) Наука о материјалима - Апстракти c)
Наноматеријали - Апстракти

COBISS.SR-ID 215704332

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MECHANICAL PROPERTIES OF POROUS CERAMIC MONOLITHS BASED ON DIATOMITE

Maja Kokunešoski¹, Jelena Majstorović², Jovana Ružić¹, Branko Matović¹, Svetlana Ilić¹, Adela Egelja¹, Aleksandra Šaponjić¹

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Diatomite from surface coal mine Kolubara, Serbia, was used as a silica source. Firstly, diatomite was purified from organic and inorganic impurities by the heat and chemical treatments. Secondly, boric acid was used as a sintering aid up to 2 wt%. So, after using different pressures of 40, 60, and 80 MPa, the compacted samples were sintered at 850, 1000, 1150, and 1300 °C for 4 h in air. A relatively high porosity in the range of 60-70% is obtained for the samples pressed at the applied pressures and sintered at 1000 °C. The relations between mechanical properties like Young modulus, Poisson ratio, and compressive strength versus content of boric acid in the investigated samples were studied and discussed. Young modulus increases with lowering porosity in the samples sintered at 1300 °C, while Young modulus of the samples sintered at 1150 °C are assumed to be almost linear function of the forming pressure. Compressive strength of the observed samples is higher for the samples sintered at 1300 °C in comparison with the samples sintered at 1150 °C.

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STRUCTURAL AND ELECTRONIC PROPERTIES OF PSEUDOBROOKITE

Zorka Z. Vasiljević¹, Maria V. Nikolic², Obrad S. Aleksic², Nebojsa Labus¹, Miloljub D. Lukovic², Smilja Markovic¹, Pantelija M. Nikolic¹

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Starting powders of TiO₂ (anatase) and Fe₂O₃ (hematite) were mixed in the molar ratio 1:1. Pseudobrookite powder was obtained by a combined milling/calination procedure. Particle size distribution was analyzed on a laser