PACS 82.45.Aa, 81.05.Uw, 82.47.Uv ISSN 1729-4428

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**Promising Cathode Material for Supercapacitors LaFe0.5Cr0.5O3 Perovskite Nanoparticles**

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In this work, the Perovskite powders LaFe0.5Cr0.5O3 with space group of P m -3 m was obtained by sol-gel method. The specific surface area of synthesized material is 14 m2/g. Moreover, the nanoscale powders of LaFe0.5Cr0.5O3 have been tested as a cathode material for electrochemical supercapacitors. The CVA and charge-discharge curves were obtained at 0.5 mV/s to 16 mV/s and 0.5 mA/s to 16 mA/s scan rates accordingly. Thus, the cathode material LaFe0.5Cr0.5O3 demonstrates the specific capacity up to 16 F/g at discharge scan rate 0.5mV/s. Additionally, the maximum of the specific capacity was calculated and it is determined that C is 29.26 F/g, and specific capacity of double electric layer CDEL is 3.44 F/g. It was determined that, the contribution of the redox reactions in specific capacity is 88%. Consequently, the Nyquist plots and Mott-Schottky plots for LaFe0.5Cr0.5O3 were obtained, which consist of two parts with different conductivity type, received the values of ﬂat band potential Efb is -1V for n type and Efb is 0.16V for p type of conductivity.

**Keywords:** sol-gel method,perovskite structure, supercapacitor, impedance, spectroscopy, specific capacity

**Introduction**

Due to the rapid development of electronics the number of personal gadgets is increasing such as mobile phones, smart watches, fitness bracelets, wireless headphones, mini cameras, smart glasses, etc. Moreover, every year they become more complex and more functional, which leads to higher energy consumption. Consequently, against this background there is a need for development new energy storage devices with better capacity, less weight, cheaper, more reliable and environment friendly [1].

Thus, supercapacitor is one of the alternative types of energy storage. The advantages of supercapacitor’s compare to classic batteries are low toxicity, high efficiency (more than 95%), less weight, etc. All this makes supercapacitors very promising in the future. Therefore, research new materials for supercapacitors with predefined properties is relevant today [2].

The perovskite is a complex oxide with the general formula АВО3. It can be realized in a big number of oxide systems, and crystals of perovskites distributed on Earth. Moreover, ABO3attract enormous interest of researchers due to their unique electrical and magnetic properties [2]. Additionally, the complex oxides with perovskite structure have been widely used in various fields, such as cathodes in solid oxide fuel cells, active oxidation catalysts, environmental monitoring films, active materials for chemical sensors for the detection of humidity, alcohols and gases, and so forth [3,4,5].

In this work, the nano-sized LaFe0.5Cr0.5O3 withperovskite structure has been synthesized. The structure and size of the obtained material were determined. Moreover, obtained material was applied as a cathode material for an electrochemical supercapacitor. The specific capacity was determined at different scan rate. The value of maximum of specific capacity was calculated. The conductivity type of tested material at different applied potentials was determined.

**Experiment technique**

The nano-sized powder of perovskite structure LaFe0.5Cr0.5O3 was obtained by sol-gel method. Thus, for the synthesis was used the nitrates crystal hydrates of corresponding metals and citric acid dissolved in distilled water. The level of PH=7 was controlled by adding 25 % ammonia solution. After that, the obtained solution was dried with access to air until it became solid. Dried xerogel heated to a temperature of 230°C to activate auto-burning [6, 7].

Phase composition analysis and determination of crystal structure of obtained materials was performed by DRON-3 diffractometer with Cu (Kα) radiation and X-rays focusing on the Bragg-Brentano. The experimental diffractograms were analyzed with software “FullProf”.

The specific surface of the samples was measured by chromatographic method in gas sorption analyzer *NOVA Quantachrome 2200е*. The method consists in determination of the volume of adsorbed (desorbed) nitrogen by the samples at a temperature of liquid nitrogen and further calculation of the specific surface by BET method.

The electrochemical investigation of obtained material were measured using a three-electrode system with 6 M KOH electrolyte. The working electrode was prepared as a composition of active material and acetylene soot in a proportion of 9 to 1 wrapped in nickel mesh, for counter electrode was used platinum wire, and reference electrode (Ag/AgCl).

Galvanostatic charge-discharge curves, cyclic voltammetry performances, and electrical impudence spectroscopy in the frequency range 105-10-2 Hz were carried on *Autolab PGSTAT/FRA2* workstation with GPES and FRA-2 software [8].

**Results and Discussion**

The powders of complex oxide LaFe0.5Cr0.5O3 with perovskite structure were obtained by sol-gel method with auto-burning. The detailed synthesis of this material described in the previous article [9]. The figure 1 shows the diffraction patterns of dried xerogel (Ammonium nitrate) (a), and synthesized nano-sized powder of the perovskite structure LaFe0.5Cr0.5O3 (b). Also, fig 1 contains theoretical approximation by Rietveld method of experimental diffraction patterns, calculated in software “FullProf”.

The synthesized material has space group P m -3 m with lattice size *а* = 3,904 Å and cell volume V = 59,5 Å3, calculated density ρ = 6,724 g/cm3. Figure 2 shows the transmission electron microscope images in scale 50 nm (a), and 10 nm (b). The average size of the particles is about 40-60 nm.

 

a b

Fig 1. Diffractogram of dried xerogel (a), and synthesized nano-sized powder of perovskite structure LaFe0.5Cr0.5O3 (b).

 

a b

Fig 2. TEM pictures of synthesized nano-sized powder of complex oxide perovskite structure LaFe0.5Cr0.5O3 in scale 50 nm (a), and 10 nm (b).

The typical asymmetric CVA curves of LaFe0.5Cr0.5O3 electrode in 6 M KOH electrolyte in an enlarged potential range of 0.0 to 0.5 V under scan rate range of 0.5 to 16 mV/s, are showed in fig.3. The CVA curves of obtained LaFe0.5Cr0.5O3 electrode are demonstrating the deviation from default electrochemical double layer behavior and redox peaks at ≈ 0.28V confirms the pseudocapacitive characteristics. There is a visible rise in peak current densities with linear increase of scan rates, which demonstrates the sufficient rate of ionic and electronic transportation during the redox reaction in electrolyte with good proficiency rate. Thus, in between scan rate from 0.5 to 16 mV/s, there is no redox peak-shift was observed, and it once again shows the excellent stability and low internal resistance [1, 10]. The peaks near 0.35V on the oxidation branch corresponds to the oxidation of Fe2+ to Fe3+ due to the intercalation of oxygen ions in an oxygen vacancy [1].



Fig 3. CVA of the LaFe0.5Cr0.5O3/KOH system for scan rates from 0.5 to 16 mV/s.



Fig 4. Specific capacity of the LaFe0.5Cr0.5O3/KOH system for scan rates from 0.5 to 16 mV/s.

The figure 4 shows the dependence of the specific capacity on the scan rate of the research material LaFe0.5Cr0.5O3.We can see the non-linear decrease of specific capacity with increasing scan rate. The maximum of the specific capacity is 16.3 F/g at scan rate of 0.5 mV/s. The specific capacity (C) consists of two parts, the first part is the capacity of the double electric layer (CDEL) and the second is capacity of the redox reactions (CF) [10]. From CVA curves view on the fig 3.a we can make an assumption that the main part of LaFe0.5Cr0.5O3 capacity contributes from the redox reactions.



Fig 5. Dependence of specific capacity on s-1/2 for LaFe0.5Cr0.5O3.



Fig 6. Dependence of C-1 on s1/2 for LaFe0.5Cr0.5O3.

From the kinetic model [11], diffuse component (CF) depends on scan rate due to it is a function of the reaction time, and scan rate is inverse to the time of diffusion. In case of linear diffusion, we can use the equation: $C=C\_{s=\infty }+a\sqrt{s}$, where $C\_{DEL}=C\_{s=\infty }$ and $a$ is a constant value. Figure 5 shows the linearly dependency of specific capacity (C) on s-1/2, we can determine the CDEL by the extrapolation of this line to the Y axis. Nevertheless, the capacity of the double electric layer CDEL is 3.44 F/g. Additionally, we can extrapolate the dependency of the specific capacitance on the scan rate to the other side s is 0. 1/C decrease linearly with s1/2, so we can use such equation $\frac{1}{C}=\frac{1}{C\_{x=0}}=b\sqrt{s}$, where b is a constant value [10]. The figure 6 shows the dependency of C-1 on s1/2 for obtained LaFe0.5Cr0.5O3, extrapolation of this dependency to the Y axis made it possible to get the maximum of the specific capacity C is 29.26 F/g. The contribution of the redox reactions in specific capacity is 88%, which confirms our assumption. Figure 7 shows the charge-discharge curves of the electrochemical system based on LaFe0.5Cr0.5O3 electrode. There are linear sections on discharge curves with different tilt angles, they indicate different mechanisms of electric energy storage. To calculate the amount of specific capacitance (Cm) we can use the formula: $C\_{m}=\frac{C}{m}=\frac{I∆t}{∆Um}$, where I (mA) is the discharge current, Δt (s) is the discharge time, ΔU (V) is the potential during the discharge, and m (g) is the mass of the active material inside the electrode[10].



Fig 7. The charge-discharge curves of LaFe0.5Cr0.5O3 in the potential window from 0 to 0.4 V at various currents. 

Fig 8. The specific capacitance density of LaFe0.5Cr0.5O3, measured from discharge part of the charge discharge curves.

The specific capacitance (Cm) values are presented at the fig 8. We can see that the capacity decreases when discharge current increases. It can be explained by the fact that there is not enough time for ions to completely infiltrate into the electrode and enter vacancies. The maximum of specific capacity 10.1 F/g is obtained at the discharge current of 0.5 mA.

Thus, the electron impedance spectroscopy in the frequency range 105-10-2 Hz measurements were made to determine the electrical properties of the nano-sized complex oxide perovskite structure LaFe0.5Cr0.5O3. The Nyquist plots at different values of potential are shown in fig 9.



Fig 9. Nyquist plots of LaFe0.5Cr0.5O3



Fig 10. Mott-Schottky plots of LaFe0.5Cr0.5O3

The Nyquist plot at the electrode potential -1V consists of small semicircle on high frequencies structure and straight line on low frequencies. Semicircle is the result of energy consumption for ion transportation through perovskite structure and straight line corresponds to double electric layer capacity. The electrode potential changing from -1V to -0.3V leads to semicircle straightening and resistance increase. During further potential increasing from -0.3V to 0.5V resistance decrease and plots return to original view with two parts.

Therefore, the Poisson equation can be modified with Boltzmann distribution for electrons distribution in the region of space charge and Gauss's law for electric field on the border of charge to obtain Mott-Schottky equation [12]:

$$\frac{1}{С^{2}}=\frac{2}{εε\_{0}A^{2}eN\_{D}}\left(V-V\_{fb}-\frac{k\_{B}T}{e}\right),$$

where ε and ε0 are dielectric constant and vacuum permittivity, e is an electron charge, A is BET surface area, ND is a carrier concentration, E and Efb are electrode potential and ﬂat band potential values, T – an absolute temperature, kB – Boltzmann constant. The linear fitting of the low-potential part allows calculating the ﬂat band potential (Efb), and the slope of the curve corresponds for the type of conductivity. The Mott-Schottky plots are presented in fig 10, they can be divided into two parts with different type of conductivity: from -1V to -0.3V and -0.2V to 0.5V with the same point of division as Nyquist plots. First area corresponds for n type of conductivity with Efb is -1V, and p type for the second area with Efb is 0.16V.

**Conclusions**

The nano-sized powder of perovskite structure LaFe0.5Cr0.5O3 was successfully obtained by sol-gel method, which confirms by X-rays analysis and transmission electron microscope images.

Moreover, the LaFe0.5Cr0.5O3 was used as cathode material and demonstrates the specific capacity up to 16 F/g at discharge scan rate 0.5mV/s.

It was determined that, the calculated maximum of the specific capacity C is 29.26 F/g, where 88% is specific capacity of redox reactions.

LaFe0.5Cr0.5O3 shows different types of conductivity depends on applied potential, received the values of ﬂat band potential Efb is -1V and Efb is 0.16V for n type and p type of conductivity accordingly.

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**Перспективний катодний матеріал для супер конденсаторів нанорозмірний перовскитний матеріал LaFe0.5Cr0.5O3**

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У цій роботі, порошки перовскиту LaFe0.5Cr0.5O3 з просторовою групою P m -3 m були отримані золь-гель методом. Площа питомої поверхні синтезованого матеріалу 14 м2/г. Також, нанорозмірні порошки LaFe0.5Cr0.5O3 були випробувані як катодний матеріал для електрохімічних супер конденсаторів. Отримано CVA криві при швидкостях сканування від 0,5 мВ/с до 16 мВ/с та заряд розрядні криві при швидкостях сканування від 0,5 мA/с до 16 мA/с. Катодний матеріал LaFe0.5Cr0.5O3 демонструє питому ємність 16 Ф/г при швидкості розряду 0,5 мА/с. розрахований максимум питомої ємності C становить 29.26 Ф/г, там питома ємність подвійного електричного шару CDEL становить 3.44 Ф/г. Було визначено, що вклад окисно-відновних реакцій у питому ємність складає 88%. Отримано діаграми Найквіста та Мотта-Шотки для LaFe0.5Cr0.5O3, котрі складаються х двох частин з різним типом провідності, отримані значення потенціалів плоскої зони Efb становить -1В для n типу та Efb становить 0.16V для p типу провідності.

**Ключові слова**: золь-гель метод, структура перовскиту,супер конденсатор, імпеданс, спектроскопія, питома ємність.