Thermoelectric Properties of Inhomogeneous Ceramics Based on the Layered Calcium Cobaltate

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Abstract The effect of cationic composition and sintering conditions on the electrotransport and thermoelectric properties of Ca₃CoₙO₉₊δ (x = 3.8, 4.0, and 4.2) had been investigated. It had been found that increase of cobalt oxide content in the samples increase their electrical conductivity, creation of phase inhomogeneity improves their thermo-EMF coefficient, and sintering above temperature of peritectoid decomposition increase their apparent density, which, in the whole, improves thermoelectric properties of ceramics based on the layered calcium cobaltate Ca₃Co₄O₉₊δ. So, power factor values of phase inhomogeneous ceramics Ca₃Co₄.2O₉₊δ, sintered in air below and above temperature of peritectoid decomposition of Ca₃Co₄O₉₊δ, at 800°C were equal 255 and 273 μW/(m⋅K²) respectively, which was 2.1 and 2.4 times larger than for the Ca₃Co₄O₉₊δ. It had been also found that sintering of phase inhomogeneous ceramics both in oxidizing or reducing atmospheres resulted in improving of its functional properties. So, power factor values of Ca₃Co₃.8O₉₊δ (sintered in oxygen at 970°C) and Ca₃Co₄.2O₉₊δ (sintered in nitrogen at 920°C) at 800°C were equal 422 and 378 μW/(m⋅K²) respectively, which was 3.5 and 3.1 times larger than for the Ca₃Co₄O₉₊δ.

Keywords Layered Calcium Cobaltate, Phase Inhomogeneity, Thermoelectrics, Sintering Conditions, Electrical Conductivity, Thermo-EMF, Power Factor

1. Introduction

The growth of global energy consumption and the negative environmental impact of many modern energy conversion technologies (for example, the burning of hydrocarbon fuels) have led to increased activity in the search for alternative energy sources. A huge renewable source of energy is the heat evolved into the environment during the work of industrial enterprises, vehicles, as well as various plants and units. This heat can be directly and effectively converted into electrical energy using thermoelectric generators, the creation of which requires so-called thermoelectric materials (thermoelectrics), which possesses high values of electrical conductivity and thermo-EMF coefficient and have low thermal conductivity [1]. Chalcogenides of heavy metals are traditional thermoelectrics, but these materials are expensive, toxic and unstable in air at high temperatures.

The indicated drawbacks are largely absent for oxide thermoelectrics, including materials based on the layered calcium cobaltate Ca₃Co₄O₉₊δ, which in recent years are considered as a promising basis for the development of p-branches of high-temperature thermoelectric generators [2, 3]. The functional (thermoelectric) characteristics of ceramics based on the Ca₃Co₄O₉₊δ can be essentially improved by using instead of the traditional ceramic methods the “soft” low-temperature synthesis methods [4, 5], using special techniques for sintering ceramics – hot pressing [6], spark plasma sintering [7–9], and also by partial substitution of calcium ions in it by bismuth [3, 10–12] or rare earth elements ions [3, 13, 14] or of cobalt ions by 3d-metal ions [3, 15, 16]. It was shown in [11, 17–19] that thermoelectric characteristics of oxide ceramics can be improved by creating phase inhomogeneity in it.

In ceramics based on layered calcium cobaltate, phase inhomogeneity can be created in various ways – by introducing a second, impurity phase into the mixture at the stage of synthesis or sintering [20–22], by varying the cationic stoichiometry of the initial mixture so that the target composition is outside the Ca₃Co₄O₉₊δ homogeneity region [23] (according to [24], in air, layered calcium cobaltate can be obtained in the composition range
Ca₃Co₃.8O₉δ – Ca₃Co₄.2O₉δ), as well as by thermal treatment (annealing) of ceramics at temperatures exceeding the temperature of peritectoid decomposition of Ca₃Co₆O₁₅ (T_p = 926°C in air [24]) by the reaction Ca₃Co₄O₉δ → Ca₃Co₂O₆ + (Co, Ca)O. In this work, the last two approaches were used to create phase inhomogeneity in ceramics based on layered calcium cobalate.

The aim of this work was to study the influence of the cationic composition and thermal prehistory of ceramics based on Ca₃Co₂O₉δ on its phase composition, physicochemical and functional properties, as well as to study the possibility of improving the thermoelectric (functional) characteristics of materials based on the layered calcium cobalate by creating of phase inhomogeneity in it.

2. Materials and Methods

Ceramics with a nominal composition of Ca₃Co₃.8O₉δ, Ca₃Co₄O₉δ, and Ca₃Co₄.2O₉δ, was obtained by the citrate method according to the procedure described in [14, 16] using Ca(NO₃)₂·4H₂O, Co(NO₃)₂·6H₂O (analytical grade), and citric acid C₆H₈O₇ (analytical grade) as the starting reagents. At the final stage, the samples were sintered for 6 h in air (A serie) at temperatures of 920°C (T<T_p) and 970°C (T>T_p), in oxygen (O serie) at temperatures of 970°C (T<T_p) and 1020°C (T>T_p), and in nitrogen (N serie) at temperatures of 890°C (T<T_p), 920°C and 970°C (T>T_p) followed by slow cooling to room temperature.

The phase composition of the samples and the crystal structure parameters of predominant phases in them were determined using X-ray diffraction analysis (XRD) (Bruker D8 XRD Advance X-ray diffractometer, CuKα-radiation, Ni filter).

The microstructure of sintered ceramics and its chemical composition was studied by means of JSM–5610 LV scanning electron microscope with EDX JED–2201 chemical analysis system (JEOL, Japan) using a reflected electron detector (accelerating voltage – 20 kV) in low vacuum mode (p = 1 Pa) and using Fei Company Quanta 200 instrument equipped with an EDAX attachment. The apparent density of the samples (ρ_exp) was determined on their mass and geometric dimensions.

The electrical conductivity (σ) and thermo-EMF coefficient (S) of sintered ceramics based on the layered calcium cobalate Ca₃Co₄O₉δ were studied in air in the temperature range of 300–1100 K [25]. The values of apparent activation energy of their electrical conductivity were determined from linear parts of ln(σ·T) = f(T) dependencies, taking into account that temperature dependencies of electrical conductivity of layered calcium cobalate Ca₃Co₄O₉δ and its derivatives above room temperature usually obey equation σ = (σ₀/T)·exp[−E₄/(k_BT)], where σ₀ is temperature independent constant, E₄ is apparent activation energy of electrical conductivity, k_B is Boltzmann’s constant, and T is absolute temperature [5, 23]. The power factor (P) values were calculated by the formula P = S²·σ.

3. Results and Their Discussion

Taking into account the phase diagram of the CaO–CoO quasibinary system in air [24], it was expected that all samples of A serie, except for Ca₃Co₄O₉δ ceramics, sintered at temperature of 920°C, will be non-monophase (Table 1). According to the XRД data (Figure 1, a, Table 1), samples of Ca₃Co₃.8O₉δ, Ca₃Co₄O₉δ, and Ca₃Co₄.2O₉δ compositions after annealing in air at 920°C, and sample of Ca₃Co₃.8O₉δ composition after annealing in air at 970°C were mono-phase, within the accuracy of XRД method, and had the structure of the layered calcium cobalate Ca₃Co₄O₉δ [26] with the lattice constants given in the Table 1. As can be seen from the Table 1, creation both excess and deficiency of cobalt oxide in the layered calcium cobalate leads to the increasing of δ parameter (repeatability period in the ab-plane of CoO₂-layers of Ca₃Co₄O₉δ crystal structure [26]. The samples of Ca₃Co₃O₉δ and Ca₃Co₄.2O₉δ compositions annealed in air at 970°C were non-monophase and consisted of calcium cobaltite Ca₃Co₂O₆ having a quasi-one-dimensional structure [27, 28] and CoO₄ oxide [29], with predominant phase of Ca₃Co₂O₆ in both cases (Table 1).
Taking into the account on the results of the XRD we can make two conclusions. Firstly, the impurity phases (Ca₃Co₂O₆, (Co,Ca)O) in the samples Ca₃Co₄.2O₉+δ, Ca₃Co₃.8O₉+δ sintered at 920°C are in amounts lower than the sensitivity of the XRD method. Secondly, the formation of layered calcium cobaltate Ca₃Co₄O₉+δ, solid solution (Co,Ca)O in the samples sintered at 920°C at their cooling to the room temperature in mixtures containing an excess of CaO occurs much faster than in the mixtures of stoichiometric composition (Ca: Co = 3: 4) and in the mixtures containing an excess of cobalt oxide.

The apparent density of ceramics varied within 2.72–3.20 g/cm³ (Table 1), increased at increasing of their sintering temperature, and the highest values of ρ_{app} were observed for samples with a lack of cobalt oxide (excess of calcium oxide) having Ca₃Co₃.8O₉+δ composition.

Taking into account the phase diagram of the CaO–CoO quasibinary system in an oxygen atmosphere [24], it was expected that all samples of O serie, except for Ca₃Co₄O₉+δ ceramics, sintered in an oxygen atmosphere at 970°C, will be non-monophase (Table 2). According to the XRD data (Figure 1, b, Table 2), all the obtained samples after annealing were, within the accuracy of XRD method, monophase and had the structure of layered calcium cobaltate Ca₃Co₄O₉+δ [26] with the crystal structure parameters given in the Table 2. Interestingly, that creation both excess and deficiency of cobalt oxide in the layered calcium cobaltate results in the similar changes of its lattice constants: increasing of b₂ parameter for the samples sintered at 970°C in oxygen (below T_p) and increasing of a parameter and decreasing of b₁ and b₂ parameters for the samples sintered at 1020°C in oxygen (above T_p).

Taking in the account the results of the XRD we can assume, that the impurity phases (Ca₃Co₂O₆, (Ca,Co)O) in the samples Ca₃Co₃.8O₉+δ, Ca₃Co₄.2O₉+δ, which had been sintered in an oxygen atmosphere at 970°C and 1020°C present in amounts less than the sensitivity of the XRD.

Table 1. Apparent density (ρ_{app}), phase composition and lattice constants (a, b₁, b₂, c, β) of predominant phase in the ceramics based on the layered calcium cobaltate obtained in air (A serie) at different temperatures T

| Composition | T, °C | ρ_{app}, g/cm³ | Phase composition | Predominant phase | Lattice constants of predominant phase |
|-------------|-------|----------------|-------------------|-------------------|--------------------------------------|
| Ca₃Co₃.8O₉+δ | 920   | 2.72           | Ca₃Co₄O₉+δ ¹       | Ca₃Co₄O₉+δ       | a = 4.829(7), b₁ = 4.566(8), b₂ = 2.846(9), c = 10.85(1), β = 98.13(8) |
| Ca₃Co₃.8O₉+δ | 970   | 3.01           | Ca₃Co₄O₉+δ, (Co, Ca)O | Ca₃Co₄O₉+δ       | a = 4.846(7), b₁ = 4.567(8), b₂ = 2.832(9), c = 10.84(1), β = 97.93(8) |

¹According to the X-ray data
²According to [24], at 970°C in air this phase is thermodynamically unstable, but can form at slow cooling of the sample to the room temperature due to the peritectoid reaction of Ca₃Co₄O₉+δ + (Co, Ca)O → Ca₃Co₄O₉+δ.

Figure 1. X-ray powder diffractograms (CuKα-radiation) of Ca₃CoO₉+δ (1, 2, 7, 8), Ca₃Co₃O₉+δ (3, 4, 9, 10), and Ca₃Co₄O₉+δ (5, 6, 11, 12) samples, annealed in air (a) and oxygen (b) at 920°C (1, 3, 5), 970°C (2, 4, 6, 7, 9, 11), and 1020°C (8, 10, 12). The Miller indices of Ca₃Co₄O₉+δ phase are indicated. Symbols * and # depict the reflexes of Ca₃Co₂O₆ and Co₃O₄ phases respectively.
method.

The apparent density of ceramics slightly depended on their composition and sintering temperature and varied within 2.53–2.86 g/cm³ (Table 2); the highest value of ρ_{app} was observed for the sample with an excess of cobalt oxide (calcium oxide deficiency) having Ca₃Co₄.2O₉+δ composition, which had been annealed at 1020°C in oxygen.

Samples of the Ca₃Co₃.0O₉δ+, Ca₃Co₄O₉δ+, and Ca₃Co₄.2O₉δ+ composition, sintered in a nitrogen atmosphere, were non-monophase, and, according to the XRD results, consisted of CoO, CaO, and Ca₃Co₄O₉δ+ phases, with cobalt oxide had being the predominant phase (Figure 2, lines 1–3, Table 3). After measuring of electrical conductivity (in air within the temperature range of 25–800°C), the phase composition of the samples changed significantly: the most intense peaks corresponded to the phase of layered calcium cobaltate Ca₃Co₄O₉δ+ (Figure 2, lines 4–6, Table 3), formed by reaction:

\[
3\text{CaO} + 4\text{CoO} + (1 + 0.5\delta)\text{O}_2 \rightarrow \text{Ca}_3\text{Co}_4\text{O}_{9+\delta}.
\]

Lattice constants of layered calcium cobaltate having both excess and deficiency of cobalt oxide in the samples sintered in nitrogen after their heat treatment in air similarly differed from the lattice constants of basic Ca₃Co₄O₉δ+ phase (Table 3): their \(a, b_1,\) and \(b_2\) parameters were essentially smaller, but \(c\) one was larger. On the diffraction patterns of these samples, in addition to the peaks of the main phase (Ca₃Co₄O₉δ+), there were weakly expressed peaks of additional phases (CaO and CoO), from which we can conclude that in this case too, the ceramic is a chemically and phase inhomogeneous composite material.

The apparent density of ceramics sintered at 920°C, slightly depended on their composition and varied within 2.72–2.78 g/cm³ (Table 3).

According to the EPMA results (Figure 3), chemical composition of the samples after their heat treatment at different conditions was corresponded, in the whole, to their nominal composition.

As can be seen from Figure 4, \(a,\) ceramics having Ca₃Co₄.0O₉δ+ composition, which had been sintered at 920°C in air contained only anisometric particles, the sizes of which in different directions varied within one to six micrometers (Table 4). Micrographs of ceramics of the same composition annealed at 970°C (Figure 4, \(b\)) showed crystallites of two types – anisometric large particles for the main Ca₃Co₄O₉ phase and small practically isometric inclusions of the Co₄O₉ phase.

### Table 2. Apparent density (ρ_{app}), phase composition and lattice constants \((a, b_1, b_2, c, \beta)\) of predominant phase in the ceramics based on the layered calcium cobaltate obtained in oxygen \((O\ serie)\) at different temperatures \(T\)

| Composition | \(T, ^°C\) | ρ_{app}, g/cm³ | Phase composition | Lattice constants of predominant phase |
|-------------|-------------|----------------|------------------|---------------------------------------|
| Ca₃Co₂O₆ | 970 | 2.73 | Ca₃Co₄O₉δ+ | \(a = 4.831(7)\ A, b_1 = 4.586(8)\ A, b_2 = 2.845(9)\ A, c = 10.85(1)\ Å, \beta = 98.08(8)\) |
| Ca₃Co₂O₆ | 970 | 2.69 | Ca₃Co₄O₉δ+ | \(a = 4.826(7)\ A, b_1 = 4.554(8)\ A, b_2 = 2.823(9)\ A, c = 10.83(1)\ Å, \beta = 98.10(8)\) |
| Ca₃Co₂O₆ | 970 | 2.53 | Ca₃Co₄O₉δ+ | \(a = 4.822(7)\ A, b_1 = 4.555(8)\ A, b_2 = 2.836(9)\ A, c = 10.83(1)\ Å, \beta = 98.13(8)\) |
| Ca₃Co₂O₆ | 970 | 2.73 | Ca₃Co₄O₉δ+ | \(a = 4.866(6)\ A, b_1 = 4.544(7)\ A, b_2 = 2.794(9)\ A, c = 10.85(1)\ Å, \beta = 98.86(7)\) |
| Ca₃Co₂O₆ | 1020 | 2.62 | Ca₃Co₄O₉δ+ | \(a = 4.843(7)\ A, b_1 = 4.575(8)\ A, b_2 = 2.821(9)\ A, c = 10.87(1)\ Å, \beta = 98.45(8)\) |
| Ca₃Co₂O₆ | 1020 | 2.86 | Ca₃Co₄O₉δ+ | \(a = 4.858(7)\ A, b_1 = 4.537(8)\ A, b_2 = 2.799(9)\ A, c = 10.85(1)\ Å, \beta = 98.83(8)\) |

*According to the X-ray data

*According to [24], at 1020°C in oxygen this phase is thermodynamically unstable, but can form at slow cooling of the sample to the room temperature due to the peritectoid reaction of Ca₃Co₂O₆ + (Co, Ca)O → Ca₃Co₄O₉δ+.
Table 3. Apparent density ($\rho_{app}$), phase composition and lattice constants ($a$, $b_1$, $b_2$, $c$, $\beta$) of predominant phase in the ceramics based on the layered calcium cobaltate obtained in nitrogen (N serie) at 920°C before (nitrogen) and after (nitrogen+air) measurements of their electrotransport properties in air within 25–800°C

| Composition       | Prehistory | $\rho_{app}$/g/cm$^3$ | Phase composition | Predominant phase | Lattice constants of predominant phase |
|-------------------|------------|------------------------|-------------------|-------------------|----------------------------------------|
| $\text{Ca}_3\text{Co}_3.8\text{O}_{9+\delta}$ | Nitrogen   | 2.76                   | CaO, CoO          | CoO               | $4.278(3)$ $-$ $-$ $-$ $-$            |
| $\text{Ca}_3\text{Co}_4\text{O}_{9+\delta}$ | Nitrogen   | 2.75                   | CaO, CoO          | CoO               | $4.282(3)$ $-$ $-$ $-$ $-$            |
| $\text{Ca}_3\text{Co}_4\text{O}_{9+\delta}$ | Nitrogen+air | 2.77               | CaO, CoO          | $\text{Ca}_3\text{Co}_4\text{O}_{9+\delta}$ | $4.824(7)$ $-$ $-$ $-$ $-$ $-$ $-$ |
| $\text{Ca}_3\text{Co}_3.8\text{O}_{9+\delta}$ | Nitrogen+air | 2.78               | CaO, CoO          | $\text{Ca}_3\text{Co}_4\text{O}_{9+\delta}$ | $4.838(6)$ $-$ $-$ $-$ $-$ $-$ $-$ |
| $\text{Ca}_3\text{Co}_3.8\text{O}_{9+\delta}$ | Nitrogen+air | 2.75               | CaO, CoO          | $\text{Ca}_3\text{Co}_4\text{O}_{9+\delta}$ | $4.804(9)$ $-$ $-$ $-$ $-$ $-$ $-$ |

*According to the X-ray data

According to the results of EMPA of the cleaved surfaces, all phases are not individual compounds, but substitution solid solutions (Table 4). The fact that the spinel phase is a solid solution of calcium oxide in cobalt oxide (II, III) ($\text{Co}_{0.9}\text{Ca}_{0.1})_3\text{O}_4$ is consistent with the phase diagram of the CaO–CoO quasibinary system [24]. The excess of cobalt in layered calcium cobaltate and quasi-one-dimensional calcium cobaltite is most likely due to the partial segregation of cobalt oxide between the volume and surface of the ceramic grains. In other words,
the ceramics we obtained is not only non-monophase, but also chemically inhomogeneous.

The crystallites of ceramics of $\text{Ca}_3\text{Co}_4\text{O}_{9+\delta}$ composition, which had been sintered in oxygen at a temperature of 1020°C were larger and more anisometric (Figure 4, c) than for ceramics of the same composition sintered in air at 920°C, and had an average particle size of about $5.8\times2.7\times1.0$ μm (Table 4). According to the EMPA results, the particles were enriched in cobalt oxide (Table 4), which, as in the previous case, is most likely explained by its partial segregation between the volume and surface of the ceramic grains.

Micrographs of ceramics annealed at 920 °C in a nitrogen atmosphere showed crystallites of two types: anisometric large particles, which are heterogeneous aggregates consisting of $\text{Ca}_3\text{Co}_4\text{O}_{9+\delta}$, $\text{CaO}$, and $\text{CoO}$ phases, and small practically isometric inclusions of the $\text{CoO}$ phase (Figure 4, d). After additional heat treatment in air (measurements of electrical conductivity), as a result of a change in the phase composition of the ceramic, a noticeable change in its microstructure was occurred, consisting in a change in the shape and size of the ceramic grains (Figure 4, e). According to the EMPA results of the cleaved surfaces, all phases are not individual compounds, but substitution solid solutions, and, as in the case of ceramics sintered in air and in oxygen, phase crystallites based on layered calcium cobaltate contained an excess amount compared to stoichiometric cobalt oxide (Table 4).

As can be seen from the data given in the Figure 5 (upper block), the materials based on the layered calcium cobaltate, which had been sintered in air, were $p$-type ($\sigma > 0$) semiconductors ($\partial\sigma/\partial T > 0$), while $\text{Ca}_3\text{Co}_4\text{O}_{9+\delta}$ ceramics sintered at 970°C was characterized by the highest electrical conductivity, and the highest values of the coefficient of thermo-EMF was observed for ceramics $\text{Ca}_3\text{Co}_4\text{O}_{9+\delta}$ sintered both at 920°C and at 970°C (Table 5), which is most likely due to its phase heterogeneity (see Figure, 1, a, Table 1). The values of electrical conductivity and coefficient of thermo-EMF of ceramics, in the whole, were increased at increasing of the cobalt oxide content in the samples (Table 5). The values of the apparent activation energy of electrical conductivity of the samples of A serie varied within 0.069–0.081 eV, which is typical for ceramics based on the layered calcium cobaltate [12, 14, 16] (excluding the samples having $\text{Ca}_3\text{Co}_4\text{O}_{9+\delta}$, $\text{Ca}_3\text{Co}_4\text{O}_{9+\delta}$ composition, which had been sintered at 970°C, whose $E_A$ values were equal to 0.274 and 0.125 eV respectively and were abnormally high, which was probably due to the peculiarities of their microstructure).

The power factor values of sintered in air ceramics had been increased at temperature increasing, and for sample having $\text{Ca}_3\text{Co}_4\text{O}_{9+\delta}$ composition, containing an excess of cobalt oxide, were significantly higher than for the basic monophase layered calcium cobaltate $\text{Ca}_3\text{Co}_4\text{O}_{9+\delta}$ (Figure 5, upper block, Table 5). So, for $\text{Ca}_3\text{Co}_4\text{O}_{9+\delta}$ ceramics sintered at 920°C and 970°C, $P$ values at 800°C were equal to 255 and 273 μW/(m·K²), respectively, which was 2.1 and 2.3 times higher than for the basic layered calcium cobaltate $\text{Ca}_3\text{Co}_4\text{O}_{9+\delta}$ at the same temperature.

Samples which were sintered in oxygen were also $p$-type ($\sigma > 0$) semiconductors ($\partial\sigma/\partial T > 0$) (Figure 5, middle block), although their electrical conductivity values at temperature gaining increased less significant than for the samples, sintered in air.

Figure 4. Electron micrographs of surface of $\text{Ca}_3\text{Co}_4\text{O}_{9+\delta}$ ceramics, annealed in air at 920°C (a) and 970°C (b), in oxygen at 1020°C (c), in nitrogen at 920°C before (d) and after (e) measurements of its electrotransport properties in air within 25 – 800°C.
Table 4. The results of analysis of surface of cleavages of ceramics based on the layered calcium cobaltate sintered at different conditions ($T$, atmosphere) according EM and EMPA

| Sintering conditions | Air 920°C | 970°C | Oxygen 1020°C | Nitrogen 920°C (+25–800°C) | Nitrogen (+air) 920°C (+25–800°C) |
|----------------------|------------|--------|----------------|---------------------------|----------------------------------|
| Phase                | Ca$_3$Co$_2$O$_{9+δ}$ | Ca$_3$Co$_2$O$_{9+δ}$ | Co$_3$O$_4$ | Ca$_3$Co$_4$O$_{9+δ}$ | CoO | Ca$_3$Co$_4$O$_{9+δ}$ |
| $x_{\text{Ca}}$, at. % | 36.35 | 48.65 | 9.98 | 33.35 | 15.01 | 32.62 |
| $x_{\text{Co}}$, at. % | 63.65 | 51.35 | 90.02 | 66.65 | 84.99 | 67.38 |
| $x_{\text{CaO}}$, wt. % | 29.94 | 41.48 | 7.66 | 27.24 | 11.68 | 26.60 |
| $x_{\text{CoO}}$, wt. % | 70.06 | 58.52 | 92.34 | 72.76 | 88.32 | 73.40 |
| Phase composition    | (Ca$_{0.85}$Co$_{0.15}$)$_3$Co$_4$O$_{9+δ}$ | (Ca$_{0.85}$Co$_{0.15}$)$_3$Co$_4$O$_{9+δ}$ | (Co$_3$/Ca$_3$)$_3$O$_4$ | (Ca$_{0.82}$Co$_{0.18}$)$_3$Co$_4$O$_{9+δ}$ | Co$_3$Co$_{0.8}$O$_{3}$ | (Ca$_{0.89}$/Ca$_{0.11}$)$_3$Co$_4$O$_{9+δ}$ |
| Average crystallite size (given in the $x\times y\times z$ directions), $\mu$m | $2.2 \times 1.7 \times 0.5$ | $5.2 \times 3.4 \times 1.7$ | $1.6 \times 1.6 \times 1.6$ | $5.8 \times 2.7 \times 1.0$ | $1.7 \times 1.7 \times 1.7$ | $2.3 \times 1.8 \times 1.7$ |
Figure 5. Temperature dependences of electrical conductivity ($\sigma$), thermo-EMF coefficient ($S$), and power factor ($P$) of Ca$_3$Co$_4$O$_9$+$\delta$ (1, 2, 3, 4, 7, 8, 13, 14), Ca$_3$Co$_3.8$O$_9$+$\delta$ (3, 4, 9, 10, 15, 16), and Ca$_3$Co$_4.2$O$_9$+$\delta$ (5, 6, 11, 12, 17, 18) ceramics, annealed in air (1 – 6), oxygen (7 – 12) and nitrogen (13 – 18) at 890°C (13), 920°C (1, 3, 5, 15, 17), 970°C (2, 4, 6, 7, 9, 11, 14, 16, 18), and 1020°C (8, 10, 12).
The highest electrical conductivity was demonstrated by ceramics having Ca₃Co₃.8O₉+δ composition which had been sintered at 1020°C, and the highest value of the thermo-EMF coefficient of ceramics containing an oxide (Ca₃Co₄.2O₉+δ) were higher than for the basic Ca₃Co₄O₉+δ phase (Table 5). It is interesting to note that the values of the apparent activation energy of electrical conductivity (Eₐ, eV) and the power factor (P₉₀₀, µW/(m²K)) of ceramics, based on the layered calcium cobaltate, sintered at different conditions (T, atmosphere) are close each other, excluding sample Ca₃Co₃.8O₉+δ sintered in nitrogen increased with increasing temperature (T, atmosphere).

The values of the apparent activation energy of electrical conductivity of ceramics having Ca₃Co₃.8O₉+δ composition which had been sintered at 1020°C, and the highest value of the thermo-EMF coefficient of ceramics containing an oxide (Ca₃Co₄.2O₉+δ) were higher than for the basic Ca₃Co₄O₉+δ phase with stoichiometric relation of cations, which conductivity had a semiconducting character (0 < δσ/δT < 0) (except for a sample of Ca₃Co₄.2O₉+δ composition, which conductivity had a semiconducting character (0 < δσ/δT < 0) and showed the highest values in this series) (Figure 5, lower block, Table 5). Interestingly, that increasing of sintering temperature of ceramics resulted in decreasing of their electrical conductivity values, probably, due to the more pronounced of phase heterogeneity as well as due to the lowering of cobalt oxidation degree in them leading to the decreasing of concentration of main charge carriers – “holes”.

The values of the thermo-EMF coefficient of ceramics sintered in nitrogen increased with increasing temperature and were close each other, excluding sample Ca₃Co₃.8O₉+δ sintered at 920°C and sample Ca₃Co₄.2O₉+δ sintered at 970°C, which thermo-EMF coefficient were approximately 25% higher (Table 5). The maximal values of thermo-EMF coefficient (like for the samples which were sintered in oxygen) was observed for chemically inhomogeneous samples, which contained an excess of calcium oxide (Ca₃Co₃.8O₉+δ) or cobalt oxide (Ca₃Co₄.2O₉+δ) (Table 5), which is another confirmation of the fact that inhomogeneity of layered calcium cobaltate improve its

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Table 5. Values of apparent density (ρₚₐₚ), electrical conductivity (σₑₑₑ), thermo-EMF coefficient (Sₑₑₑ), apparent activation energy of electrical conductivity (Eₐ), and power factor (Pₙₙₙ) of ceramics, based on the layered calcium cobaltate, sintered at different conditions (T, atmosphere)

| Sintering conditions | Composition    | ρₚₐₚ, g/cm³ | σₑₑₑ, S/cm | Sₑₑₑ, µV/K | Eₐ, eV | Pₙₙₙ, µW/(m²K) |
|----------------------|---------------|-------------|------------|------------|--------|----------------|
| **Air**              |               |             |            |            |        |                |
| 920°C                | Ca₃Co₃.8O₉+δ | 2.87        | 29.4       | 170        | 0.069  | 85.1           |
|                      | Ca₃Co₄O₉+δ   | 2.77        | 44.8       | 164        | 0.081  | 121            |
| 970°C                | Ca₃Co₃.8O₉+δ | 3.20        | 32.6       | 137        | 0.274  | 60.8           |
|                      | Ca₃Co₄O₉+δ   | 3.01        | 37.9       | 171        | 0.125  | 111            |
| **Oxygen**           |               |             |            |            |        |                |
| 970°C                | Ca₃Co₃.8O₉+δ | 3.11        | 52.0       | 229        | 0.064  | 273            |
| 1020°C               | Ca₃Co₃.8O₉+δ | 2.73        | 31.6       | 366        | 0.080  | 422            |
|                      | Ca₃Co₄O₉+δ   | 2.69        | 25.2       | 323        | 0.065  | 262            |
| **Nitrogen***        |               |             |            |            |        |                |
| 890°C                | Ca₃Co₄O₉+δ   | 2.63        | 37.3       | 219        | 0.060  | 179            |
| 920°C                | Ca₃Co₃.8O₉+δ | 2.76        | 34.5       | 288        | 0.050  | 286            |
|                      | Ca₃Co₄O₉+δ   | 2.75        | 50.7       | –          | 0.062  | –              |
| 970°C                | Ca₃Co₃.8O₉+δ | 2.95        | 17.3       | 229        | 0.064  | 91.1           |
|                      | Ca₃Co₄O₉+δ   | 2.73        | 48.4       | 230        | 0.059  | 256            |

*All the samples sintered in nitrogen atmosphere, were light-brown in color, but after measurements of their electrical conductivity or thermo-EMF in air their color was changing into black, obviously, due to their oxidation and change in phase composition (from mixture of CoO and CaO into ceramics based on the Ca₃Co₄O₉+δ phase)
thermoelectric properties.

The values of the apparent activation energy of electrical conductivity of the samples of N serie varied within 0.050–0.072 eV, which is typical for ceramics based on the layered calcium cobaltate [12, 14, 16], slightly depended on the cationic composition of the samples and temperature of their sintering, and were less than for the samples which had been sintered in air or in oxygen.

The power factor values of ceramics sintered in nitrogen increased at temperature increasing (Figure 5, lower block), while the character of the $P = f(T)$ dependences of the samples was determined mainly by the character of the temperature dependences of their thermo-EMF coefficient ($S = f(T)$). As can be seen (Figure 5, Table 5), the highest values of the power factor at 1100 K had the samples containing excess of cobalt oxide $\delta$ – $Ca_{3}Co_{2}O_{9+\delta}$, sintered at temperatures of $920°C \ (P_{800} = 378 \ \mu W/(mK^2)$ and $970°C \ (P_{800} = 356 \ \mu W/(mK^2))$, which is 3.1 and 2.9 times higher than for the basic monophase ceramics of layered calcium cobaltate $Ca_{3}Co_{4}O_{9+\delta}$.

Comparing the results obtained for the ceramics having various cationic composition and sintered at different conditions we can make some conclusions:

a) electrical conductivity of the samples was increased after they had been sintered both in oxidizing (oxygen) and reducing (nitrogen) atmospheres; in the first case it is due to the increasing of average oxidation degree of cobalt ions in their crystal structure leading to the increasing of concentration of main charge carriers – “holes”, in the second case the heat post-treatment in air atmosphere of samples sintered in nitrogen (during measurements of their electrical conductivity) led to their oxidation and, consequently, to increasing of charge carriers concentration;

b) the values of apparent activation energy of electrical conductivity of the samples, in the whole, were close each other, and varied within 0.05–0.08 eV; so, mechanism of electrical conductivity in all the samples studied is the same, and charge transfer in them is took place through the $Ca_{3}Co_{4}O_{9+\delta}$ phase (both in homogeneous and heterogeneous ceramics);

c) values of thermo-EMF coefficient are larger for the samples containing an excess both cobalt oxide and calcium oxide, which was more pronounced for the samples sintered in oxygen and nitrogen; thus, creation in the ceramics based on the layered calcium cobaltate of phase and chemical inhomogeneity essentially improves its thermoelectric (functional) properties.

4. Conclusions

Ceramic samples of $Ca_{3}Co_{x}O_{9+\delta}$ ($x = 3.8, 4.0, \text{and} 4.2$) composition were prepared using citrate method and then sintered in different atmospheres (oxygen, air, nitrogen) below and above temperature of peritectoid decomposition of layered calcium cobaltate $Ca_{3}Co_{4}O_{9+\delta}$.

Their crystal structure, microstructure, electrical conductivity, thermo-EMF coefficient, and power factor in air within 25–800°C were investigated. It had been found that increase of cobalt oxide content in the samples led to the increasing of their electrical conductivity, the creation of phase inhomogeneity in ceramics (by both varying of their cationic composition and by heat treatment at temperatures above temperature of peritectoid decomposition) improves its thermo-EMF coefficient, which, in the whole, improves thermoelectric properties of ceramics based on the layered calcium cobaltate $Ca_{3}Co_{4}O_{9+\delta}$. So, power factor values of phase and chemically inhomogeneous ceramics $Ca_{3}Co_{2.5}O_{9+\delta}$, sintered in air below and above temperature of peritectoid decomposition of layered calcium cobaltate, at 800°C were equal 255 and 273 $\mu W/(mK^2)$ respectively, which was 2.1 and 2.4 times larger than for the $Ca_{3}Co_{4}O_{9+\delta}$.

It had been also found that sintering of phase inhomogeneous ceramics both in oxidizing or reducing atmospheres led to the improving of its functional properties. So, power factor values of $Ca_{3}Co_{3.0}O_{9+\delta}$ (sintered in oxygen at 970°C) and $Ca_{3}Co_{2.5}O_{9+\delta}$ (sintered in nitrogen at 920°C) at 800°C were equal 422 and 378 $\mu W/(mK^2)$ respectively, which was 3.5 and 3.1 times larger than for the $Ca_{3}Co_{4}O_{9+\delta}$.

Abbreviations

EMF electromotive force
XRD x-ray diffraction analysis
EDX energy-dispersive x-ray analysis
EM electron microscopy
EMPA electronic micro probe analysis

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