Technical University of Denmark



High-temperature Thermoelectric Properties of Ca0.9Y0.1Mn1-xFexO3 (0 x 0.25)

Le, Thanh Hung; Van Nong, Ngo; Han, Li; Minh, Dang Le; Borup, Kasper A.; Iversen, Bo B.; Pryds, Nini; Linderoth, Søren *Published in:*

Journal of Materials Science

Link to article, DOI: 10.1007/s10853-012-6834-z

Publication date: 2013

Link back to DTU Orbit

Citation (APA):

Le, T. H., Van Nong, N., Han, L., Minh, D. L., Borup, K. A., Iversen, B. B., ... Linderoth, S. (2013). Hightemperature Thermoelectric Properties of Ca0.9Y0.1Mn1-xFexO3 (0 x 0.25). Journal of Materials Science, 48(7), 2817-2822. DOI: 10.1007/s10853-012-6834-z

DTU Library Technical Information Center of Denmark

General rights

Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

- Users may download and print one copy of any publication from the public portal for the purpose of private study or research.
- You may not further distribute the material or use it for any profit-making activity or commercial gain
- You may freely distribute the URL identifying the publication in the public portal

If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

ENERGY MATERIALS & THERMOELECTRICS

High-temperature thermoelectric properties of $Ca_{0.9}Y_{0.1}Mn_{1-x}Fe_xO_3$ ($0 \le x \le 0.25$)

4 Le Thanh Hung · Ngo Van Nong · Li Han ·

5 Dang Le Minh · Kasper A. Borup · Bo B. Iversen ·

6 Nini Pryds · Søren Linderoth

Received: 25 May 2012/Accepted: 23 August 2012 © Springer Science+Business Media, LLC 2012

9 **Abstract** Polycrystalline compounds of $Ca_{0.9}Y_{0.1}Mn_{1-x}$ Fe_xO_3 for $0 \le x \le 0.25$ were prepared by solid-state 10 11 reaction, followed by spark plasma sintering process, and 12 their thermoelectric properties from 300 to 1200 K were 13 systematically investigated in terms of Y and Fe co-doping 14 at the Ca- and Mn-sites, respectively. Crystal structure 15 refinement revealed that all the investigated samples have 16 the O'-type orthorhombic structure, and the lattice param-17 eters slightly increased with increasing Fe concentration, 18 causing a crystal distortion. It was found that with 19 increasing the content of Fe doping, the Seebeck coefficient 20 of Ca_{0.9}Y_{0.1}Mn_{1-x}Fe_xO₃ tended to increase, while the 21 tendency toward the electrical conductivity was more 22 complicated. The highest power factor was found to be 2.1×10^{-4} W/mK² at 1150 K for the sample with 23 24 x = 0.05 after annealing at 1523 K for 24 h in air. Thermal 25 conductivity of the Fe-doped samples showed a lower 26 value than that of the x = 0 sample, and the highest 27 dimensionless figure of merit, ZT was found to be 28 improved about 20 % for the sample with x = 0.05 as 29 compared to that of the x = 0 sample at 1150 K.

- 30
- A1 L. T. Hung (🖂) · N. V. Nong · L. Han · N. Pryds · S. Linderoth

A2 Department of Energy Conversion and Storage, Technical

- A3 University of Denmark, DTU Risø Campus, 4000 Roskilde,
- A4 Denmark
- A5 e-mail: lthh@dtu.dk
- A6 D. L. Minh
- A7 Solid State Department, Faculty of Physics, Hanoi University
- A8 of Science, Vietnam National University of Hanoi, Hanoi,
- A9 Vietnam
- A10 K. A. Borup · B. B. Iversen
- A11 Centre for Materials Crystallography, Department of Chemistry
- A12 and iNANO, University of Aarhus, 8000 Aarhus C, Denmark

Introduction

With increasing the global energy demand, thermoelectric 32 materials have recently gained much interest in both the 33 theoretical and technological aspects due to the potential 34 use of these materials in converting waste heat into elec-35 tricity [1, 2]. In general, for a single thermoelectric mate-36 rial, the conversion efficiency can be evaluated by the 37 dimensionless figure of merit ($ZT = \sigma S^2 T / \kappa$, where σ , S, T, 38 κ are the electrical conductivity, the Seebeck coefficient, 39 the absolute temperature, and the thermal conductivity, 40 respectively). The requirements for practical application of 41 high thermal-to-electrical energy conversion place on 42 finding suitable thermoelectric materials, and are not easily 43 satisfied. They should not only possess good thermoelectric 44 performance, they must also be stable at high temperatures 45 and be composed of nontoxic and low-cost elements, but 46 also must be able to be processed and shaped cheaply. For 47 this purpose, metal oxide-based materials are considered as 48 49 good candidates.

50 CaMnO₃, which is a perovskite oxide with orthorhombic structure at room temperature, has also been considered as 51 52 a promising thermoelectric n-type material for hightemperature application [3–9]. Many attempts have been 53 made in order to improve the thermoelectric performance 54 of this type of material, mainly to enhance the electrical 55 conductivity, reduce further the thermal conductivity, 56 57 while avoiding degradation of the Seebeck coefficient. 58 Most of these studies have been focused on doping, for example, Yb at Ca-site [4-7] or Nb at Mn-site [3, 8], while 59 only few reports performed the research on dually doping, 60 e.g., Sr and Yb at Ca-site [9]. Previous reports have showed 61 that the substitution of Y for Ca resulted in a significant 62 improvement in the thermoelectric performance of Ca_{1-x} 63 Y_x MnO₃ system in a wide temperature region, and the 64



Journal : Large 10853	Dispatch : 29-8-2012	Pages : 6
Article No. : 6834		□ TYPESET
MS Code : JMSC29016	🗹 СР	🗹 DISK

<u>Author Proo</u>

7

8

1

72

73

74

75

76

77

78

79

80

81

optimum doping level was found to be around x = 0.1 [5, 10]. Similar to other multi-valence systems such as cobaltites [11] or titanates [12], the interrelation between Mn³⁺ and Mn⁴⁺ should be responsible for the transport mechanism in the CaMnO₃ material. Therefore, doping of trivalent ions such as Fe³⁺ or Co³⁺ at the Mn-site would probably influence the transport properties of this material.

In this study, we have prepared the $Ca_{0.9}Y_{0.1}Mn_{1-x}$ Fe_xO₃ system with $0 \le x \le 0.25$, in which Ca-site was substituted with Y at a fixed concentration and Mn-site was partly replaced by Fe. The structural and the thermoelectric properties of these set of materials were investigated in detail. The influence of Y and Fe doping at Ca- and Mnsites, respectively, on the crystal structure was carefully studied by the Rietveld refinement analysis. The correlation between the crystal structures and the thermoelectric properties are discussed.

82 Experimental

83 Polycrystalline samples of Ca_{0.9}Y_{0.1}Mn_{1-x}Fe_xO₃ with 84 x = 0, 0.05, 0.15, 0.2, and 0.25 were synthesized by a 85 solid-state reaction. A mixture of commercially available 86 CaCO₃ (98 %), MnO₂ (99.9 %), Fe₂O₃ (99.9 %), and Y₂O₃ 87 (99.9 %) precursors were thoroughly mixed by ball milling 88 with ethanol for 24 h. The mixtures were dried and then 89 calcined at 1273 K for 24 h in air with an intermediate 90 grinding procedure. The densify processing was carried out 91 using a spark plasma sintering (SPS) system (SPS Syntex 92 Inc., Japan). The samples were heated to 1123 K, while a 93 uniaxial pressure of 50 MPa was applied for 8 min in Ar 94 atmosphere. During the experiment, the temperature, 95 applied pressure, and displacement of the sample were 96 recorded continuously. The as-prepared samples were then 97 polished in order to remove the graphite foil used during 98 the SPS processing. The pellets were then cut into bar 99 $(3.5 \times 3.5 \times 12 \text{ mm}^3)$ and plate $(10 \times 10 \times 1.4 \text{ mm}^3)$ 100 shapes for the thermoelectric properties and thermal con-101 ductivity measurements, respectively. XRD analysis was 102 carried out on the powders after calcining and after the SPS 103 processing using a Bruker robot diffractometer with Cu-Kα 104 radiation. Structural refinement was carried out using the 105 Rietveld method with TOPAS 4.1. Microstructures of the 106 samples were observed using scanning electron microscopy 107 (SEM) with a Hitachi TM-1000 system. The electrical 108 resistivity and the Seebeck coefficient were measured 109 simultaneously from room temperature to 1200 K using an 110 ULVAC-RIKO ZEM3 measurement system in a low-111 pressure helium atmosphere. The thermal conductivity, κ , 112 was determined from the measured thermal diffusivity, α , the heat capacity, C_{p} , and the density, d, using the formula: 113 $\kappa = d \times \alpha \times C_{p}$. The densities of the samples were 114

🖉 Springer

117

measured by an AccuPyc-1340 pycnometer. The thermal 115 diffusivity was measured by a LFA-457 laser flash system. 116

Results and discussion

Figure 1 shows powder X-ray diffraction (XRD) spectra 118 measured at room temperature for pure CaMnO₃ and for 119 $Ca_{0.9}Y_{0.1}Mn_{1-x}Fe_xO_3$ samples with x = 0, 0.05, 0.1, 0.15, 1200.2, and 0.25 after they were calcined at 1273 K for 24 h in 121 air. All the visible XRD peaks can be indexed as the pure 122 phase of CaMnO₃, indicating that all the investigated 123 samples are single phase. 124

Figure 2 displays XRD patterns of a typical sample with 125 x = 0.05 for the calcined powder (a), spark plasma sintered 126 pellet (b), and the SPS sample after further heat treatment 127 at 1523 K for 24 h in air (c). As indicated by this figure 128 regardless of heat treatment, the structure remained the 129 same. The structure refinement for the calcined powders 130 $Ca_{0.9}Y_{0.1}Mn_{1-x}Fe_xO_3$ system was conducted using Topas 131 4.1 Rietveld refinement software with input parameters 132 which were taken from Poeppelmeier et al. [13] using 133 space group Pnma (No.62), and the refined results are 134 summarized in Table 1. The profile R value (R_p) , weighted 135 profile R-factor (R_{wp}) , and Goodness of fit (GOF) values 136 obtained in this analysis are of high quality, and is clearly 137 illustrated in Fig. 2 for a typical $Ca_{0.9}Y_{0.1}Mn_{1-x}Fe_xO_3$ 138 sample with x = 0.05 as an example. This result implies 139 that Y and Fe most likely substituted on the Ca- and Mn-140 sites of CaMnO₃, respectively. It can be judged from the 141 data in Table 1, that the lattice parameters follow a relation 142 of $c/\sqrt{2} \le a \le b$, confirming that the polycrystalline 143



Fig. 1 XRD patterns of CaMnO₃ and Ca_{0.9}Y_{0.1}Mn_{1-x}Fe_xO₃ with x = 0, 0.05, 0.1, 0.15, 0.2, 0.25 samples after calcining at 1273 K for 24 h in air

~	Journal : Large 10853	Dispatch : 29-8-2012	Pages : 6
	Article No. : 6834	□ LE	□ TYPESET
	MS Code : JMSC29016	🖌 СЬ	🗹 DISK



Fig. 2 XRD patterns of a typical sample Ca_{0.9}Y_{0.1}Mn_{0.95}Fe_{0.05}O₃: (a) Rietveld refinement profile of the calcined powder, (b) pellet sample sintered by SPS at 1173 K under pressure 50 MPa for 8 min under Ar atmosphere, (c) SPS sample after annealing at 1523 K in air for 24 h

144 compounds $Ca_{0.9}Y_{0.1}Mn_{1-x}Fe_xO_3$ of our samples have O'-145 type orthorhombic structure [9, 14].

146 The dependence of the lattice parameters and the cell 147 volumes of $Ca_{0.9}Y_{0.1}Mn_{1-x}Fe_xO_3$ on the amount of Fe 148 substituent are presented in Fig. 3. The result shows that 149 the lattice parameters slightly increased with the increasing 150 Fe concentration, resulting in an expansion in the unit cell 151 volume. The increase in lattice parameters may be associated with the substitution of Fe³⁺ with a larger ionic 152 radius (0.55 Å) for smaller Mn^{4+} (0.53 Å) ion [15]. In the 153 case if Fe^{3+} substitutes for Mn^{3+} (0.58 Å), which has 154 155 larger ionic radius, one would expect a slight contraction of 156 the unit cell volume. The geometric distortion of ABO₃-157 type perovskites can be explained by Goldsmith tolerance 158 factor, which is defined as

$$t = (r_{\rm A} + r_{\rm O})/\sqrt{2}(r_{\rm B} + r_{\rm O})$$
(1)

160 where r_A , r_B , and r_O are the ionic radii of A, B, and O 161 atoms, respectively [15]. For $Ca_{0.9}Y_{0.1}Mn_{1-x}Fe_xO_3$ com-162 pounds, calculation of Goldsmith tolerance factors 163 (t) showed that the highest t value was 0.988 in the case of Fe^{3+} substitutes for Mn⁴⁺ and the smallest *t* value was 164

0.963 with Fe^{3+} substitutes for Mn^{3+} . It implies that the 165 orthorhombic structure is stable for all Ca_{0.9}Y_{0.1} 166 $Mn_{1-x}Fe_xO_3$ compounds. 167

Figure 4 depicts the temperature dependence of the 168 electrical conductivity for $Ca_{0.9}Y_{0.1}Mn_{1-x}Fe_xO_3$ with 169 x = 0, 0.05, 0.1, 0.15, 0.2, and 0.25 spark plasma sintered 170 samples. The result points out that the electrical conduc-171 172 tivity of the entire samples exhibit a semiconducting-like behavior over the whole measured temperature range. 173 174 However, the electrical conductivity of the spark plasma sintered samples does not show a clear tendency with the 175 increase of Fe doping concentration. σ tends to decrease 176 with increasing Fe concentration for x > 0.1, while 177 increases for the samples with x < 0.1. It should be noted 178 here that those samples were sintered under a high pressure 179 at high temperature and in inert gas atmosphere. The 180 oxygen content or even the microstructure may be varied 181 from the samples, causing the different behaviors of the 182 183 electrical conductivity as a result.

Temperature dependence of the Seebeck coefficient 184 (S) for the spark plasma sintered samples of $Ca_{0.9}Y_{0.1}$ 185 $Mn_{1-x}Fe_xO_3$ with x = 0, 0.05, 0.1, 0.15, 0.2, and 0.25 are 186 shown in Fig. 5. S of all investigated samples show a 187 188 negative values over the whole measured temperature range, indicating n-type conduction. Contrastingly to the 189 electrical conductivity, the absolute S values increase with 190 increasing Fe concentration, and the effect was more sub-191 stantial in low temperature region. 192

In order to understand further the influence of the Fe 193 doping on the thermoelectric properties, four SPS samples 194 195 with x = 0, 0.05, 0.1, and 0.15 were selected and further 196 annealed at 1523 K for 24 h in air. Figure 6a shows the electrical conductivity and the Seebeck coefficient as a 197 198 function of temperature after annealing at 1523 K for 24 h in air. As seen from Fig 6a, a clear tendency showing the 199 decrease of σ , while S increases with increasing Fe con-200 centration from x = 0, 0.05, 0.1 to 0.15. The value of 201 electrical conductivity was found to increase more than two 202 times as compared with post samples, while the Seebeck 203 coefficient remained almost the same. 204

In general, the conduction mechanism of CaMnO₃ can 205 be interpreted by hopping conduction [16] where hopping 206

Table 1 Structural refinement factors, lattice parameters, and cell volumes of	Compositions (x)	0	0.05	0.1	0.15	0.2	0.25
$Ca_{0.9}Y_{0.1}Mn_{1-x}Fe_{x}O_{3}$	$R_{ m wp}$ (%)	8.42	9.33	9.39	11.27	10.62	10.62
	$R_{\rm p}~(\%)$	6.68	6.70	6.42	7.67	7.45	6.63
	GOF	1.73	1.90	1.71	1.83	1.92	1.85
	a (Å)	5.28233(2)	5.27480(3)	5.3018(3)	5.3004(1)	5.29596(2)	5.29999(3)
	<i>b</i> (Å)	7.46185(3)	7.45797(4)	7.4846(6)	7.4806(2)	7.48990(3)	7.49067(5)
	<i>c</i> (Å)	5.26841(2)	5.27498(5)	5.2899(2)	5.3035(3)	5.28405(2)	5.28344(3)
	$V(\text{\AA}^3)$	207.659(2)	207.514(2)	209.914(2)	210.284(3)	209.598(3)	209.755(2)



1	Journal : Large 10853	Dispatch : 29-8-2012	Pages : 6
	Article No. : 6834	□ LE	□ TYPESET
	MS Code : JMSC29016	🗹 СР	🖌 disk



Fig. 3 Lattice parameters and cell volume of $Ca_{0.9}Y_{0.1}Mn_{1-x}Fe_xO_3$ as function of Fe content (x)



Fig. 4 Temperature dependence of the electrical conductivity for $Ca_{0.9}Y_{0.1}Mn_{1-x}Fe_xO_3$ with x = 0, 0.05, 0.1, 0.15, 0.2, 0.25 spark plasma sintered samples; Inset, the activation energies were fitted from experimental data



Fig. 5 Temperature dependence of the Seebeck coefficient for $Ca_{0.9}Y_{0.1}Mn_{1-x}Fe_xO_3$ with x = 0, 0.05, 0.1, 0.15, 0.2, and 0.25 spark plasma sintered samples

🖉 Springer

140 (a) -80 130 -100 120 110 -120 100 90 -140 σ (Ω⁻¹cm⁻¹ 80 S (µVIK) -160 70 -180 40 -200 -220 20 v=0 -240 x = 0.05S x = 0.1S -260 S x = 0.150 400 600 800 1000 1200 T (K) (b) Anealing @ 1523 K 2.0 0 x = 0. 0.05 0 P (10⁻⁴ Wm⁻¹K⁻²) 1.5 1.0 0.5



x = 0, 0.05, 0.1, 0.15, 0.2, 0.25 and selective samples with x = 0, 0.05, 0.1, 0.15 after annealing at 1523 K for 24 h in air

207 of the charge carriers is thermally activated with the activation energy $E_{\rm a}$, the temperature dependence of the 208 electrical conductivity is given as 209

$$\sigma = C/T \exp(-E_{\rm a}/k_{\rm B}T) \tag{2}$$

211 where T is absolute temperature, $k_{\rm B}$ is the Boltzmann constant, and C is a constant depending on the charge 212 carrier concentration. The activation energy could be esti-213 mated from the Arrhenius plot of LnT versus 1/T as shown 214 in the inset in Fig. 4. The calculated activation energy, $E_{\rm a}$ 215 is listed in Table 2 for all investigated samples, showing 216 217 that $E_{\rm a}$ is linearly increasing with the increase of Fe substituent. However, the relationship between and E_a is only 218 obeyed the hopping conduction's equation at temperatures 219 below 700 K, as shown in Fig. 4 inset as well as in 220 Table 2. As for the x = 0 and 0.05 samples, the LnT versus 221 1/T curve showed two different slopes in the temperature 222 regions of T < 700 and T > 700 yielding two activate 223 energies (see Table 2), which is similar to the observation 224 225 by Vecherskii et al. [17] on the oxygen non-stoichiometry CaMnO_{3- δ} system. 226

~	Journal : Large 10853	Dispatch : 29-8-2012	Pages : 6
	Article No. : 6834	□ LE	□ TYPESET
	MS Code : JMSC29016	🖌 СР	🗹 disk

Author Proof

Table 2 Relative densities and electrical characteristics of $Ca_{0.9}Y_{0.1}$ $Mn_{1-v}Fe_vO_3$

1 4 4 5						
Compositions (x)	0	0.05	0.1	0.15	0.2	0.25
Relative density (%)	94.36	94.86	97.40	96.65	95.60	93.39
$E_{\rm a}~({\rm meV})$	117.46	146.15	155.28	181.33	189.69	227.47

For an extrinsic *n*-type semiconductor with negligible hole conduction, the thermoelectric power can be given by [18, 19]:

$$S(T) \approx -\frac{k_{\rm B}}{e} \left[\ln\left(\frac{N_{\rm v}}{n}\right) + A \right]$$
 (3)

231 where e is the electric charge of the carrier, $k_{\rm B}$ the Boltz-232 mann constant, $N_{\rm V}$ the density of states (DOS), *n* the carrier 233 concentration, and A is a transport constant. Equations 2 234 and 3 clearly show that the decrease in carrier concentra-235 tion (n) will result in an increase in the thermoelectric 236 power (S) and vice versa. This can well explain the ten-237 dency of the Seebeck coefficient and the electrical con-238 ductivity as a function of temperature observed for the 239 investigated samples after annealing with the increasing Fe 240 concentration (see Fig. 6). Increasing the Fe content 241 decreases the conductivity and increases the Seebeck 242 coefficient which is also related to the carrier concentration 243 via Eq. 3. However, further investigation on the carrier 244 density and the mobility by means of the Hall measure-245 ments is currently ongoing to evidently support this 246 interpretation.

247 Figure 6b shows the power factor (PF) as a function of 248 temperature for all the spark plasma sintered samples and 249 the selected ones after annealing. It is obvious that the PF 250 values were remarkably improved by further heat treatment 251 in air. The x = 0.05 sample showed the highest PF values 252 over the whole measured temperature region, and the 253 maximum PF attained was 2.1×10^{-4} W/mK² at about 1150 K. 254

255 To understand the reason which led to the finding 256 interesting effect on the thermoelectric properties of the 257 samples after heat treatment, the microstructure of the 258 samples after SPS and after further annealing was studied 259 using SEM, and the results are shown in Fig. 7a, b. Fig-260 ure 7 shows an obvious difference in the grain size before and after the annealing. The small grains size structure 261 observed for the spark plasma sintered sample means to be 262 263 more grain boundaries, leading to the increase in electron 264 scattering at the grain boundaries, and thus decreasing the electrical conductivity. This result well explained the 265 behavior of the electrical conductivity for the samples 266 267 before and after heat treatment.

Figure 8 shows the total thermal conductivity (κ_{total}) for all investigated samples. It can be seen that κ decreases



Fig. 7 SEM images from fractured surfaces of a typical $Ca_{0.9}Y_{0.1}$ $Mn_{1-x}Fe_xO_3$ with x = 0.05 sample: **a** sample was sintered by SPS, **b** sample was annealed at 1523 K for 24 h in air flow



Fig. 8 the total thermal conductivity (κ_{total}), the electronic and phonon components (κ_e and κ_{ph}) of Ca_{0.9}Y_{0.1}Mn_{1-x}Fe_xO₃ samples with x = 0, 0.05, 0.1, 0.15 as a function of temperature

 Journal : Large 10853	Dispatch : 29-8-2012	Pages : 6
Article No. : 6834	□ LE	□ TYPESET
MS Code : JMSC29016	🖌 СР	🖌 DISK



Fig. 9 The dimensionless figure of merit (ZT) as a function of temperature for $Ca_{0.9}Y_{0.1}Mn_{1-x}Fe_xO_3$ with x = 0, 0.05, 0.1, 0.15selective SPS samples after heated treatment at 1523 K for 24 h in air

270 with increasing temperature. The substitution of Fe at 271 Mn-sites generally decreases the thermal conductivity. The 272 total thermal conductivity can be expressed by the sum of a 273 lattice component ($\kappa_{\rm ph}$) and an electronic component ($\kappa_{\rm e}$), 274 i.e., as $\kappa_{\text{total}} = \kappa_{\text{ph}} + \kappa_{\text{e}}$. In this case, the contribution of κ_{e} 275 to κ_{total} , estimated from the Wiedemann–Franz relation, is 276 small, indicating the major contribution of the phonon term 277 $\kappa_{\rm ph}$, as clearly illustrated in Fig. 8. Finally, using the 278 measured thermoelectric data, the dimensionless figure of 279 merit of these compositions was calculated. Figure 9 pre-280 sents the dimensionless figure of merit, ZT, versus tem-281 perature for the x = 0, 0.05, 0.1, and 0.15 samples, 282 showing that ZT increased for the x = 0.05 and then 283 decreased again with increasing x over the whole temper-284 ature range. The maximum ZT value reached a value of 285 0.11 at about 1150 K for the x = 0.05 samples.

286 Conclusion

The effect of Fe substitution on the structure and the high-287 temperature thermoelectric properties of $Ca_{0.9}Y_{0.1}Mn_{1-x}$ 288 289 Fe_xO_3 (x = 0, 0.05, 0.1, 0.15, 0.2, 0.25) was investigated in 290 details. Structural analysis shows that lattice parameters 291 slightly increase with increasing amount of Fe substituent, 292 which originates from the difference in the ionic radii 293 between Fe and Mn ions. The thermoelectric properties 294 were found to be improved for the Fe-doped samples with x < 0.1, particularly for the SPS samples with further 295 296 annealing mainly due to the increase in the Seebeck coefficient that could overcome the simultaneous decrease 297 of the electrical conductivity. The thermal conductivity 298 was suppressed by the substitution of Fe for Mn. The 299 maximum PF attained was 2.1×10^{-4} W/mK² for the 300 x = 0.05 sample at 1150 K giving a maximum ZT = 0.11, 301 which is about 20 % higher than the x = 0 sample. Further 302 study should be performed with finer Fe substituent tuning 303 304 with x < 0.1 in order to optimize these compounds high-305 temperature thermoelectric properties.

306 Acknowledgements The authors would like to thank the Programme 307 Commission on Energy and Environment (EnMi) which is part of the 308 Danish Council for Strategic Research (Contract No. 10-093971) for 309 sponsoring this work via the OTE-POWER research work.

References

310

319

320

321

322

323

324

333

337

338

339

- 1. Rowe DM (ed) (2006) Thermoelectric handbook: macro to nano. 311 312 CRC/Taylor & Francis, Boca Raton 313
- 2. Snyder GJ, Toberer ES (2008) Nat Mater 7:105
- 314 3. Bocher L, Aguirre MH, Logvinovich D, Shkabko A, Robert R, Trottmann M, Weidenkaffet A (2008) Inorg Chem 47:8077 315
- 316 4. Kosuga A, Urate S, Kurosaki K, Yamanaka S, Funahashi R 317 (2008) Jpn J Appl Phys 47(8):6399 318
- 5. Wang Y, Sui FanH, Wang X, Su Y, Su W, Liu X (2009) Chem Mater 21:4653
- 6. Flahaut D, Mihara T, Funahashi R, Nabeshima N, Lee L, Ohta H, Koumoto K (2006) J Appl Phys 100:084911
- 7 Wang Y, Sui Y, Su W (2008) J Appl Phys 104:093703
- 8. Bocher L, Aguirre MH, Robert R, Logvinovich D, Bakardjieva S,
- Hejtmanek J, Weidenkaff A (2009) Acta Mater 57:5667 9. Kosuga A, Isse Y, Wang Y, Koumoto K, Funahashi R (2009) J
- 325 326 Appl Phys 105:093717 327
- 10. Thuy NT, Minh DL, Nong NV, Bahl CRH and Pryds N, 7-9 328 November, Ho Chi Minh city, Vietnam (2011), Proceedings of 329 the solid state physics and materials science symposium (in press)
- 330 11. Nong NV, Pryds N, Linderoth S, Ohtaki M (2011) J Adv Mater 331 23(21):2484 332
- 12. Wang HC, Wang CL, Su WB, Liu J, Sun Y, Peng H, Mei LM (2011) J Am Ceram Soc 94(3):838
- 334 13. Poeppelmeier KR, Leonowicz ME, Scanlon JC, Longo JM (1982) 335 J Solid State Chem 45:71 336
- 14. Kostogloudis GC, Fertis P, Ftikos C (1999) Solid State Ionics 118:241
- 15. Shanon RD (1976) Acta Crystallogr A A32:751
- 16. Karim DP, Aldred AT (1979) Phys Rev B 20:2255
- 340 17. Vecherskii SI, Konopel'ko MA, Esina NO and Batalov NN 341 (2002) Inorg Mater 38(12):1491 342
- 18. Jonker GH (1968) Philips Res Rep 23:131
- 343 19. Ohtaki M, Tsubota T, Eguchi K, Arai H (1996) J Appl Phys 344 79:1816 345

Author

🖉 Springer



Journal : Large 10853	Dispatch : 29-8-2012	Pages : 6
Article No. : 6834	□ LE	□ TYPESET
MS Code : JMSC29016	🖌 СР	🗹 disk