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ABSTRACTS

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Structural analysis of LiNi_{1/3}Mn_{1/3}Co_{1/3}O₂, Li_{0.96} Na_{0.94}Ni_{1/3}Mn_{1/3}Co_{1/3}O₂ and Li_{0.96}K_{0.04}Ni_{1/3}Mn_{1/2}Co_{1/3}O₂ materials synthesized by Pechini method

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Layered tri-transition metal oxides, specially LiNi_{1/2}Co_{1/2}Mn_{1/2}O₂ (NMC 333), have become a promising alternative to LiCoO2 electrode material in the rechargeable Lithium-Ion Battery (LIB). The electrochemical performances of NMC 333 mainly depend on its crystallographic structural properties including lattice parameters, the unit-cell, c/a ratio, volume, crystallite size (D), dislocation density (δ), and lattice strain. This study aims to synthesize LiNius Mnus CoursO2, Lines Nanos Nius Mnus CoursO2, and Line-Kon Ni, a Mn, a Co. a On materials and study their structural properties. The Pechini method was used for powder synthesis in this study. The synthesized materials were characterized using X-ray diffraction (XRD), X-ray characterization confirmed the formation of only the single-phase layered hexagonal lattice (α-NaFeO2-type) structure without any impurity phase for all these prepared materials. Interestingly, while confirming the formation of layered structures, a better splitting of the (006)/(102) and (108)/(110) peaks appeared for LieukKeneNii/2Mni/2Coi/2O2 than that of LiNi 1/2 Mn 1/2 Co 1/2 O2 and Lio 26 Naco Ni 1/2 Mn 1/2 Co 1/2 O2 in the diffractograms. The lattice parameters, i.e. a, c, c/a, the unit-cell volume, the crystallite size (D), and dislocation density(8) are 2.8641(Å), $14.2143(\text{\AA}), 4.9629, 100.979(\text{\AA}^2), 77.45 \text{ nm}, 1.666 \times 10^{16} \text{ m}^{-2}, \text{ for LiNi}_{1/2}\text{Mn}_{1/2}\text{Co}_{1/2}\text{O}_{3}$ While they are 14.2317(Å), 101.347(Å²), $1.382 \times 10^{14} \text{ m}^{-2}$ 2.8675(Å), 4.9630, 85.06 for LioneNanouNi, Mn, CO, CO, 202 and 2.869 (Å), 14.2421(Å), 4.9641, 101.528(Å2), 128.38 nm, 0.606 × 1016 m-2 for Lings Kons Ni 13 Mn 13 Co 13 Op. respectively. It is also observed that the lattice parameters, the unit-cell volume, c/a, and the crystallite size are increased with the substitution of Li+ by Na+ and K. It may be due to the radii of Na+ and K- are bigger than that of Li+ and that will have the way for increasing the interlayer space of the substituted materials with the substitution of bigger ions. The c/a ratio constitutes a direct indication of the cation mixing, Linea Nana Nita Mnita Cona 20 and Liona Kooa Niug Mnug Coug Oz exhibit higher c/a values than LiNiug Mnug Coug Oz, supporting the observation that the substituting bigger ions such as Na+ and K+ into LiNi, 2Mn, 2Co, 2O2 suppresses the cation mixing and forms a well-defined layered structure. The micro-strain calculated for the LiNi1/2Mn1/2C01/2O2, LinseNano4Ni1/2Mn1/2C01/2O2, and LinseKno4Ni1/2Mn1/2C01/2O2 are 1.38 × 10⁻², 2.17 × 10-2 and 1.46 × 10-2, respectively. This implies a slight difference in the crystallinity of the materials, as the micro-strain was slightly affected by substituting Na+ and K+. Crystallite size (D) was 77.45 nm, 85.06 nm, and 128.38 nm for LiNi, Mn, CO, CO, D. Lio, Naco, Ni, Mn, CO, CO, and Lione Koon Nil 27 Mn 1/2 Co 1/2 O 2 respectively. It exhibits an increment of crystallite size, indicating a lowering of the dislocation density with the substitution of bigger ions. Altogether, this study reveals that substituting Li+ with bigger ions of Na+ and K+ is improving the structural stability of NMC 333.

Keywords: NMC materials, Na doping, K doping, Li-ion battery, Pechini method

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