

Supplementary Materials

Synergistic internal-external disorder to activate electrochemically inert maricite NaFePO₄

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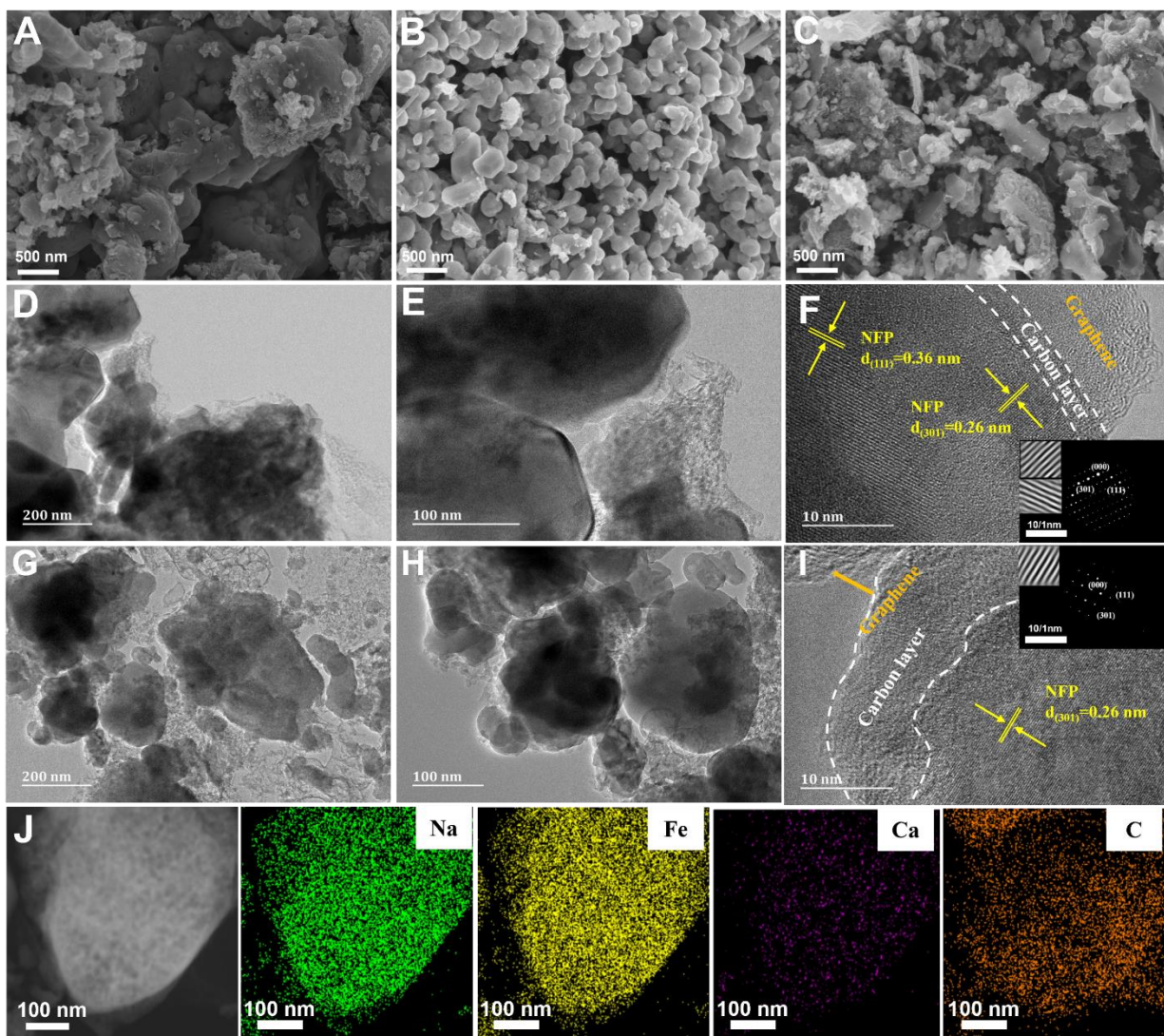


Figure S1. SEM images of NFP/C (A), NFP/C-H (B) and NFP/C-T_{0.01} (C) to NFP/C-T_{0.01} (B), NFP/C-H (C); (D-F) TEM and HRTEM images of NFP/C-T_{0.01}; (G-I) TEM and HRTEM images of NFP/C-H; (J) EDS images of NFP/C-T_{0.01}.

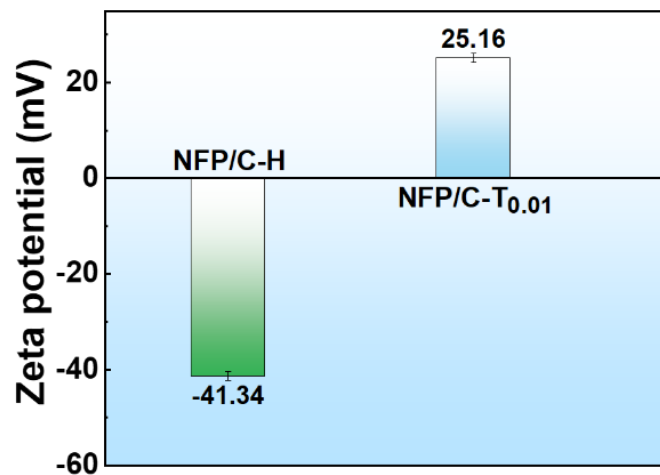


Figure S2. Zeta potential values of the NFP/C, NFP/C-H and NFP/C-T_{0.01} samples powder dispersed in H₂O.

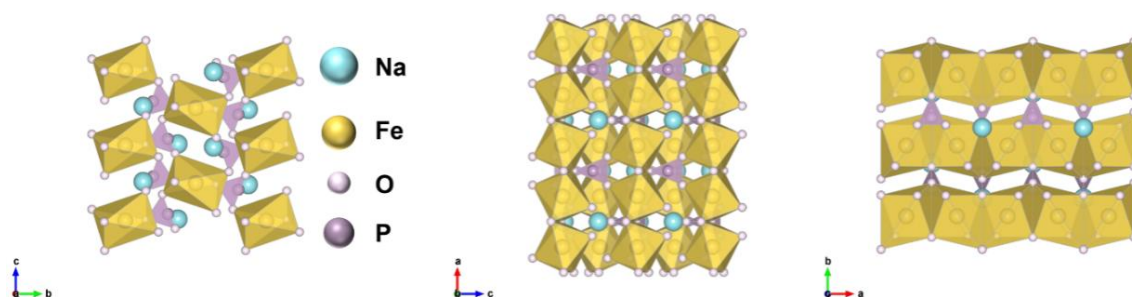


Figure S3. Crystal structure schematics of NFP along different orientations.

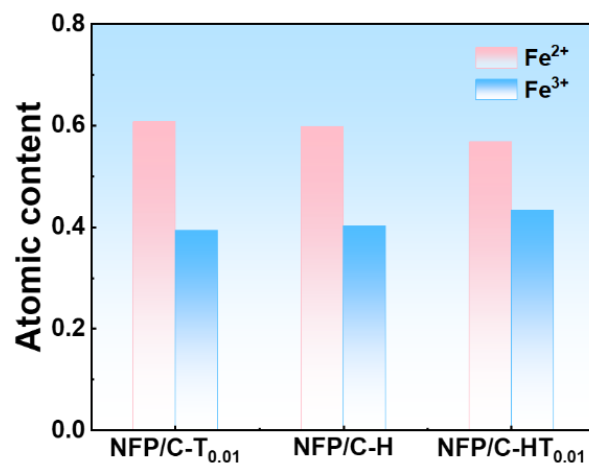


Figure S4. Ratio of Fe²⁺ to Fe³⁺ in NFP/C- T_{0.01}, NFP/C-H, and NFP/C-HT_{0.01}.

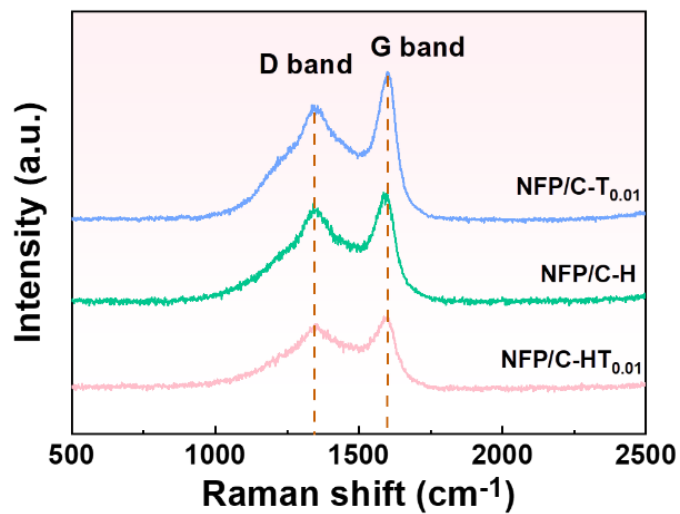


Figure S5. Raman spectra of NFP/C- $T_{0.01}$, NFP/C-H and NFP/C-HT $_{0.01}$ samples.

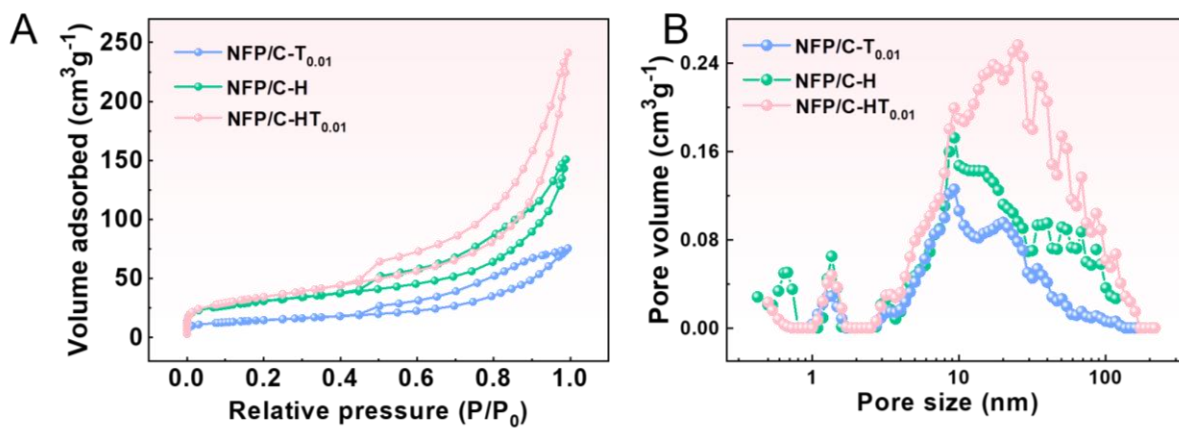


Figure S6. (A) N_2 adsorption-desorption isotherm and (B) pore size distribution curve of NFP/C-H, NFP/C- $T_{0.01}$ and NFP/C-HT $_{0.01}$ samples.

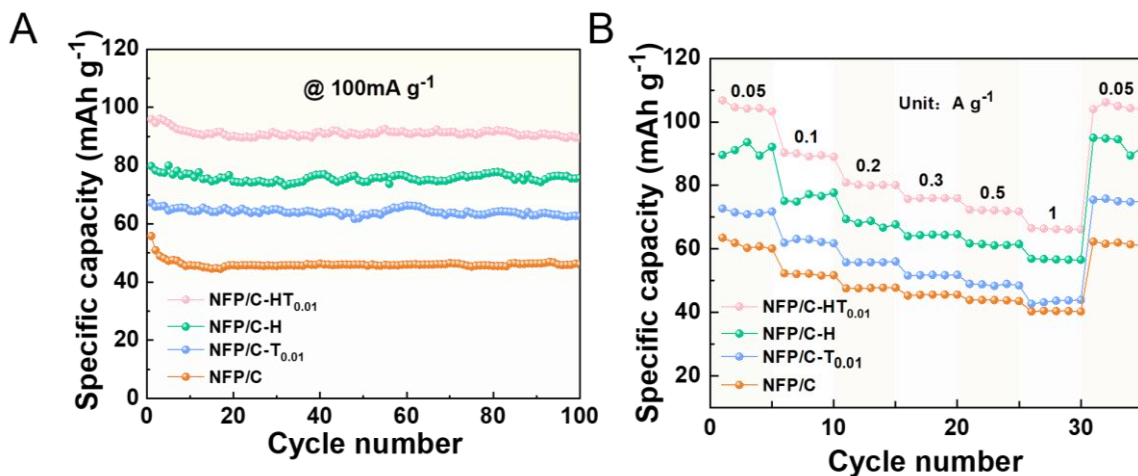


Figure S7. (A) Cycle performance at a current density of 100 mA g⁻¹ and (B) rate performance of NFP/C, NFP/C-H, NFP/C-T_{0.01} and NFP/C-HT_{0.01} samples.

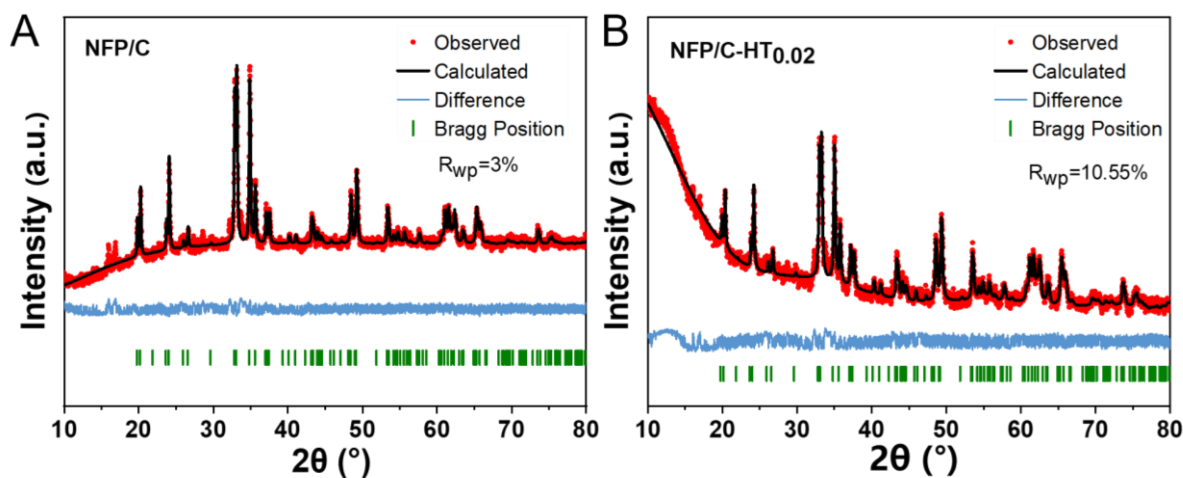


Figure S8. Rietveld refinements of NFP/C (A) and NFP/C-HT_{0.02} (B).

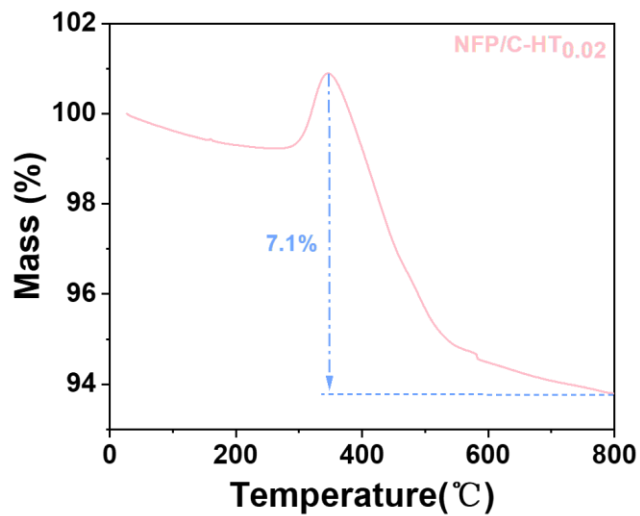


Figure S9. TGA of NFP/C-HT_{0.02} in air atmosphere.

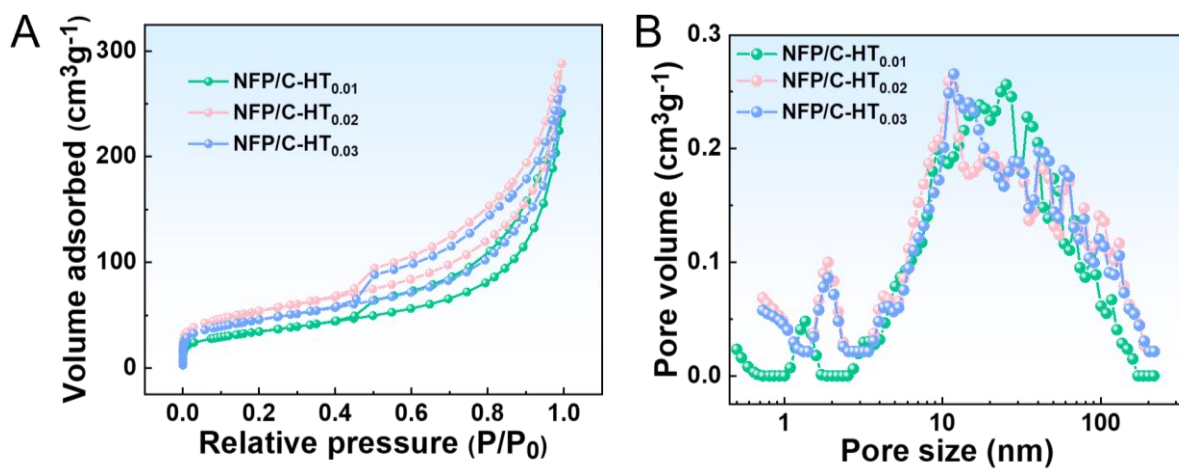


Figure S10. (A) N₂ adsorption-desorption isotherm and (B) pore size distribution curve of NFP/C-HT_{0.01}, NFP/C-HT_{0.02} and NFP/C-HT_{0.03} samples.

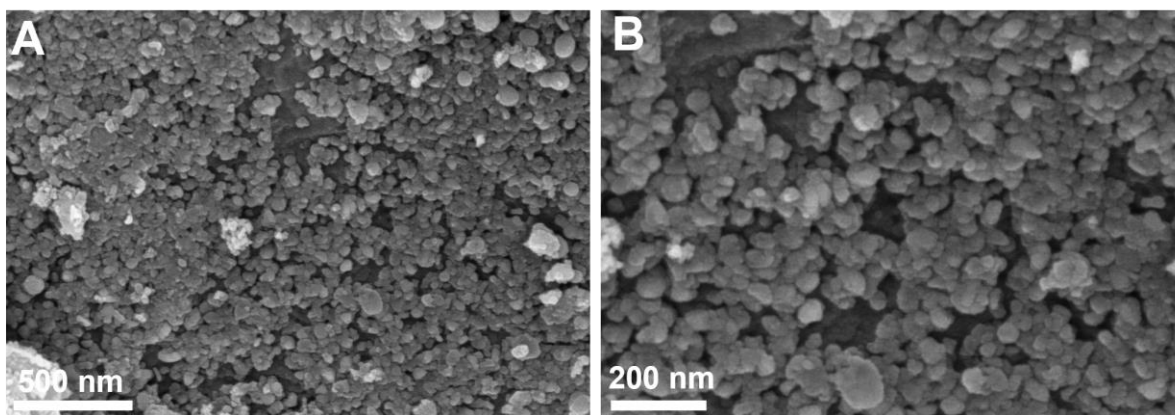


Figure S11. (A, B) SEM images of NFP/C-HT_{0.03}.

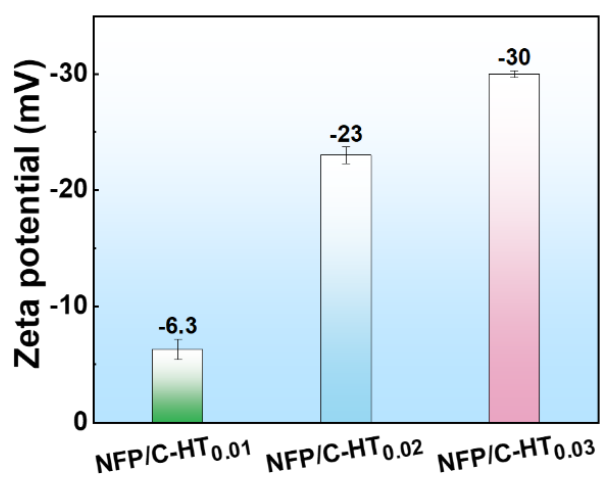


Figure S12. Zeta potential values of the NFP/C-HT_{0.01}, NFP/C-HT_{0.02} and NFP/C-HT_{0.03} samples powder dispersed in H₂O.

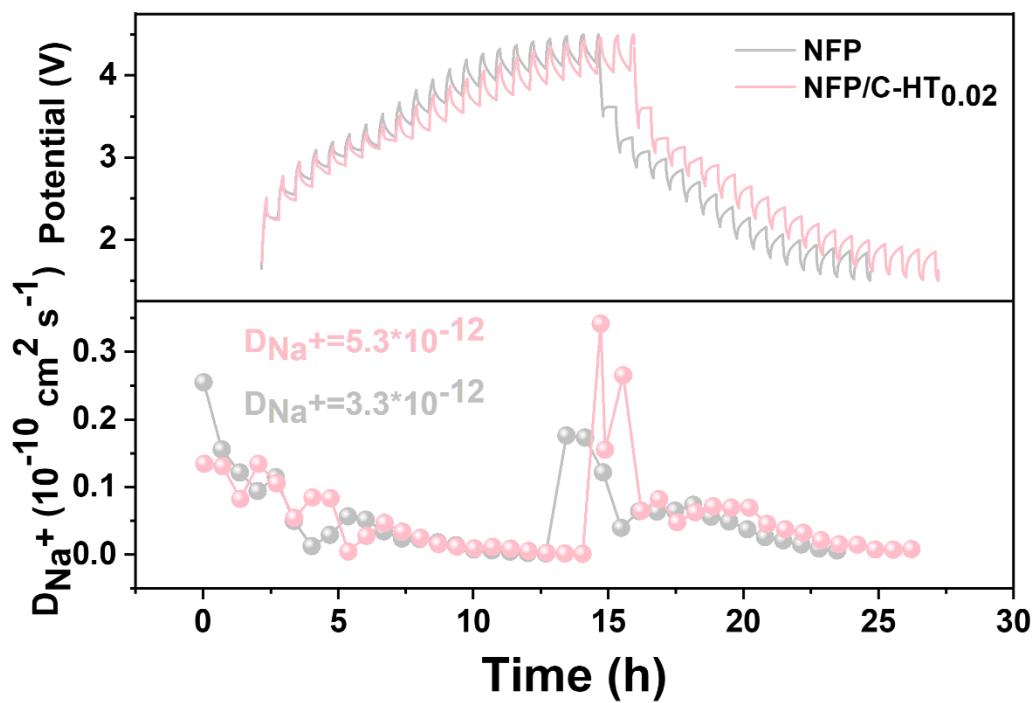


Figure S13. GITT curves and Na^+ diffusion coefficients of NFP/C-HT_{0.02} and NFP.

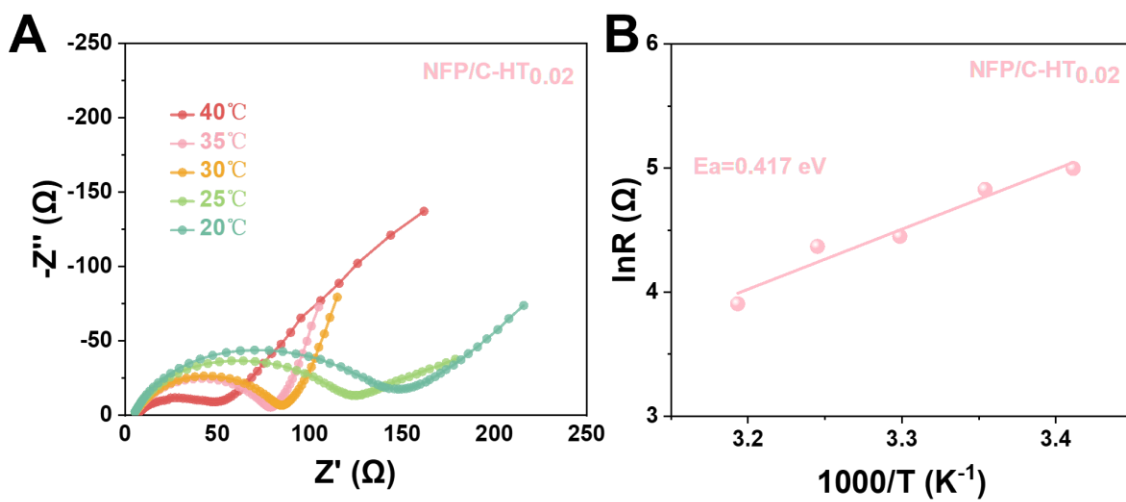


Figure S14. Nyquist plots of NFP/C-HT_{0.02} at different temperatures (A), and the temperature dependence of the corresponding resistances (B).

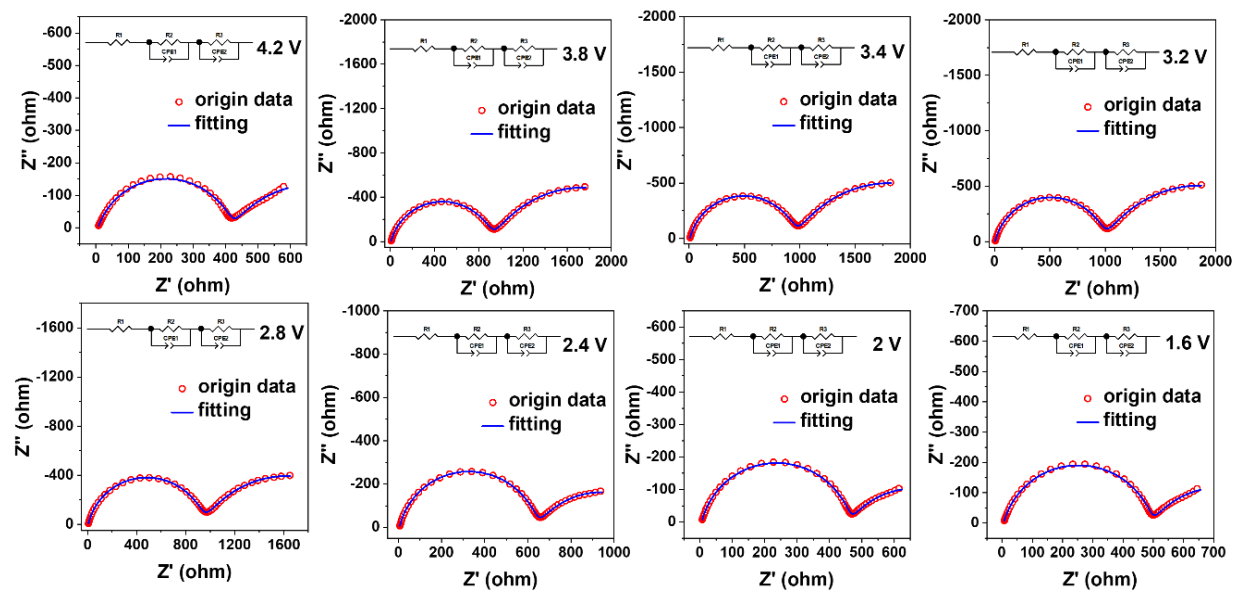


Figure S15. EIS of NFP/C-HT_{0.02} at Various Discharge Voltages.

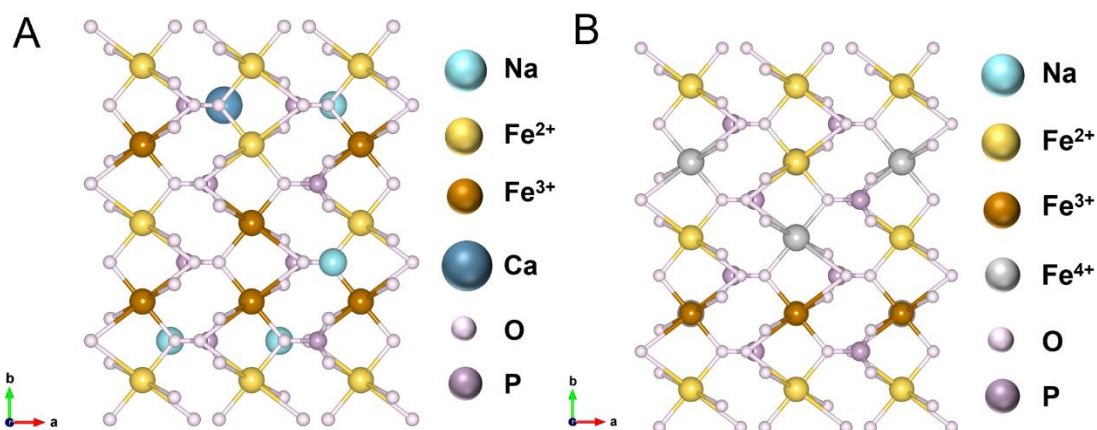


Figure S16. Crystal structure schematic of NFP featuring Fe³⁺ (A) and Fe⁴⁺ (B).