

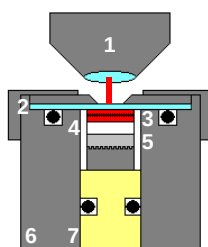


Combined *in situ* Raman and IR microscopy of electrode materials for lithium-ion batteries

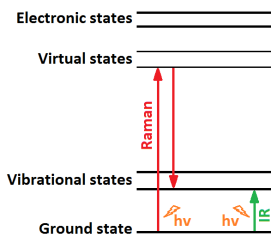
P. Lanz, S. Pérez-Villar, C. Villevieille and P. Novák

Paul Scherrer Institute, Electrochemistry Laboratory, CH-5232 Villigen PSI, Switzerland
patrick.lanz@psi.ch, +41 56 310 24 74

In situ cell / Principle / Approach



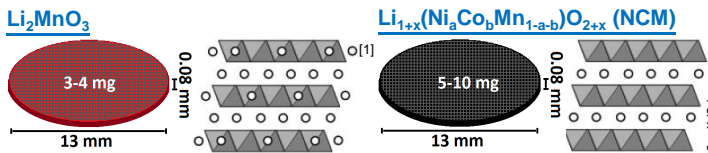
- 1 Objective (Raman or IR)
- 2 Window (CaF_2)
- 3 Working electrode (stainless steel mesh)
- 4 Separator (glass fibre)
- 5 Counter/reference electrode (Li)
- 6 Cell body (stainless steel)
- 7 Current collector (Ti in PEEK) (PEEK = polyether ether ketone)



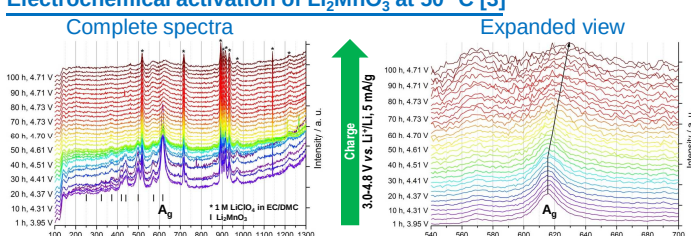
Combined *in situ* microscopic approach

- **Raman:** Particularly sensitive to structural changes in the electrode material
- **IR:** Particularly sensitive to the interface with the organic electrolyte
- ***In situ:*** Avoids relaxation & determines the exact potential of electrochemical processes
- **Microscopy:** Allows lateral resolution
- **Automation:** Switches between Raman & IR

Positive electrodes (Raman microscopy)



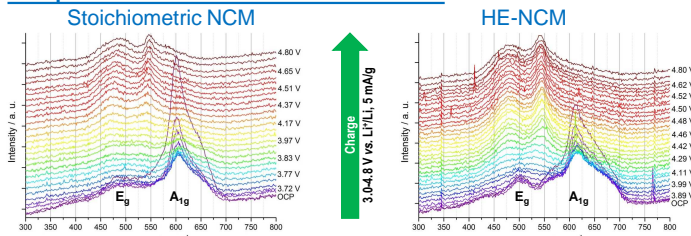
Electrochemical activation of Li_2MnO_3 at 50 °C [3]



➤ Observation: Shift of A_g to higher wavenumbers starting at 4.4 V

➤ Electrochemical activation of Li_2MnO_3 above 4.4 V
In agreement with the potential plateau observed in HE-NCM

Comparison of stoichiometric and HE-NCM

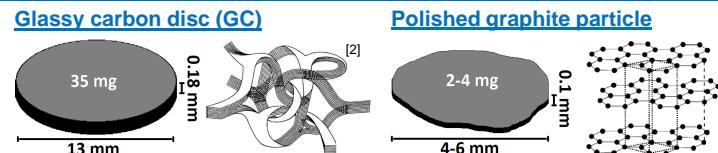


➤ Observation 1: Disappearance of A_{1g} (reversible, due to intercalation)

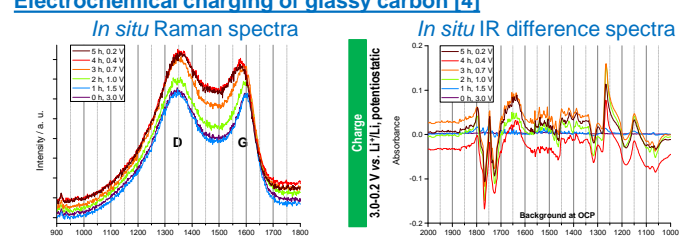
➤ Observation 2: New band at 545 cm⁻¹ (more pronounced in HE-NCM)

➤ New band at 545 cm⁻¹ possibly due to activation products (e.g. Li_2O) of Li_2MnO_3 integrated in the NCM structure
(Small amounts of Li_2MnO_3 may be present in stoichiometric NCM)

Negative electrodes (Combined microscopy)



Electrochemical charging of glassy carbon [4]



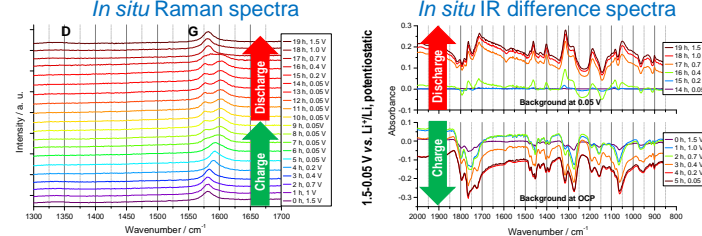
➤ Raman: No significant D and G shifts/splitting

➤ No significant lithium intercalation

➤ IR: Bipolar bands (separated by a few cm⁻¹) & positive band at 1650 cm⁻¹

➤ Increase in free and decrease in coord. EC/DMC & SEI formation

Electrochemical charging of a polished graphite particle



➤ Raman: G shifts/splitting (due to staging)

➤ Reversible Li-intercalation

➤ IR 1: Decrease/increase in free and coord. EC/DMC on charge/discharge

➤ IR 2: Pronounced jump in absorbance at the transition 0.4 V / 0.7 V

➤ Potential-dependent solvation effects & Li-intercalation

Free = not coordinated to Li^+ , coord. = coordinated to Li^+

Conclusions & Outlook

- Li_2MnO_3 is activated at 4.4 V vs. Li^+/Li when charged at 50 °C. Charging of NCM leads to the evolution of a new band at 545 cm⁻¹ (stronger in HE-NCM).
- Unlike GC, graphite intercalates Li. Strong solvation effects in both carbons. SEI products were detected on GC (band at 1650 cm⁻¹ due to EC reduction).
- Further *in situ* IR experiments to identify SEI products on polished graphite particles are planned.
- Combined *in situ* Raman and IR microscopy is a powerful method for the characterization of Li-ion batteries (electrode materials and electrolyte).

Acknowledgements

- Financial support from BASF SE is gratefully acknowledged.
- The authors are also grateful to Hermann Kaiser and Christoph Junker for technical assistance, for designing the *in situ* cells and for building the automation mechanism.

References

- [1] M.M. Thackeray et al., J. Mater. Chem., 2007, 17, 3115
- [2] G.M. Jenkins, K. Kawamura, Nature, 1971, 231, 175
- [3] P. Lanz et al., Electrochim. Acta, 2013, 109, 426-432
- [4] S. Perez-Villar et al., Electrochim. Acta, 2013, 106, 506-515

Scan me!

