

素子材料特論  
第3授業

## Li-ion電池負極(II)

### ハードカーボン系負極

#### Charge-Discharge Curves of Various Carbon Materials

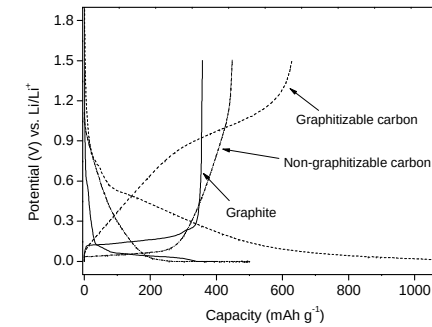


Figure 1-4. Charge-discharge profiles of representative carbon materials

#### Anodic performance and their related factors

| Performances                       | Factors                                                                                                                                                                                                    |
|------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Capacity                           | <ul style="list-style-type: none"> <li>Sites for Li incorporation</li> </ul>                                                                                                                               |
| Potential for charge and discharge | <ul style="list-style-type: none"> <li>Reversibility of charge and discharge</li> <li>Over potential</li> <li>Non-electrochemical reaction</li> </ul>                                                      |
| charge and discharge rate          | <ul style="list-style-type: none"> <li>Diffusivity of Li</li> </ul>                                                                                                                                        |
| Non-dischargeable charge           | <ul style="list-style-type: none"> <li>Reactivity of electrolyte</li> <li>Reactivity of anode, hetero atomic groups, terminal C-H, edge carbon</li> <li>Irreversible sites for Li incorporation</li> </ul> |
| Cycle ability                      | <ul style="list-style-type: none"> <li>Irreversible charge in structure</li> </ul>                                                                                                                         |
| Safety                             | <ul style="list-style-type: none"> <li>Stability of charged Li</li> <li>Li-Carbon intercalation</li> <li>Thermal stability of SEI</li> <li>Reactivity of electrolyte</li> </ul>                            |

#### Mechanisms for Lithium Insertion in Carbonaceous Materials

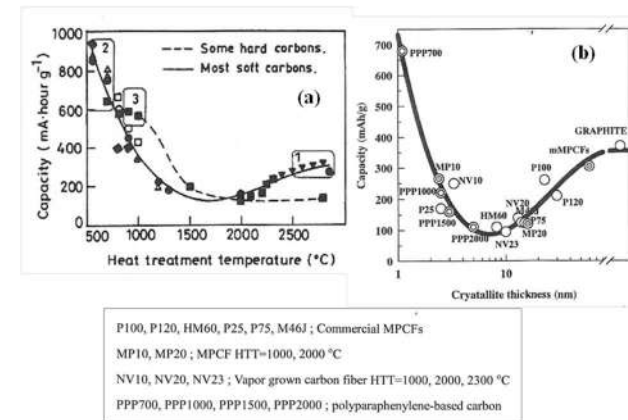


Figure 1-5. (a) Plot of reversible capacity for Li vs. HTT for a variety of carbon samples (□ hard carbon, ■ soft carbon), (b) Charge capacity as a function of the height of stacking (Lc002)

J. R. Dahn,\* Tao Zheng, Yinghu Liu, J. S. Xue SCIENCE, 270, 27 OCTOBER 1995

## Characteristics of various materials

|                                                  | Precursor                                                                                        | Advantages                                                                                 | Disadvantages                                                                                   |
|--------------------------------------------------|--------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------|
| Graphite<br>(over 2800°C)                        | Natural graphite<br>Artificial graphite<br>MCMB<br>Needle cokes<br>VGCF                          | Low discharge<br>potential<br>(around 0.2V)<br>Long cycle life                             | Low discharge capacity<br>(372 mAh/g)<br>High cost                                              |
| Graphitizable<br>carbon<br>(600~800°C)           | MCMB<br>Meso phase pitch<br>Green cokes                                                          | High capacity<br>(700~1000mAh/g)<br>Low cost                                               | High discharge potential<br>(around 1.0V)<br>High irreversible capacity<br>Poor cycle stability |
| Non-<br>graphitizable<br>carbon<br>(1000~1400°C) | Thermosetting polymer<br>Glassy carbon<br>Coal<br>Organic material<br>Stabilized isotropic pitch | High capacity<br>(400~700mAh/g)<br>Low discharge<br>potential<br>(around 0.1V)<br>Low cost | High irreversible capacity                                                                      |

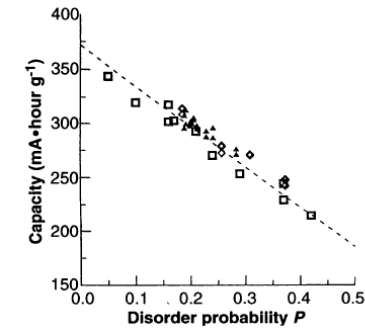


Fig. 3. Reversible capacity of region 1 carbons plotted as a function of the probability P of turbostratic disorder between adjacent carbon sheets. The line is the relation  $Q = 372(1 - P)$ , where Q is the capacity. For the purposes of this plot, samples corresponding to different symbols are equivalent.

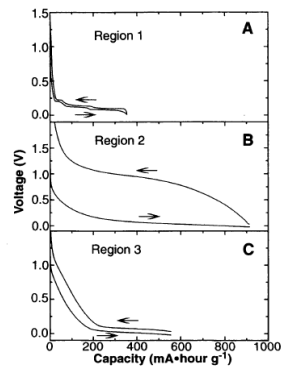
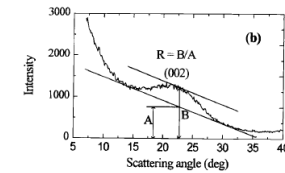
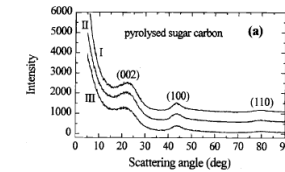


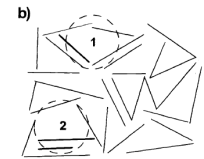
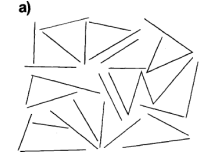
Fig. 2. Plots of voltage versus reversible capacity for the second charge-discharge cycle of representative carbon samples from regions 1, 2, and 3 of Fig. 1. (A) Synthetic graphite (Johnson-Matthey); (B) petroleum pitch (Crowley Tar Co.) heated to 550°C; (C) resole resin (Occidental Chemical Co.) heated to 1 000°C. Arrows designate the directions the curves are traversed as Li is added to (to the right) or removed from (to the left) the carbon samples.

## ハードカーボンの製造

- 原料: 難黒鉛化性前駆体
  - Biomass: Husk, Cellulose, Sugar, Rignin, Tree, Crab (Chichin), ...
  - Polymer: Phenol Resin, Unsaturated Resin, Epoxy Resin
  - Pitch: Isotropic Pitch and Coke
- 熱処理温度
  - 800~1400°C
- 構造
  - 難黒鉛化性カーボン

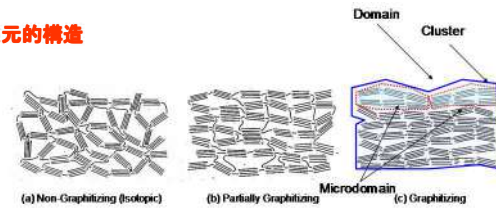


### FALLING CARDS MODEL

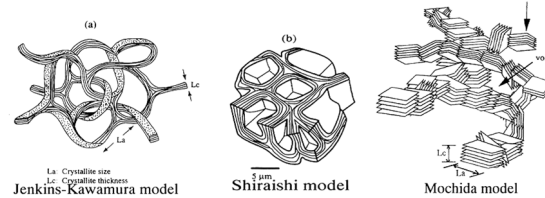


## ハードカーボンの構造

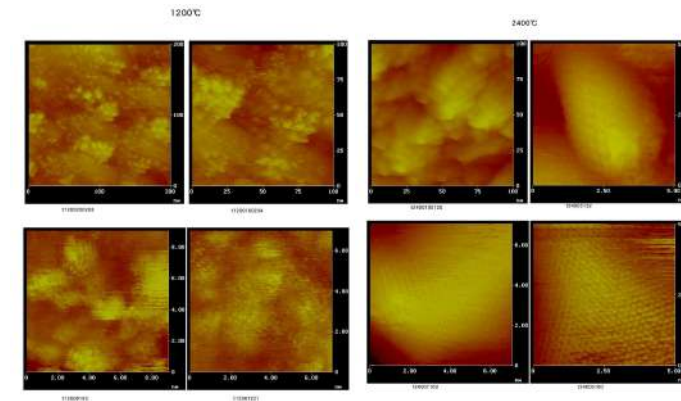
### 2次元の構造



### 3次元の構造



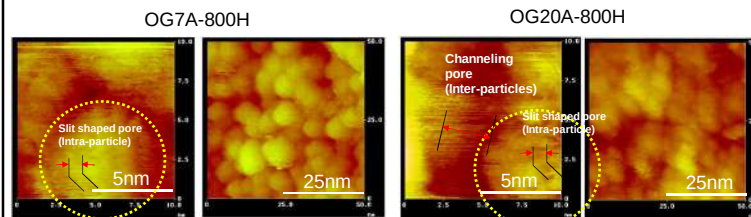
## ガラス炭素の微細構造



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## 活性炭素繊維の微細構造

In order to remove oxygen containing functional groups for removing the heterogeneous effect of STM, OG7A and OG20A were heat-treated at 800°C in a hydrogen atmosphere ( $H_2/He = 1/4$ ).



♣ Vacant spaces between the two domains of OG20A are larger than that of OG7A.

♣ Domain size of OG20A is a little smaller than that of OG7A.

♣ Slit type pores were observed in domains of OG7A and OG20A.

♣ It can be presumed that almost pores larger than 2nm nucleated by the inter-particle mechanism.

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## ハードカーボンの長所と短所

- 長所
  - 高容量が可能
  - 安価(?)
  - 資源が豊富(?)
  - High rate property
  - Hybrid系材料が可能
  - 低温特性がよい。
  - 電解液のPC使用可能
- 短所
  - 低密度
  - 放電プロファイルがNon-platten
  - 不純物の除去が困難
  - 低1st cycle Coulombic efficiency
  - 低Cycle性
  - ...

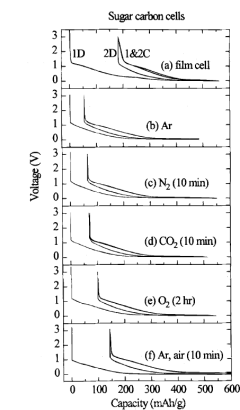


Fig. 2. Voltage profiles of super carbon/I cells: (a) film electrode prepared in air and (b-f) tablet electrodes exposed to different gases as indicated.

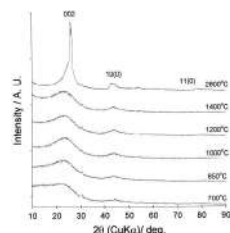


Fig. 1. Powder X-ray diffraction patterns of MNP-derived carbonaceous materials. The sharp (002) diffraction lines appearing at  $2\theta \approx 26$  and  $26.5^\circ$  in the 2800°C material come from the small portion of graphitized carbon.

Carbon 38 (2000) 995–1001

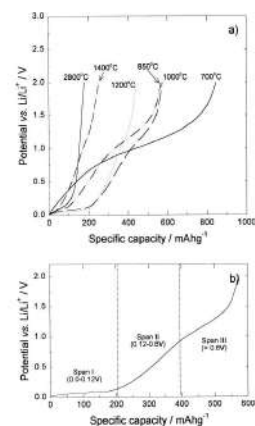


Fig. 2. Galvanostatic discharge curves of MNP-derived carbons. The profile obtained with the 1000°C-prepared material is separately displayed in (b). Before the tests, the carbon electrodes were galvanostatically charged down to 0 V and further short-circuited with the Li counter electrode for 96 h. The current density was  $30 \text{ mA g}^{-1}$  ( $0.15 \text{ mA cm}^{-2}$ ). Note the change of slope at three different potential regions in (b).

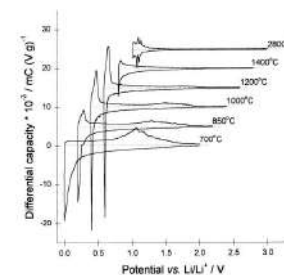


Fig. 4. EVS (electrochemical voltage spectroscopy) profiles obtained with the MNP-derived carbons. In each profile, the upper traces correspond to discharging ( $\text{Li}^+$  de-storage) and the bottom ones to charging ( $\text{Li}^+$  storage).  $i_{\text{discharge}} = 0.01 \text{ mA}$  and voltage step =  $5 \text{ mV}$ .

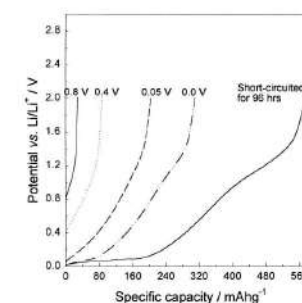
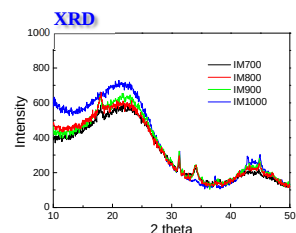


Fig. 5. Galvanostatic discharge curves recorded with the 1000°C material. The charging end is indicated. The experimental conditions were the same as for Fig. 2.

## Hard Carbon from Indonesian Mangrove Char

### Elemental analysis & Surface area

|         | Elemental analysis (wt%) |     |      |      |     | S.A.<br>( $\text{m}^2/\text{g}$ ) |
|---------|--------------------------|-----|------|------|-----|-----------------------------------|
|         | C                        | H   | N    | O    | Ash |                                   |
| IM      | 70.6                     | 3.6 | 0.38 | 23.7 | 1.7 | -                                 |
| IM 700  | 89.4                     | 1.0 | 0.72 | 582  | 3.1 | 318                               |
| IM 800  | 90.0                     | 0.8 | 0.74 | 5.5  | 3.0 | 19                                |
| IM 900  | 91.0                     | 0.6 | 1.17 | 4.3  | 3.0 | 3                                 |
| IM 1000 | 91.5                     | 0.5 | 1.13 | 4.0  | 2.9 | 54                                |



### STM image of IM1000

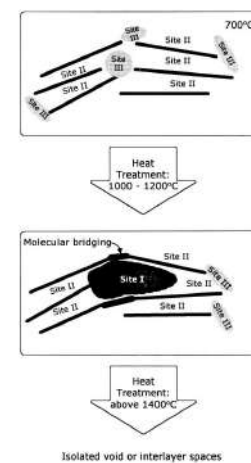
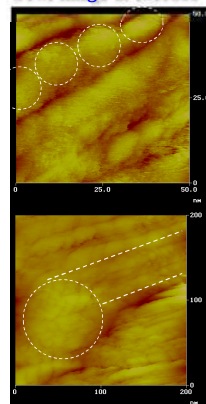


Fig. 7. A schematic illustration for the evolution of site I with heat-treatment temperature.

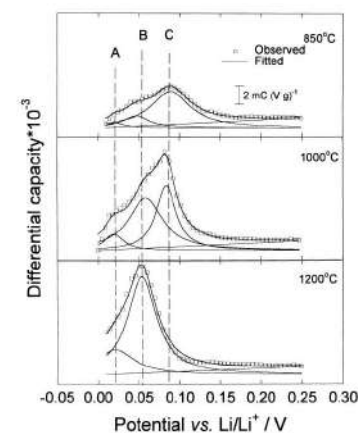
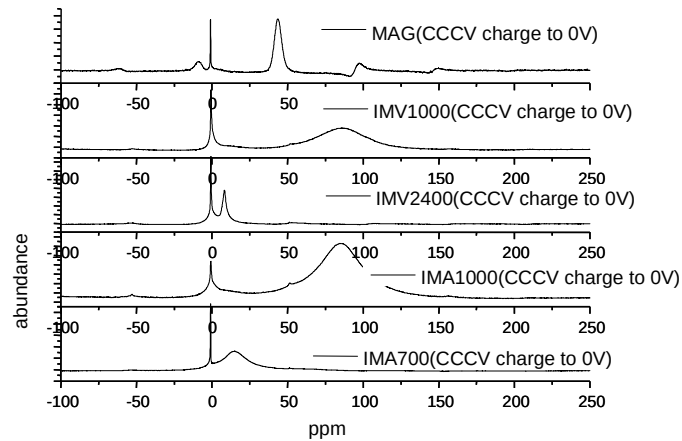


Fig. 8. Expanded view of the discharging EVS profiles near 0 V. The curves were fitted with three Lorentzian peaks. Note the evolution of peaks B and C with the heat treatment temperature.

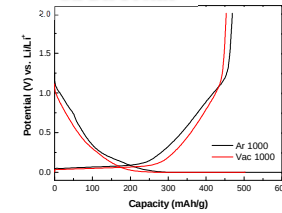
"Li+ storage sites in non-graphitizable carbons prepared from methylnaphthalene-derived isotropic pitches(MNIP)", *Carbon*, 38, 995 - 1001 (2000), C. W. Park, S. I. Lee, S.H. Yoon, S. M. Oh

## Li-NMR of Various Carbons



## Hard Carbon from Indonesian Mangrove Char

### Ch-Dis Profile

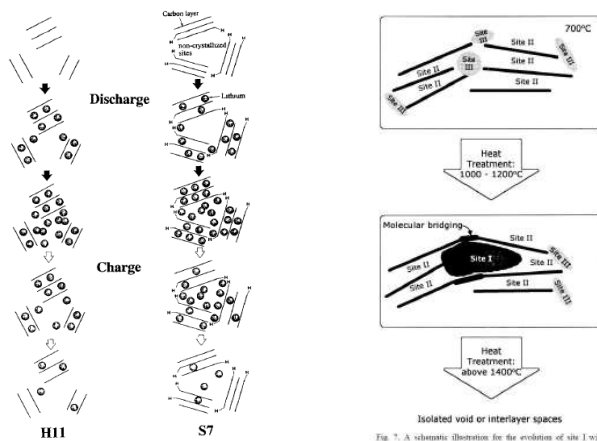


|    | Condition | Ch<br>(mAh/g) | Dis<br>(mAh/g) | Efficiency<br>(1cy, %) | Cap.avec.<br>(mAh/g) |
|----|-----------|---------------|----------------|------------------------|----------------------|
| IM | Ar 1000   | 628           | 469            | 74.7                   | 159                  |
|    | Vac 1000  | 503           | 453            | 90.1                   | 50                   |

IM V1000 showed the better initial efficiency than that of IM Ar1000

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## ハードカーボンの充放電機構



Low-crystallized carbon materials for lithium-ion secondary batteries,  
Hayato Higuchi, Keiichi Uenae, Akira Kawakami  
JOURNAL OF POWER SOURCES, 68, 1997

## 理想的なハードカーボン

- 高密度: 1.9g/cm<sup>3</sup>以上
- 高容量: 650mAh/cc
- 高レート性: 90%以上 (5C/0.2C: Half cell, 20C/0.2C: Single cell)
- Low Impurity: 100ppm以下
- 安価: 800円/Kg以下
- 高1st Coulombic efficiency
- 高低温特性
- 高サイクル性
- 豊富な原料からの調製
- マイルドな炭化条件
- Hybrid系が可能な材料: SnOxまたはSiとのハイブリッド化⇒高容量化

⇒ 達成するためには、各々因子に対応する原因把握が要求。

⇒ まだ、原因説明が明らかになっていない。

⇒ 今後の研究に期待する。

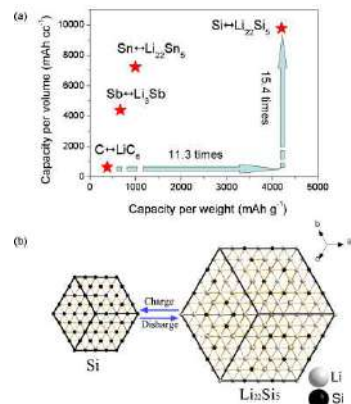


Figure 1-15. Typical high performance materials as anode ;  
(a) Comparison of graphite, Si, Sn and Sb  
(b) Volume expansion model of Si during charge

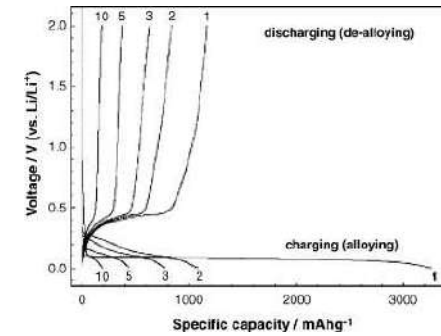


Fig. 2. Galvanostatic charge-discharge profiles for micro-Si (10 μm) anode.

J.H. Ryu, J.W. Kim, Y.-E. Sung, S.M. Oh  
Electrochem. Solid State Lett., 7 (2004), p. A306

## High performance material & their problems

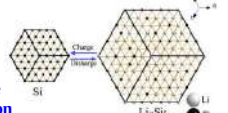
- **SiO, Si and Sn** (SiO 2100, Si 4200, Sn 931 mAh/g) are very promising materials as anodic materials of LIB for their large theoretical capacities, however, they have poor cycle performances because of internal crack in particles caused by large volumetric expansion in charge process.

### Li-Si system

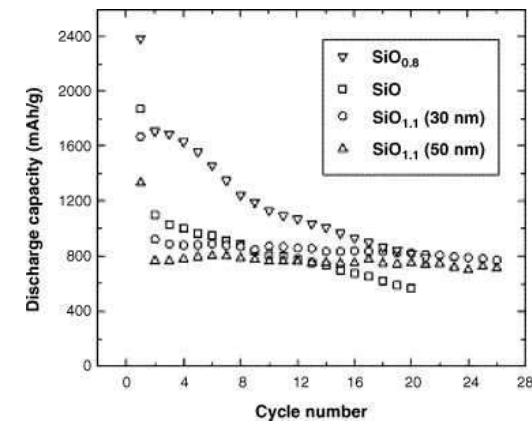
| Compound                         | Structure    | Unit cell vol. (Å <sup>3</sup> ) | Vol. / Si atom (Å <sup>3</sup> ) |
|----------------------------------|--------------|----------------------------------|----------------------------------|
| Si                               | Cubic        | 160.2                            | 20.0                             |
| Li <sub>12</sub> Si <sub>7</sub> | Orthorhombic | 243.6                            | 58.0                             |
| Li <sub>14</sub> Si <sub>6</sub> | Rhombohedral | 308.9                            | 51.5                             |
| Li <sub>13</sub> Si <sub>4</sub> | Orthorhombic | 538.4                            | 67.3                             |
| Li <sub>22</sub> Si <sub>5</sub> | Cubic        | 659.2                            | 82.4                             |

Volume expansion (over 400%)

Ref.) J. Power Sources 192 (2) (2009) 644-651

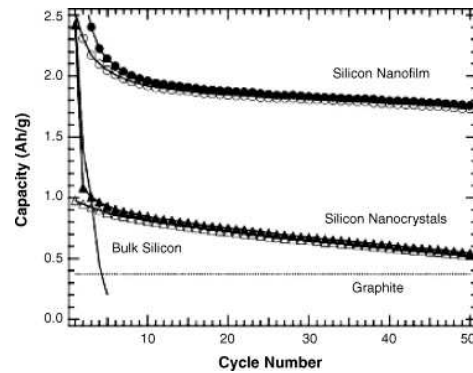


Ref. : A. John Appleby and et al., J Power Sources 163 (2007) 1003-1039



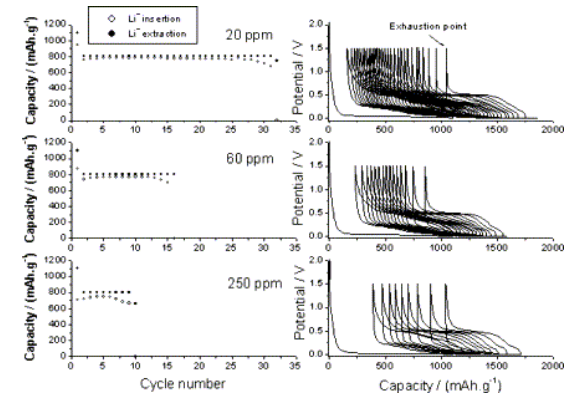
Cycling behavior of SiO<sub>x</sub> with different oxygen content and particle size

J. Yang, Y. Takeda, N. Imanishi, C. Capiglia, J.Y. Xie, O. Yamamoto  
Solid State Ionics, 152-153 (2002), p. 125



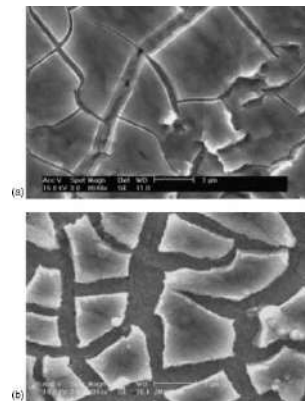
Specific capacity vs. cycle number for nano-crystalline Si and nano-amorphous Si thin film anodes prepared by thermal vapor deposition. Specific capacity of graphite and bulk-silicon anodes also shown

J. Graetz, C.C. Ahn, R. Yazami, B. Fultz  
Electrochem. Solid-State Lett., 6 (2003), p. A194



Cycling performance of carbon-coated silicon anodes in 1 M  $\text{LiPF}_6$  EC:DMC (1:2) at different water contents present in the electrolyte. In all cases, the 'exhaustion point' (last extraction peak) is centered around 1000 mA h/g

[Electrochimica Acta](#)  
Volume 48, Issue 11, 15 May 2003, Pages 1579–1587



SEM morphology of 250 nm a-Si film on Cu cycled at C/2.5 for (a) 1 cycle, and (b) 30 cycles

J.P. Maranchi, A.F. Hepp, A.G. Evans, N.T. Nuhfer, P.N. Kumta  
J. Electrochem. Soc., 153 (2006), p. A1246

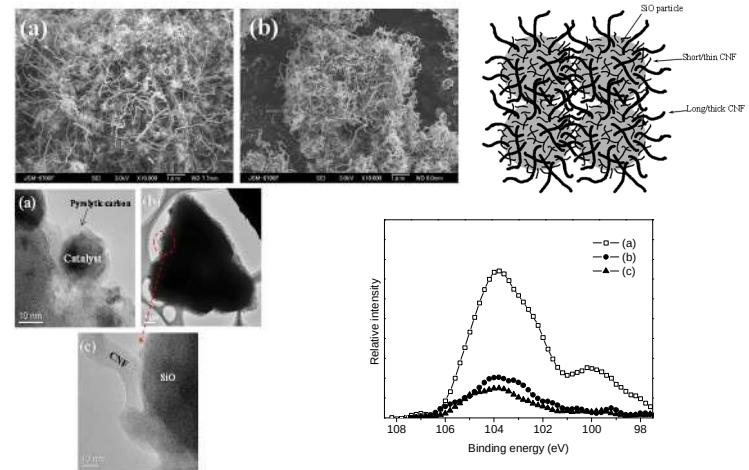


Figure Schematic model of cocoon shaped composite of silicon monoxide and surficial carbon nanofiber (SiO-CNF composite)



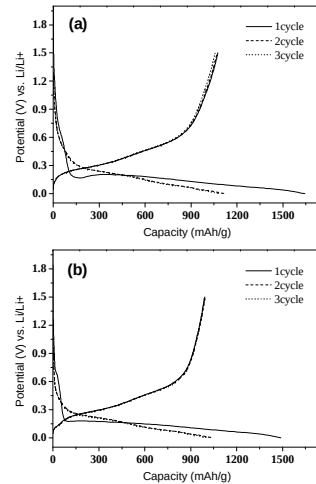


Figure 2-11. Cycle performances with SiO-CNF composite until 3 cycles according to the various amount of CNFs with Ni (a) or Fe catalyst (b).

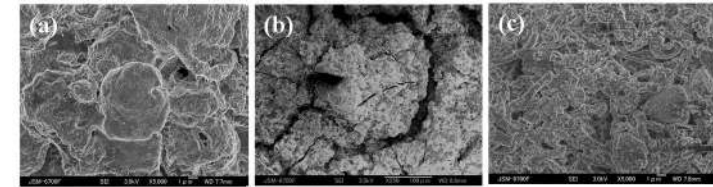
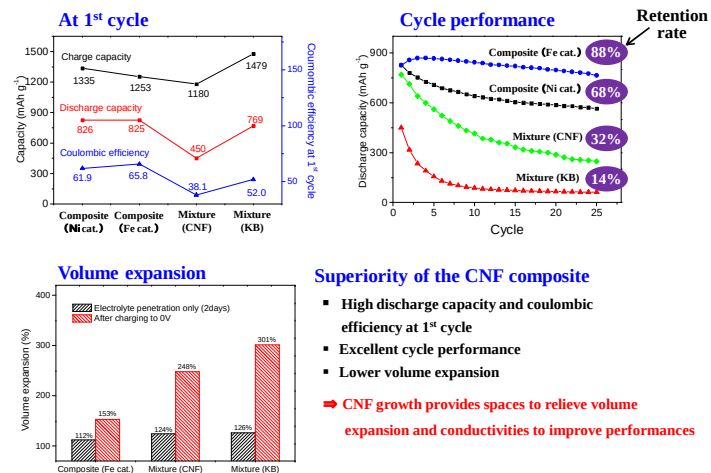


Figure 2-12. SEM images of electrode surface after 3 cycles : (a) SiO-CNF composite with Fe catalyst, mixture of SiO-KB (b) or SiO-CNF (c).

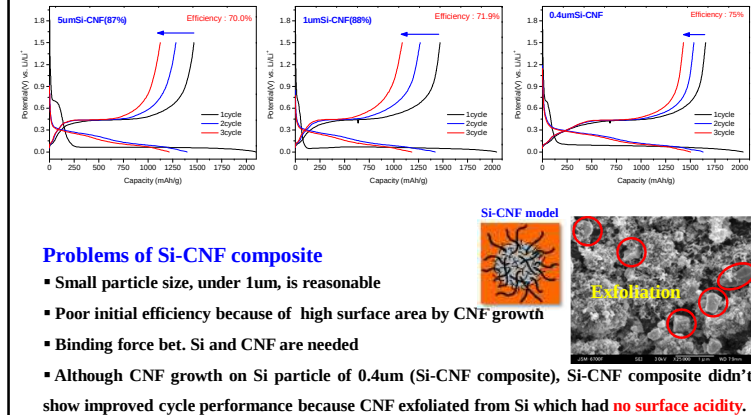
### Comparison bet. Composite and Mixture



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### Problem in Si-CNF composite

Si-CNF composite : CNF growth directly on Si(5um, 1um, 0.4um)



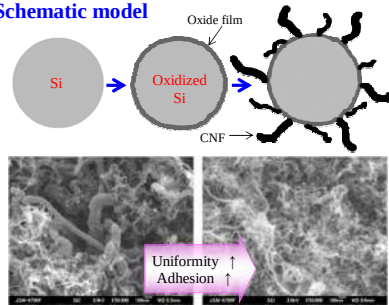
32



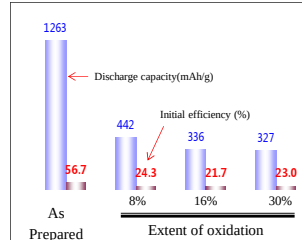
### Oxidized Si-CNF composite

To improve surface acidity,  
Si particle was oxidized at 700~900°C for 3 hrs with  $\text{H}_2\text{O}_\text{g}$ .

#### Schematic model



#### Performance test



Adhesive power and uniformity of CNF was improved by surface oxidized Si, where as charge-discharge performance became poor.

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### Preparation of PCSi-CNF composite

|            | Code     | PC/ or CNF amount (wt. %) |      |               | Condition                                        |
|------------|----------|---------------------------|------|---------------|--------------------------------------------------|
|            |          | PC                        | CNF  | PC re-coating |                                                  |
| Samples    | Step I   | Si-PC                     | 6 %  | -             | PC coating (900°C-CH <sub>4</sub> /He- 30min)    |
|            | Step II  | Si-PC-CNF                 | 6 %  | 93 %          | CNF growth on PCSi (580°C-CO/He- 30min)          |
|            | Step III | Si-PC-CNF-RC              | 6 %  | 93 %          | Catalyst removal by HCl                          |
|            | Step IV  | Si-PC-CNF-RC-PC           | 6 %  | 93 %          | PC re-coating (900°C-CH <sub>4</sub> /He- 30min) |
| Comparison | Si-CNF   | -                         | 98 % | -             | CNF growth directly on Si surface                |

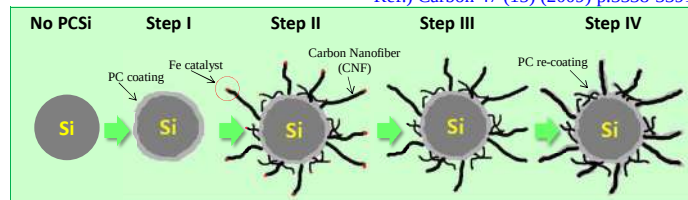
35

### Composite of PC coated Si and CNF

As a solution, Pyrolytic carbon (PC) coating on Si particle is suggested.

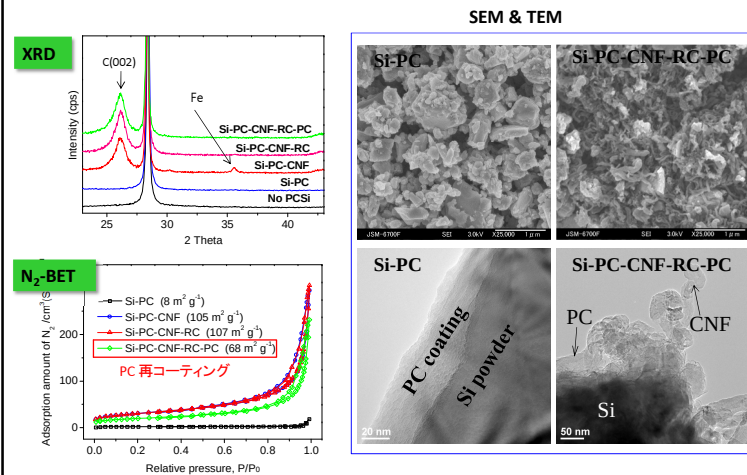
- Step I : PC is coated on Si particle to improve adhesion between Si and CNF (PCSi).  
⇒ The coated PC provide conductivity with Si particles as well as adhesion strength.
- Step II : CNF is grown on PCSi to improve cycle performance
- Step III : Removal catalyst by HCl treatment
- Step IV : PC is re-coated on PCSi-CNF composite to improve initial efficiency by decreasing surface area.

Ref.) Carbon 47 (15) (2009) p.3338-3391

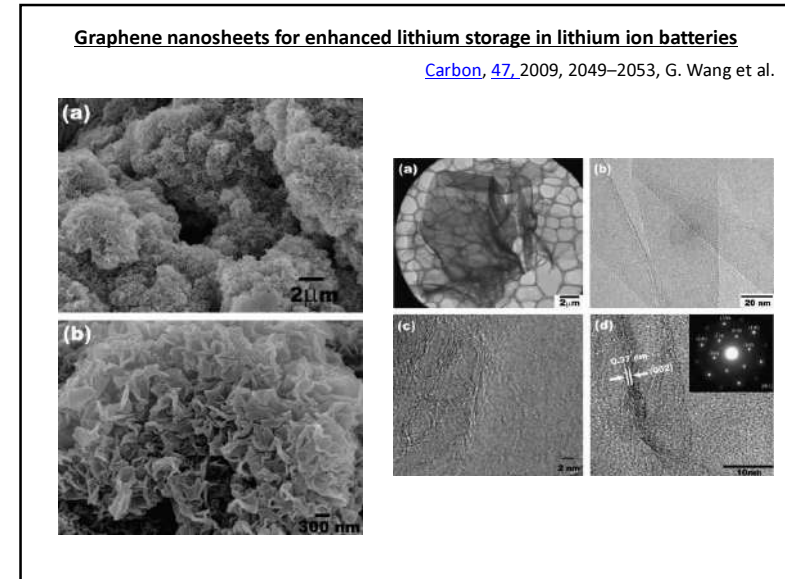
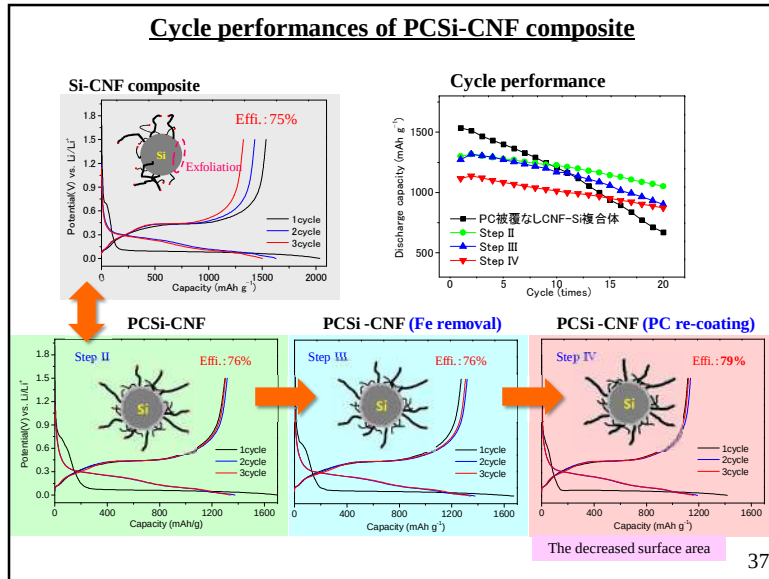


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### Analysis of PCSi-CNF composite



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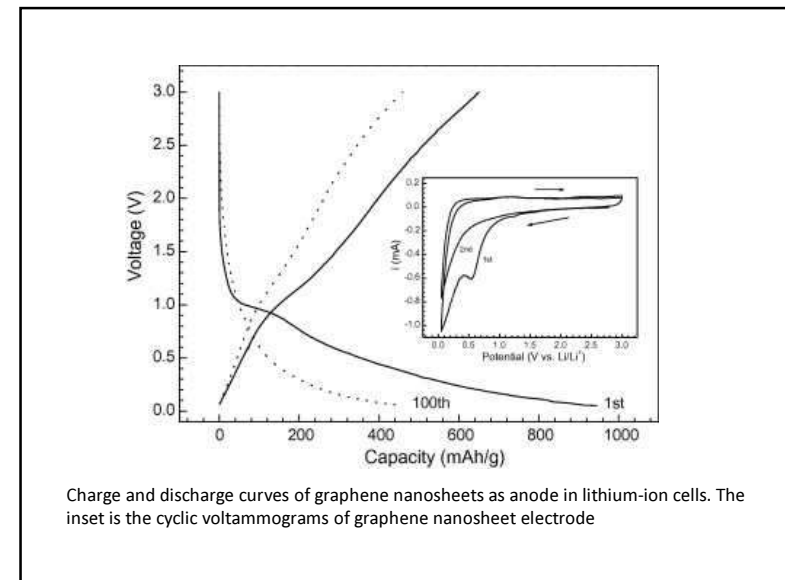
### Cycle performances of PCSi-CNF composite

|            |                 | Capacity at 1 <sup>st</sup> cycle (mAh/g) |      |       |
|------------|-----------------|-------------------------------------------|------|-------|
|            |                 | Ch                                        | Dis  | 効率(%) |
| Samples    | Si-PC-CNF       | 1709                                      | 1299 | 76.0  |
|            | Si-PC-CNF-RC    | 1674                                      | 1272 | 76.0  |
|            | Si-PC-CNF-RC-PC | 1415                                      | 1115 | 78.8  |
| Comparison | Si-CNF          | 2037                                      | 1535 | 75.4  |

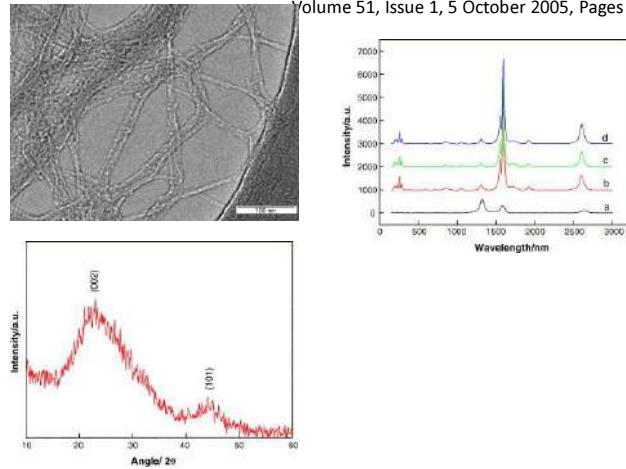
|            |                 | Dis.(max)<br>(mAh / g) | At 20 cycle (mAh / g) |                    |
|------------|-----------------|------------------------|-----------------------|--------------------|
|            |                 |                        | Dis                   | Retention rate (%) |
| Samples    | Si-PC-CNF       | 1317                   | 1051                  | 80                 |
|            | Si-PC-CNF-RC    | 1318                   | 903                   | 69                 |
|            | Si-PC-CNF-RC-PC | 1136                   | 873                   | 77                 |
| Comparison | Si-CNF          | 1535                   | 670                   | 44                 |

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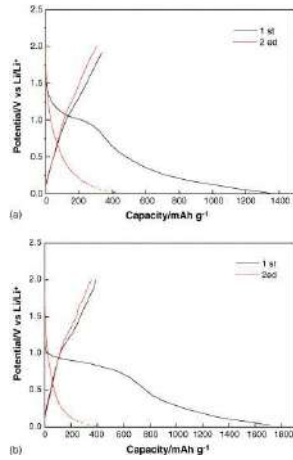
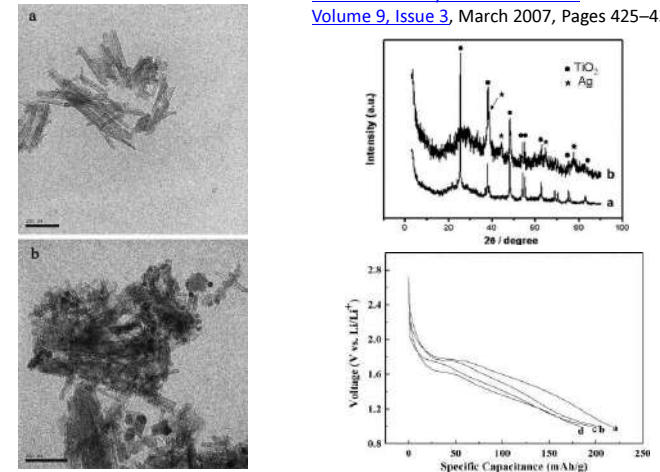
# Single wall carbon nanotube paper as anode for lithium-ion battery

Electrochimica Acta  
Volume 51, Issue 1, 5 October 2005, Pages 23–28



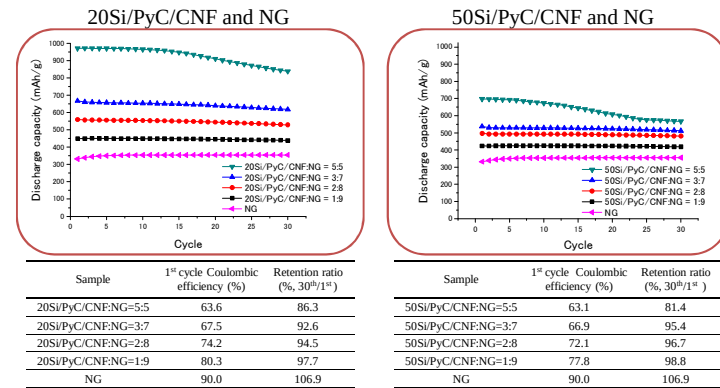
# Preparation and electrochemical properties of Ag-modified TiO<sub>2</sub> nanotube anode material for lithium-ion battery

Electrochimica Acta  
Volume 51, Issue 1, 5 October 2005, Pages 23–28



The charge/discharge profiles of SWNT electrodes: (a) conventional slurry coated electrode and (b) "Free standing" electrode.

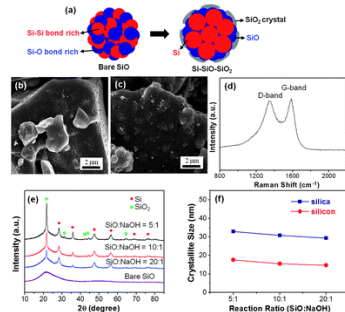
# Hybridization of Si/PyC/CNF and NG



✓ The hybrids of the Si/PyC/CNF and NG showed better cycle-ability than the hybrids of Si/PyC and NG.

### Highly stable Si-based multicomponent anodes for practical use in lithium-ion batteries

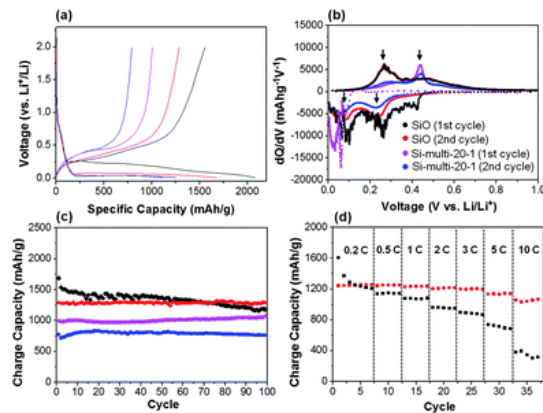
Jung-In Lee, Nam-Soon Choi and Soojin Park  
*Energy Environ. Sci.*, 2012, 5, 7878-7882  
 Interdisciplinary School of Green Energy, Ulsan National  
 Institute of Science and Technology, Ulsan, Korea 689-798



Synthesis of an Si-based multicomponent from bulk SiO particles via thermal annealing in the presence of NaOH. (a) Schematic illustration for the conversion of bare SiO to Si-SiO-SiO<sub>2</sub> three-components, (b) SEM image of Si-based multicomponents, SEM image (c) and Raman spectrum (d) of carbon-coated Si-based multicomponents, (e) XRD patterns of Si-based multicomponents as a function of NaOH amount, and (f) calculation of silica and silicon crystallite size as a function of NaOH amount

### まとめ

- Bulk Carbon以外は、長所と共に短所も持っており、まだLi-ion電池用負極としては商品化されていない。
- 今後の研究によって短所が解決できれば、電池の特性はより改善できる。



Electrochemical performances of c-SiO and c-Si-SiO-SiO<sub>2</sub> three-component electrodes. (a) Voltage profiles of c-SiO (black), c-Si-multi-20-1 (red), c-Si-multi-10-1 (pink), and c-Si-multi-5-1 (blue). (b) dQ/dV plots of c-SiO and c-Si-multi-20-1 in the first and second cycles. (c) Cycle performances of c-SiO (black), c-Si-multi-20-1 (red), c-Si-multi-10-1 (pink), and c-Si-multi-5-1 (blue) at 0.1 C rate. (d) Rate capabilities of c-SiO and c-Si-multi-20-1 electrodes. The discharge rate was fixed at a rate of 0.1 C