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**Theoretical Study of Novel Nanostructured Materials for Lithium-Ion Batteries**

**Mario Sanchez-Vazquez**  
**CENTRO DE INVESTIGACION EN MATERIALES AVANZADOS, S.C.**

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**Final Report**

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# Theoretical study of $\text{Si}_x\text{Ge}_y\text{Li}_z^-$ ( $x=4-10$ , $y=1-10$ , $z=0-10$ ) clusters for designing of novel nanostructured materials to be utilized as anodes for Lithium-ion batteries

## *Final Report*

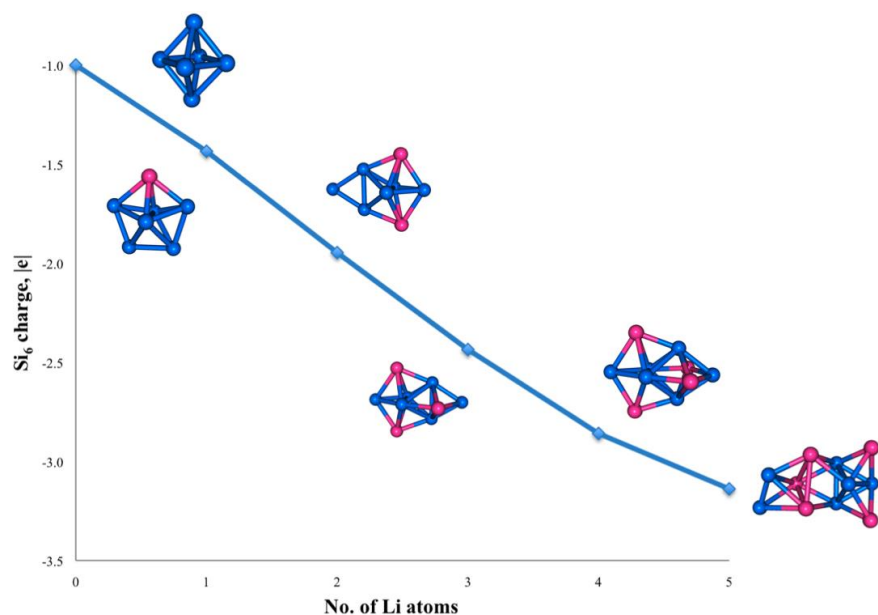
Nancy Perez-Peralta and Mario Sanchez-Vazquez

### **Abstract**

In order to find out if silicon nanostructured materials resist lithium cations insertion and extraction under the presence of germanium atoms, we have explored theoretically the potential energy surface of  $\text{Si}_6\text{Ge}^-$ ,  $\text{Si}_6\text{GeLi}_5^-$ ,  $\text{Si}_6\text{GeLi}_7^-$ ,  $\text{Si}_6\text{GeLi}_{10}^-$ ,  $\text{Si}_6\text{Ge}_2^-$ ,  $\text{Si}_6\text{Ge}_2\text{Li}_5^-$ , and  $\text{Si}_{10}\text{Ge}_{10}\text{Li}_{10}^-$  clusters. This study was performed using the Kick Coalescence method coupled to Gaussian 09 program, and re-optimization all structures with the B3LYP/def2-TZVPP method. Our results confirm that incorporating germanium atoms to silicon clusters improves and prevents fragmentation.

### **Introduction**

Rechargeable Li-ion batteries are the most promising energy storage devices for hybrid, plug-in hybrid electric, and all-electric vehicles. Lithium is an ideal material for batteries: it is the lightest metal in addition to having a high electric potential. Charging a Li-ion battery usually means moving lithium cations from cathode into anode. There is an increasing interest in developing rechargeable lithium batteries with higher energy capacity and longer cycle life for applications in portable electronic devices and electric vehicles. Currently, the graphite anode is the most commercially used due to its good capacity ( $372 \text{ mA h g}^{-1}$ ) together with its rate capability and long life. Silicon has recently become very popular as a potential anode material for lithium batteries because it has a low discharge potential and the highest known theoretical charge capacity (which could be 10x that of graphite).<sup>1-4</sup> However, silicon anodes have limited applications because of the large volume change upon lithium cations insertion or extraction. Silicon nanowires have been shown to be promising as high-performance lithium battery anodes because they can accommodate large strains derived from lithium charging or discharging.<sup>3</sup>



**Figure 1.** Si<sub>6</sub> skeleton charges for every global minimum structure vs. the number of lithium atoms are plotted. As one can see, the addition of five lithium atoms splits the Si<sub>6</sub> skeleton into two fragments: a S<sub>4</sub> kernel and a two-silicon fragment.

Very recently, an *ab initio* study on the lithiation of the Si<sub>4</sub><sup>-</sup> cluster was reported. Results of this study revealed that the maximum formal charge transfer from alkali metals to Si<sub>4</sub> system is four and that the Si<sub>4</sub><sup>-</sup> tetrahedral kernel is a robust building block.<sup>5</sup> However, a study on the lithiation of the Si<sub>6</sub><sup>-</sup> shows that lithium cations are able to split the silicon skeleton into two fragments: the S<sub>4</sub> kernel and a two-silicon fragment (see Figure 1). This result actually gives an explanation to the large volume change of silicon anodes upon lithium cations insertion. The latter study suggests that even though the tetrahedral Si<sub>4</sub><sup>4-</sup> structure is resistant, any material designed upon it will be broken up by lithium cations.<sup>6</sup> Song et al. have recently reported that Si/Ge double-layered nanotubes (Si/Ge DLNT) are promising materials as anodes for Li-ion batteries. Compared to silicon nanotubes, Si/Ge DLNT improve both cyclability and rate capability.<sup>7</sup> It is apparent from the last results that germanium atoms play a key role in the silicon structure stabilization upon lithium cations insertion and extraction. Therefore, we have proposed to investigate both qualitatively and quantitative germanium effects on silicon skeletons.

### Computational details

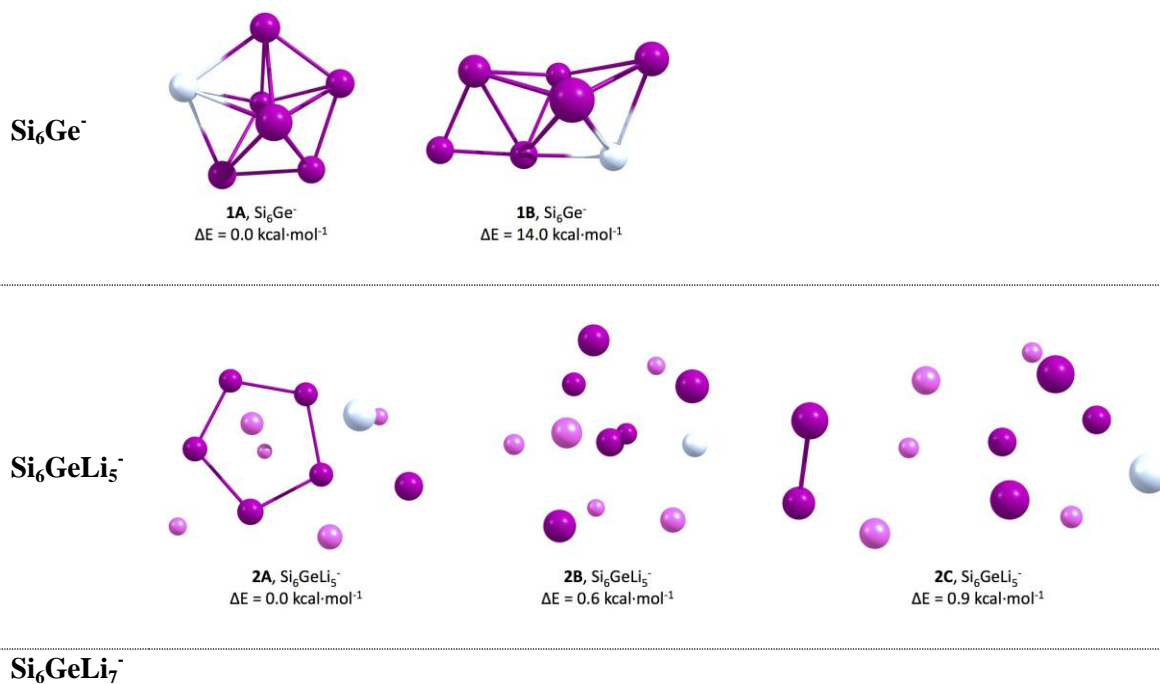
In order to find out if silicon nanostructured materials resist lithium cations insertion and extraction under the presence of germanium atoms, we have been studying theoretically the lithiation process of the Si<sub>x</sub>Ge<sub>y</sub><sup>-</sup> (x=4-10, y=1-10) clusters. At the moment, we have explored the potential energy surfaces of the Si<sub>x</sub>Ge<sub>y</sub>Li<sub>z</sub><sup>-</sup> (x=6, y=1-2, z=0-10) clusters throughout global minimum search techniques, i.e., Kick Coalescence method.<sup>8</sup>

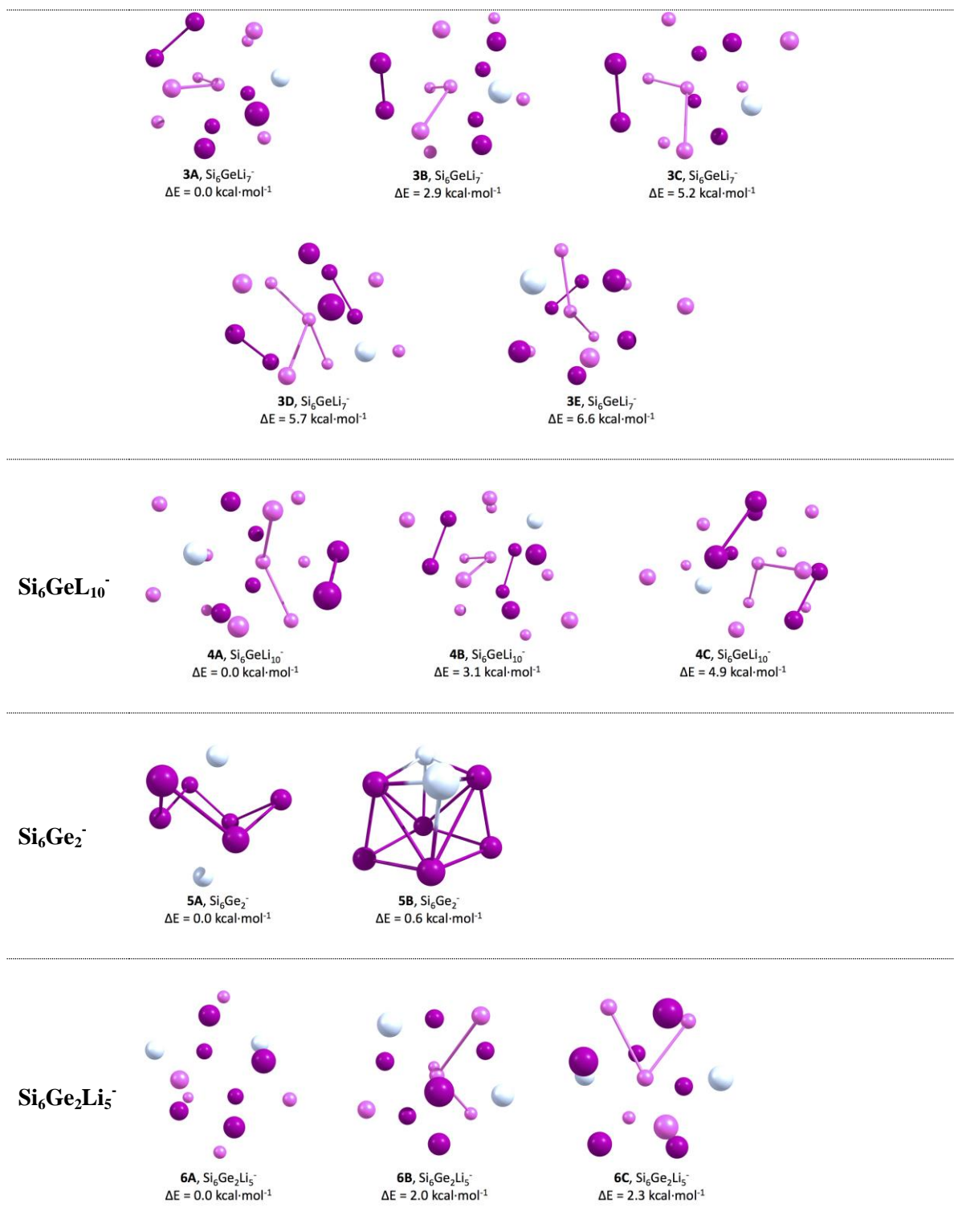
In order to find the global minimum structure of any cluster, a comprehensive scanning on the corresponding potential energy surface has been performed using the Kick Coalescence

method. Kick Coalescence (KC) method, is particularly powerful given that it combines the accuracy of *ab initio* and DFT approximations with a very fast scanning of any potential energy surface; in addition, it is unbiased. In this method, a very large population of structures that are generated randomly is subjected to a coalescence procedure with all atoms being pushed gradually to the molecular center of mass to avoid fragmented structures. Subsequently, these structures are optimized to the nearest local minimum using an electronic structure package. Currently, the KC method is coupled to Gaussian 09; but it can be easily coupled to any software that calculates electronic structure of atoms and molecules. At this stage, a low level of theory is required in order to get a fast scanning. In previous works on silicon clusters, reliable results have been got using the combination of the hybrid functional B3LYP with the 3-21G basis set. This optimization process was followed by a reoptimization and frequency calculation at a higher level of theory; B3LYP/def2-TZVPP has been used in the present research.

## Results

At the moment, we have explored the potential energy surface of  $\text{Si}_6\text{Ge}^-$ ,  $\text{Si}_6\text{GeLi}_5^-$ ,  $\text{Si}_6\text{GeLi}_7^-$ ,  $\text{Si}_6\text{GeLi}_{10}^-$ ,  $\text{Si}_6\text{Ge}_2^-$ ,  $\text{Si}_6\text{Ge}_2\text{Li}_5^-$ , and  $\text{Si}_6\text{Ge}_{10}\text{Li}_{10}^-$  clusters. Low-lying isomers structures are depicted in Figure 2. As is can be seen,  $\text{Si}_6$  skeleton is distorted in order to accommodate the Ge atom in  $\text{Si}_6\text{Ge}^-$ . The global minimum structure, **1A**, corresponds to a planar five-member ring shaped by four Si atoms and one Ge atom, with other two Ge atoms located above and below the ring. The second most stable isomer, **1B**, is  $14.0 \text{ kcal}\cdot\text{mol}^{-1}$  less stable than **1A**.



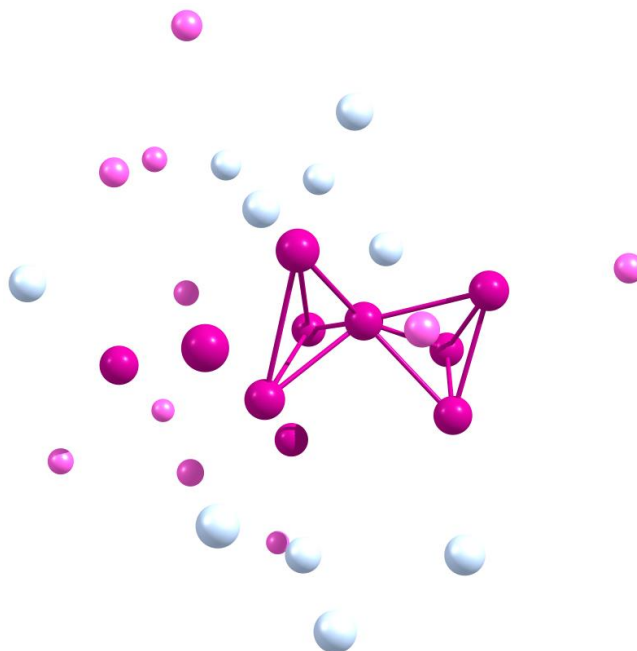


**Figure 2.** Low-lying isomers of Si<sub>6</sub>Ge<sup>-</sup>, Si<sub>6</sub>GeLi<sub>5</sub><sup>-</sup>, Si<sub>6</sub>GeLi<sub>7</sub><sup>-</sup>, Si<sub>6</sub>GeLi<sub>10</sub><sup>-</sup>, Si<sub>6</sub>Ge<sub>2</sub><sup>-</sup>, and Si<sub>6</sub>Ge<sub>2</sub>Li<sub>5</sub><sup>-</sup> clusters calculated at the B3LYP/def2-TZVPP level. Si atoms are represented by dark-pink spheres, Ge by white spheres, and Li by light-pink spheres.

Given that  $\text{Si}_6^-$  is fragmented when five Li atoms are added, we decided to explore the insertion of five Li atoms into the  $\text{Si}_6\text{Ge}^-$  cluster. The global minimum for  $\text{Si}_6\text{GeLi}_5^-$  corresponds to structure **2A**, which consists of a planar five-member Si ring surrounded by the remaining Si atoms, one Ge atom and five Li atoms. It is noticeable that  $\text{Si}_6$  skeleton is not longer fragmented by the five Li atoms. However, there are a number of isomers lying in the range of  $0 - 1 \text{ kcal}\cdot\text{mol}^{-1}$ , where the  $\text{Si}_6$  skeleton is actually fragmented. These results suggest that even though one Ge atom helps to prevent Si fragmentation, it is not enough. In order to corroborate our conclusion, we decided to explore  $\text{Si}_6\text{GeLi}_7^-$ , and  $\text{Si}_6\text{GeLi}_{10}^-$ . As it can be seen in Figure 2, silicon-germanium skeleton is fragmented by Li atoms in both cases.

Subsequently, we decided to explore the effect of the addition of two Ge atoms to  $\text{Si}_6^-$  cluster. Only two low-lying isomers were found: **5A** and **5B**. Isomer **5A** resembles chair conformation of cyclohexane with the Ge atoms located above and below the Si ring. The second most stable isomer, **5B**, can be seen as two Si tetrahedrons bridged by the Ge atoms. The relative energy between both isomers is only  $0.6 \text{ kcal}\cdot\text{mol}^{-1}$ . When five Li atoms are added to the  $\text{Si}_6\text{Ge}_2^-$  cluster, the resulting minimum structure corresponds to **6A**. This structure can be seen as a silicon aggregate surrounded by Ge and Li atoms. However, the second most stable isomer, **6B**, is a silicon-fragmented structure. Nonetheless, it is noticeable, that the difference in energy is  $2.0 \text{ kcal}\cdot\text{mol}^{-1}$  now. This result suggests that incrementing the Ge atoms into the silicon cluster actually prevents its fragmentation. Because of that, we decided to study the biggest system proposed,  $\text{Si}_{10}\text{Ge}_{10}\text{Li}_{10}^-$ .

After exploring the potential energy surface of  $\text{Si}_{10}\text{Ge}_{10}\text{Li}_{10}^-$ , the structure depicted in Figure 3 was found to be the global minimum. This structure accommodates two distorted silicon tetrahedra connected by a silicon atom, surrounded by other three silicon atoms as well as the germanium fragment divided into two skeletons that also corresponds to two distorted tetrahedra. The lithium atoms are accommodated around the Si-Ge skeleton. There is no fragmentation due to lithium atoms in this structure. The last result confirms that actually germanium atoms prevent fragmentation of the silicon structure, as well as the tetrahedral skeleton of silicon atoms can be used as building-block fragment that stabilizes these materials.



**Figure 3.** Global minimum structure of  $\text{Si}_{10}\text{Ge}_{10}\text{Li}_{10}^-$  (**7A**).

The future work consists of studying materials based on the tetrahedral skeleton of silicon atoms, surrounded by tetrahedral fragments of germanium atoms. Such a material should resist fragmentation and be adequate to be used as anode in Li-ion batteries.

Finally, the xyz Cartesian coordinates are described below in order to provide better detail of the studies in this work. These Cartesian coordinates can be displayed in programs such as Mercury, ArgusLab, etc, or some other program that supports this type of coordinates.

### Cartesian coordinates

#### 1A

14	-0.308418000	2.114693000	-0.000392000
14	2.136519000	-1.302871000	0.000211000
14	-0.308292000	-2.114602000	0.000156000
14	2.136153000	1.302904000	-0.000109000
14	0.346930000	0.000601000	1.581198000
14	0.346973000	-0.000395000	-1.581184000
32	-1.903066000	-0.000145000	0.000052000

#### 1B

14	0.923207000	-1.121609000	0.000373000
14	3.284098000	-0.788374000	-0.000194000
14	1.920375000	1.422610000	-0.000149000
14	-0.183384000	0.900110000	1.408206000

14	-0.183205000	0.900655000	-1.407640000
14	-2.162027000	1.439057000	0.000007000
32	-1.574590000	-1.204196000	-0.000264000

## 2A

14	-0.855176000	-0.809783000	-1.950081000
14	-0.614792000	0.020476000	2.143840000
14	1.290649000	-1.585002000	1.518766000
14	0.616227000	0.336506000	-0.135972000
14	2.108441000	2.298260000	-0.455511000
14	1.135764000	-2.089137000	-0.985130000
32	-2.034480000	0.496835000	-0.013688000
3	-1.105135000	-1.952559000	0.434554000
3	0.520906000	1.586451000	-2.391274000
3	3.000332000	-0.065140000	-1.077247000
3	-0.169569000	2.651477000	0.743433000
3	2.276060000	1.014034000	1.802286000

## 2B

14	-2.434596000	-1.930313000	0.062926000
14	2.996343000	0.941621000	0.559125000
14	-0.080422000	-1.671137000	0.759330000
14	-1.208136000	2.055372000	-0.119941000
14	0.532625000	0.820943000	1.106595000
14	-3.132954000	0.365925000	-0.404870000
32	1.714097000	-0.596282000	-0.951193000
3	-0.789212000	-0.261490000	-1.359055000
3	2.177464000	-1.212471000	1.762285000
3	-3.588246000	2.787280000	-0.227890000
3	-1.793569000	0.122072000	1.815605000
3	1.236513000	2.207029000	-1.006320000

## 2C

14	0.030405000	2.351388000	-0.006100000
14	-2.308553000	1.354346000	0.233693000
14	1.978367000	-0.910138000	1.279456000
14	3.768550000	-0.000281000	-0.109531000
14	0.115700000	-1.627579000	-0.460908000
14	1.430144000	0.553795000	-0.973014000
32	-2.324403000	-1.150724000	0.063380000
3	-2.021391000	3.776448000	-0.081293000
3	-0.973193000	0.428225000	-1.775405000
3	-0.358778000	0.142646000	1.441730000
3	2.286452000	1.848798000	1.182631000
3	2.459019000	-1.955538000	-1.273827000

## 3A

14	-1.048663000	0.263839000	-2.074080000
14	1.973708000	-2.200255000	0.000538000
14	-1.049119000	0.262461000	2.074391000
14	3.216861000	-0.386178000	-0.000855000
14	0.242559000	2.161981000	-1.264560000
14	0.242428000	2.161246000	1.266219000
32	-1.904236000	-0.964795000	-0.000641000
3	-0.304889000	-2.201025000	1.519550000
3	-0.304746000	-2.199558000	-1.522063000
3	1.646155000	0.176952000	2.214502000
3	1.646729000	0.179979000	-2.214083000
3	2.482449000	2.191393000	0.000214000
3	0.442444000	-0.027153000	0.000492000
3	-1.992566000	1.609449000	0.000511000

### 3B

14	1.695931000	-0.262055000	1.553608000
14	0.290749000	0.036911000	-2.146641000
14	0.190714000	2.308993000	-1.125496000
14	-3.258816000	0.261937000	-0.057688000
14	1.050969000	2.130464000	1.142586000
14	-2.575085000	-1.209640000	1.408031000
32	1.259457000	-1.636739000	-0.533387000
3	-0.123943000	-2.154513000	1.704901000
3	2.330172000	0.776433000	-0.762649000
3	2.086672000	1.451606000	3.291122000
3	-1.455049000	-1.625999000	-1.213779000
3	-1.979699000	1.307149000	-2.093858000
3	-1.684707000	2.285042000	0.850402000
3	-0.448488000	0.174650000	0.299458000

### 3C

14	2.819111000	0.767989000	-1.556171000
14	0.358045000	-0.143175000	2.386157000
14	-0.797964000	-1.876761000	1.029750000
14	0.021614000	2.106877000	1.464186000
14	-1.351757000	1.802654000	-0.555321000
14	2.936046000	-1.209143000	-0.634345000
32	-1.880308000	-0.705967000	-0.884308000
3	0.478904000	0.093665000	-0.188639000
3	-2.777996000	1.040876000	-2.410386000
3	1.099060000	2.687171000	-0.946203000
3	0.474737000	-2.170785000	-1.286444000
3	1.733542000	-2.143745000	1.497693000
3	2.479960000	0.868916000	1.279221000
3	-2.028690000	0.394823000	1.527515000

**3D**

14	-2.404316000	-1.728573000	-1.151701000
14	-0.209617000	2.399673000	0.683226000
14	0.324081000	0.438911000	2.136354000
14	-3.280774000	-0.239986000	0.187355000
14	1.635321000	-0.041632000	-1.710694000
14	0.597868000	2.143218000	-1.646129000
32	1.490149000	-1.211542000	0.676134000
3	2.129037000	1.320365000	0.454328000
3	-0.477117000	0.057733000	-0.319937000
3	0.136169000	-2.171857000	-1.636767000
3	3.316106000	-1.690868000	-0.946520000
3	-2.144725000	1.323258000	1.950848000
3	-1.201953000	-1.578029000	1.506875000
3	-2.077727000	1.794996000	-1.213508000

**3E**

14	2.815489000	0.469652000	-1.514798000
14	-0.993648000	-1.815055000	1.083465000
14	3.011209000	-1.319696000	-0.277563000
14	0.104355000	-0.091702000	2.536879000
14	-1.200279000	1.573886000	-0.928583000
14	0.022358000	2.032765000	1.322962000
32	-1.884452000	-0.845239000	-1.059072000
3	-2.119495000	0.526252000	1.249380000
3	2.407457000	0.793416000	1.435235000
3	1.256035000	2.461976000	-0.997015000
3	-1.261893000	3.831107000	0.143956000
3	0.312362000	-0.175358000	-2.505651000
3	0.502740000	-0.106916000	-0.040780000
3	1.459352000	-2.280560000	1.640620000

**4A**

14	2.256670000	-2.022338000	-0.774851000
14	-0.639851000	0.285443000	2.226476000
14	3.141155000	-0.315040000	0.356103000
14	0.153766000	2.238268000	-1.325841000
14	0.291786000	2.317604000	1.148711000
14	-1.000823000	0.028500000	-1.997123000
32	-1.547579000	-1.279515000	0.319655000
3	-0.091676000	-2.353723000	-1.552904000
3	-3.149327000	-0.303006000	2.058508000
3	-3.449747000	-0.378501000	-1.230993000
3	-1.871359000	1.374604000	0.155878000
3	1.968392000	0.769458000	2.436200000
3	0.809601000	-1.875262000	1.491535000
3	-2.127870000	2.222859000	-2.598226000

3	0.587047000	0.045852000	-0.024784000
3	2.556471000	2.181931000	-0.176366000
3	1.663367000	0.145904000	-2.258056000

#### 4B

14	1.175110000	-0.018069000	-2.027720000
14	-3.061936000	0.805698000	-0.160782000
14	0.437841000	1.773519000	1.810923000
14	0.915837000	-0.574613000	2.136786000
14	0.630697000	2.261724000	-0.840198000
14	-2.539722000	-1.467884000	0.132328000
32	1.406655000	-1.714865000	-0.126379000
3	-1.835811000	0.186962000	2.122752000
3	-1.300077000	1.130636000	-2.249525000
3	2.354652000	2.071546000	-2.647316000
3	-0.691687000	-1.958214000	-1.728108000
3	-1.608124000	2.734646000	0.486235000
3	2.337528000	0.644161000	0.295064000
3	-0.613132000	-2.532650000	1.373250000
3	-0.430279000	0.110023000	0.015583000
3	1.711407000	3.706551000	0.867366000
3	-3.531992000	-0.776852000	-2.093509000

#### 4C

14	-0.214868000	2.138200000	-1.555114000
14	-0.280395000	2.442446000	0.947751000
14	0.838497000	0.560235000	2.151857000
14	0.974901000	0.106916000	-2.138179000
14	-2.005487000	-2.162762000	-0.414015000
14	-3.137281000	-0.175045000	0.124760000
32	1.806022000	-1.021830000	0.204497000
3	0.236785000	-2.317055000	-1.513307000
3	3.310177000	-0.659982000	-1.746028000
3	-1.783811000	-0.167270000	-2.184991000
3	-2.550194000	2.235152000	-0.407324000
3	-1.859943000	0.773005000	2.258397000
3	-3.152679000	-2.028309000	1.850850000
3	1.901309000	1.555916000	-0.107074000
3	-0.492450000	0.037971000	-0.018430000
3	-0.218906000	-1.910287000	1.677915000
3	3.193764000	-0.199578000	2.129070000

#### 5A

14	0.000068000	-1.080734000	1.254187000
14	-1.431212000	1.430198000	0.294129000
14	1.429697000	1.430981000	0.294841000
14	-0.000436000	2.335791000	-1.538769000

14	0.000595000	-0.159196000	-1.646361000
14	-0.000877000	1.082392000	2.437043000
32	2.026427000	-1.101774000	-0.239297000
32	-2.025480000	-1.102978000	-0.239796000

**5B**

14	1.154075000	2.327987000	-0.000229000
14	-0.697501000	-1.670797000	0.000236000
14	0.652044000	-1.340103000	2.031767000
14	-0.329155000	0.950059000	-1.452032000
14	0.654051000	-1.341499000	-2.030159000
14	-0.330245000	0.950978000	1.451235000
32	-2.410068000	0.172668000	-0.000641000
32	1.927388000	-0.118691000	0.000283000

**6A**

14	0.624761000	-0.733039000	-1.647412000
14	-0.748956000	2.136761000	1.162102000
14	0.168358000	-1.709867000	1.135875000
14	-0.872951000	1.349944000	-1.475552000
14	1.430981000	1.806176000	0.024208000
14	-1.602874000	-0.112929000	1.953131000
32	-1.837916000	-0.977746000	-0.589241000
32	2.362850000	-0.588581000	0.258485000
3	1.521328000	1.550585000	-2.499168000
3	0.944768000	0.486135000	2.222179000
3	-0.199935000	3.688783000	-0.756870000
3	-0.270870000	-3.056037000	-0.938908000
3	-2.924741000	1.265141000	0.123191000

**6B**

14	1.200013000	-0.160550000	2.322537000
14	-2.048754000	1.467690000	0.312541000
14	1.308928000	1.134272000	-1.703933000
14	0.261642000	-1.307364000	-1.782288000
14	-1.245593000	-0.348351000	2.027744000
14	-0.103133000	2.783334000	-0.495122000
32	-1.989356000	-1.120434000	-0.466004000
32	2.189759000	-0.713156000	0.018039000
3	-1.292561000	0.907152000	-2.150048000
3	2.162476000	1.814923000	0.684776000
3	0.169748000	-2.234277000	0.799537000
3	0.026322000	0.311185000	0.127338000
3	-0.278099000	2.103840000	2.136454000

**6C**

14	-1.079847000	0.675598000	2.100388000
----	--------------	-------------	-------------

14	-1.068367000	-1.860034000	-1.443054000
14	1.081525000	-0.678713000	2.098375000
14	0.995137000	-2.164854000	-0.128486000
14	1.066977000	1.861941000	-1.441993000
14	-0.994268000	2.164954000	-0.125471000
32	-2.395838000	-0.105942000	-0.132672000
32	2.395595000	0.106367000	-0.133167000
3	1.216387000	1.952095000	1.320722000
3	-1.216879000	-1.953397000	1.318127000
3	-1.241219000	0.724007000	-2.327962000
3	1.239178000	-0.721563000	-2.329105000
3	-0.000277000	-0.000509000	-0.091705000

### 7A

14	-0.115042000	3.096381000	-1.225232000
14	-0.649485000	-3.094037000	1.792278000
14	1.017738000	-1.266001000	-1.268817000
14	-2.193980000	-2.290292000	0.092447000
14	-1.077238000	1.692851000	0.540410000
14	-0.421502000	-0.632928000	0.562040000
14	0.347253000	0.806245000	2.509680000
14	-0.383818000	-3.524855000	-1.008768000
14	-1.242096000	2.324515000	3.472236000
14	-2.860892000	0.834607000	-3.771554000
32	2.178435000	3.830716000	-1.477292000
32	-4.642127000	-1.724956000	-0.103207000
32	5.360294000	-0.348154000	0.670459000
32	-3.576706000	-0.158653000	-1.915350000
32	6.035191000	0.543553000	-1.327734000
32	-3.144496000	1.523499000	1.983169000
32	2.947394000	-1.075178000	0.347284000
32	4.109931000	-1.091408000	2.817559000
32	2.796800000	1.021802000	1.955362000
32	-4.686008000	-0.326328000	2.178105000
3	1.873680000	-3.543860000	0.374346000
3	-2.691937000	2.697858000	-1.691818000
3	-0.316956000	0.637134000	-2.685120000
3	2.161279000	1.072247000	-0.986518000
3	-1.961386000	-0.887968000	2.981444000
3	1.385545000	3.121609000	1.072669000
3	4.234286000	2.632518000	0.223602000
3	0.375433000	4.649317000	-3.145985000
3	1.414987000	-1.709448000	2.691336000
3	-5.218972000	0.156502000	-4.406860000

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1.

**1. Report Type**

Final Report

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**Organization / Institution name**

Centro de Investigacion en Materiales Avanzados, S.C.

**Grant/Contract Title**

The full title of the funded effort.

Theoretical study of SixGeLi<sub>z</sub>- (x=4-10, y=1-10, z=0-10) clusters for designing of novel nanostructured materials to be utilized as anodes for Lithium-ion batteries

**Grant/Contract Number**

AFOSR assigned control number. It must begin with "FA9550" or "F49620" or "FA2386".

FA9550-13-1-0175

**Principal Investigator Name**

The full name of the principal investigator on the grant or contract.

Mario Sanchez-Vazquez

**Program Manager**

The AFOSR Program Manager currently assigned to the award

Mr. James Fillerup

**Reporting Period Start Date**

04/01/2013

**Reporting Period End Date**

12/31/2014

**Abstract**

In order to find out if silicon nanostructured materials resist lithium cations insertion and extraction under the presence of germanium atoms, we have explored theoretically the potential energy surface of Si<sub>6</sub>Ge-, Si<sub>6</sub>GeLi<sub>5</sub>-, Si<sub>6</sub>GeLi<sub>7</sub>-, Si<sub>6</sub>GeLi<sub>10</sub>-, Si<sub>6</sub>Ge<sub>2</sub>-, Si<sub>6</sub>Ge<sub>2</sub>Li<sub>5</sub>-, and Si<sub>10</sub>Ge<sub>10</sub>Li<sub>10</sub>- clusters. This study was performed using the Kick Coalescence method coupled to Gaussian 09 program, and re-optimization all structures with the B3LYP/def2-TZVPP method. Our results confirm that incorporating germanium atoms to silicon clusters improves and prevents fragmentation.

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