Isolobal Series of Chemical Fragments

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ABSTRACT

During the study of carbonyl clusters using 14n and 4n rules, it was discovered that fragments and atoms could be categorized. This paper presents the categorization of isolobal fragments, molecules and ions using the 14n and 4n rules. In the process of categorization, the conventional periodic table was recreated. A new method of categorizing clusters using fragments and series has been introduced for the first time and this immensely simplifies the categorization of clusters from simple to more complex ones including boranes, carboranes, transition metal carbonyls and metalloboranes. The paper also introduces a simple algebra of cluster series.

Key words: Chemical fragments and series, 4n and 14n rules, Wade-Mingos rules, Hoffman’s isolobal principle, cluster series, Rudolph correlation system, cluster number k value, Isolobal series

INTRODUCTION

The arrival of Hoffman’s isolobal principle greatly assisted the advancement of chemical knowledge of linking up the inorganic, organic and organometallic fields1-2. A combination of isolobal principle and Wade-Mingos rules3-4 laid a strong base for cluster chemistry of boranes, carboranes and transition metal carbonyls. It has recently been found that the organic, and many inorganic and organometallic compounds and clusters follow the 4n and 14n series5-6. In continuing to examine the application of the series in analyzing chemical structures, it has been discovered that the isolobal principle also follows the 4n and 14n series. This revelation is presented in this paper.

RESULTS AND DISCUSSION

In the study of osmium carbonyl clusters7, it was found that some of the series follow a pattern in which one carbonyl ligand is removed at every step until a given cluster is completely stripped naked. In the Hoffmann’s isolobal principle the initial fragment molecule used is methane, CH4. The stripping off one H atom creates CH3 fragment. The next stages produce the fragments CH2, CH and finally the naked C is the final product. Using the 4n rule of classification1-2 the CH4 molecule or fragment belongs to 4n+4 series, while the CH3 belongs to 4n+3 series, CH2 is a member of 4n+2 series, CH is a member of 4n+1 series and C belongs to 4n series. The series give the value of the electron
content of the fragment. For example in CH₄, n = 1, 
the series value S = 4n+4 = 4x1+4 = 8, CH₃, S = 
4n+3, n =1, S = 7, CH₂, S = 4n+2, n= 1, S = 6, CH, S 
= 5, and S, C, S=4. In terms of series, what is found 
interesting is that these values correspond to the 
respective valence content of the fragments. The 
CH₃ fragment was found to be isolobal with Mn(CO)₅ 
fragment. Also applying the 14n rule of classification, 
Mn(CO)₅ fragment belongs to 14n + 3 series. The 
14n +3 series and 4n+3 series while they are not 
identical, they are similar. If n =1 for the fragment 
Mn(CO)₅, then S = 14n+3 = 14x1+3 =17, the valence 
content of the fragment (7+5x2=17). When 10 n is 
subtracted from 14n+3 it becomes 4n+3. This means 
if 10n = 10x1 = 10 is removed from 17, then 7 is left 
which is the same as the valence content of CH₃. 
Thus, the two fragments CH₃ and Mn(CO)₅ are 
terrelated by series. Accordingly, the fragments 
are truly isolobal CH₃ and Mn(CO)₅. In this paper, 
14n + q and 4n+q will be used interchangeably as 
needed.

The recreation of the periodic table

What is fascinating and exciting is that the 
use of 14n and 4n rule the Periodic Table of Elements 
and Fragments has been created. For the ordinary 
elements of the periodic table they have been placed 
precisely into their usual places while the isolobal 
fragments have been classified and placed side by 
side with the ordinary elements into equivalent 
periodic groups. This information is summarized in 
Table1. This table has implications. For example, if 
CH fragment may be regarded as equivalent to N 
atom, it implies that the H atom acted as an electron 
donor to the carbon atom(C) making it have 5 
valence electrons in the (CH) fragment like the 
nitrogen atom (N) and hence placing it into group 5 
or 15 like the nitrogen atom. In the same manner, in 
the fragment [Cr(CO)] when the CO ligand donates 
the two electrons to Cr which is in group 6 of the 
transition elements it makes it behave as if it is an 
iron atom(Fe) atom of group 8. On the basis of the 
series, it appears that the basis of the isolobal 
principle is the electron content of the sum valence 
electrons around the skeletal atom. In the case of 
CH₃ fragment, the carbon has 4 valence electrons 
and the three hydrogen atoms donate three more 
electrons making the carbon atom have a 
surrounding environment of 7 valence electrons. 
Thus, it acquires the status of a group 7 element 
such as fluorine. Group 7 has other family members 
such as Cl, Br, and I. Moving along the period from 
C the next element is nitrogen atom(N). The addition 
of 2H atoms converts it NH₂ which also has a total 
valence content of 7. The NH₂ fragment will also be 
isolobal to CH₂. Since N is in group 5, other family 
members namely, PH₃, AsH₃, SbH₃, and BiH₃ will 
also be isolobal to CH₂. In case of group 17, those 
elements with carboylns will also be isolobal to CH₂. 
Since we are actually comparing parallel series 
14n+3 with 4n+3, if we remove 10n from 14n+3 we 
will convert the carbonyl series to main group series 
of 4n+3. Futhermore, since we are dealing with a 
single skeletal element, then 10n is equivalent to 
10x1 = 10. Hence, if we subtract 10 from 17 we will 
get 7 the valence electron content of CH₃ fragment 
and group 7 elements. The carbonyl fragments 
include the hypothetical and real ones such as 
V(CO)₆, Nb(CO)₆, Ta(CO)₆, Mn(CO)₅, Tc(CO)₅, 
Re(CO)₅, Co(CO)₄, Rh(CO)₄, Ir(CO)₄, Cu(CO)₃, 
Ag(CO)₃ and Au(CO)₃. The fragments from the main 
group elements and transition metal fragments 
which are isolobal to CH₂ and CH fragments 
corresponding to 6 and 5 valence electrons can 
similarly be identified. Again the removal of 10 
electron units from the carbonyl fragments will yield 
the corresponding 6 and 5 valence electron 
contents. It should be pointed out that the hydrogen 
atom can be replaced by an alkyl group, R or aryl 
group Ar. The selected isolobal extended family 
members are indicated in Tables 2 to 4. In searching 
for possible isolobal fragments for CH₄ from main 
group elements, numbers could act as a guide. For 
example since we are looking for seven valence 
electrons, we may use sets such as (4+3) this can 
lead us to group 4 elements combined with 3 
hydrogen atoms. Hence, we generate, CH₃, SiH₃, 
GeH₃, SnH₃, and PbH₃. In the case of transition 
metals, we want to generate 17 electron systems. 
The set of numbers we may need for example could 
include (5+12). This set will give us the carbonyl 
fragments such as V(CO)₆, Nb(CO)₆, and Ta(CO)₆, 
while the set (7+10) could give us the carbonyls 
M(CO)₅, M = Mn, Tc, and Re. The combination (9+8) 
will generate M(CO)₄ where M = Co, Rh, and Ir. The 
other possible combination of numbers to give 17 
is (8+5+4). This set will generate the fragments such 
M(Cp)(CO)₂ where M = Fe, Ru and Os and Cp = η⁵-
C₅H₅.
### Table 1. Classification of Isolobal Fragments into Series

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<thead>
<tr>
<th>14n-9</th>
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References:
Arising from the arranging isolobal fragments and elements according to series as indicated in Table 1, it became apparent that the 14n and 4n series were in fact recreating the Periodic Table of Elements as is currently known. A portion of this is portrayed in Table 5. Since we are considering one skeletal element in this case, n = 1 and this generates the numbers in the first row of the table. These findings are fascinating.

Assembling the fragments

As can be seen from Table 1, atoms and fragments can be categorized according to series. In actual fact, what has happened in Table 1, the 14n and 4n series have created or reproduced a Periodic Table of elements and fragments. Just as atoms can be assembled according to valence rules to produce molecules, fragments can do the same. Scheme 1 gives some few illustrative examples. Fragments can be combined to generate molecular structures. Let us illustrate this concept starting with the simplest fragment CH3. This fragment belongs to 4n+3 series. Adding two of these fragments is equivalent to 2[CH3(4n+3)] = C2H6 (4n+6). Thus, the

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**Table 2: Selected CH3 Isolobal Family**

<table>
<thead>
<tr>
<th>GROUP 17</th>
<th>GROUP 7</th>
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<tr>
<td><strong>14n+3 SERIES</strong></td>
<td><strong>4n+3 SERIES</strong></td>
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<tr>
<td>Cu(CO)3</td>
<td>Co(CO)4</td>
</tr>
<tr>
<td>Ag(CO)3</td>
<td>Rh(CO)4</td>
</tr>
<tr>
<td>Au(CO)3</td>
<td>Ir(CO)4</td>
</tr>
<tr>
<td>11+6 = 17</td>
<td>9+8 = 17</td>
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<tr>
<td>17-10 = 7</td>
<td>17-10 = 7</td>
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**Table 3: Selected CH2 Isolobal Family**

<table>
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<th>GROUP 6</th>
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<tr>
<td><strong>14n+2 SERIES</strong></td>
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<td>Zn(CO)2</td>
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<tr>
<td>Cd(CO)2</td>
<td>Pd(CO)3</td>
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<tr>
<td>Hg(CO)2</td>
<td>Pt(CO)3</td>
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<td>12+4 = 16</td>
<td>10+6 = 16</td>
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<tr>
<td>16-10 = 6</td>
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**Table 4: Selected CH Isolobal Family**

<table>
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<th>GROUP 15</th>
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<tbody>
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<td><strong>14n+1 SERIES</strong></td>
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<td>Cu(CO)2</td>
<td>Co(CO)3</td>
</tr>
<tr>
<td>Ag(CO)2</td>
<td>Rh(CO)3</td>
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<td>Au(CO)2</td>
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<tr>
<td>11+4 = 15</td>
<td>9+6 = 15</td>
</tr>
<tr>
<td>15-10 = 5</td>
<td>15-10 = 5</td>
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multiplication affects the digit component of the series \(2 \times 3 = 6\) but not the \(4n\) component. In actual fact the \(4n\) represents the number of the valence electrons contributed by the carbon skeletal atoms and the digital part represents the hydrogen atoms contribution to the valence account. In this result, when \(n = 2\), the value of the series \(S = 4n+6 = 4x2+6 = 14\), which the sum of the valence electrons of \(\text{C}_2\text{H}_6\). This means that in series, the rule is that the addition or multiplication does not affect the \((4n)\) component of the series. The digit portion is the one to be added or subtracted whatever the case maybe. This makes sense because \(4n\) for \(n = 1\) gives us 4 while \(4n\) for \(n = 2\) gives us 8. This gives us a simple algebraic rule of adding series. Let us take \(\text{CH}_2\) fragments for illustrating the concept combining series. The fragment \(\text{CH}_2\) belongs to \(4n+2\) series.

The value \(k = 2n-x\) where \(= 0, 1, 2, 3, 4\) and son comes from a simple correlation of the empirical observation of the number of skeletal atoms involves and the corresponding digit of the molecular fragment series formed. Thus, in the case of \(\text{C}_2\text{H}_6\), we know it has a single C-C bond and 2 skeletal atoms. In order to create a simple mathematical relationship of a single bond with the series \(S = 4n+6\), it is common sense that \(k = 2n-3\). The digit 3 is just one half of 6 and \(n\) represents the number of the skeletal atoms. Naturally for \(\text{N}_2\) with \(S = 4n+2\), the \(k\) value is given by \(2k-1 = 3\), the triple bond of dinitrogen as we know it. In this way, a simple mathematical relationship between skeletal bonds or linkages and the corresponding number of skeletal atoms is developed. Let us try one example to illustrate the concept of assembling fragments. We can combine 4 fragments of \(\text{Ir(CO)}_3\) as follows.

\[
4[\text{Ir(CO)}_3] = \text{Ir}_4(\text{CO})_{12}.
\]

\[
4[14n+1, n = 1] = 14n+4 (n = 4); k = 2n-2 = 2x4-2 = 6.
\]

\[
4[14X1+1 = 15] = 14X4+4 = 60 = V.
\]

This the number of valence electrons of the cluster and \(k = 6\) is the cluster number in this case. It also corresponds to the number of linkages of the skeletal atoms which have a tetrahedral geometry \((T_d)\) as sketched in Figure 1. Since the 14n series run parallel to the 4n series, the removal of 10 \(n\) electrons from \(V = 60\) will give us the number...
Scheme 1: Combination of two Isolobal Fragments
of valence electrons characteristic to corresponding cluster of the main group elements. In this case 60-10x4 = 20. This number of valence electrons can give us a (CH)_4, P_4 or (BH_2)_4 cluster with the ideal T_d symmetry.

Unique simplification and structure prediction using isolobal principle and series

The most fascinating and exciting aspect of isolobal principle in combination with series is the way they can simplify an apparently imposing cluster to a bare simple number from which its structure (shape) can be predicted with high accuracy. Let us consider the carbonyl cluster, Ru_4(C)(CO)_15 which has a square pyramid geometry. Applying the isolobal principle and series, we can split up the cluster into simpler monoskeletal fragments as follows. The splitting of molecule into fragment, the nature of monoskeletal fragments produced does not matter as long as you can readily work out their corresponding fragmental series. Let us represent the molecular formula with F = Ru_4(C)(CO)_15 = 5[Ru(CO)_3] + C. For Ru(CO)_3 fragment; V = 8 +3x2 = 14 valence electrons, and n =1, hence 14n =14. Therefore the fragment belongs to S = 14n +0 series. The approach is to determine the overall cluster series (SF) of the formula F. If regard the C atom as a donor of its valence electrons and not part of the skeletal cluster fragment, then the overall series of the formula F is obtained in a straightforward manner. 5[Ru(CO)_3] ; SF = 5[14n+0] + 4 = 14n +4 where n = 5. Therefore the series SF = 14n +4 tells us that the cluster is a member of the NIDO family. A nido structure is just one step away from being a closo (the ideal of full or standard) geometry. In this case, an M-5 skeletal fragment is one step away from an ideal M-6 octahedral (closo) symmetry. For the main group elements, this series is the same as that of SF = 4n+4. Furthermore, it also gives us the valence content of the cluster if we substitute n =5 in the formula series. Just like the 4n+4 series, the cluster k value is obtained from k =2n-2 = 2x5-2 = 8. This implies that an M-5 skeletal cluster with k = 8 has a characteristic shape of a square pyramid sketched in Figure 3. The shape is flattened for ease of visibility. The borane cluster B_5H_9 is known to have an ideal square pyramid shape. We can readily determine its series by splitting up into monoskeletal fragment as we please and deduce the series of each fragment. Then finally sum up the series to get the formula series of B_5H_9. This is illustrated below.

B_5H_9 = [BH]+ 4[BH_3]

[BH], V = 3+1 = 4, n = 1, 4n = 4, S = 4n + 0
[BH_3], V = 3+2 = 5, n = 1, 4n = 4, S = 4n +1.

$SF = (4n+0) + 4(4n+1) = 4n + 4$, for $n = 5$, $S = 24$, the number of valence electrons of $\text{B}_5\text{H}_9$. The $k$ value is $2n-2 = 2 \times 5 - 2 = 9$. Hence, this is an $M-5$ cluster with $k=8$, and with an ideal square pyramid shape as is already known.

Let us now tackle the $[(\text{Cp})\text{Co}][\text{B}_4\text{H}_9]$ cluster. The cluster can be split into fragments as follows.

$[(\text{Cp})\text{Co}][\text{B}_4\text{H}_9] = [\text{CpCo}] + 4[\text{BH}_3]$, $[\text{CpCo}], V = 5 + 9 = 14$, $n = 1$, $14n = 14$. Hence, $S = 14n + 0 = 4n + 0$. $[\text{BH}_3], V = 3 + 2 = 5$, $n = 1$, $4n + 1$. Hence, $4[\text{BH}_3] = 4[4n+1], n = 1] = 4n + 4 (n = 4)$. $SF = (4n+0) + (4n+4) = 4n + 4 (n = 5)$. Therefore the formula, for the series $SF = 4n + 4 (n = 5) = 24$ if Co is regarded as main group element. Since the main group skeletal elements are more than the coordinated metal atoms in the metalloborane complexes, it is best to regard to consider them as if they belong to the main group elements. With $SF = 4n + 4$ for the $[(\text{Cp})\text{Co}][\text{B}_4\text{H}_9]$ cluster, then $k = 2n - 2 = 2 \times 5 - 2 = 8$. Since the cluster has 5 skeletal elements $M-5$ and has $k = 8$, its ideal shape will be a square pyramid. The presence of a metal atom, makes it possible for the $M-5$ cluster to have two possible isomers as sketched in Figure 4. Again the square pyramid geometry has been flattened to allow a better visibility. The larger point in the sketch represent CpCo fragment and Cp = $C_5H_5$.

Let us try a few more examples to demonstrate the power of fragments and series in predicting the geometry by considering the example $(\text{Cp})_3\text{Co}_3\text{B}_3\text{H}_5$ cluster. The cluster can be fragmented into $3[\text{CpCo}] + [\text{BH}] + 2[\text{BH}_3]$ units. The series for each fragment can be worked out and the total series for the cluster easily obtained. This is shown in Scheme 2. With a cluster of $M-6$ and $k = 11$ belonging to the $4n + 2$ series (CLOSO), the geometry is predicted to be an octahedral which is the case. A possible skeletal shape is sketched in Figure 5.

Consider the cobalt cluster $(\text{CpCo})_2(\text{C}_2\text{B}_6\text{H}_{10})$. Its structure can readily be deduced by use of fragments and series approach. The formula of the cluster can be re-written as follows $(\text{C}_2\text{Co})_2(\text{BH})_8 \times 4$. The $[\text{CpCo}]$ fragment with 14 electrons belong to $14n$ ($= 4n + 0$) series. The 6 BH fragments also belong to the $4n$ series. However the two CH fragments contribute $2[4n+1] = 4n + 2$ towards the series. The net series of the cluster $SF = 2[4n+0] + 6[4n+0] + 2[4n+1] = 4n + 2$ series for $n = 10$. Hence, the final series is $SF = 4n + 2$. This means the cluster belongs to CLOSO series. Since $n = 10$, it will have a shape similar to that of $\text{B}_{10}\text{H}_{10}^{2-}$. This skeletal structure is given in Figure 6 with two of the ten atoms being cobalt atoms carrying the Cp $(C_5H_5)$ rings. Since, the use of series to predict shapes is not common, let us briefly restate them. The $S = 4n + 2$ is closo, $4n + 4$ (nido), $4n + 8$ (archachno), $4n + 10$ (klapo) and so on. But for $4n + 0$ (monocapped), $4n - 2$ (bicapped), $4n - 4$ (tricapped), $4n - 6$ (tetracapped) and so on. In the case of the capped series the $k$ value takes a positive number after the $n$, thus $k = 2n + 1$ for bicapped series, $k = 2n + 2$ for tricapped series and so on.
Revisiting the rudolph correlation system of clusters

A lot of work has been done on boranes and its relatives mainly carboranes, and metalloboranes as well as transition metal carbonyl clusters\(^1\)-\(^2\). The prediction of shapes of these clusters is based mainly on the well known Revisiting the Rudolph correlation system also based on known shapes \(^1\)-\(^2\). For example, if an M-5 skeletal system is an arachno, then it is traced to an M-7 closo system whose shape is known (pentagonal bipyramid) in order to derive its shape by correlation. However, if M-5 system is a Nido one, it is related to the next M-6 (octahedral) closo system. This correlation system is presented in formula form in Table 6. It is quite clear that the Rudolph correlation system is a DECAPPING SYSTEM which is characterized by change in \(k\) value by 3. For example from \(B_6H_6 \rightarrow B_7H_8\) of \(k = 11\) to \(B_7H_9\) \((k = 8)\) to \(B_7H_{10}\) \((k = 5)\). On the other hand if we know the \(k\)-value of a given M-x cluster, we can use that as a guide to predict the shape. For instance \(B_5H_{11}\) \((M=5)\) with \(k = 7\) and of \(4n+6\) series, will be a Nido and with a shape deduced from the \(k\) value or correlation with \(k = 11\) an octahedral closo shape. On the other hand if \(B_5H_{9}\) \((M=5, k = 9)\). This predicted to be a trigonal bipyramid. It must be pointed out that the 4\(n+2\) series are known to exist in ionic form. For example, \(B_5H_7\) exists as \(B_5H_5\) and \(B_6H_8\) as \(B_6H_6\) \(^2\). If the type of cluster series of a given formula is known, then the \(k\) value is readily calculated. For instance, \(B_9H_{13}\) if we write it as \((BH_2)_4(BH)_5\) fragments, we can readily deduce its series. Thus; \(4[4n+1]+5[4n+0] = 4n+4\). In the algebra of series, the \(4n\) remains constant regardless of the multiplier. What changes is the value of the digit after \(4n\) (or \(14n\)). Also additions of \(4n\) do not change it. Hence, \(4[4n+1]\) simply becomes \(4n+4\) as the multiplier \(4\) does not affect it. However, the multiplier affects the digit and so we get, \(4x1 = 4\). Similarly, \(5[4n+0]\) becomes \(4n+0\) as the multiplier 5 does not affect \(4n\) and \(5x0 = 0\). When \((4n+4)\) is added to \((4n+0)\), the result becomes \(4n+4\) as the addition does not affect the \(4n\) term but the digits \(4+0 = 4\). The net result is \(S = 4n+4\) the series for \(B_5H_9\) borane cluster. If \(n = 5\) for a 5-skeletal cluster, the value of \(S\) becomes \(S = 4x5+4 = 24\). This the sum of valence electrons for \(B_5H_9\). This can be verified from \(5x3+9 = 24\) as derived from the formula. In order to get more familiar with the power...
of deducing the series of a given molecular formula, let us take $B_6H_{10}$ cluster. There are several ways of splitting it up into fragments. But let us consider one of them given in Scheme 4. The $B_6H_{10}$ cluster was split into $(BH_2)(BH_2)_4$, and then the series of each fragment was determined. Hence, the overall series of the cluster easily deduced. It is amazing and delightful to see how the combination of fragments and series work together to readily decompose a seemingly complex formula into its family of series. Once the cluster series has been deduced, it becomes quite easy to predict its shape in the usual way. The ideal skeletal shape of $B_6H_{10}$ is shown in Figure 7.

### Table 6. Rudolph Correlation Scheme

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<td>NIDO</td>
<td>ARACHNO</td>
</tr>
<tr>
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<td>$B_6H_5$</td>
<td>$B_6H_5$</td>
</tr>
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</tbody>
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### Scheme 4. Using Fragments and their Series to Categorize $B_6H_{10}$ into its Family Series

#### CONCLUSION

This paper is a climax of intense efforts in an attempt to fully understand and be able to explain structures of clusters of the main group elements and transition metal carbonyls in much simpler way than is currently presented. The work started with the introduction of cluster number ($k$ value) which led to the establishment of $14n$ and $4n$ rules\textsuperscript{14-15}. It was demonstrated that clusters could easily be categorized by using the $14n$ and $4n$ rules\textsuperscript{5-6}. Further investigations into the nature of carbonyl clusters...
with respect to capping series led to the revelation that molecules, fragments, atoms could be categorized by 14n and 4n rules. This sparked great interest in applying the 14n and 4n rules to Hoffmann’s isolobal fragments. This has proved a tremendous success as it was discovered that isolobal fragments, molecules were interlinked by 14n and 4n rules and that they could be categorized in the same way as elements in the periodic table. Furthermore, what is interesting was that the placement of fragments and elements resulted in recreating the modern periodic table. It became easier to appreciate why a fragment such as Os(CO)$_4$ of a transition behaved like CH$_2$ of the main group elements. The reason being Os(CO)$_4$ fragment has a CH$_2$ fragment belongs to the 4n system. If 10 electrons are removed from 16 electrons of the Os(CO)$_4$ fragment the remainder is 6 electrons corresponding to those of the CH$_2$ fragment. It is as if another CH$_2$ fragment is hiding within Os(CO)$_4$ fragment. Finally, when the concept of fragments and series are applied to clusters they are easily reduced to simple series from which cluster structure can be predicted in the usual way. What is much more heart-warming and pleasant was that not only does the 14n and 4n rules reproduce Mandeleev’s periodic table of elements, categorize Hoffmann’s fragments into their positions in the periodic table, but it also provides us with the basis for OCTET RULE AND EIGHTEEN ELECTRON RULES that have played vital roles in science. If we look carefully at Table 1 column 18/8, we can discover all those fragments listed there possess an element that obeys either octet rule or 18-electron rule. For instance, HF, OH$_2$, NH$_3$, CH$_4$ and BH$_3$ obey the octet rule while Ne, Cr(CO)$_6$, Mo(CO)$_6$, W(CO)$_6$, Fe(CO)$_5$, Ru(CO)$_5$, Os(CO)$_5$, Ni(CO)$_5$, Zn(CO)$_3$ and Cd(CO)$_3$ in principle obey the 18-electron rule. Since atoms, fragments and molecules can be categorized into series and they can also be combined to produce other fragments and molecules and in turn can be split up into fragments, this implied that the corresponding series of fragments can somehow also be maneuvered. Indeed a simple algebra of series has briefly been introduced to simplify the categorization of clusters. In the author’s reflections and meditations, seeing the beauty and smooth flow of using simple numbers to generate these results with no heavy computational calculations, D. Mendeleev(1834-1907), I. Langmuir(1881-1957), G.N. Lewis(1875-1948) and even R. Hoffmann (1937-) would smile broadly at how nature is so wonderful. This work complements the existing knowledge on clusters whose skeletal elements obey the 18 or 8 electron rules.

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