# Chemical Trees Enumeration Algorithms \*

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Received: date / Revised version: date

Abstract In the chemical community the need for representing chemical structures within a given family and of efficiently enumerating these structures suggested the use of computers and the implementation of fast enumeration algorithms. This paper considers the isomeric acyclic structures focusing on the enumeration of the alkane molecular family. For this family, Trinajstić et al. ([12]) devised an enumeration algorithm which is the most widely known and utilized nowadays. Kvasnička and Pospichal ([9]) have proposed an algorithmic scheme which, from the computational complexity point of view, can prove to be more efficient than the Trinajstić one, nevertheless, this algorithm, to the best of our knowledge, has never been implemented. Indeed an efficient implementation requires the introduction of non trivial data structures and other computational tricks. The main contribution of this paper consists of the definition of the implementation details of Kvasnička-Pospichal's algorithm, in a comparison of Trinajstić's, Kvasnička-Pospichal's and two new algorithms, proposed here, in terms of both computational complexity analysis and running times.

## 1 Introduction

The enumeration of the alkane molecular family is a problem whose roots go back to the nineteenth century. Alkanes are chemical compounds with identical molecular formulae and weights but differing at least in some physical and/or chemical properties leading to different structures. Alkanes belong to the general class of isomeric acyclic structures. The term isomerism and the definition of isomers were introduced in 1830 by Berzelius ([3]).

 $<sup>^{\</sup>star}\,$  Part of this work has been developed during a visit of the first two authors at the EPFL of Lausanne

The alkane molecular family is partitioned into classes of homologous molecules, that is molecules with the same numbers of carbonium and hydrogen atoms; the *n*-th class is characterized by the formula  $C_nH_{2n+2}$ , n = 1, 2, ...

An alkane molecule is usually represented by indicating the carbonium atoms and their (primary) links, omitting to represent hydrogen atoms (see Figure 1), whose placement can be obtained automatically. When n is suffi-



**Fig. 1**  $C_3H_8$  propane and  $C_4H_{10}$  butanes

ciently large  $(n \ge 4)$ , to the same formula correspond molecules with different structures. Take as an example butane  $(C_4H_{10})$ : there are two possible molecular structures (nbutane and isobutane) as depicted in Figure 1.

Alkanes can be represented by acyclic connected graphs whose nodes correspond to carbonium atoms and edges correspond to the primary links. There is a one-one correspondence between elements in the family of alkanes and elements in the family of the connected acyclic graphs (i.e. trees) whose nodes have degree less than or equal to 4, therefore an alkane molecule can be represented by such a degree constrained tree.

At the end of the 19-th century, Cayley was the first to realize the importance of tree mathematical theory in the enumeration of isomeric acyclic structures ([4]). He enumerated manually the alkane isomers and alkyl radicals with up to n = 13 carbonium atoms, though introducing some counting errors. This enumeration had a considerable impact on chemists of that time.

Subsequently, the development took a turn to computer-oriented methods for the enumeration and generation of isomeric structures: the development of techniques for the generation of graphs (chemical structures) by computing devices has made possible the direct enumeration of isomeric acyclic structures by way of enumerating algorithms. The pioneering contributions in this area were made by Lederberg and his group ([10]) and Corey and his group ([5]).

One of the possible applications of chemical tree enumeration algorithms is the investigation of different chemical and physical properties on massive quantities of molecules through the computation of topological indices, e.g. the efficient computation of Hyper-Wiener index ([1]).

For these enumeration purposes, the algorithm most widely used by chemists is that devised by Trinajstić et al. in [12]. In 1991, Kvasnička and Pospichal ([9]) proposed an algorithmic scheme which in principle can be more efficient than the other one, depending on its actual implementation. However the details described in [9] are not sufficient to implement a computer code. Here we propose, to our knowledge, the first implementation of the Kvasnička and Pospichal algorithm. Moreover, in this paper we introduce two new algorithms for the generation of the alkanes molecular family based on Reverse Search ([2]) which is a technique to avoid duplicated trees in the enumeration.

In section 2 we introduce the molecular codes used to encode alkanes, the two algorithms proposed in literature ([12,9]) are described in section 3, and our new algorithms are presented in section 4. The details needed to implement our algorithms as well as that in [9] are also reported. The computational complexity analysis of all algorithms is carried out in section 5. Section 6 reports running times comparison of the algorithms applied to molecules with up to n = 27 carbonium atoms. Some conclusions are drawn in section 7.

Hereafter, we denote by n-tree and n-rooted respectively a tree and a rooted tree with n nodes. For the sake of simplicity, we use the expression "tree" or "alkane" as a synonymous with "degree constrained tree".

### 2 Tree codes

In order to efficiently enumerate degree constrained trees, data must be suitably encoded: the motivation is that a code can be easily handled by computer program. In [11], the author lists some desirable properties for a molecular code such as being linear, unique, well-defined, brief, simple, easily comprehensible to chemists, and efficiently encodable and decodable.

The codes used in this paper are the N-TUPLE code ([11,12]) and the *centered* N-TUPLE code proposed in [6] hereafter called CN-TUPLE code for short. We define the N-TUPLE code for a rooted tree, then we extend this definition to an unrooted tree. The definition of CN-TUPLE code for a tree is a variant of that for the N-TUPLE.

First we recall that a sequence of integers  $C = c_1 c_2 \dots c_l$  is *lexicographically* larger than a sequence  $C' = c'_1 c'_2 \dots c'_{l'}$  if i) there exists an integer j,  $1 \leq j \leq \min\{l, l'\}$ , such that  $c_i = c'_i$  for all  $i = 1, \dots, j - 1$  and  $c_j > c'_j$  or ii) l > l' and  $c_i = c'_i$  for all  $i = 1, \dots, l'$ .

The code of a *n*-rooted tree, rooted in r,  $T_r$  is recursively defined as follows: the *N*-TUPLE of a pendant node is 0; let g be the number of nodes  $(v_1, v_2, \ldots, v_g)$  adjacent to r. In order to obtain the *N*-TUPLE of tree  $T_r$ , rooted in r, we delete from  $T_r$  the root r together with its incident edges; we compute the *N*-TUPLES of the subtrees rooted in  $v_1, v_2, \ldots, v_g$ ; then we concatenate these *N*-TUPLES in such a way as to obtain a lexicographically maximum sequence S. The *N*-TUPLE code of  $T_r$  is the concatenation of gand S.

For an unrooted tree T, all nodes of maximum degree are chosen successively as roots and the corresponding N-TUPLES are computed. The N-TUPLE code of T is defined as being the lexicographically maximum N-

TUPLE among these. For example, the N-TUPLE codes for the butanes in Figure 1 are 2100 and 3000, respectively.

To obtain the CN-TUPLE code, we first identify the *center* (or the centers) of a tree T. We delete simultaneously all pendant nodes of T together with their incident edges, thus obtaining a smaller tree; we repeat this operation until a tree with a single node (or two adjacent nodes) is obtained which is, by definition, the center (or the centers) of T. The CN-TUPLE code for a tree T is the N-TUPLE code of the rooted tree with the root in the center (or the the lexicographically maximum N-TUPLE of the two rooted trees rooted in the centers).

The computation of the N-TUPLE code and the CN-TUPLE code for a n-tree requires respectively  $O(n^2)$  and  $O(n \log n)$  time as reported in [6].

#### **3** Previous work

In this section we report two different enumeration algorithms: the algorithm proposed by Trinajstić et al. ([12]), briefly referred to as TNKMS, and that outlined by Kvasnička and Pospichal ([9]), briefly referred to as KP.

The algorithm TNKMS is based on the following idea: a *n*-tree can be obtained by adding one node (and one edge) to a (n-1)-tree in such a way that the degree constraint still holds. We refer to this class of algorithms as *one-to-one enumeration*.



Fig. 2 Example of one-to-one enumeration

One-to-one enumeration does not guarantee avoiding the generation of multiple copies of the same tree. An example of enumeration is reported in Figure 2: each arrow represents how the tree can be derived and the label is the number of different ways in which it can be obtained. The multiple generation of a tree starting from the same tree or from different trees is the crucial point emerging from this class of enumeration algorithms. Therefore the introduction of some rules which guarantee the uniqueness of enumerated trees is required. Note that TNKMS as described in [12] does not introduce any such rules.

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The algorithm KP, also called *constructive enumeration* algorithm, is based on the following observation: an *n*-tree can be decomposed into one node and one or more rooted trees; then, each rooted tree can be decomposed in the same way, and so on. Therefore, a *n*-tree can be enumerated joining one node together with one or more rooted trees in such a way that the sum of nodes in the rooted trees is equal to n - 1. An example is given in Figure 3:  $T_1$  is generated by joining the blank node with two 2-rooted and one 1-rooted trees;  $T_2$  is generated by joining three 1-rooted and one 2-rooted trees.



Fig. 3 Examples of constructive enumeration

In [9] a more formal description of this idea is reported. In particular, this description is based on the rules defined by Jordan's theorem ([8]) which guarantee the unique enumeration of trees.

**Theorem 1 (Jordan 1869)** Let T be a n-tree, the following three different cases should be separately considered:

- (odd case) For an odd n = 2k + 1 there exist a unique node, called the **centroid**, such that all (two or more) incident subtrees are composed, at most, by k nodes.
- For an even n = 2k there exist either
- (even case i) a unique node centroid such that all (three or more) incident subtrees are composed of less than k nodes, or
- (even case ii) a unique edge, called the **bicentroid**, such that the incident two subtrees are composed exactly of k nodes.

Note that to generate a degree constrained tree, we have to take into account that we are not allowed to combine more than four rooted trees. For example, the set of 10-trees can be generated joining one node together with three or four rooted trees in such a way that the sum of nodes is equal to 9 (even case i). Moreover, we can combine two rooted trees with 5 nodes (even case ii).

Jordan's theorem defines a way to compose rooted trees, e.g. one 4rooted and one 5-rooted for a 9-tree. It is clear that different trees can be obtained by composing different rooted trees with the same cardinality. Figure 3 shows the trees obtained by combining all the pairs of 3-rooted trees, i.e.  $T_3$ ,  $T_4$  and  $T_5$ . Finding the possible combinations of t rooted trees with no more than k nodes is equivalent to finding all the possible combinations of t numbers smaller than or equal to k in such a way that their sum is equal to n-1. Note that t has to be greater than or equal to 2 according to Jordan's theorem and less than or equal to 4 according to the degree constraint.

KP generates all *n*-trees computing all the possible combinations of t rooted trees and, for each combination, combining the rooted trees with the same cardinality in all possible ways. We have found neither computational results for this approach nor any computer code implementing it, therefore we devised our own implementation which required to specify data structures and clarify the details missing in [9].

Note that both TNKMS and KP encode the enumerated trees by using the  $N\mbox{-}\ensuremath{\mathsf{TUPLE}}$  code.

#### 3.1 Implementation details of KP algorithm

KP algorithm requires computing of all the sets of *p*-rooted trees, say  $R_p$ , for  $p = 1, \ldots, \lfloor \frac{n}{2} \rfloor$  to generate the set of *n*-trees according to the rules defined by Jordan's theorem and to the degree constraint.

As discussed above, a combination of t rooted trees having no more than p nodes to build an n-tree is equivalent to a combination of t integer numbers smaller than or equal to p whose sum is equal to n-1. We denote such a combination with

$$(a(x), b(y), c(z), d(w)) \tag{1}$$

which means that x a-rooted trees and y b-rooted trees (and so on) are used to build one n-tree and  $ax + by + \ldots = n - 1$ .

Moreover, each combination of numbers generating n-tree is subject to the following conditions ([9]) according to Jordan's theorem and to the degree constraint.

(Condition 1) The combination (a(x), b(y), c(z)) yielding a *p*-rooted tree must be such that:

1. ax + by + cz = p - 1;

2.  $1 \le x + y + z \le 3$ , (degree constraint);

3.  $a, b, c \in \{1, \dots, p-1\}, x, y, z \in \{0, \dots, 3\};$ 

(Condition 2) The combination (a(x), b(y), c(z), d(w)) yielding an *n*-tree (with *n* odd) must be such that:

1. ax + by + cz + dw = n - 1;

2.  $x + y + z + w \ge 2$ , (Jordan's theorem, odd case);

3.  $x + y + z + w \le 4$ , (degree constraint);

4.  $a, b, c, d \in \{1, \dots, \lfloor n/2 \rfloor\}$ , (Jordan's theorem, odd case);

5.  $x, y, z, w \in \{0, \dots, 4\};$ 

(Condition 3) The combination (a(x), b(y), c(z), d(w)) yielding an *n*-tree (with *n* even) must be such that:

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1. ax + by + cz + dw = n - 1;

2.  $x + y + z + w \ge 3$ , (Jordan's theorem, even case i);

3.  $x + y + z + w \le 4$ , (degree constraint);

4.  $a, b, c, d \in \{1, \ldots, n/2\}$ , (Jordan's theorem, even case i);

5.  $x, y, z, w \in \{0, \dots, 4\}$ 

(Condition 4) The combination (k(2)) yielding an *n*-tree (with *n* even) must be such that  $k = \frac{n}{2}$ , (Jordan's theorem, even case ii);

Referring to algorithm KP summarized in Figure 4, we first compute all sets (R[p]) of rooted trees with up to  $\lfloor \frac{n}{2} \rfloor$  nodes generating all combinations (a(x), b(y), c(z)) such that Condition 1 holds. Then, the set of *n*-trees (T) is generated: for odd *n*, **KP** generates all combinations (a(x), b(y), c(z), d(w)) such that Condition 2 holds and the corresponding *n*-trees are generated by function **Compose**; in the case of even *n*, the *n*-trees are generated from all combinations verifying Condition 3 and all combinations of two rooted trees with  $\frac{n}{2}$  nodes verifying Condition 4.

```
 \begin{array}{l} \textbf{Procedure KP} (n) \left\{ \\ k := \lfloor n/2 \rfloor; R[1..k] : \text{sets of rooted trees}; T : \text{set of trees}; \\ R[1] := \{0\}; NofAlk := 0; T := \emptyset; \\ \textbf{for } p = 2 \ \textbf{to} \ k \ \textbf{do} \left\{ & \{\text{rooted tree generation}\} \\ R[p] := \emptyset; \\ \textbf{for each} [a(\mathbf{x}), \mathbf{b}(\mathbf{y}), \mathbf{c}(\mathbf{z})] \ \textbf{s.t. Condition 1 holds } \textbf{do} \\ R[p] := R[p] \cup \{\textbf{Compose}(R[a], x, R[b], y, R[c], z, \emptyset, 0)\}; \\ \}; \\ \textbf{if} (Odd(n)) \left\{ & \{\text{odd n-tree generation}\} \\ \textbf{for each} [a(\mathbf{x}), \mathbf{b}(\mathbf{y}), \mathbf{c}(\mathbf{z}), \mathbf{d}(\mathbf{w})] \ \textbf{s.t. Condition 2 holds } \textbf{do} \\ T := T \cup \{\textbf{Compose}(R[a], x, R[b], y, R[c], z, R[d], w)\}; \\ \} \ \textbf{else} \left\{ & \{\text{even n-tree generation}\} \\ \textbf{for each} [a(\mathbf{x}), \mathbf{b}(\mathbf{y}), \mathbf{c}(\mathbf{z}), \mathbf{d}(\mathbf{w})] \ \textbf{s.t. Condition 3 holds } \textbf{do} \\ T := T \cup \{\textbf{Compose}(R[a], x, R[b], y, R[c], z, R[d], w)\}; \\ T := T \cup \{\textbf{Compose}(R[a], x, R[b], y, R[c], z, R[d], w)\}; \\ T := T \cup \{\textbf{Compose}(R[a], x, R[b], y, R[c], z, R[d], w)\}; \\ Nof Alk := \textbf{CardinalityOf}(T); \\ \textbf{return}(NofAlk, ElapsedTime, T); \\ \}. \end{array}
```



#### 4 Two new enumeration algorithms

In this section we present two new enumeration algorithms whose main difference with respect to TNKMS algorithm is the introduction of rules avoiding the multiple generation of trees. The two algorithms, namely AHM and HPVK ([7]), have the same structure but use different codes to represent trees: AHM uses the CN-TUPLE code while HPVK uses the N-TUPLE code.

As already discussed in section 3 (see Figure 2), the enumeration scheme suggests the use of some rules guaranteeing the uniqueness of the enumerated trees. A first attempt can be carried out by implementing a list of already enumerated trees: a new tree is accepted if and only if it does not belong to the list. The use of a list of codes (ordered or not) requires a large amount of computational resources, in terms of time and memory, as we will show in section 6.

In order to avoid the use of such a list, we introduced two techniques which prevent the generation of multiple copies of the same tree, namely *Reverse Search* (RS) and *Symmetry Detection* (SD).

Let  $T_o, T_d$  be two trees such that  $T_d$  can be generated adding one pendant node to  $T_o$  in such a way that the degree constraint holds. Let the tree generation function  $G_T : T_o \mapsto T_d$  be the function describing such operation. Moreover, let  $D(T_o) = \{T_d : G_T(T_o) = T_d\}$  be the multiset of trees generated from  $T_o$  and  $O(T_d) = \{T_o : G_T^{-1}(T_d) = T_o\}$  be the multiset of trees from which  $T_d$  can be generated. Note that multiple copies of the same tree may belong to  $D(T_o)$  and  $O(T_d)$ . For example:

 $D(200) = \{2100, 2100, 3000\}, O(31000) = \{3000, 3000, 3000, 2100, 2100\}.$ 

Given a tree and a root r, for any node v let  $l_r(v)$  be the distance (i.e. the number of edges) between v and r.

RS has been introduced by Avis and Fukuda ([2]) for efficiently enumerating polyhedra's vertices and it can be considered as a particular search on a graph.

We summarize the basic idea of Reverse Search in the following, referring to [2] for a more formal and detailed description. Let G be a connected graph whose vertices are the objects to be enumerated, and suppose that we have an objective function to be maximized over all vertices of G. A local search algorithm on G is a deterministic procedure which explores the neighborhood of a given vertex of G to find a vertex improving the objective function value. The search is repeated until there exist no better neighboring vertex. For simplicity, suppose that the considered local search avoids cycling and there is only one optimal vertex  $x^*$ . Consider now the digraph T having the same vertices of G and the edges (x, x') corresponding to the moves from x to x' performed by the local search algorithm starting from each vertex of G. Since  $x^*$  is the unique global optimum, T turns out to be a spanning arborescence having  $x^*$  as sink. Thus if we trace backward T from  $x^*$  systematically, e.g. by depth first search, or, in other words, reversing the local search, we can enumerate all vertices. Note that RS does not require storing any information about visited vertices.

By applying RS to our problem with the lexicographical order as the objective function, we derive the following *enumeration rule*.

**Enumeration Rule 1 (RS)** An enumerated tree  $T_d = G_T(T_o)$  is accepted if and only if  $T_o$  is the lexicographically maximum among the codes in  $O(T_d)$ . For example,  $T_d = 31000$  is accepted if  $T_o = 3000$  and it is not accepted if  $T_o = 2100$ .

Let us now illustrate the Symmetry Detection technique, that is the way to recognize identical subtrees or *symmetries* within a given  $T_o$ . Consider  $T_o$  as composed by two or more identical subtrees; clearly, it is possible to generate two or more identical  $T_d$  by adding a pendant node to any of the identical subtrees of  $T_o$ . For example,  $T_d = 2100$  can be generated twice from  $T_o = 200$  (see Figure 5).

Consider  $T_o$  with root r and two subtrees  $S_1$  and  $S_2$  of  $T_o$  with roots vand w respectively.  $S_1$  and  $S_2$  are symmetrical if and only if i) the code of  $S_1$  is equal to that of  $S_2$  and ii)  $l_r(v) = l_r(w)$ . The enumeration rule is:

**Enumeration Rule 2 (SD)** Given  $S_i$  symmetrical subtrees of  $T_o$  with  $i = 1, \ldots, q, 2 \le q \le 4$ , the trees generated from  $T_o$  are those obtained expanding  $S_q$  thus skipping  $S_1, \ldots, S_{q-1}$ .



We observe that the enumeration rule 2 can be applied recursively to all subtrees of  $S_q$  and to all non symmetrical subtrees of  $T_o$ . Note that the choice of  $S_q$  instead of  $S_1$  is only motivated by implementation reasons (see §4.1).

The use of RS and SD techniques guarantees the uniqueness of enumerated trees. Enumeration rule 1 avoids multiple enumeration of trees from different  $T_o$  while enumeration rule 2 avoids multiple enumeration of trees from the same  $T_o$ .

#### 4.1 Implementation details

We report only the main details concerning the implementation of algorithm AHM. These observations can be extended to HPVK in a straightforward manner considering the fact that this algorithm uses *N*-TUPLE codes, hence it requires the use of simpler data structures.

In particular AHM makes use of the *code tree* data structure (CTREE for short) which is exemplified in Figure 6.

CTREE is a rooted tree with root in r which is decomposed into levels where a *level* i is the set of nodes v such that  $l_r(v) = i$ .



**Fig. 6** A tree with the indication of  $l_r(v)$  and the corresponding CTREE

Starting from a pendant node whose code is 0, the code of a node v is obtained concatenating the number of subtrees incident in v with the subtree codes in lexicographical order.

CTREE drives the code computation of a new enumerated tree and the application of the enumeration rules 1 and 2.

First of all, we observe that the subtree code  $C = c_1 c_2 \dots c_l$  can be easily manipulated: if a pendant node is added to the root of subtree the new code is  $C' = (c_1 + 1)c_2 \dots c_l 0$ ; otherwise, if a pendant node is deleted from the root, the code is  $C' = (c_1 - 1)c_2 \dots c_{l-1}$ .

Figure 7 shows how the CTREE of  $T_o$  can be used to compute a new code  $T_d$ : first we compute the new code of the right subtree where a pendant node is added (200); then the codes at same level are lexicographically ordered (200, 110) to correctly compose the subtree code at the upper level (2 200 110); these operations (*ordering* and *composing* steps) are repeated until level 0 is reached, that is until the new code  $T_d$  is computed.



Fig. 7 New code computation

The same procedure can be utilized to compute  $T_o$  code from  $T_d$ . The new subtree code is obtained by deleting a pendant node, instead of adding it. Then, the ordering and composing steps are repeated in the same way as before until level 0 is reached.

The enumeration rule 2 can be implemented by performing a *breadth first search* (BFS) on CTREE : finding a symmetry is equivalent to finding two nodes at the same level with the same code.

The enumeration rule 1 can be implemented by generating all possible  $T_o$  codes and checking if the lexicographically largest one corresponds to the code of the current  $T_o$ . Note that the enumeration rule 2 can be also used to compute  $T_o$  trees.

Procedure **GenSetD** (see Figure 8) taking as input a tree  $T_o$ , generates the set  $D(T_o)$  by applying the enumeration rules 1 and 2. In particular, the CTREE corresponding to the input tree is generated and the codes belonging to level 1 are inserted into a queue **Q**. Following the BFS search, at each iteration a code u is extracted from **Q**; if u is equal to the first code in **Q** then it is skipped (enumeration rule 2); otherwise, if the degree constraint holds  $(adj(u) \leq 3)$ , the procedure tries to add a pendant node to u and recomputes the code corresponding to the new tree, if the enumeration rule 1 applies. The pseudo-code of **GenSetD** is depicted in Figure 8.

```
Procedure GenSetD (T, D) {
    D, O : sets of trees; Q : queue;
    \mathbf{MakeCTreeOf}(T); \ D := \emptyset; \ Q := \{v : v = succ(root(T))\};
    while Q \neq \emptyset
         u = \mathbf{GetFirst}(Q); \ Q := Q \setminus \{u\};
         if (code(u) \neq code(next(u))) {
                                                               {enum. rule 2}
             \begin{array}{l} Q:=Q\cup\{v:v=succ(u)\};\\ \text{if }(adj(u)\leq 3) \end{array} \}
                  t := \mathbf{NewCode}(u, l(u));
                  m := \mathbf{Compute} \mathbf{Max}(O(t));
                                                                  {enum. rule 1}
                  if (T = m) D := D \cup \{t\};
             };
         };
    };
    return(D);
}.
```



The procedure **OneToOneEnumeration** (see Figure 9) generates the set of all *n*-trees by calling procedure **GenSetD**. The pseudo-code reported is amenable to both AHM and HPVK algorithms.

## 5 Complexity of algorithms

In this section, we compare the computational complexities of AHM, HPVK and KP algorithms. The complexity of TNKMS is not reported since the algorithm structure is not described precisely in the original paper ([12]).

```
\begin{array}{l} \textbf{Procedure OneToOneEnumeration } (n) \ \{ \\ Trees[1..n] : sets of trees; \\ Trees[1] := \{10\}; \ NofTrees[1] := 1; \\ Trees[2..n] := \emptyset; \ NofTrees[2..n] := 0; \ node := 2; \\ \textbf{while } ( \ node \leq n \ ) \ \{ \\ \textbf{while } ( \ Trees[node - 1] \neq \emptyset \ \{ \\ t := \textbf{GetAndDel}(Trees[node - 1] \ ); \\ \textbf{GenSetD}( \ t, \ Trees[node] \ ); \\ NofTrees[node] := \textbf{CardinalityOf}(Trees[node]); \\ \}; \\ node := node + 1; \\ \}; \\ \textbf{return}( \ NofTrees[n], \ ElapsedTime, \ Trees[n] \ ); \\ \}. \end{array}
```

Fig. 9 AHM and HPVK algorithms

#### 5.1 KP algorithm

The complexity of procedure **KP** depends on the number of combinations (1) subject to Conditions 1-4. We recall that the problem of finding all possible combinations of t rooted trees with no more than k nodes is equivalent to finding all possible combinations of t numbers less than or equal to k in such a way that their sum equals n - 1.

Let  $C_c$ ,  $N_t$  and  $N_q$  be respectively the cost of composing a new code, the number of triplets verifying Condition 1 and the number of quadruplets verifying both Conditions 2 and 3. Let  $N_c$  be the number of possible combinations of rooted trees associated with each triplet or quadruplet. Note that Condition 4 holds only for one combination. The complexity of cycles in **KP** (see Figure 4) is given by

$$O(k \times C_c \times N_c \times N_t) \tag{2}$$

for the generation of p-rooted trees, and

$$O(C_c \times N_c \times N_q) \tag{3}$$

for the generation of n-trees. The proof of the following Lemma is straightforward.

**Lemma 1** The number of triplets (a(x), b(y), c(z)) verifying condition 1 are  $O(n^2)$ . The number of quadruplets (a(x), b(y), c(z), d(w)) verifying both conditions 2 and 3 are  $O(n^3)$ .

Let  $K_p$  be the cardinality of *p*-rooted set. The number of possible combinations of *m p*-rooted trees  $(p = 1, .., \lfloor \frac{n}{2} \rfloor)$  is  $O(H^m)$  where *H* is the maximum cardinality of rooted trees sets used to generate the *p*-rooted set in procedure **Compose**, that is  $H = \max\{K_a, K_b, K_c\}$  with ax+by+cz = p-1according to Condition 1. Note that the function **Compose** in **KP** generates a new code by composing the subtree codes in lexicographical order in linear time. Therefore,  $C_c$  is O(n). According to (2) and (3), the rooted tree generation cycle is  $O(n^4 K^3_{\lfloor \frac{n}{2} \rfloor})$  while the *n*-tree generation cycle is  $O(n^4 K^4_{\lfloor \frac{n}{2} \rfloor})$ . Moreover, for even *n*, the generation of bicentroids is  $O(nK^2_{\lfloor \frac{n}{2} \rfloor})$ . Therefore, we have

**Theorem 2** Algorithm KP is  $O(n^4 K_{\lfloor \frac{n}{2} \rfloor}^4)$ .

#### 5.2 AHM and HPVK algorithms

AHM and HPVK have the same computational complexity since the only difference between the two algorithms is that HPVK does not use the CTREE data structure to drive the implementation of the enumeration rules 1 and 2.

The complexity of AHM depends on the time required to compute the code of a new enumerated tree  $T_d$  (procedure **NewCode**) and to compute  $T_o$  codes when the algorithm checks the enumeration rule 1 (procedure **ComputeMax**). We show that this computation can be carried out in linear time. As depicted in Figure 7, this computation consists of the subtree code computation and the consequent ordering and composing steps.

The code of a modified subtree (i.e. obtained by adding or removing a pendant node) is computed in constant time (see  $\S4.1$ ).

The ordering step consists of checking the lexicographical order of two codes which can be carried out in linear time in the code length d. The worst case arises when from a code we generate a complete CTREE with L levels. Let  $d_j$  be the length of a subtree code at level j = 0, ..., L, therefore we have:

$$d_0 = n, \quad d_1 = \frac{n-1}{4}, \quad d_L = 0.$$
  
 $d_j = \frac{d_{j-1}-1}{3}, \quad j = 2, \dots, L-1,$ 

In fact, by definition of CTREE,  $d_0 = n$  and  $d_L = 0$ . Since the CTREE is complete, the root has 4 subtrees whose code lengths are equal to  $\frac{n-1}{4}$ .

At level  $j \ge 2$ , the length of the 3 subtree codes is  $\frac{d_{j-1}-1}{3}$ . The number of codes compared are 2 at level  $j = 2, \ldots, L-1$ , and 3 at level j = 1. Each comparison requires  $O(d_i)$  steps. Hence the overall number of steps is:

$$3d_1 + \sum_{j=2}^{L-1} 2d_j \le 3\sum_{j=1}^{L-1} d_j = 3\left[\frac{n-1}{4} + \frac{n-1}{4} - \frac{1}{3} + \dots\right]$$
$$\le 3\left[\frac{n-1}{4} + \frac{n-1}{43} + \dots\right] \le 3\sum_{j=1}^{L-1} \frac{n-1}{43^j} = \frac{3}{4}(n-1)\sum_{j=1}^{L-1} \frac{1}{3^j}$$
$$\le \frac{3}{4}(n-1)\sum_{j=1}^{\infty} \frac{1}{3^j} = \frac{3}{4}(n-1)\frac{1}{2} = \frac{3}{8}(n-1)$$

which is O(n).

Finally, the concatenation of two or more subtree codes is O(n). Therefore, the computation of a new code after adding or deleting a pendant node is O(n).

Procedure **GenSetD** (Figure 8) visits once all n-1 nodes of CTREE in the worst case. For each one, the application of enumeration rule 2 is O(n), the encoding of a new enumerated tree  $T_d$  is O(n) and the application of enumeration rule 1 is  $O(n^2)$ : in fact, the codes of all n-1 possible  $T_o = G_T(T_d)$  are obtained in O(n). This means that **GenSetD** is  $O(n^3)$ .

By consequence, the overall complexity of procedure **OneToOneEnu**meration (Figure 9) is given by

$$\sum_{i=1}^{n-1} i^3 S_{i-1}.$$

where  $S_i$  is the number of *i*-trees. Since the following inequalities hold

$$\sum_{i=1}^{n} i^{3} S_{i} \leq \sum_{i=1}^{n} i^{3} S_{n} \leq S_{n} \sum_{i=1}^{n} i^{3} = S_{n} \left( \sum_{i=1}^{n} i \right)^{2} = S_{n} \frac{n^{2} (n+1)^{2}}{4} \quad ,$$

we have theorem 3.

**Theorem 3** Algorithms AHM and HPVK are  $O(n^4S_n)$ .

$$\diamond$$

Table 1 Computational complexities of reported algorithms

Enumeration algorithms								
KP	HPVK	AHM	TNKMS					
$O(n^4 K^4_{\lfloor \frac{n}{2} \rfloor})$	$O(n^4 S_n)$	$O(n^4 S_n)$						

Table 1 summarizes the complexity of algorithms. The reported complexities depend on the number of carbonium atoms n, that is the size of each generated tree, and the size of the set of trees (rooted tree in the case of KP algorithm) used by the algorithms to generate the required class of alkanes. For example, for n = 20,  $K_{10} = 879$  while  $S_{20} = 251731$  (see table 2). Looking at these results, it is not possible to select a priori which is the best algorithm since it is difficult to clearly evaluate any single contribution to the complexity functions. Therefore, running time comparisons are also required.

## 6 Running time comparisons

This section reports some computational results based on our running time experiences made on Silicon Graphics Indingo 4000, processor Risc IP20 100



Fig. 10 List vs RS with SD

MHz, 48 Mbytes of main memory. AHM, HPVK and KP are implemented in Pascal while TNKMS in Fortran77, as reported in [12].

Figure 10 plots the running times performances obtained by comparing the use of Reverse Search and Symmetry Detection techniques and the use of a list of codes in order to avoid multiple generations. The joint use of these techniques outperforms the use of an unordered list. Moreover, they allow to save memory since they do not require storing all generated codes. Therefore, in principle, these techniques make it possible to enumerate classes of alkanes with larger numbers of carbonium atoms.

Table 2 reports the running times of the algorithms presented in the course of the paper. TNKMS appears the slowest algorithm to generate the alkane molecular family. This might be due to the lack of rules avoiding multiple enumerations of trees. AHM and HPVK have similar running times and their gaps depend on the different computer implementations. KP is surprisingly the best algorithm: with n = 21, KP is about 4, 5 times faster than HPVK and AHM and about 1320 times faster than TNKMS.

The main reason seems to be the fact that KP builds the alkane molecular family employing only  $\lfloor \frac{n}{2} \rfloor$ -rooted trees. On the contrary, the other algorithms need to generate all the alkane molecular families up to n-1. In other words, KP generates 879 rooted trees to enumerate all the 366319 20-trees while AHM, HPVK and TNKMS need 251731 trees.

One may observe that contrary to the KP algorithm, AHM, HPVK and TNKMS algorithms give as output not only the set of all trees of a given cardinality n, but they also give as a byproduct all the sets of trees of cardinality m with m < n. However, in order to achieve the same result also utilizing algorithm KP we may run it once for each possible value of m. The overall running time would be the sum of running times of each run. It is clear from table 2 that even in this case KP proves to be much faster than the other algorithms. Moreover, an optimized algorithm computes once the rooted trees needed to enumerate all alkane families saving further computational time.

n	$S_n$	HPVK	AHM	TNKMS	KP	$\sum_{p=1}^{\lfloor \frac{n}{2} \rfloor} K_p$
16	10359	0"	0"	45"	0"	161
17	24894	2"	2"	2'43''	0"	161
18	60523	6"	7"	9'50"	2"	372
19	148284	14"	16"	35'44''	4"	372
20	366319	37"	42"	2h11'20''	15"	879
21	910726	1'38"	1'52''	8h04'39''	22"	879
22	2278658	4'17"	4'57"		1'34"	2117
23	5731580	11'11"	13'09"		2'25''	2117
24	14490245	29'22''	31'39"		10'18"	5174
25	36797588	1h18'43"	1h24'11"		15'58''	5174
26	93839412	3h33'50"	3h49'33''		1h06'44''	12813
27	240215803	9h09'35"	9h36'55''		$1\mathrm{h}46'25"$	12813

Table 2 KP, HPVK, AHM and TNKMS running time

In conclusion, KP appears the best algorithm to enumerate an alkane molecular family with n carbonium atoms.

## 7 Conclusions

We have presented four different algorithms to enumerate the alkane molecular family  $C_n H_{2n+2}$  for any given n.

Reading the alkane enumeration literature, we have noted that remarkable work on this topic has been done by researchers not deeply involved with algorithmic aspects. They often avoid studying or reporting on the computational complexity and the running times. For these reasons, we reported a detailed complexity study.

One-to-one enumeration requires some rules which avoid duplications. We have shown how Reverse Search and Symmetry Detection techniques are useful to avoid multiple enumeration and to save memory allowing the enumeration of larger families. Moreover, they speed up the running time of the algorithms HPVK and AHM in comparison to TNKMS. Jordan's Theorem guarantees the uniqueness in the constructive enumeration algorithm. As shown in section 6, KP is surprisingly the best algorithm.

#### Acknowledgments

The authors are indebted to the Editor and to the anonymous referee for their comments and suggestions which improved the presentation.

#### References

 Aringhieri R., Hansen P. and Malucelli F., A Linear Algorithm for The Hyper Wiener Index of Chemical Trees. J. Chem. Inf. Comput. Sci., 2001, 41(4), 958-963.

- Avis D.; Fukuda K., Reverse Search for enumeration. Discrete Applied Mathematics 1996, 65, 21-46.
- 3. Berzelius J. J., Ann. Phys. Chem. 1830, 19, 305-.
- Cayley A., On the Analytical Figures called Trees in Mathematics and Their Applications to the Theory of Chemical Combinations. Ber. Dtsch. Chem. Ges. 1875, 8, 1056-1059.
- 5. Corey E. J.; Wipke W. T., Science 1969, 166, 178-.
- Hansen P.; Jaumard B.; Labatteux C.; Zeng M., Coding Chemical Trees with the Centered N-Tuple Code. J. Chem. Inf. Comput. Sci., 1994, 34, 782-790.
- 7. Hansen P.; Pitteloud G.; Van der Klink R., Enumeration d'Alcanes par le Code N-Tuple et la Recherche Inversée. *Travail de semestre* 1995, DMA, EPFL, Lausanne.
- 8. Jordan C., Reine Angew. Math. 1869, 185-.
- Kvasnička V.; Pospichal J., Constructive Enumeration of Acyclic Molecules. Collect. Czech. Chem. Commun. 1991, 56, 1777-1802.
- Lederberg J.; Sutherland J. L.; Buchanan B. G.; Feigenbaum E. A.; Lindsay R. K., Application of Artificial Intelligence for Organic Chemistry: the DENDRAL project. McGraw-Hill: New York, 1980.
- Read R. C., The Coding of Various Kind of Unlabelled Trees in *Graph Theory* and *Computing*; Ed Read R. C., Academic Press: New York, **1972**, Chapter 12, 153-.
- 12. Trinajstić N.; Nicolić S.; Knop J. V.; Müller W. R.; Szymanski K., Computational Chemical Graph Theory. Ellis Horwood: New York. **1991**.