New Topological Information Indices Based on the Full Neighborhood of All Atoms

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Introduction

We present a new family of topological information indices based on the full neighborhood of all atoms.

In previous definitions, mostly certain partitions of atoms have been used for this purpose [1-3]. We consider each atom of a molecular structure as a sub-system.

For each atom the complete neighborhood is characterized by an information functional [4], considering the number of atoms in all possible spheres around the atom.

An appropriate weighting scheme combines the number of atoms in the different spheres resulting in a characteristic property of the atom.

The properties of all atoms are normalized to give "probabilities for the sub-systems" necessary for the computation of an entropy measure.

In the current version only skeletons of the chemical structures are considered, with all atoms equal and all bonds equal.
Method

A chemical structure is represented by a graph.
The graph consists of \( n \) subsystems corresponding to the \( n \) atoms of the structure.
For each subsystem the value \( f_i \) of an invariant is calculated based on the complete neighborhood.
We call \( f_i \) a special information functional.
The values of the invariants are normalized to give "probabilities" \( p_i \) that are combined to an entropy measure \( E \), defining a molecular descriptor.

\[
\begin{align*}
  f_i &= c_1 s_{i1} + c_2 s_{i2} + \ldots + c_d s_{id} \\
  s_{ik} &= \text{number of atoms in sphere } k \text{ of atom } i \\
  c_k &= \text{weight for sphere } k \text{ (e.g. linearly decreasing with increasing } k; c_1 = d; c_2 = d - 1; \ldots, c_d = 1) \\
  d &= \text{topological diameter of the structure/graph} \\
  p_i &= \frac{f_i}{\sum_{j=1}^{n} f_j} \\
  E &= a \left( \ln n + \sum_{i=1}^{n} p_i \ln p_i \right)
\end{align*}
\]

- \( a \) is a scaling constant, e.g. 1000.
- If all atoms are topologically equivalent (vertex transitive), \( E = 0 \). Examples: rings, prism, tetraeder, cube
- \( E \) increases with increasing "neighborhood-diversity" of the atoms
  Examples: chain structures
Results

Relationship to other molecular descriptors

A PCA loading plot is used to characterize the multivariate similarity of molecular descriptors, including the new information index $E$.

**Data**

$n = 3943$ chemical structures, randomly selected from a spectroscopic database [5].

$m = 211$ molecular descriptors calculated by software *Dragon* [6] from 2D H-depleted structures.

**PCA loading plot**

Calculated from autoscaled descriptors. Descriptors most similar to $E$ - and some others are marked.
Results

Distribution of $E$ for various structures

3,943 randomly selected structures from a spectroscopic database
16,979 (all) graphs with 12 vertices, containing 2 rings [7]
3,232 (all) graphs with 12 vertices, containing 1 rings [7]
355 (all) tree graphs with 12 vertices [7]

<table>
<thead>
<tr>
<th>Structure</th>
<th>Minimum $E$</th>
<th>Maximum $E$</th>
</tr>
</thead>
<tbody>
<tr>
<td>trees</td>
<td>1.8413</td>
<td>11.6165</td>
</tr>
<tr>
<td>one ring</td>
<td>0</td>
<td>12.4312</td>
</tr>
<tr>
<td>two rings</td>
<td>0.1678</td>
<td>12.7809</td>
</tr>
<tr>
<td>spec database</td>
<td>0</td>
<td>11.6431</td>
</tr>
</tbody>
</table>

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Summary

- $E$ characterizes the diversity of the atoms in terms of neighborhood, and thereby a special type of structural complexity and inner symmetry.
- In contrary to previously defined information indices, $E$ uses each atom separately (and not in groups), and the neighborhood of atoms considers the whole molecule.
- Extensions of the basic measure $E$ for colored graphs (different atoms and different bonds) is under development.
- We generalized the classical information indices because our measure is parameterized and allows the incorporation of various information functionals. Thus, these molecular descriptors can be optimized by machine learning techniques using appropriate data sets.

References