Abstract. Main topoenergetic principles are reviewed in view to emphasize the UNIVERSAL relationships between time conversion of a system in transformation and one of its governing potentials. It results that: (i) any quantity can be calibrated in unit of time and (ii) any quantity can be calibrated by another one.

1 Short history of topoenergetic principles

Topoenergetic principles were initiated by calorimetric studies of polyethylene morphologies created by different thermal, mechanical and/or chemical treatments [1,2]. The basic observation was that all systems in transformation have composite structure because at least two components interact between them and exchange energy (heat) with the surrounding medium (Figure 1). Calorimetry was the most appropriate analytical technique allowing development of topoenergetic principles [1,3]. In view to quantitatively describe the kinetics of the transforming process by its components, I considered the energy circuit in similar terms like electric circuits, i.e. as composed by components with elementary behaviours: resistors, capacitors, energy sources, etc. This idea was launched in the same period of time by Oster and Auslander [4], but unfortunately they considered on the one hand very complicated cases and on the other hand that these components are spatially distributed in the transforming system. In this way mathematical description became unrealistic.

The first kinetic equation established in topoenergetic terms was a modified Arrhenius equation applied for crystallization of polyethylenes by using a so called “drop calorimetry” [5]. The experimental principle of this technique consists in keeping first the calorimetric cell with the specimen at a temperature over melting point, Tm, and dropping it in the calorimeter already prepared at crystallization temperature, T. Two exothermal flows appear, namely: one immediately at the contact with calorimeter (at time considered zero) given by inert component, Cin, denoted as win, and the flow delayed at a specific time, ti, given by component in transformation, Ctr (wtr), in this case the phase which crystallizes. The kinetic equation is:

\[ \ln(ti*T) = - \frac{E}{R*T} + K \]

where \( E \) – activation energy, \( R \) – the gas constant, T in Kelvin and the free term

\[ K = \ln(Cin*R*E/R) \]

Diagram:

- Cin – inert component
- Ctr – transforming component
- ctr – kinetic unit

Figure 1. Composite structure of a test specimen in transformation.

* Corresponding author: dragan_gdf@yahoo.com, gdf.dragan@gmail.com

This is an Open Access article distributed under the terms of the Creative Commons Attribution License 2.0, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Article available at http://cfmetrologie.edpsciences.org or http://dx.doi.org/10.1051/metrology/201317001
where $\mathbf{R}$ is the dissipative (resistive) element specific to the experimental device. It is important to mention that if we repeat experiments with the same sample, but using different specimen size and/or calorimeters, the parameters ($t_i$, $E$, $K$) may be different. In this experiment temperature is the potential governing the transforming process and $T_m$ is the threshold value above which the crystallization does not occur. In other words, the calorimetric experiment reveals the evolution of the test specimen between two equilibrium states located on the both sides of the threshold value. The same equation was proved to be valid for processes occurring by increasing temperature, i.e. the process occurs at a temperature over the threshold value (for instance melting [6], curing-polymerization [7], etc.). By considering more transforming processes driven by temperature by using different measuring systems (analytical techniques), but the same principle of stepwise perturbation crossing a threshold value, kinetic equation remains valid [8, 9].

Table 1. ARRHENIUS representation:

$$\ln(t_i^*T) = -\frac{E}{(R^*T)} + K;$$
$$K = n_1^*E + m_1$$

Table 2. UNIVERSAL representation:

$$\ln(t_i) = N*\ln|U - U_o| + M$$

$$M = n_1*N + m_1$$

Another important parameter is the process polarity, $P$, which was initially defined as a trivial parameter for thermally driven processes as revealed in calorimetric systems (Table 1). UNIVERSAL representation allows defining also the coupling strength (CS) between Ctr and Cin (Table 2). For groups of samples showing different nature of transforming processes, it is possible to establish higher phylogenies [1,6], so the overall experimental data have a pyramidal structure (Table 3). For instance energy structure of atoms and all atomic spectra can be described by a unique pyramid of topoenergetic parameters [10].

2 Standard Experimental Conditions (SEC)

Figure 2 suggestively presents the experimental procedure in view to define the ontogeny of a system according to topoenergetic principles. There are represented three sections: the bellow sections represent the stepwise perturbation on the governing potential $U$ by increasing (1) or decreasing (2) crossing the threshold value $U_o$. In the upper section the triggered process of transformation is revealed by the time conversion of a response function, $\Theta$, between the two equilibrium values $\Theta_{1,2}$, or by its derivative, $d\Theta/dt$, defining the induction time, $t_i$ as the eigenvalue from the considered response function. In general the effect of Cin can be removed and/or neglected. It is important to mention that in view to define the ontogeny parameters and subsequently the higher phylogenies, all experiments must be performed in the same Standard Experimental Conditions (SEC), namely:

(i) Conditions of repeatability;
(ii) the same size & shape of tested specimens;
(iii) by applying an increasing or decreasing stepwise perturbation of governing potential, $U$, crossing the threshold value $U_o$ specific to the transforming process;
(iv) by considering the same response function, its eigenvalue, governing potential and by keeping unchanged the units for all quantities during all measurements;
(v) linear relationships can be established between parameters obtained in two different repeatable conditions [11].

17001-p.2
3 Metrological aspects

The UNIVERSAL kinetic equation establishes the univocal relationship between \( t_i \) and \( U \) specific for a transforming process which can be considered as a calibration curve of \( U \) units in time units. Furthermore, \( t_i \) is a particular case of eigenvalue from time conversion of the response function. It can be replaced by other specific quantities; for instance, electric resistance of NTC-thermistors as a function of temperature, a.c. electric conductivity quantities as a function of solute concentration in electrolyte solutions, etc. [10]. Topoenergetic principles can solve many problems of actual metrology (for instance viscosity).

Table 3. Pyramidal structure of data bank obtained according to topoenergetic principles.

<table>
<thead>
<tr>
<th>(ni, mi)</th>
<th>higher phylogeny parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>(n1, m1)</td>
<td>first phylogeny parameters</td>
</tr>
<tr>
<td>(E, K), (N, M, Uo)</td>
<td>ontogeny parameters</td>
</tr>
<tr>
<td>(t1, U)</td>
<td>basic experimental data obtained in accurate SEC</td>
</tr>
</tbody>
</table>

References

[10] GDF Databanks Bull. (ISSN 1453-1674) publishes applications of topoenergetic principles. It is registered at Biblioteca Nationala a Romaniei and National Library of Australia and it is also posted on the website www.gdfdatabanks.ro