Opinion: Chemical --Mesoscopics..for the Mesoparticles Reactivity Explanation

V.I. Kodolov¹,²* V.V. Kodolova-Chukhontseva³

1. Basic Research - High Educational Centre of Chemical Physics and Mesoscopics, Udmurt Federal Research Centre, UD, RAS, Izhevsk, Russia
2. M.T. Kalashnikov Izhevsk State Technical University, Izhevsk, Russia
3. Institute Macromolecular Compounds, RAS, StPetersburg, Russia

ARTICLE INFO

Article history
Received: 6 September 2020
Accepted: 30 October 2020
Published Online: 30 November 2020

Keywords:
Kolmogorov-Avrami equation
Relative energetic and volume characteristics
Semi empiric Taft constants
Reaction series notion in chemical processes

ABSTRACT

The estimation of chemical particles reactivity and the determination of chemical reactions direction are the actual theme in new scientific trend - Chemical Mesoscopics. Paper includes the proposal about the using the theory of free energy linear dependence from physical organic chemistry and their applications for prognosis of reactions flowing. The semi-empiric constants is given according to mesoscopic physics definitions as well as the transformed Kolmogorov-Avrami equation is discussed. It is the development of Chemical Mesoscopics for organic reactivity estimation including nanostructures reactivity.

1. Introduction

Chemical Mesoscopics is based on the notions about self organization and self similarity [1].

According to the fractal theory [1] any system can be presented as aggregate of elements similar to whole system. These elements have own energetic and geometric (volume) parameters owing to which they are found within the system. The change of these parameters because of the action of external factors leads to disturbance of system balance. At this case the system is destructed or transformed. The estimation of these changes is possible with the using of the relative parameters in which the energetic and volume values are compared with the definite standard values for the correspondent elements (fragments) in the definite reaction series. This approach to reactivity consideration is near to Taft and Pal’m theoretical works [2]. For the relative energetic parameters the following formula \((\varepsilon - \varepsilon_0)/\varepsilon_0\) is proposed [3], where \(\varepsilon_0\) corresponds to the surface energy for standard chemical fragment. In turn, analogous relative parameters are proposed [4] for the volume characteristics \((V - V_0)/V_0\).

*Corresponding Author:
V.I. Kodolov,
Basic Research - High Educational Centre of Chemical Physics and Mesoscopics, Udmurt Federal Research Centre, UD, RAS, Izhevsk, Russia;
M.T. Kalashnikov Izhevsk State Technical University, Izhevsk, Russia;
Email: vkodol.av@mail.ru
2. Discussion

The development of Chemical Mesoscopics in this direction is connected with the research of the size and energetic characteristics of chemical particles. The size of mesoparticles is denoted as approximately 10 nm, and the motion freedom of nanostructures (mesoparticles) is limited by the vibration with high frequency and electron transport across them. The peculiarities of mesoparticles consist in the radiation of energy quants of negative or positive charges. This radiation is the main reason of the stimulation of chemical processes. At the imposition of the negative charge quants the interference takes place and the chemical bonds are formed. In turn the imposition of the negative and positive quants together the phenomenon of annihilation is created. At this case the direct electromagnetic field is appeared that leads to the stimulation of negative charge quants moving and the growth of chemical bonds formation. The phenomena of interference and annihilation are reasons of start for self organization process with reservation conformation order, which determine the finished product structure. For the process explanation the equation of Kolmogorov-Avrami\(^1\) can be used -

\[ W = 1 - \exp (-k\tau^n), \]

where \(W\) - the part of obtained product (for instance, polymer), \(k\) - the process rate constant, \(\tau\) - the duration of process, \(n\) - the fractal dimension (for one measured process \(n = 1\)).

For the comparative estimation of reagents (or nanostructures) in one reaction series it’s possible the application of the theory of free energy linear dependences. In this case the reactions are considered with using one of reagent as the standard compound for which \(W\) is fixed \(W_0\). The estimation of reactivity can be proposed on the difference \(W_0-W\), where \(W\) is calculated on formula 2, and \(W_0\) is defined on analogous formula with changes \(k_0\tau_0^n\). It’s noted, the fractal dimension \(n\) do not change because the comparison is carried out for one type of reactions. The following equation for difference \(W-W_0\) can be written -

\[ W - W_0 = \exp (-k_0\tau_0^n) - \exp (-k\tau^n) \]

and after eq. 2 transformation -

\[ W_0 = k/k_0 (\tau/\tau_0)^n - 1 \]

If \(\lg k/k_0\) is defined as -

\[ \lg k/k_0 = -2.3 \text{ RT } \left\{ \left[ (\varepsilon_0 - \varepsilon)/\varepsilon_0 \right] a + \left[ (V_0 - V)/V_0 \right] b \right\} \]

and this expression after transformation stand in the eq. 3, than the equation 5 is received -

\[ \lg (W - W_0) = (\tau/\tau_0)^n \exp \left\{ \left[ (\varepsilon_0 - \varepsilon)/\varepsilon_0 \right] a + \left[ (V_0 - V)/V_0 \right] b \right\} \]

where values \(a\) and \(b\) - parameters, which correct the influence polar and steric (spaced) effects on reactivity in polymerization, the relation \((\varepsilon_0 - \varepsilon)/\varepsilon_0\) corresponds to Taft constant \(\sigma\) (polarity constant), and the relation \((V_0 - V)/V_0\) - Taft constant \(E_s\) (steric or spaced constant)\(^3\).

3. Conclusion

The application of above notions for the Chemical Mesoscopics development is very perspective because stimulates the mathematic apparatus creation for the chemical processes flowing direction prediction.

References

[1] V.I. Kodolov, V.V. Kodolova-Chukhotzeva. Fundamentals of Chemical Physics. Izhevsk: M.T. Kalashnikov IzhevSTU, 2019: 218.
[2] V.A. Palm. The introduction in theoretical organic chemistry. M.: Publ. “High School”, 1974: 446.
[3] V.I. Kodolov. Possibilities of modeling in organic chemistry. Organic Reactivity, 1965, 2(4-6): 11-17.
[4] V.I. Kodolov, S.S. Spasskiy. Parameters in Alfrey-Price and Taft equations. Vysokomol. Soed., 1976, 18(9): 1986-1992.