Further remarks on the new criterion differentiating between Non-Redox and Redox Electrolytic Systems

Abstract

The new general criterion, distinguishing between non-redox and redox electrolytic systems, is based on the properties of the linear combination \( f_i = 2\pi(O) - \pi(H) \) of elemental balances: \( f_i = \pi(H) \) for \( Y_i = H \), and \( f_i = \pi(O) \) for \( Y_i = O \). The \( f_i \) is the primary form of the Generalized Electron Balance (GEB), completing the set of balances needed to formulate a redox system, according to Generalized Approach to Electrolytic Systems (GATES). In the redox system, \( f_i \) is linearly independent on charge \( f_j \) and other elemental/core balances \( f_k = \pi(Y_k) \) \((k=3,...,K)\), i.e., \( f_0, f_1, f_2, ..., f_F \) form the set of independent balances needed to formulate the redox system. In a non-redox system, \( f_i \) is linearly dependent on the balances \( f_0, f_1, f_2, ..., f_F \) and is resolved with use of iterative computer program MATLAB [35,36]. The problem of oxidation numbers (ONs) calculation on the step of formulation of D+T mixture and its subsystems (D, T) is also considered.

Keywords: Electrolytic redox systems; GATES/GEBS; Approach I to GEB; Approach II to GEB; Potentiometric titration

Introduction

Electrolytic systems, of different degree of complexity, can be resolved according to Generalized Approach to Electrolytic Systems (GATES) [1-3], principles, perceived as the best thermodynamic approach to equilibrium and metastable [4-6] systems, where all attainable physicochemical knowledge on a system tested can be involved. The advantages of GATES are particularly related to redox systems, where the Generalized Electron Balance (GEB) is formulated within GATES/GEBS [1-3,6-34].

All the inferences made within GATES/GEBS are based on firm, algebraic foundations. It allows to understand far better all physicochemical phenomena occurring in a system in question, and improve some methods of analysis. All the facts confirm the huge potency of simulated calculations made according to GATES/GEBS principles, with all attainable and preselected physicochemical knowledge involved therein.

This paper provides an example of a dynamic electrolytic redox system realized according to titrimetric mode, where \( V \) mL of titrant T is added, up to a given point of the titration, into \( V_0 \) mL of titrand D, and \( V_0 + V \) mL of D+T mixture is thus obtained, if the additivity of the volumes is pre-assumed/valid. For modelling purposes, the titration is considered as the isothermal process realized in the closed system, separated from the environment by diathermal walls. The related system will be formulated according to the GATES principles, and resolved with use of iterative computer program MATLAB [35,36]. The problem of oxidation numbers (ONs) calculation on the step of formulation of D+T mixture and its subsystems (D, T) is also considered.

Preliminary notations

According to GATES/GEBS principles, any species \( X_i \) in an electrolytic system can be perceived in its natural form, i.e., as a hydrate \( X_i \cdot n_{H_2O} \) in aqueous (W=H\( _2 \)O) solution, where \( z_i \) \((z_i = 0, \pm 1, \pm 2,...)\) is the external charge of \( X_i \), expressed in terms of elementary charge unit \( e = F/N_a \) (F – Faraday constant, \( N_a \) – Avogadro’s number), and \( n_{H_2O} = n_{H_2O} (\geq 0) \) is the mean number of water \((W = H_2O)\) molecules attached to \( X_i \). The known chemical formulas of \( X_i \) (\( i = 1, ..., I \)...)
provide the information necessary/sufficient to formulate the respective balances.

As were stated above, the terms: components and species are distinguished, for the balancing purposes. In the notation applied here, \( N_j \) (\( j=1,2,\ldots,L \)) be the number of molecules of a component of \( j \)-th kind, composing a static (\( D,T \)) or dynamic \( D+T \) system, whereby the \( D \) and \( T \) are composed separately, from defined components, including water. The mono- or two-phase electrolytic system thus obtained involves \( N \) molecules of \( H_2O \) and \( N \) species of \( i \)-th kind, \( X^i_{1} \cdot n_{iW} \) (\( i=2,3,\ldots,L \)), denoted briefly as \( X^i_{1} \) (\( N,n \)), where \( n_{iW}=n_{iW}(H_2O) \); then we have: \( H^+ (N,n) \), \( OH^- (N,n) \), \( \ldots \), for ordering purposes. Molar concentration of the species \( X^i_{1} \cdot n_{iW} \) in \( D+T \) system is denoted as \( [X^i_{1}] \).

The GEB is recognized as the law of Nature \([1, 41]\), as the hidden connection of physicochemical laws, and as the breakthrough in thermodynamic theory of electrolytic redox systems. The GEB was discovered by Michałowski: as the Approach I to GEB (1992), and as the Approach II to GEB (2005). In the Approach I to GEB, perceived according to ‘card game’ principle, electron-active elements are perceived as ‘players’, electron-non-active elements as ‘fans’, and electrons as ‘money’; the knowledge of oxidation numbers is needed here. The Approach II introduces the balance \( f_{x_i} = 2 f_{x_j} - f_{x_k} \) as the combination of elemental balances: \( f_{x} = f(Y) \) for \( Y = H \) and \( f_{x} = f(O) \) for \( Y = O \). Charge balance (\( f_{x} = ChB \) and other, elemental and/or core balances \( f_{x} = f(Y_k) \) for \( Y = H \), \( O; k=3,\ldots,K \)) are also considered within GATES. A core is a cluster of different atoms with defined composition (expressed by chemical formula), structure and external charge, unchanged in the system considered. In a redox system, \( f_{x_i} \) is linearly independent on the balances \( f_{x_1}, f_{x_2}, \ldots, f_{x_k} \), i.e., a redox system is formulated with use of \( K \) independent balances \( f_{x_1}, f_{x_2}, \ldots, f_{x_k} \) that form the complete set of algebraic equations needed for the solution of the redox system, according to GATES/GEB principles.

In a non-redox system, \( f_{x_i} \) is linearly dependent on the balances: \( f_{x_1}, f_{x_2}, \ldots, f_{x_k} \), i.e., a non-redox system is formulated with use of \( K-1 \) independent balances \( f_{x_1}, f_{x_2}, \ldots, f_{x_k} \). The balances \( f_{x_i} \) and \( f_{x_j} \), and then \( f_{x_k} \), are not used for the solution of a non-redox system. The linear dependency or independency of \( f_{x_i} \) from \( f_{x_1}, f_{x_2}, \ldots, f_{x_k} \) is then the general property distinguishing between non-redox and redox systems, of any degree of complexity \([40,41]\).

Other, general properties are also valid here, within GATES. Among others, oxidation number (ON) is the derivative concept, resulting from linear combination of the balances. Application of controversial electronegativity (EN) concept, where a highly doubtful (artificial) qualification of ionic bonds (practiced hitherto in literature) is made, is thus avoided. There is no need to pre-assign the roles of oxidants and reducers to individual components and species; oxidant and reductant are derivative concepts within GATES/GEB.

Formulaion of GEB according to Approach II needs none prior knowledge of ONs of elements in all components forming a system and in all species present in the system. For a redox system with \( K-K \) ‘players’, \( f_{x_i} \) is linearly independent on \( f_{x_1}, f_{x_2}, \ldots, f_{x_k} \), i.e., the redox system is described by \( K \) independent balances \( f_{x_1}, f_{x_2}, \ldots, f_{x_k} \). For a non-redox system (\( K=K \)), \( f_{x_i} \) is linearly dependent on \( f_{x_1}, f_{x_2}, \ldots, f_{x_k} \), i.e., a non-redox system is described by \( K-1 \) independent balances \( f_{x_1}, f_{x_2}, \ldots, f_{x_k} \).

Consequently, the linear combination \( \sum_{k=3}^{K} d_k f_k \) applied to a redox system does not give the identity, also after further combination with \( K-K \) balances for ‘players’. The linear combination \( \sum_{k=1}^{K} d_k f_k - f_{0} \) for a redox system is composed only of components and species, where ‘players’ are involved. These regularities are confirmed also in the examples presented below. On this basis, we express/confirm our conviction that these regularities are valid for electrolytic systems of any degree of complexity, with biological systems included \textit{a priori}.

The charged/ionic species \( X^i_{1} \cdot n_{iW} \), i.e., the species with \( z > 0 \) (\( z > 0 \) for cations, \( z < 0 \) for anions), are involved in charge balance, \( f_{x} = ChB \).

\[ f_{0} = \sum_{i=2}^{L} z_i N_i = 0 = \sum_{i=2}^{L} z_i [X^i_{1}] = 0 \]  

The terms: charge balance will be used to both forms of this relation, in accordance with the Ockham razor principle; this should not lead to ambiguity, in the right context. The same viewpoint is referenced to generalized electron balance (GEB). The elemental/core balances, when expressed in terms of molar concentrations, are named as concentration balances.

Free water particles, and water bound in the hydrates \( X^i_{1} \cdot n_{iW} \), are included in balances: \( f_{x} = f(H) \) and \( f_{x} = f(O) \):

\[ f_{1} = f(H) = 2 N_i - \sum_{i=1}^{L} (a_{i1} + 2 n_{iW}) \cdot N_i = \sum_{i=1}^{L} b_{i1} \cdot N_{0j} = 0 \]  

\[ f_{2} = f(O) = 2 N_i - \sum_{i=1}^{L} (a_{i2} + 2 n_{iW}) \cdot N_i - \sum_{i=1}^{L} b_{i2} \cdot N_{0j} = 0 \]  

Then the balance

\[ f_{2} = 2 \cdot f_{2} - f_{1} - \sum_{i=1}^{L} (a_{i2} - a_{i1}) \cdot N_i - \sum_{i=1}^{L} (2 \cdot b_{i2} - b_{i1}) \cdot N_{0j} = 0 \]  

is formulated.

The elemental/core balances: \( f_{y_1}, \ldots, f_{y_k} \) interrelating the numbers of atoms/cores \( Y_k \) \( \in H, O \) in components and species, are as follows

\[ f_{y_k} = f(Y_k) = \sum_{i=1}^{L} a_{ki} \cdot N_i - \sum_{i=1}^{L} b_{ki} \cdot N_{0j} = 0 \quad (k=3,\ldots,K) \]  

where \( a_{ki} \) and \( b_{ki} \) are the numbers of elements/cores \( Y_k \) in \( X^i_{1} \cdot n_{iW} \), and in the \( j \)-th component of the system, resp. For example, \( N \) species/ions HSO\(_4\)\(^{-}\)\( n_{iW}H_2O \) involve \( N_1(1+2n_i) \) atoms of \( H \) (where \( a_{n_i}=1 \)), \( N_2(4+n_i) \) atoms of \( O \) (where \( a_{n_i}=4 \)),
and N atoms of S; N_{o4} molecules of FeSO_{4·7H2O} as a component involve 14N_{o4} atoms of H, 11N_{o4} atoms of O, N_{o4} atoms of S and N_{o4} atoms of Fe.

Formulation of linear combinations is applicable to check the linear dependency or independency of the balances \( f_1, f_2, \ldots, f_K \). For this purpose we will try, in all instances, to obtain the simplest form of the linear combination of these balances. A very useful/effective manner for checking/stating the linear dependency or independency of the balances is the transformation of their linear combination to the identity, \( 0 = 0 \). For a redox system, the proper linear combination of the balances gives the simplest form of the linear combination procedure, \( w \), which is the transformation of their linear combination to the identity, \( 0 = 0 \). For this purpose we will try, in all instances, to obtain the simplest (most desired) form of the related linear combination (Equation 7), as will be explained in examples presented below.

In Equation 4 and then in Equation 7, the terms involved with water, i.e., \( n_{10}, n_{11} \) for related to H O as the component, also as hydrating water, and \( n_{12} = n_{13} \) are not involved. The necessity of prior knowledge of \( n_{14} \) values in the balancing is thus avoided, already at the stage of \( f_1 \) formulation.

Consequently, the set of K independent balances: \( f_1, f_2, \ldots, f_K \) is related to a redox system, whereas \( f_1, f_2, \ldots, f_K \) form the set of K–1 independent balances related to a non-redox system, where \( f_1, \ldots, f_K \) is the set of K–2 elemental/core balances \( f_i = f(Y_i) \) for \( k = 3, \ldots, K \), i.e., for \( Y_i \neq H, O \) (Equation 5). The balancing is necessary for computer simulation of titrimetric procedure according to GATES principles.

The elemental/core balances, expressed in terms of particular units: \( N_{o4} \) for components and \( N \) for species, are the basis to formulation of Generalized Electron Balance (GEB), charge balance (ChB) and concentration balances, expressed in terms of molar concentrations, see Equation 1.

### Formulation of electrolytic D+T system

#### Preliminary information:

As an example let us consider the dynamic D+T system, where \( V \) mL of titrant (D) containing FeSO_{4·7H2O} (C_{o4} mol/L), H_{2}C_{2}O_{4·2H2O} (C_{o5} mol/L), H_{2}SO_{4} (C_{o6} mol/L) and CO_{2} (C_{o7} mol/L) is titrated with \( V \) mL of KMnO_{4} (C_{o8} mol/L) and FeSO_{4·7H2O} (C_{o4} mol/L) + H_{2}SO_{4} (C_{o6} mol/L) + CO_{2} (C_{o7} mol/L) + H_{2}O (C_{o8} mol/L) mL of D. This system involves the following species:

\[
\begin{align*}
\text{H}_2\text{O} & \quad \text{H}^{+1} \quad \text{OH}^{-1} \quad \text{K}^{+1} \quad \text{HSO4}^{-2} \quad \text{Fe}^{+2} \quad \text{FeO}^{+2} \quad \text{Fe}^{+3} \quad \text{FeOH}^{+2} \quad \text{Fe(OH)}^{+2} \quad \text{Mn}^{+2} \quad \text{MnO}^{+2} \quad \text{MnO}_4^{-2} \\
\text{FeSO}_4 \quad \text{H}_{2}\text{C}_{2}\text{O}_{4} \quad \text{H}_{2}\text{SO}_4 \quad \text{CO}_2 \quad \text{H}_{2}\text{O} \\
\end{align*}
\]

It is assumed that the D+T system is composed of KMnO_{4} (N_{o4} molecules), FeSO_{4·7H2O} (N_{o4} molecules) + CO_{2} (N_{o7} molecules) + FeSO_{4·7H2O} (N_{o4} molecules) + H_{2}O (N_{o8} molecules) in \( V \) mL of T and FeSO_{4·7H2O} (N_{o4} molecules) + H_{2}C_{2}O_{4·2H2O} (N_{o5} molecules) + H_{2}SO_{4} (N_{o6} molecules) + CO_{2} (N_{o7} molecules) + H_{2}O (N_{o8} molecules) in \( V \) mL of D. This system involves the following species:

\[
\begin{align*}
\text{Fe}^{2+} \quad \text{Fe}^{3+} \quad \text{FeOH}^{+2} \quad \text{Fe(OH)}^{+2} \\
\text{Mn}^{2+} \quad \text{MnO}^{2+} \quad \text{MnO}_4^{-2} \quad \text{Mn}_{2}\text{O}_4^{-2} \\
\text{H}_{2}\text{C}_{2}\text{O}_{4} \quad \text{H}_{2}\text{SO}_4 \quad \text{CO}_2 \quad \text{H}_{2}\text{O} \\
\end{align*}
\]

The molecules of the (pre-assumed) precipitates of the oxalates are written in bold: FeC_{o6}O_{o4} and MnC_{o6}O_{o4}; \( a \neq 1 \), if the precipitate indicated (\( i = 1, 2 \)) is in the equilibrium solid phase in the system; otherwise, \( a = 0 \).

#### Formulation of balances for D+T system according to approach I

The charge and elemental/core balances, formulated on the basis of the data specified above for electron-non–active elements (‘fans’) are as follows:

\[
f_1 = \text{ChB}
\]

\[
N_1 - N_4 + N_5 - N_6 - 2N_9 - N_{11} - 2N_{13} - N_{15} - 2N_{16} + 3N_{17} + 2N_{18} \neq 0\]

\[
N_1 - N_4 + 3N_{19} + 2N_{20} - N_{21} - 2N_{22} - 2N_{23} - N_{24} + 2N_{25} + 2N_{26} + N_{27} - 2N_{28} - 4N_{29} + 3N_{30} \neq 0\]

\[
2N_{31} + N_{32} + 4N_{33} + N_{34} - N_{35} + N_{36} - 3N_{37} = 0\]

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On the basis of charge and elemental/core balances, we formulate the linear combinations:

$$\begin{align*}
\text{Linear combinations of the balances} \\
&= \sum \text{charges on elements} \\
&= \sum \text{elementalrepid} \\
&= \sum \text{corerepid} \\
\end{align*}$$

Applying atomic numbers: $Z_c = 6$ for C, $Z_{Fe} = 26$ for Fe and $Z_{Mn} = 25$ for Mn, from Equations (13) – (15) and (17a) we have, by turns:

$$\begin{align*}
Z_c f_C &= Z_{Fe} f_C(C) \\
Z_c \left(2(N_7 + N_8 + N_9) + (N_{10} + N_{11} + N_{12}) + 2N_{17} + 4N_{18} + 6N_{19} + 7N_{20} + 4N_{21} + 4N_{24} + 6N_{25} + 4N_{26} + 6N_{29} + 6N_{30} + 7N_{01} + 4(N_{02} + N_{07}) + 2N_{03} + 2N_{04} + 6N_{05} + 4N_{06} + 2N_{07} + 6N_{08}
\end{align*}$$

$$\begin{align*}
\text{Z_{Fe} f_Fe} &= Z_{Fe} f_C(Fe) \\
Z_{Fe} \left(N_{15} + N_{16} + N_{17} + N_{18} + N_{19} + N_{20} + N_{21} + N_{22} + N_{23} + N_{24} + N_{25} + N_{26} + N_{27} + N_{28} + N_{29} + N_{30} + N_{31} + N_{32} + N_{33} + 2N_{34} + 2N_{35} + 3N_{36} + 3N_{37}
\end{align*}$$

$$\begin{align*}
\text{Z_{Mn} f_Mn} &= Z_{Mn} f_C(Mn) \\
Z_{Mn} \left(N_{13} + N_{14} + N_{15} + N_{16} + N_{17} + N_{18} + N_{19} + N_{20} + N_{21} + N_{22} + N_{23} + N_{24} + N_{25} + N_{26} + N_{27} + N_{28} + N_{29} + N_{30} + N_{31} + N_{32} + N_{33} + 2N_{34} + 3N_{35} + 3N_{36} + 3N_{37} + 3N_{38} + 3N_{39} + a_1N_{40} + a_2N_{41}
\end{align*}$$

$$\begin{align*}
\text{Z_{C} f_C} &= Z_{Fe} f_C(Fe) \\
Z_{C} \left(2(N_7 + N_8 + N_9) + (N_{10} + N_{11} + N_{12}) + 2N_{17} + 4N_{18} + 6N_{19} + 7N_{20} + 4N_{21} + 4N_{24} + 6N_{25} + 4N_{26} + 6N_{29} + 6N_{30} + 7N_{01} + 4(N_{02} + N_{07}) + 2N_{03} + 2N_{04} + 6N_{05} + 4N_{06} + 2N_{07} + 6N_{08}
\end{align*}$$
Formed in the system by C, Fe and Mn. In effect:

\( \text{N}_1 \) molecules of \( \text{H}_2\text{C}_2\text{O}_4\cdot\text{n}_1\text{H}_2\text{O} \) carries 2(Z_c – 3)N_c carbon electrons;

\( \text{N}_4 \) ions of \( \text{HC}_2\text{O}_4^{-}\cdot\text{n}_4\text{H}_2\text{O} \) carries 2(Z_c – 3)N_b carbon electrons;

\( \text{N}_6 \) ions of \( \text{C}_2\text{O}_4^{2-}\cdot\text{n}_6\text{H}_2\text{O} \) carries 2(Z_c – 3)N_b carbon electrons;

\( \text{N}_11 \) molecules of \( \text{H}_2\text{CO}_3\cdot\text{n}_11\text{H}_2\text{O} \) carries (Z_c – 4)N_c carbon electrons;

\( \text{N}_11 \) ions of \( \text{HCO}_3^{-}\cdot\text{n}_11\text{H}_2\text{O} \) carries (Z_c – 4)N_b carbon electrons;

\( \text{N}_11 \) ions of \( \text{CO}_3^{2-}\cdot\text{n}_11\text{H}_2\text{O} \) carries (Z_c – 4)N_b carbon electrons;

\( \text{N}_3 \) ions of \( \text{MnO}_4^{-}\cdot\text{n}_3\text{H}_2\text{O} \) carries (Z_mn – 7)N_mn manganese electrons;

\( \text{N}_3 \) ions of \( \text{MnO}_4^{-}\cdot\text{n}_3\text{H}_2\text{O} \) carries (Z_mn – 6)N_mn manganese electrons;

\( \text{N}_4 \) ions of \( \text{MnOH}^+\cdot\text{n}_4\text{H}_2\text{O} \) carries (Z_mn – 3)N_mn manganese electrons;

\( \text{N}_12 \) ions of \( \text{MnC}_2\text{O}_4^{-}\cdot\text{n}_12\text{H}_2\text{O} \) carries (Z_mn – 3)N_mn manganese electrons and 2(Z_c – 3)N_c carbon electrons; together (Z_mn+2Z_c – 9)N_c electrons;

\( \text{N}_19 \) ions of \( \text{MnC}_2\text{O}_4^{-}\cdot\text{n}_19\text{H}_2\text{O} \) carries (Z_mn – 3)N_mn manganese electrons and 4(Z_c – 3)N_b carbon electrons; together (Z_mn+4Z_c – 15)N_b electrons;

\( \text{N}_13 \) ions of \( \text{MnC}_2\text{O}_4^{-}\cdot\text{n}_13\text{H}_2\text{O} \) carries (Z_mn – 2)N_mn manganese electrons and 6(Z_c – 3)N_b carbon electrons; together (Z_mn+6Z_c – 21)N_b electrons;

\( \text{N}_13 \) ions of \( \text{Mn}^2+\cdot\text{n}_13\text{H}_2\text{O} \) carries (Z_mn – 2)N_mn manganese electrons;

\( \text{N}_13 \) ions of \( \text{MnOH}^+\cdot\text{n}_13\text{H}_2\text{O} \) carries (Z_mn – 2)N_mn manganese electrons;

\( \text{N}_13 \) molecules of \( \text{MnSO}_4\cdot\text{n}_13\text{H}_2\text{O} \) carries (Z_mn – 2)N_mn manganese electrons;

\( \text{N}_13 \) molecules of \( \text{MnC}_2\text{O}_4\cdot\text{n}_13\text{H}_2\text{O} \) carries (Z_mn – 2)N_mn manganese electrons and 2(Z_c – 3)N_c carbon electrons; together (Z_mn+2Z_c – 8)N_c electrons;

\( \text{N}_13 \) ions of \( \text{MnC}_2\text{O}_4\cdot\text{n}_13\text{H}_2\text{O} \) carries (Z_mn – 2)N_mn manganese electrons and 4(Z_c – 3)N_b carbon electrons; together (Z_mn+4Z_c – 14)N_b electrons;

\( \text{N}_13 \) ions of \( \text{MnC}_2\text{O}_4\cdot\text{n}_13\text{H}_2\text{O} \) carries (Z_mn – 2)N_mn manganese electrons and 6(Z_c – 3)N_b carbon electrons; together (Z_mn+6Z_c – 21)N_b electrons.

\( \text{N}_13 \) ions of \( \text{Fe}^{2+}\cdot\text{n}_13\text{H}_2\text{O} \) carries (Z_mn – 2)N_mn iron electrons;

\( \text{N}_13 \) ions of \( \text{Fe}^{2+}\cdot\text{n}_13\text{H}_2\text{O} \) carries (Z_mn – 2)N_mn iron electrons;

\( \text{N}_13 \) ions of \( \text{Fe}^{3+}\cdot\text{n}_13\text{H}_2\text{O} \) carries (Z_mn – 2)N_mn iron electrons;

\( \text{N}_13 \) ions of \( \text{Fe}^{3+}\cdot\text{n}_13\text{H}_2\text{O} \) carries (Z_mn – 2)N_mn iron electrons;

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Equivalency of Approaches I and II to GEB

The Approach I to GEB needs prior knowledge of ONs for the ‘players’, i.e., C, Fe and Mn, in different components and species formed by them, in the D+T system considered. It is assumed there, that both ‘players’ as components introduce, to the common pool, a certain number of their own electrons, assumed there, that both ‘players’ as components introduce, species formed by them, in the D+T system considered. It is

Equation 19 will be compared with the one obtained according to Approach I to GEB, introduced initially (as such) in the papers [42–47] and applied for modelling of titrations curves, realized according to modified Gran I and II methods, in numerous versions [48–52], proposed by Michałowski.

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\[
\text{In V mL of T, } N_m \text{ molecules of KMnO}_4 \text{ introduce } (Z_mn – 7)N_m \text{ manganese electrons, } N_{c2} \text{ molecules of CO}_2 \text{ introduce } (Z_c – 4)N_{c2} \text{ carbon electrons;}
\]

In V mL of D, N_{c2} molecules of FeSO_{4} \cdot 7H_{2}O introduce (Z_{c2} – 2)N_{c2} iron electrons, N_{c3} molecules of H_{2}C_{2}O_{4} \cdot 2H_{2}O introduce 2(Z_{c2} – 3)N_{c3} carbon electrons, and N_{c4} molecules of CO_{3} introduce (Z_{c2} – 4)N_{c4} carbon electrons.

Then the total number of electrons introduced by the ‘players’ is

\[
(Z_{m} – 7)N_m + 2(Z_c – 3)N_{c2} + (Z_c – 4)(N_{c3} + N_{c4}) + (Z_{c2} – 2)N_{c4}
\]

These electrons are ‘dissipated’ between different species.
N34 molecules of FeSO4 \( \times n_{34}H_2O \) carries \((Z_{Fe} - 3)N_{34}\) iron electrons; 
N35 ions of Fe(OH)+1 \( \times n_{35}H_2O \) carries \((Z_{Fe} - 3)N_{35}\) iron electrons; 
N36 ions of Fe(OH)2+1 \( \times n_{36}H_2O \) carries \((Z_{Fe} - 3)N_{36}\) iron electrons; 
N37 ions of Fe+3 \( \times n_{37}H_2O \) carries \((Z_{Fe} - 3)N_{37}\) iron electrons; 
N38 ions of Fe2(OH)2+1 \( \times n_{38}H_2O \) carries \((Z_{Fe} - 3)N_{38}\) iron electrons; 
N39 ions of Fe(OH)3+1 \( \times n_{39}H_2O \) carries \((Z_{Fe} - 3)N_{39}\) iron electrons; 
N40 molecules of FeC2O4 \( \times n_{40}H_2O \) carries \(a_1(Z_{Fe} - 2)N_{40}\) iron electrons and \(2a_1(Z_{C} - 3)N_{40}\) carbon electrons; together \(a_1(Z_{Fe}+2Z_{C} - 8)N_{40}\) electrons; 
N41 molecules of MnC2O4 \( \times n_{41}H_2O \) carries \(a_2(Z_{Mn} - 2)N_{41}\) manganese electrons and \(2a_2(Z_{C} - 3)N_{41}\) carbon electrons; together \(a_2(Z_{Fe}+2Z_{C} - 8)N_{41}\) electrons;

Balancing the electrons brought by ‘players’ in components and those dissipated between ‘players’ in the species, we obtain Eq. 19. This way, the equivalence of the Approaches I and II to GEB is proved here.

**Linearity properties involved with T and D subsystems**

The **T subsystem** (V): Applying the notations from (9), we have the following balances:

\[ f_0 = \text{ChB} \]
\[ N_1 - N_2 - N_3 - N_4 - 2N_5 - 2N_6 - 2N_7 - N_{10} = 0 \]
\[ f_1 = f(H) \]
\[ 2N_1 + N_2(1+2n_2) + N_3(1+2n_3) + N_5(1+2n_5) + 2N_6n_6 + N_7(2+2n_7) + N_8(1+2n_8) + 2N_9n_9 + N_{10}(2+2n_{10}) + N_{11}(1+2n_{11}) + 2N_{12}n_{12} + 2N_{26}n_{26} + N_{27}(1+2n_{27}) + 2N_{28}n_{28} + 2N_{29}n_{29} + 2N_{30}n_{30} = 14N_04 + 6N_05 + 2N_06 + 2N_08 \]
\[ f_2 = f(O) \]
\[ N_1 + N_2n_2 + N_3(1+n_3) + N_4(4+n_4) + N_6(4+n_6) + N_7(4+n_7) + N_8(4+n_8) + N_{10}(4+n_{10}) = N_{12}(4+n_{12}), \]
\[ f_{12} = 2f_2 - f_1 \]
\[ -N_1 + N_2 + 4N_{10} + 5N_8 + 6N_9 + 8N_{11} = 8N_0 + 4N_{02} \]
\[ f_{1}_2 = -f_1 \]
\[ -f_3 = -f(K) \]
\[ N_0 = N_4 \]
\[ 4N_{02} = 4N_{0} + 4N_{11} + 4N_{12} \]
\[ 7f_7 = 7f(Mn) \]
\[ 7N_{40} = 7N_{11} \]
\[ f_{13} - f_0 - f_4 - 7f_7 = 0 \]
\[ f_{4} = 0 \]

As we see, the linear combination (19) for T, considered as a non-redox system, is transformed into identity (20a) [32].

**Formulation of the D subsystem (V0)**

Applying again the notations from (9), we have the following balances:

\[ f_0 = \text{ChB} \]
\[ N_1 - N_2 - N_3 - 2N_5 - N_6 - 2N_7 - 2N_9 - N_{10} = 0 \]
\[ f_1 = f(H) \]
\[ 2N_1 + N_2(1+2n_2) + N_3(1+2n_3) + N_5(1+2n_5) + 2N_6n_6 + N_7(2+2n_7) + N_8(1+2n_8) + 2N_9n_9 + N_{10}(2+2n_{10}) + N_{11}(1+2n_{11}) + 2N_{12}n_{12} + 2N_{26}n_{26} + N_{27}(1+2n_{27}) + 2N_{28}n_{28} + 2N_{29}n_{29} + 2N_{30}n_{30} + 2a_1N_{40}n_{40} = 14N_04 + 6N_05 + 2N_06 + 2N_08 \]
\[ f_2 = f(O) \]
\[ N_1 + N_2n_2 + N_3(1+n_3) + N_4(4+n_4) + N_6(4+n_6) + N_7(4+n_7) + N_8(4+n_8) + N_{10}(4+n_{10}) + N_{12}(4+n_{12}) = N_{13}(4+n_{13}), \]
\[ f_{13} = 2f_2 - f_1 \]
\[ -N_1 + N_2 + 7N_8 + 8N_9 + 7N_9 + 8N_{10} + 5N_{11} + 6N_{12} + 5N_9 + 8N_{10} + 16N_9 = 32N_03 + 6a_1N_{40}n_{40} = 8N_04 + 6N_06 + 4N_{07} + 7N_{08} \]
\[ -f_3 = -f(Mn) \]
\[ f_{12} = 2f_2 - f_1 \]
As we see, the linear combination (20) for D, considered as a non-redox system, is transformed into identity (21a) [32].

The balances for the D+T system expressed in terms of concentrations

Applying the notations specified above, we have the relations:

\[
\begin{bmatrix}
X^0 + V \\
\end{bmatrix} = 10^3 \begin{bmatrix}
N_1 \\
\end{bmatrix}
\]

\[
C_{01} V_0 = 10^{\frac{2 N_01 + N_02}{N_A}}
\]

\[
C_{02} V_0 = 10^{\frac{2 N_04}{N_A}}
\]

\[
C_{03} V_0 = 10^{\frac{2 N_05}{N_A}}
\]

\[
C_{04} V_0 = 10^{\frac{2 N_06}{N_A}}
\]

\[
= 10^{\frac{2 N_07}{N_A}}
\]

The balances (10) – (15), (17a) written in terms of molar concentrations are as follows:

\[\begin{align*}
[H^+1] & = \text{OH}^{-1} + \text{K}^{+1} - [\text{HSO}_4^{2-}] - 2[\text{SO}_4^{2-}] - [\text{HC}_2\text{O}_4^{-}] - 2[\text{CO}_3^{2-}] - [\text{HCO}_3^{-}] \\
-2[\text{CO}_3^{2-}] & - [\text{Mn}^{2+}] - 2[\text{MnO}_4^{2-}] + 3[\text{Mn}^{3+}] + 2[\text{MnOH}^{2+}] + [\text{MnC}_2\text{O}_4^{2-}] - [\text{Mn(C}_2\text{O}_4)_3^{-}] \\
-3[\text{Mn(C}_2\text{O}_4)_3^{-}] & + 2[\text{Mn}^{2+}] + [\text{MnOH}^{2+}] - 2[\text{Mn(C}_2\text{O}_4)_2^{-}] - 2[\text{Mn(OH)}^{2+}] + 2[\text{Fe}^{2+}] + [\text{FeOH}^{2+}] \\
-2[\text{Fe}^{2+}] & - 4[\text{Fe(C}_2\text{O}_4)_3^{-}] + 3[\text{Fe}^{3+}] + 2[\text{FeOH}^{2+}] + [\text{Fe(OH)}^{2+}] + 4[\text{Fe(OH)}^{3+}] \\
+ [\text{FeSO}_4^{2-}] & - [\text{Fe(SO}_4)_2^{2-}] - [\text{Fe(C}_2\text{O}_4)_3^{-}] - [\text{Fe(C}_2\text{O}_4)_2^{-}] - 3[\text{Fe(C}_2\text{O}_4)_3^{-}] = 0
\end{align*}\]

\[\text{K}^{+1}] = CV/(V \text{+} V)
\]

\[\begin{align*}
[\text{HSO}_4^{2-}] & + [\text{SO}_4^{2-}] + [\text{MnSO}_4] + [\text{FeSO}_4] + 2[\text{Fe(SO}_4)_2^{2-}] - (C_{01} \text{c}_{02} V_0)/(V \text{+} V) = 0
\end{align*}\]

\[\begin{align*}
2[\text{H}_2\text{CO}_3] & + [\text{HC}_2\text{O}_4^{-}] + [\text{HCO}_3^{-}] + \text{C}_2\text{O}_4^{-} + \text{CO}_3^{2-} + \text{MnC}_2\text{O}_4^{2-} + 4[\text{MnC}_2\text{O}_4] + 4[\text{MnC}_2\text{O}_4] + 6[\text{MnC}_2\text{O}_4] + 2[\text{Mn(OH)}^{2+}] + 4[\text{MnC}_2\text{O}_4] + 6[\text{MnC}_2\text{O}_4] + 4[\text{FeC}_2\text{O}_4^{-}] + 6[\text{Fe(C}_2\text{O}_4)_2^{-}] + [\text{FeC}_2\text{O}_4] + 2a_2[\text{FeC}_2\text{O}_4] + 2a_2[\text{MnC}_2\text{O}_4] - (C_{03} V_0 + C_{04} V_0)/(V \text{+} V) = 0
\end{align*}\]
(Z_{\text{min}} + 2Z_{\text{C}} - 20)[\text{Mn}(\text{C}_2\text{O}_4)^{2-}] + (Z_{\text{min}} + 1\dfrac{3}{2}Z_{\text{C}} - 9)[\text{MnC}_2\text{O}_4]^{2-} + [\text{Mn}(\text{C}_2\text{O}_4)^{3-}] + (Z_{\text{min}} + 2\dfrac{3}{2}Z_{\text{C}} - 15)[\text{Mn}(\text{C}_2\text{O}_4)^{3-}] + (Z_{\text{min}} + 3\dfrac{3}{2}Z_{\text{C}} - 21)[\text{Mn}(\text{C}_2\text{O}_4)^{3-}] - (2Z_{\text{C}} - 3)C V_{\text{C}} + (Z_{\text{min}} - 4)[\text{C}_2\text{O}_4]^{2-} + (Z_{\text{min}} - 7)CV)/(V + V) = 0 \quad (19a)

Some regularities involved with linear combinations of the balances

Equations (16b), (17b) and (19a) are equivalent forms of GEB. Equation (16b) is obtained from the balance (16b), considered as the primary form of GEB, pr-GEB [1]. Equation (19a), obtained also on the basis of prior knowledge of ONs for C, Fe and Mn, is also called as the ‘short’ version of GEB [1] for this system. Equation (17b) contains components and species composed only of ‘players’, i.e., where ‘fans’ are not involved. These ‘fans’ are cancelled after further combination with balances related to other ‘fans’. The relation (11a), where only one species is involved, is considered as equality, not equation; it can enter immediately the charge balance (10a). The volume V added from the start up to a defined point of the titration, is the current parameter (not variable), from the viewpoint of the calculation procedure.

Equation (17b) and, consequently, Equation (19a) includes only the components and species where ‘players’ are involved. The equivalent forms: (16b), (17b), (19a) of GEB have different notation lengths. One can, however, reduce the length of this notation, making further, linear combination of $f_x + f_y - f_z - 6f_x$ (Equation 17a) with $f_x$, $f_y$ and $f_z$ (Equations 13–16, resp.). After addition of (17a) to the balances (13b) – (15b):

$-4f_x = -4f(C)$

$8N_{29} + 4(N_{25} + N_{30}) = 8(N_{13} + N_{14} + N_{15} + N_{16}) + 8N_{17} + 16N_{28} + 8N_{29} + 8N_{30}$

$+ 16N_{24} + 24N_{25} + 16N_{29} + 24N_{30} + 8N_{31} + 8a_1N_{29} + 8a_2N_{30}$

$-3f_y = -3f(Fe)$

$3N_{23} = 3(N_{25} + N_{26} + N_{27}) + 3N_{30} + 3N_{35} + 3(N_{39} + N_{40} + N_{41}) + 2N_{15} + N_{16} + 3N_{28} + a_1N_{23}$

$-2f_z = -2f(Mn)$

$2N_{45} = 2N_{13} + 2N_{14} + 2N_{15} + 2N_{16} + 2N_{17} + 2N_{18} + 2N_{19} + 2(N_{20} + N_{31} + N_{32}) + 2N_{23} + 2N_{24}$

$2N_{15} + a_1N_{41}$

and further rearrangements the terms, we get the simplest (in this respect) combination

$f_x + f_y - f_z - 6f_x - 4f_y - 2f_z - 2f_2$ (23)

By the way, a new criterion applying only the components and species where ‘players’ are involved, presents the following calculation algorithm:

On the calculation procedure

Concentrations of the species in Equations (10a) – (15a), (19a) are also interrelated in the set of independent expressions for equilibrium constants, found e.g. in [53–55]. Altogether, 29 independent equilibrium constants are involved in the computer program, presented in Appendix for the D+T system with 41 species. Among others, the relationships for solubility products of the precipitates: $\text{FeC}_4\text{O}_4$ and $\text{MnC}_4\text{O}_4$, pre-assumed in the model are as follows:

$$K_{\text{sol}} = [\text{Fe}^{2+}][\text{C}_2\text{O}_4^{2-}] \quad (pK_{\text{sol}} = 6.7), \quad K_{\text{so2}} = [\text{Mn}^{2+}][\text{C}_2\text{O}_4^{2-}] \quad (pK_{\text{so2}} = 5.3).$$

The complete set of independent equilibrium constants provides all quantitative knowledge on the system in question. Some qualitative knowledge on the system in question is also valuable; this kind of knowledge is particularly desired in the case of metastable systems.

Six independent variables:

$x(1) = \text{pH}, \quad x(2) = E,$

$x(3) = -\text{log}[\text{Mn}^{2+}]$ or $x(3) = -\text{log}[\text{MnC}_2\text{O}_4]$,

$x(4) = -\text{log}[\text{Fe}^{2+}]$ or $x(4) = -\text{log}[\text{FeC}_2\text{O}_4]$,

$x(5) = -\text{log}[\text{H}_2\text{C}_2\text{O}_4]$.

are involved in the set of six independent equations: (9a), (11a), (12a), (13a), (14a) and (17b) in the computer program. The volume V of titrant T is the current (operating) parameter in the related algorithm.

Concentrations of particular species are presented there without parentheses. Some exemplary concentrations are denoted as follows:

$$[\text{Fe}^{2+}] = \text{Fe}^3, \quad [\text{MnOH}^{+} + 2] = \text{MnOH}, \quad [\text{MnO}_4^{2-}] = \text{MnO}_4, \quad [\text{MnO}_4^{2-}] = \text{MnO}_4, \quad [\text{FeC}_2\text{O}_4]^{2-} = \text{FeC}_2\text{O}_4,$$

$$[\text{pr}_1] = [\text{FeC}_4\text{O}_4], \quad [\text{pr}_2] = [\text{MnC}_4\text{O}_4], \quad \text{logr}_1 = \text{log}[\text{FeC}_4\text{O}_4].$$

At the pre-assumed concentrations of components in D and T,
Conclusions

The example presented above provides some general regularities, obligatory for all electrolytic systems, in aqueous media, and extended on non-aqueous and mixed-solvent media, with amphiprotic (co)solvent(s) included. These regularities can be listed from different viewpoints.

The criterion distinguishing between redox and non-redox systems

The simplest/shortest form of GEB (Equations 23a, 23b) related to D+T mixture as the redox system is different from identity, \( 0 = 0 \). This way, the linear independency of \( f_{12} \) from other balances: \( f_0, f_3, f_5, f_6, f_7 \) is proved here. The identity property, \( 0 = 0 \), was stated for T (19, 19a) and D (20, 20a), considered as non-redox (sub)systems, i.e., \( f_{12} \) is linearly dependent on:

- \( f_0, f_3, \ldots, f_k \) for all non-redox systems, of any degree of complexity; consequently, non-redox systems are fully described with use of \( K - 1 \) balances \( f_0, f_3, \ldots, f_k \), i.e., the balances for H and O: \( f = f(H) \) and \( f = f(O) \) and then \( f_{13} = 2f_2 - f_1 \) are not applied for description of non-redox systems;

- \( f_{12} \) is linearly independent on \( f_0, f_3, \ldots, f_k \) for all redox systems, of any degree of complexity; consequently, redox systems are fully described with use of \( K \) balances \( f_0, f_{12}, f_3, \ldots, f_k \); the balance \( f_{13} = 2f_2 - f_1 \) is the basis to formulation of GEB, completing the set of independent equations needed for description of redox systems;

the GEB completing the set of equations needed for description of a redox system can be obtained according to two equivalent approaches, named as Approach I to GEB and Approach II to GEB.

Figure 1: The (a) \( E = E(V) \), (b) \( pH = pH(V) \) relationships and dynamic speciation diagrams for (c) Mn- and (d) Fe- species, plotted at \( V_0 = 100, C_{01} = 0.02, C_{02} = 0.01, C = 0.02, C_{03} = 0.5, C_1 = C_{ss} = 0.001 \).
the linear independency or dependency of \( f_{u} \) from \( f_{v}, f_{w}, ... \), \( f_{u} \) is the general criterion distinguishing between redox and non-redox systems.

**Oxidation numbers (ONS)**

The Approach I to GEB needs prior knowledge of ONs for ‘players’ involved in components and species of the system; ONs of elements in ‘fans’ are not taken into account there;

Within the Approach I to GEB, all components and species with ‘players’ are distinguished and collected; ‘fans’ are not considered on the step of GEB formulation according to Approach I;

The terms oxidant(s) and reductant(s) are not applied (are derivative terms) on the step of formulation of a redox system according to Approaches I and II. This fact is of capital importance, when redox equilibria involved e.g., with complex organic species, are considered.

The Approach II to GEB needs none prior knowledge of ONs for elements in components and species; the terms: ‘players’ and ‘fans’, oxidant and reductant are not indicated; full ‘democracy’ is provided within the Approach II to GEB. Known (or pre-assumed) composition of a species, expressed by its formula, together with external charge of this species, provides an information sufficient to formulate the related balances.

**The linear combination of balances as the source of ONs**

Linear combination of charge and elemental/core balances suggested/practiced above was aimed at finding the simplest form of a linear combination, in the sense of the minimizing the number of kinds of different components.

Let us refer to linear combinations of linear combinations of balances related to the non-redox systems (D, T) and the redox system (D+T). Applying the notation suggested in (4), (5), (6a) and (6b), we rewrite (20) and (21) as follows.

For non-redox subsystem T

\[
\begin{align*}
\text{f}_{at} + f_{o} - f_{s} - 4f_{k} - 7f_{s} &= 0 & \Rightarrow & & 1(+1)f_{at} + 1(-2)f_{o} + 1(+4)f_{s} + 1(+7)f_{k} = f_{o} \\
1(+1)(H) + 1(-2)(O) + 1(+1)(K) + 1(+4)(CO_{3}) + 1(+7)(MnO_{4}) &= \text{ChB} \\
\end{align*}
\]

For redox subsystem D

\[
\begin{align*}
1(+1)f_{at} + 1(-2)f_{o} + 1(+6)f_{s} + 2(+3)f_{k} + 1(+4)f_{s} + 1(+2)f_{s} &= f_{o} \\
1(+1)(H) + 1(-2)(O) + 1(+6)(SO_{4}) + 2(+3)(C_{2}O_{4}) + 1(+4)(CO_{3}) + 1(+2)(Fe) &= \text{ChB} \\
\end{align*}
\]

In both subsystems, all ‘fans’ are involved. The coefficients at particular balances in the related linear combinations are equal to the product of: a number of elements in the related entities (component, species) and the ON of the element in these entities.

For the redox D+T system

Linear combination of balances for ‘fans’ of this system (Equation 17) can be rearranged as follows:

\[
\begin{align*}
1(+1)f_{at} + 1(-2)f_{o} + 1(+4)f_{s} + 1(+7)f_{k} = f_{o} \\
1(+1)(H) + 1(-2)(O) + 1(+6)(SO_{4}) + 2(+3)(C_{2}O_{4}) + 1(+4)(CO_{3}) + 1(+2)(Fe) &= \text{ChB} \\
\end{align*}
\]

**A remark.** Some subsystems of other D + T systems may have redox properties. This is the case, for example, in D of D + T system, where (a) NaOH solution as T is added to: Br₂ or HBrO, solution [23,24], as D, or (b) HCl solution as T is added into NaIO solution [23,24]. Redox properties result there from disproportionating of the solute (Br₂, HBrO, NaIO) dissolved in water.

**The elemental versus core balances**

Unlike the D + T system, where oxalate species are transformed into carbonate species, in D they exist side by side, without mutual transformation. Therefore, oxalate and carbonate species in D can be placed in separate balances, only for an illustrative consideration. Writing the common balance \( f_{at} = 2f_{at} + f_{at} \) is a matter of choice for D, but the necessity for D+T.

The element S enters the related system only as a core \( SO_{4} \), and then the balances: \( f(S) \) and \( f(SO_{4}) \) for the related systems are identical.

**Further reference to the ‘card game’**

Let us recall again the ‘card game’ with ‘players’, ‘fans’ and ‘money’, presented in ref [1]. (pp. 41–43).
As usually happens in the card game practice, the players devote to the game only a part of their cash resources. Similarly, in redox reactions may participate electrons from the valence shells of atoms of electron–active elements; the electrons from the valence shell of the reductant atoms are transferred onto the valence shell of the oxidant atoms. However, this restriction to the valence electrons is not required here. For example, one can replace \( Z_{i} \) by \( \zeta_{i} (Z_{i} < \zeta_{i}) \), \( Z_{p} \) by \( \zeta_{p} (Z_{p} < \zeta_{p}) \) and \( Z_{mn} \) by \( \zeta_{mn} (Z_{mn} < \zeta_{mn}) \) in Equation 19. In particular, we can put in there \( \zeta_{i} = \zeta_{p} = 0 \) for \( X = C, Fe, Mn \). Obviously, we get here the relation identical with Equation 17a.

This way, we recall the card game without ‘live cash’ but with ‘debt of honor’ – in not accidental reference to the title of the thriller novel by T. Clancy; btw, the “Debt of honor” was published in 1994, like the papers [43–45].

**Final comments**

A keystone for the overall, thermodynamic knowledge on electrolytic systems is the linear combination 2·(H)–(O) of elemental balances: \( f(H) \) for H and \( f(O) \) for O, that can be formulated both for non–redox and redox systems, in aqueous, non–aqueous and mixed–solvent systems, with amphiprotic (co)solvent(s) involved. In any redox system, the 2·(H) – (f(H)) is linearly independent on ChB and other, elemental/core balances, for For any non–redox system, 2·(O) – (f(H)) is linearly dependent on those balances. The equation for 2·(O) – (f(H)) considered as the primary form of GEB, \( \mathbf{P} = \Delta \mathbf{E}_{\text{B}} = 2 \cdot f(O) – f(H) \), is the basis of GEB formulation for redox systems according to Approach II to GEB. Then the linear independency/dependency of 2·(O) – (f(H)) on the other balances is the general criterion distinguishing between redox and non–redox systems.

The number of electron–active elements (‘players’) in a redox system, considered according to GATES/GEB principles, is practically unlimited; among others, the systems with one, two, three or four ‘players’ were considered.

All earlier (dated from 1960s) trials made towards formulation of electrolytic redox systems were only clumsy attempts of resolution of the problem in question, as stated in [10–12]. Those approaches were slavishly related to the stoichiometric reaction notations, involving only two pairs of indicated species participating in redox reaction; there were usually minor species of the system considered. The species different from those involved in the reaction notation were thus omitted in considerations. What is more, the charge balance and concentration balances for accompanying substances were also omitted. Theoretical considerations were related to virtual cases, not to real, electrolytic redox systems. The computer simulation realized within GATES with use of iterative computer programs, e.g. MATLAB, provides quite a new quality in knowledge gaining. It enables to follow the details of the process, registered with use of measurable quantities, such as pH and/or potential E.

All the inferences made within GATES/GEB are based on firm, mathematical (algebraic) foundations, not on an extremely “fragile” chemical notation principle that is only a faint imitation of a true, algebraic notation [21,30]. The approach proposed allows to understand far better all physicochemical phenomena occurring in the system in question and improve some methods of analysis. All the facts testify very well about the potency of simulated calculations made, according to GATES, on the basis of all attainable physicochemical knowledge. Testing the complex redox and non–redox systems with use of iterative computer programs deserves wider popularization among physicochemists and chemists–analysts.

**Epilogue**

In the fourteenth century, William of Ockham has formulated the parsimony (economy of thinking) [50], principle that “entities should not be multiplied unnecessarily” (“Pluralitas non est ponenda sine necessitate”), known briefly as “Ockham’s razor”. In reference to science, it means that simpler theories are generally better than more complex ones. GATES, based on fundamental rules of the matter conservation, acts according to the Ockham razor principle.

The Approach II to GEB shows immediately that the equivalent equations for GEB are derived from the common root of the elements conservation and then GEB is fully compatible with charge and concentration balances, like “the lotus flower, lotus leaf and lotus seed come from the same root” [7]. This compatibility is directly visible from the viewpoint of the Approach II to GEB. The GEB, based on a reliable law of the matter conservation, is equally robust as equations for charge and concentration balances. The complementarity of the GEB (Approaches I and II) to other balances is regarded as the expression of Harmony of Nature, and GATES/GEB as an example of excellent epistemological paradigm.

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