# A Mathematical Computer Model for Estimating Phase Weight/ Atomic Fractions for Iron Slags' System ( $\mathrm{SiO}_{\mathbf{2}}-\mathrm{Al}_{2} \mathbf{0}_{3}-\mathrm{CaO}-\mathrm{MgO}$ ) at Subsolidus Temperatures 

Tanju Yelkenci ${ }^{1} \cdot$ Z. Engin Erkmen ${ }^{2}$

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#### Abstract

In this study, the 3 dimensional description of the quaternary subsolidus slag system aluminasilicacalciamagnesia is evaluated using Mathematica programming based on previous experimental research data. The lever rule in the tetrahedra is then applied for a given arbitrary liquid composition $\% \mathrm{~A}+\% \mathrm{~B}+\% \mathrm{C}+\% \mathrm{D}=100$ in order to calculate the solid phases present (neglecting solid solutions) with their relative proportions after the solidification is completed. A novel transformation and algorithm are then developed using simple geometry and Matlab programming which can also be used for other subsolidus systems in case appropriate data are provided.


Keywords iron steel slag • mathematical model • quaternary system • subsolidus equilibrium

## 1 Introduction

Ternary/Quaternary phase equilibrium diagrams are the milestones for determining phases and their corresponding wt . (or at.)\% and their concentrations under equilibrium conditions in multicomponents, multiphase systems at given pressures and temperatures. The system $\mathrm{Al}_{2} \mathrm{O}_{3}-\mathrm{MgO}_{-} \mathrm{SiO}_{2}-$

[^0]CaO is one of the most studied in the oxide field due to its applications in blast furnace slags, bioceramics, cement production, refractories, glasses and technical ceramics. The phase compatibility model of the system is derived from the literature data and built in 3 dimensions using solid wood and cords and using Mathematica programming based on the geometrical consideration. Fifty one equilibrium quaternary phase assemblages have been established, which define the entire subsolidus phase space based on data given in Ref 1. There are so far very few studies on iron and steel slags that include quaternary descriptions. ${ }^{[2-4]}$ Vasquez et al. ${ }^{[5,6]}$ studied the crystallization volume of alumina in the $\mathrm{Al}_{2} \mathrm{O}_{3}{ }^{-}$ $\mathrm{MgO}-\mathrm{SiO}_{2}-\mathrm{CaO}$ system and the invariants points were determined in compatibility tetrahedrons $\mathrm{Al}_{2} \mathrm{O}_{3}+\mathrm{MgAl}_{2}-$ $\mathrm{O}_{4}+\mathrm{Al}_{6} \mathrm{Si}_{2} \mathrm{O}_{13}+\mathrm{CaAl}_{2} \mathrm{Si}_{2} \mathrm{O}_{8}$ (anorthite) and $\mathrm{Al}_{2} \mathrm{O}_{3}$ $+\mathrm{MgAl}_{2} \mathrm{O}_{4}+\mathrm{CaAl}_{2} \mathrm{Si}_{2} \mathrm{O}_{8}+\mathrm{CaAl}_{2} \mathrm{O}_{19}$ which are also presented in our descriptive model. A general summary of quaternary systems can be found in Hummel's ${ }^{[7]}$ and Cox et al.'s textbooks ${ }^{[8]}$ in Chapter 8. Lutsyk et al. worked on T-x-$y-z$ diagrams invariant point determination by the geometrical computation and mainly based on intersecting planes for findings invariant points to find out the compositions and claimed to predict the view of horizontal or vertical sections. ${ }^{[9]}$ The three dimensional descriptive model of $\mathrm{Al}_{2} \mathrm{O}_{3^{-}}$ $\mathrm{MgO}-\mathrm{SiO}_{2}-\mathrm{CaO}$ system in the present work excluding solid solutions is based essentially on previous research. ${ }^{[10-12]}$ Preliminary studies for determining the phase distribution were started in the second half of the previous century by Devries and Osborn ${ }^{[13]}$ and kinetic studies of the slag crystallization were performed by Öveçoğlu et al. ${ }^{[14]}$ and Lai et al. ${ }^{[15]}$ and specific compositions under specific basicity and $\mathrm{CaO} / \mathrm{SiO}_{2}$ ratio were studied by Kou et al. ${ }^{[16]}$ and Wang et al. ${ }^{[17]}$, respectively.

A join connecting the compositions of the primary crystals of two primary fields in two dimensional ternary
system or two volumes in three dimensional quaternary system having a common boundary line or surface respectively is called an Alkemade line. Thus it is essential that for an Alkemade line to exist, common phase boundaries/surfaces should exist between components.

Besides the existing thermodynamical software developed for estimating phase states and their corresponding weight/atomic fractions for the given input pressures, temperatures and concentrations, the developed model is based on pure mathematical and geometrical principles provided that the compositional data of binary and ternary compounds in any existing quaternary systems are supplied as the input. Any arbitrary quaternary composition chosen will essentially be located in one of the compatibility tetrahedra for which at the end of slow cooling, below solidus, the equilibrium phases will be those at the apices of the tetrahedron; their weight or molecular percentages will then be calculated applying the lever rule.

## 2 Method

The Mathematica programming is used to draw the Figures of the tetrahedron while the Matlab. programming is used for developing the algorithm. First, the coordinates of the 4 apices of a tetrahedron given in Fig. 1 are calculated in the Cartesian coordinate system.

$$
\mathrm{Al}_{2} \mathrm{O}_{3}=\{-50,0,0\}
$$



Fig. 1 Three dimensional model of $\mathrm{Al}_{2} \mathrm{O}_{3}-\mathrm{MgO}-\mathrm{SiO}_{2}-\mathrm{CaO}$ quaternary system
$\mathrm{MgO}=\{0,86.6,0\}$
$\mathrm{SiO}_{2}=\{50,0,0\}$
$\mathrm{CaO}=\{0,28.87,81.65\}$
Next, the mathematical direct (and reverse transformations) of the existing compositions from the quaternary system ( $\mathrm{A} \%, \mathrm{~B} \%, \mathrm{C} \%, \mathrm{D} \%$ ) to the cartesian coordinate system ( $\mathrm{x}, \mathrm{y}, \mathrm{z}$ ) are derived. Using the direct transformations the xyz coordinates of the existing compounds in quaternary system are calculated and given as:
$\mathrm{CS}=\{25.86,13.93,39.4\}, \mathrm{C} 3 \mathrm{~S} 2$
$=\{20.83,16.84,47.63\}, \mathrm{C} 2 \mathrm{~S}=\{17.45,18.79,53.16\}$,
$\mathrm{C} 3 \mathrm{~S}=\{13.15,21.27,60.16\}, \mathrm{C} 3 \mathrm{~A}$
$=\{-18.88,17.96,50.80\}, \mathrm{C} 12 \mathrm{~A} 7$
$=\{-25.75,13.99,39.58\}$,
$\mathrm{CA}=\{-32.27,10.23,28.93\}, \mathrm{CA} 2$
$=\{-39.23,6.06,17.14\}$,
CA6 $=\{-45.80,2.42,6.84\}$,
$\mathrm{A} 3 \mathrm{~S} 2=\{-21.83,0,0\}, \mathrm{CAS} 2=\{3.23,5.77,16.33\}$,
C2AS $=\{-7.5,11.83,33.48\}$,
M2A2S5 $=\{8.25,11.90,0\}$, M2S $=\{21.35,49.62,0\}$, MS $=\{29.9,34.72,0\}$


Fig. 23 Dimensional modelling of quaternary equilibrium system $\mathrm{Al}_{2} \mathrm{O}_{3}-\mathrm{MgO}-\mathrm{SiO}_{2}-\mathrm{CaO}$ system below solidus temperature


Fig. 3 The $\mathrm{Al}_{2} \mathrm{O}_{3}-\mathrm{MgO}-\mathrm{SiO}_{2}-\mathrm{CaO}$ quaternary system presented at different directions
$\mathrm{AKER}=\mathrm{C} 2 \mathrm{MS} 2=\{22.05,24.68,33.56\}$,
M4A5S2 $=\{-24.63,17.59,0\}$,
MONT $=\mathrm{CMS}=\{19,32.91,29.39\}$, C3MS2 $=$ MERV
$=\{18.35,25.25,41.69\}$,
$\mathrm{DIOP}=\mathrm{CMS} 2=\{27.7,23.5,21.05\}$,
$\mathrm{MA}=\{-35.83,24.52,0\}$,
C3MA2 $=\{-24.74,20.25,33.26\}$
in which $\mathrm{C}=\mathrm{CaO}, \mathrm{A}=\mathrm{Al}_{2} \mathrm{O}_{3}=$ corundum, $\mathrm{M}=\mathrm{MgO}$, $\mathrm{S}=\mathrm{SiO}_{2}, \quad$ AKER $=$ akermenite,$\quad$ Diop $=$ diopsite, MERV = mervinite.

With these coordinates of the binary and ternary components obtained using the originally derived transformations, the whole quaternary system in 3 dimensions including the known Alkemade lines are drawn using the commands given in Mathematica software (Fig. 2).

A novel program is then developed when an arbitrary composition $\quad \mathrm{X}_{\mathrm{A}}=\mathrm{wt} . \% \mathrm{Al}_{2} \mathrm{O}_{3}, \quad \mathrm{X}_{\mathrm{B}}=\mathrm{wt} . \% \mathrm{MgO}, \quad \mathrm{X}_{\mathrm{C}}-$ $=\mathrm{wt} . \% \mathrm{SiO}_{2}$ and $\mathrm{X}_{\mathrm{D}}=\mathrm{wt} . \% \mathrm{CaO}$ is chosen for which it enables finding the Alkemade tetrahedron that contains the composition and calculates the distances to the apices of the tetrahedron. Thereafter applying the lever rule in the particular tetrahedron where $\mathrm{X}_{\mathrm{A}}, \mathrm{X}_{\mathrm{B}}, \mathrm{X}_{\mathrm{C}}$ and $\mathrm{X}_{\mathrm{D}}$ locate, the $\mathrm{wt} . \%$ or at. $\%$ of the existing minerals can be calculated after complete solidification of the melt is achieved (Fig. 3).

The 51 tetrahedra existing in the main $\mathrm{Al}_{2} \mathrm{O}_{3}-\mathrm{MgO}-$ $\mathrm{SiO}_{2}-\mathrm{CaO}$ system were firstly defined in the program by inserting their apices coordinates as an input. Then, the program calculates the tetrahedron's volumes one by one using iterative determinant method and calculates the total volume of the quaternary system. When any composition is
arbitrarily chosen $\left(X_{A}+X_{B}+X_{C}+X_{D}=1\right)$, the program directly transforms it into Cartesian coordinates xyz and then calculates the volumes of the 4 sub-tetrahedra formed when point XYZ divides each tetrahedron into 4 ; this is repeated by iteration for 51 tetrahedra forming the main tetrahedron. Another iteration sums the subtetrahedra volumes for each of the 51 tetrahedra and compares the results with their initial volumes determined earlier. The equality holds only when the right tetrahedron in which XYZ locates is reached and therefore summation of subtetrahedra volumes equalize the volume of its parent. As an example the two tetrahedra for which the apices are anorthite, forsterite, dyopside, enstatite and anorthite, cordierite, forsterite, spinel are given in Fig. 4. The third step of the program is to define the surfaces of the tetrahedron mathematically in which XYZ locates and then calculating the length of the segments initiating from the apices, crossing XYZ and joining the faces of the tetrahedron; the lever rule for quaternary system is then applied in order to find out the molecular fractions of the components forming the apices of the tetrahedron.

### 2.1 Validation of the Program by Comparing with the Experimental Results

The quaternary system A-M-S-C has already been studied experimentally by a few researchers ${ }^{[7,9,13-17]}$ and its ternary components were already determined and published many years ago. ${ }^{[1,5,9-12]}$ Based on the early experiments, ${ }^{[1]}$ the proposed model is tested for two arbitrary ternary compositions, ( 1,2 ), and four quaternary compositions, 3,(4),5 and 6, given in Table 1.


Fig. 4 The alkemade tetrahedra showing (a) Anorthite, cordierite, forsterite, spinel and (b) Anorthite, forsterite, dyopside, enstatite systems separately

The developed code is run for the experimental data (Table 1) given as input, the results are then compared with x-ray diffraction (XRD) findings and with the lever rule calculations performed on the $\mathrm{Al}_{2} \mathrm{O}_{3}-\mathrm{MgO}-\mathrm{SiO}_{2}-\mathrm{CaO}$ ternary system. Small numerical differences are possibly due to lever rule measurements for compositions 1 and 2.

Code outputs are provided as follows:

## FOR THE LIQUID SLAG COMPOSITION 1:

Calculate which tetrahedron the liquid composition is in:
The total sum of the main compounds must be $" \mathrm{a}+\mathrm{b}+\mathrm{c}+\mathrm{d}=100 "$

Enter \% a associated with the first corner/compound Al2O3: 30

Enter \% b associated with the second corner/compound MgO: 0

Enter \% c associated with the third corner/compound SiO2: 40

Enter \% d associated with the fourth corner/compound CaO: 30

The composition point 5 is in tetrahedron of number 2, ans $=$ "Tetrahedron \#2: Gehlenite-Wollastonite-Akerman-ite-Anorthite"

The respective labels of the tetrahedron corners are:
Corner_1 = Gehlenite
Corner_2 = Wollastonite
Corner_3 = Akermanite
Corner_4 = Anorthite
Q1 being the intersection point of the line connecting the composition point 5 and the Corner_1 with the plane
containing the respective corners 2,3 and 4 , the distance between the composition point 5 and the point Q1 is:distance5Q1 $=4.6014$, distance1Q1 denoting the distance between the Corner_1 and the intersection point Q1, the ratio distance5Q1/distance1Q1 is:ratio11 $=0.2265$

Q2 being the intersection point of the line connecting the composition point 5 and the Corner_2 with the plane containing the respective corners 1,3 and 4 , the distance between the composition point 5 and the point Q2 is:distance5Q2 $=5.9553$, distance2Q2 denoting the distance between the Corner_2 and the intersection point Q2, the ratio distance5Q2/distance2Q2 is:ratio22 $=0.1854$

Q3 being the intersection point of the line connecting the composition point 5 and the Corner_3 with the plane containing the respective corners 1,2 and 4 , the distance between the composition point 5 and the point Q3 is:distance5Q3 $=0.0072$, distance3Q3 denoting the distance between the Corner_3 and the intersection point Q3, the ratio distance5Q3/distance3Q3 is:ratio33 $=2.8498 \mathrm{e}-04$

Q4 being the intersection point of the line connecting the composition point 5 and the Corner_4 with the plane containing the respective corners 1,2 and 3 , the distance between the composition point 5 and the point Q4 is:distance5Q4 $=12.6114$, distance4Q4 denoting the distance between the Corner_4 and the intersection point Q4, the ratio distance5Q4/distance4Q4 is:ratio44 $=0.5879$

FOR THE LIQUID SLAG COMPOSITION 2:
Calculate which tetrahedron the liquid composition is in:

Table 1 Experimental slag compositions and equilibrium solid phases and corresponding code output

| Liquid comp.\# | wt. $\% \mathrm{Al}_{2} \mathrm{O}_{3}$ | wt. $\% \mathrm{MgO}$ | wt. $\% \mathrm{SiO}_{2}$ | wt. \% CaO | Exp. output | Code output | Refs. |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | 30 | 0 | 40 | 30 | anorthite | anorthite | [1] |
|  |  |  |  |  | gehlenite | gehlenite |  |
|  |  |  |  |  | wollastonite 0.1857 | wollastonite |  |
|  |  |  |  |  |  | $0.1854^{+}$ |  |
| 2 | 20 | 0 | 70 | 10 | anorthite <br> mullite <br> cristobalite 0.4777 | anorthite | [1] |
|  |  |  |  |  |  | mullite |  |
|  |  |  |  |  |  | cristobalite |  |
|  |  |  |  |  |  | $0.4775^{++}$ |  |
| 3 | 11.6 | 3.2 | 39.2 | 46 | geh. aker undetermined quantitavely | geh.: 0.3931 | [14] |
|  |  |  |  |  |  | aker: 0.2165 |  |
|  |  |  |  |  |  |  |  |
| 4 | 90 | 1 | 7 | 2 | corundum, anorthite, mullite liquid $\mathrm{t}=1352^{\circ} \mathrm{C}$ <br> above solidus | corundum | 5(8)* |
|  |  |  |  |  |  | 0.7697 |  |
|  |  |  |  |  |  | anorthite |  |
|  |  |  |  |  |  | 0.10 |  |
|  |  |  |  |  |  | mullite |  |
|  |  |  |  |  |  | 0.0950 |  |
|  |  |  |  |  |  | spinel |  |
|  |  |  |  |  |  | 0.0353 |  |
| 5 | 90 | 2.5 | 4 | 3.5 | corundum | corundum | $5(4) * *$ |
|  |  |  |  |  | spinel | 0.6222 |  |
|  |  |  |  |  | $\mathrm{ca}_{6}$ | spinel |  |
|  |  |  |  |  | liquid | 0.0883 |  |
|  |  |  |  |  | $\mathrm{t}=1490{ }^{\circ} \mathrm{C}$ | ca6 |  |
|  |  |  |  |  | above solidus | 0.1971 |  |
|  |  |  |  |  |  | anorthite |  |
|  |  |  |  |  |  | 0.0924 |  |
| 6 | 15 | 8 | 38.5 | 38.5 | geh. aker. augite undetermined quantitavely | geh. 0.4053 | [15] |
|  |  |  |  |  |  | aker |  |
|  |  |  |  |  |  | 0.2653 (diopsite) ${ }^{\text {x }}$ |  |
|  |  |  |  |  |  | 0.2202 |  |

* Batch \#8 ** Batch \#4 + wt. \% wollastonite $++\mathrm{wt} . \%$ cristobalite x close to augite.

The total sum of the main compounds must be $" \mathrm{a}+\mathrm{b}+\mathrm{c}+\mathrm{d}=100 "$

Enter \% a associated with the first corner/compound Al2O3: 20

Enter \% b associated with the second corner/compound MgO: 0

Enter \% c associated with the third corner/compound SiO2: 70

Enter \% d associated with the fourth corner/compound CaO: 10

The composition point 5 is in tetrahedron of number 51 , ans $=$ "Tetrahedron \#51: Cord-Anorthite-Mullite-Cristobalite"

The respective labels of the tetrahedron corners are:
Corner_1 = Cord
Corner_2 = Anorthite
Corner_3 = Mullite

Corner_4 = Cristobalite
Q1 being the intersection point of the line connecting the composition point 5 and the Corner_1 with the plane containing the respective corners 2,3 and 4 , the distance between the composition point 5 and the point Q1 is:distance5Q1 $=0.0031$, distance1Q1 denoting the distance between the Corner_1 and the intersection point Q1, the ratio distance5Q1/distance1Q1 is:ratio11 $=1.4819 \mathrm{e}-04$

Q2 being the intersection point of the line connecting the composition point 5 and the Corner_2 with the plane containing the respective corners 1,3 and 4 , the distance between the composition point 5 and the point Q2 is:distance5Q2 $=23.4287$, distance2Q2 denoting the distance between the Corner_2 and the intersection point Q 2 , the ratio distance5Q2/distance2Q2 is:ratio22 $=0.5000$


Fig. 5 Verification of the code for initial melt composition 1 and 2 as indicated on the $\mathrm{Al}_{2} \mathrm{O}_{3}-\mathrm{SiO}_{2}-\mathrm{CaO}$ ternary equilibria [1]

Q3 being the intersection point of the line connecting the composition point 5 and the Corner_3 with the plane containing the respective corners 1,2 and 4 , the distance between the composition point 5 and the point Q3 is:distance5Q3 $=1.0912$, distance3Q3 denoting the distance between the Corner_3 and the intersection point Q3, the ratio distance5Q3/distance3Q3 is:ratio33 $=0.0224$

Q4 being the intersection point of the line connecting the composition point 5 and the Corner_4 with the plane containing the respective corners 1,2 and 3 , the distance between the composition point 5 and the point Q4 is:distance5Q4 $=24.1745$, distance4Q4 denoting the distance between the Corner_4 and the intersection point Q4, the ratio distance5Q4/distance4Q4 is:ratio44 $=0.4775$

FOR THE LIQUID SLAG COMPOSITION 4:
It is calculated in which tetrahedron the composition point is located.

The total sum of the percent values of compounds must be " $a+b+c+d=100$ "

Enter \% a associated with the first corner/compound Al2O3: 90

Enter \% b associated with the second corner/compound MgO: 1

Enter \% c associated with the third corner/compound SiO2: 7

Enter \% d associated with the fourth corner/compound CaO: 2

The composition point 5 is in tetrahedron of number 19, ans $=$ "Tetrahedron \#19: Anorthite-Corundum-Mullite-Spinel"

The respective labels of the tetrahedron corners are:
Corner_1 = Anorthite
Corner_2 = Corundum
Corner_3 = Mullite
Corner_4 = Spinel
Q1 being the intersection point of the line connecting the composition point 5 and the Corner_1 with the plane containing the respective corners 2,3 and 4 , the distance between the composition point 5 and the point Q 1 is: distance5Q1 $=5.2534$, distance1Q1 denoting the distance between the Corner_1 and the intersection point Q1, the ratio distance5Q1/distance1Q1 is:ratio11 $=0.1000$

Q2 being the intersection point of the line connecting the composition point 5 and the Corner_2 with the plane containing the respective corners 1,3 and 4 , the distance between the composition point 5 and the point Q2 is:distance5Q2 $=29.3207$, distance2Q2 denoting the distance between the Corner_2 and the intersection point Q2, the ratio distance5Q2/distance2Q2 is:ratio22 $=0.7697$

Q3 being the intersection point of the line connecting the composition point 5 and the Corner_3 with the plane containing the respective corners 1,2 and 4 , the distance between the composition point 5 and the point Q3 is:distance5Q3 $=2.0776$, distance3Q3 denoting the distance between the Corner_3 and the intersection point Q3, the ratio distance5Q3/distance3Q3 is:ratio33 $=0.0950$

Q4 being the intersection point of the line connecting the composition point 5 and the Corner_4 with the plane containing the respective corners 1,2 and 3 , the distance between the composition point 5 and the point Q4 is:distance5Q4 $=0.8724$, distance4Q4 denoting the distance between the Corner_4 and the intersection point Q 4 , the ratio distance5Q4/distance4Q4 is:ratio44 $=0.0353$

The lever rule is then applied for composition 1 in order to calculate $\mathrm{wt} . \%$ of wollastonite as indicated ratio 22 and for composition 2 to calculate wt. $\%$ of cristobalite as indicated ratio 44 given as above in output of the code. When the lever rule calculations based on geometrical measurements are observed in the Fig. 5, it is clearly seen that a nearly perfect match is obtained which necessarily validates the code.

For liquid slag compositions 3 and 6 provided in Ref. 14 and 15 based on EDS (Energy Dispersive Spectroscopy) and XRD analysis, the crystal configurations after solidification were determined to be mainly gehlenite, akermanite and gehlenite, akermanite, augite respectively which the latter is chemically close to diopsite. These results are also proved by the code when it is run except the presence of wollastonite (CS) and $\mathrm{C}_{3} \mathrm{~S}_{2}$ for composition 3 and CS for composition 4 which appeared in the output. Composition 3 is determined to be located in tetrahedron \#24 whereas composition 4 is calculated to be in tetrahedron\#14. The missing components $\left(\mathrm{CS}, \mathrm{C}_{3} \mathrm{~S}_{2}\right)$ which were not determined by XRD are most probably due to possible solid solutions for which the code does not work or to the imprecise measurements during XRD analysis for determining minor phases. For liquid slag composition 4 and 5 provided in Ref 5, a perfect match of solid phases including their relative amounts is also found (Table 1). The presence of liquid phase determined experimentally did not appear as an output due to limitation of the code which works for subsolidus case.

## 3 Conclusion

With this original algorithm developed using simple 3 D geometry and linear algebra principles, mathematical programming allows scientists to estimate the subsolidus phases in equilibrium including their relative amounts when an initial melt composition is entered into the system as an input. Before starting experimental studies and XRD/SEM characterizations, this algorithm will help reducing the costs and time consumption which is the main contribution of this research. Further studies can be done including non-stochiometric solid solutions such as $(\mathrm{MgO})_{1-x}\left(\mathrm{Al}_{2} \mathrm{O}_{3}\right)_{1+x}$ for which another loop repeating in a certain solid solution range ( $a<x<b$ interval) can be devised in the code, also including liquid range for which non planar liquidus surfaces including their boundaries may be defined using line and surface integrals.

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[^0]:    Tanju Yelkenci
    tanju.yelkenci@tau.edu.tr
    Z. Engin Erkmen
    eerkmen@marmara.edu.tr
    1 Turkish-German University, Faculty of Engineering, 34820 Istanbul, Turkey
    2 Marmara University, Faculty of Engineering, 34854 Istanbul, Turkey

