PAPER • OPEN ACCESS

Preface

To cite this article: 2020 J. Phys.: Conf. Ser. 1445 011001

View the article online for updates and enhancements.



IOP ebooks[™]

Bringing together innovative digital publishing with leading authors from the global scientific community.

Start exploring the collection-download the first chapter of every title for free.

Preface

The International Symposium on Nanoscience & Nanotechnology in Life Sciences 2017 (ISNNLS 2017) took place between 28-29 November 2017 at Hotel Santika Premiere, Surabaya, Indonesia. The symposium was organized by the Research Center for Quantum Engineering Design and Faculty of Science and Technology, Universitas Airlangga, Indonesia. ISNNLS 2017 was the fourth annual symposium that initiated and previously held by Research Center for Nanosciences and Nanotechnology (RCNN), Institut Teknologi Bandung, Indonesia.

In the last decade, nanotechnology has advanced, and nanoscale materials are used in everything from chemical catalyst to antibacterial agents. The scientific program of the symposium included many topics in the field of nanotechnology and its role in life sciences. The symposium presented keynote speakers from notable experts of nanoscience and nanotechnology, i.e., Kyle E. Cordova from University of California, USA, Prof. Yoshitada Morikawa from Osaka University, Japan, Prof. Heni Rachmawati from Institut Teknologi Bandung, Indonesia, Dr. Tommy Julianto Bustami Effendi from Universiti Teknologi MARA, Malaysia, and Mochamad Zakki Fahmi, Ph.D. from Universitas Airlangga, Indonesia. ISNNLS 2017 facilitated researchers, scientists, and engineers to exchange ideas and discuss progress in four main tracks, chapter of modeling, chapter of synthesis, chapter of treatment and chapter of supporting.

More than 100 participants took part in the symposium. We received 46 submissions to all main tracks. Papers were evaluated to the high standard. Two reviewers from Program Committee and additional reviewers were assigned to review each article. After the completion of the peer review process, 29 papers were selected for publication in the Journal of Physics: Conference Series (JPCS).

We would like to thank all authors, program committee members, reviewers, and fellow members of the symposium committee for their contribution to the symposium. We also greatly appreciated the publication support from Center for Journals Development and Scientific Publications, Universitas Airlangga, Indonesia.

Organization

1. Steering Committee

Position	Name	Institution
Head	Prof. Hermawan K. Dipojono	ITB
	Prof. Win Darmanto	UA
Member	Prof. Suprijadi	ITB
	Prof. Moh. Yasin	UA

2. Scientific Board

Position	Name	Institution
Editor-in-chief	Prof. Yoshitada Morikawa	OU
Deputy	Prof. Sulaiman W. Harun	UM
Member	Prof. Heni Rachmawati	ITB
	Dr. Tommy J. B. Effendi	UiTM
	Mochamad Z. Fahmi, Ph.D	ITB
	Andi H. Zaidan, Ph.D	UA
	Prof. Brian Yuliarto	ITB
	Mohammad K. Agusta, Ph.D	ITB
	Triati D. K. Wungu, Ph.D	ITB
	Damar R. Adhika, Ph.D	ITB
	Benny Permana, Ph.D	ITB
	Fadjar Fathurrahman, Ph.D	ITB

3. Organizing Committee

Position	Name	Institution
Program Chair	Febdian Rusydi, Ph.D	UA
Deputy	Adhitya G. Saputro, Ph.D	ITB
	Enggar Alfianto, M.Si	ITATS
Treasury	Ira Puspitasari, Ph.D	UA

Section A: Secretarial

Head	Nufida Dwi Aisyah, S.Si	UA
Member	Etika D. Susanti, S.Si	UA
	Viny V. Tanuwijaya, M.T	ITB
	Ema Rimawati	ITB
	Muhamad N. Romadhoni, S.Si	UA
	Roichatul Madinah, S.Si	UA
	Rochmatun Nisa'	UA
	Fitri N. Febriana, S.Si	UA
	Husnul Khuluq	UA
	Andri Wahyudianto	UA

Section B: Publication

Head	Rizka N. Fadilla, S.Si	UA
Member	Grendy Firmanda, S.Si	UA
	Isniar Wardani, S.Si	UA
	Binti Q. A'yuni, S.Si	UA
	Dalliyah A. Aminati	UA

Section C: Event

Head	Astrid N. Jannah, S.Si	UA
Member	Novi Irvianty	ITB
	Syifa M. Restian	ITB
	Putri A. Lestari	ITB
	Maghfira Maulidiyah, S.Si	UA
	Dian E. Candrasari, S.Si	UA
	Lafitara G. Arisha, S.Si	UA
	Winda O. D. Cahyani, S.Si	UA
	Nikmatul Khoiroh, S.Si	UA
	Muhammad C. E. Dien, S.Si	UA
	Beni Hamdani	UA
	Ella Z. Fadilah	UA
	Daysta A. Zahra	UA
	Rachma Arinsyah	UA
	Syahrul Munir, S.Si	UA
	Soleha R. Junia, S.Si	UA
	Billy Y. Wijoyo	UA
	Hakam Pranatagama, S.Si	UA
	Jeremy Pamungkas	UA
	Samuel E. P. P. Masan	UA

Abbreviation list for Institution

ITB	Institut Teknologi Bandung, Indonesia
UA	Universitas Airlangga, Indonesia
OU	Osaka University, Japan
UM	Universiti Malaya, Malaysia
UITM	Universiti Teknologi Mara, Malaysia
ITATS	Institut Teknologi Adhi Tama Surabaya, Indonesia

NOTICE: Access in China: Some users in China are being blocked by IOP's security software. Please contact china@ioppublishing.org

Table of contents

Volume 1445

January 2020

♦ Previous issue
 Next issue ▶

International Symposium on Nanoscience & Nanotechnology in Life Sciences 2017 28-29 November 2017, Surabaya, Indonesia

Accepted papers received: 24 December 2019 Published online: 27 January 2020

View all abstracts

Preface			
OPEN ACCESS			011001
Preface			
	View article	🔁 PDF	
OPEN ACCESS			011002
Peer review state	ment		
	View article	🔁 PDF	
Papers			
Modelling			
OPEN ACCESS			012001
The Perovskite Ph	nase Optimize of Ba	rium Titanate Nanoparticles	
Jan Ady, Arum Nur	pratiwi, Aliyah and Wi	inda Apriliana	
➡ View abstract	View article	🔁 PDF	
OPEN ACCESS			012002
Penetration Dept	h of Free Falling Int	ruder into a Particles Bed in Fluid-Immersed Two-	

Penetration Depth of Free Falling Intruder into a Particles Bed in Fluid-Immersed Two Dimension Spherical Particle System

S Viridi and T A Sanny

	View article	PDF	
OPEN ACCESS			012003
In vitro study of Na Substitute Associa	ano Hydroxyapatite ated- 3D printed B	e/Streptomycin -Gelatin-Based Injectable Bone one Scaffold for Spinal Tuberculosis Case	
Inten Firdhausi War	dhani, Rofi Mega Riz	zki Samudra, Katherine and Dyah Hikmawati	
	View article	PDF	
OPEN ACCESS			012004
Ultraviolet (UV) Ac Extract to <i>Streptor</i>	tivation Effect on A coccus mutans	Intibacterial Agents of Red Betel (<i>Piper Crocatum</i>)	I
Suryani Dyah Astuti Moh. Yasin	, Rio Dysan Tirtana, /	Amalia Fitriana Mahmud, Amiliyatul Mawaddah, Abdur	achman and
	View article	PDF	
OPEN ACCESS			012005
A Computational T Detected Sensor & Configuration	Theory Study of Sub based-on Fe ₃ O ₄ Na	rface Plasmon Resonance (SPR) Porcine Gelatine anoparticle—CNT with ATR Method in Kretschmanr	1
Maulina Lutfiyah, W	'ahyu Aji Eko Prabow	o and Asih Melati	
	View article	PDF	
OPEN ACCESS			012006
Implementation o Density Functiona	f Go language to ca I Theory (DFT)	alculate ground state energy of atoms based on	
Lafitiara Gita Arisha	, Enggar Alfianto and	d Febdian Rusydi	
	View article	PDF	
OPEN ACCESS			012007
Theoretical Investi using First Princip	gation of Fe and A les Calculation	I Surface Structure in the Case of H Adsorption	
N D Aisyah, D E Car	ndrasari, A Stefanus,	R Madinah, R Nisa' and A H Zaidan	
➡ View abstract	View article	🔁 PDF	
OPEN ACCESS			012008
Theoretical Study principles Calcula	on Radiationless D tion	Decay in Butadiene Isomerization Case using First-	
R N Fadilla, A N Jan	nah, F N Febriana, S	Munir and A H Zaidan	
	View article	PDF	
OPEN ACCESS			012009
Approximation Re	ctangular Function	as Potential Barrier	
I Wardani, N D Aisya	ah and A Supardi		

View abstract	View article	🔁 PDF
---------------	--------------	-------

OPEN ACCESS			012010
The Effect of Basis Isomerization	s Set on Quantum 1	unneling Probability with the Case of trans-HCOH	
E D Susanti, S R Ju	nia, R N Fadilla and A	A Supardi	
	View article	PDF	
OPEN ACCESS			012011
D-band Center Th Al(100) Surfaces:	eory for the Case of A Density Functior	f Hydrogen Atom Adsorption on Fe(100) and nal Study	
Wahyu Aji Eko Prab	owo, Nikmatul Khoir	oh, Satriyaji Wibisono and Adri Supardi	
➡ View abstract	Tiew article	PDF	
Synthesis			
OPEN ACCESS			012012
Physical Characte Aerosolization Teo	eristics of Erythropo chnique	etin Encapsulated into Alginate Polymer Using	
Dewi Melani Hariya	di, Noorma Rosita ar	nd Kamila Amalia	
	View article	🔁 PDF	
OPEN ACCESS			012013
Synthesis of Alum	ninium Nanoparticle E Kurpiawan	es Using Electrochemical Method	
 View abstract 	View article		
OPEN ACCESS			012014
Synthesis of ZnO (High Energy Milli	Nanoparticles Usin ng)	g Mechano-Chemical Method By Utilizing 3D HEM	
Siswanto and Maya	asari Hariyanto		
	View article	PDF	
OPEN ACCESS			012015
Synthesis of Hydr Substitution	oxyapatite Based o	n Nano Coral Using precipitation Method For Bone	
Siswanto, Dyah Hik	mawati, N Benecdita	and Siti Nurmala	
	Tiew article	PDF	
OPEN ACCESS			012016
Synthesis of SiO ₂ Gelatine	– PVA – Gelatine N	lanocomposite Membrane by Handling of the	

Jan Ady, Muhammad Abdul Aziz and Siti Nur Seha

View abstract	View article	🔁 PDF	
OPEN ACCESS			012017
Temperature Effect Characterization c	ct of Chemical Bath of Zinc Oxide Nanor	Deposition (CBD) to Fabrication and ods Thin Films Based Gas Sensing: Ethanol	
Adimas Ramadhan,	, Ni Luh Wulan Septia	ani, Wahyu Aji Eko Prabowo and Asih Melati	
➡ View abstract	Tiew article	🔁 PDF	
Treatment			
OPEN ACCESS			012018
Hepato-Renal Pro Extract in Streptoz	tective Effects of M cotocin-induced Dia	angosteen (<i>Garcinia mangostana</i> L.) Pericarp abetic Mice	
Saikhu Akhmad Hu	sen, Septian Hary Ka	lqutny, Arif Nur Muhammad Ansori,	
Raden Joko Kuncor	oningrat Susilo, Firas	s Khaleyla and Dwi Winarni	
View abstract	View article	PDF	
OPEN ACCESS			012020
Snedds (Self-nan Extract on Cervica	oemulsifying Drug I I Cancer Cells (HeL	Delivery System) Formulation of <i>Sarang Semut</i> a) with MTT Assay Method	012020
B H Nugroho, M R S	Syifaudin, L R Fauzi, E	Anggraini and H O Ritonga	
View abstract	View article	🔁 PDF	
OPEN ACCESS			012021
Determination of I Candidate of Phot	Infrared Laser Ener todynamic Therapy	gy Dose for Cancer Cells Inactivation as a	
Septia Kholimatuss	a'diah, Suryani Dyah	Astuti and Retna Apsari	
➡ View abstract	View article	🔁 PDF	
OPEN ACCESS			012022
Electrospun Colla	gen-based Scaffold	d as Therapeutic Agent for Ocular Chemical Inju	ry
N A F Hasbiyani, D I	Hikmawati and Siswa	into	
	View article	🔁 PDF	
OPEN ACCESS			012023
The Effect of Addit as Bone Filler	tive Substitute of M	gO Nanoparticle on the Characteristics of Expo	rts
Djony Izak Rudyardj	jo and Setiawan Wija	yanto	
	View article	PDF	
Supporting			

The Influence of Solvent Parameters along Terminal Jet Radius and Fiber Diameter in	
Electrospinning	

P M Widartiningsih, F Iskandar, M M Munir and S Viridi

View abstract	View article	🔁 PDF
---------------	--------------	-------

OPEN ACCESS			012026
Expert System for as Diagnosis Sup	Stroke Classification porting Device	on Using Naive Bayes Classifier and Certainty Factor	
Khusnul Ain, Hanik	B. Hidayati and Olivia	a Aulia Nastiti	
➡ View abstract	Tiew article	PDF	
OPEN ACCESS			012027
Design Monitoring	g Electrical Power C	onsumtion at Computer Cluster	
Enggar Alfianto, Sit	i Agustini, Syahri Muł	narom, Febdian Rusydi and Ira Puspitasari	
	View article	PDF	
OPEN ACCESS			012028
Numerical Simula	ition of Spear Motic	on as Game Items	
R R Muhima, S Mar	di, M Hariadi and I P	uspitasari	
	View article	PDF	
OPEN ACCESS			012029
Modeling Structur	re of Portable River	Bridge using Fiber – Reinforced Polymer (FRP)	
A Sa'diyah, A F Pras	setya and E Alfianto		
	View article	PDF	
JOURNAL LINKS			
Journal home			
Information for orga	anizers		
Information for aut	nors		
Search for publishe	d proceedings		
Contact us			

Reprint services from Curran Associates

PAPER • OPEN ACCESS

Implementation of Go language to calculate ground state energy of atoms based on Density Functional Theory (DFT)

To cite this article: Lafitiara Gita Arisha et al 2020 J. Phys.: Conf. Ser. 1445 012006

View the article online for updates and enhancements.



This content was downloaded from IP address 182.1.70.117 on 04/03/2020 at 13:59

Implementation of Go language to calculate ground state energy of atoms based on Density Functional Theory (DFT)

¹Lafitiara Gita Arisha, ²Enggar Alfianto, ¹Febdian Rusydi

¹Theoretical Physics Research Group, Dep. of Physics, Fac. of Science and Technology, Airlangga University, Jl. Mulyorejo, Surabaya, Indonesia 60115 ²Dep. of Computer System, Institut Teknologi Adhi Tama Surabaya, Jl Arif Rachman Hakim 100, Surabaya, Indonesia.60111

Email: rusydi@fst.unair.ac.id

Abstract. This study is using Go programming language that support parallel programming for numerical calculation. The program was created is designed for calculate ground-state energy of electron, which is based on Density Functional Theory (DFT). The basic mathematics of this program is using many basic concept of numerical mathematics (matrix calculation, Poisson solver, and standard routine of numerical mathematics).

1. Research Background

Processor is the part of the computer that acts as the brain. The function of processor is to handle the speed of processing data, executing user commands, and the ability of the computer to run multiple tasks together (multi-task). Speed and multi-task are depends on a part of the processor called core. The processor expanding the development by add number of cores that construct it.

The development of the processor and the number of cores can increase the performance of calculation and reduce calculation time. It is because the elements of the processor can do different tasks at the same time (multi-tasking). The multi-tasking process is supported by many core systems. Multi core systems are the right solution when users want to increase high processing speeds, for example in numerical calculations.

In addition to cores, supporting programming languages also affect the speed of numerical calculations [7]. There are programming languages that are often used for numerical calculations, such as the Fortran, Python, Pascal, and C programming languages. Besides the programming language, there is a new programming language called Go. The process of Go language compilation is faster than C language [6]. In addition, the Go language also supports multiple core systems, making it suitable for numerical calculations. Error! Reference source not found.

This study utilizes the Go language as a programming language in DFT-based calculation programs. The aims of the program is to calculate the ground state energy of simple atoms

International Symposium on Nanoscience & Na	notechnology in Life Sci	ences 2017	IOP Publishing
Journal of Physics: Conference Series	1445 (2020) 012006	doi:10.1088/1742	-6596/1445/1/012006

2. Calculation method

The program is designed by implementing DFT to the Go programming language. The program divided into four processes that we specify, such as Figure 1. Process A determines the initial electron density (ρ_N). Process B determines the potential energy of the electron (V). Process C determines the ground state energy (E_N). The D process determines the new electron density (ρ_{N+1}) for the iteration process. If the fourth process has finished, the program will iterate. The output in the fourth process will be input to the second process, replacing ρ_N in the first process.



Figure 1. Flow chart of the program based on DFT calculation

The flow chart that have designed is then used to build a ground state energy calculation program using the Go programming language. The program is tested on a number of selected atoms, such as: H, He, C, O, Cl, Fe, and Zn. The results of testing the calculation atoms compared with the results obtained using the C language program.

In addition to getting the energy, we also get the program calculation time. So we also compare the time needed by the Go language and the C language for the calculation of each atom.

3. Result and discussion

3.1. Flow diagram based on DFT

The flow of this program started with process A is to determine the initial electron density (ρ_N) of an atom tested. In Figure 2 shows the program flow to determine ρ_N . It is seen that to determine ρ_N requires an atomic number (Z), the maximum number of orbitals (nmax), the maximum number of angular momentum (lmax), and the number of electrons that fill each atomic skin (F [1] [n]) as input. We get this information from the atomic electron configuration tested.

Furthermore, these inputs are used in four stages. The first step to determine ρ_N is to determine the total energy for all orbitals (E). The Z and nmax values are used as inputs to calculate the energy of each orbital (En). Each En value is n = 0 until nmax is added up, then multiplied by the number of electrons filling the skin (F [1] [n]). So that we get total energy for all orbitals (E).

The second step is to calculate the total effective potential energy of the electron at all angular momentum, which we represent with V_0 . Before getting the value V_0 , we first calculate the effective potential energy value V_{eff} at each angular momentum. V_{eff} is calculated for each value of l, starting

International Symposium on Nanoscience & N	Vanotechnology in Life Sci	ences 2017	IOP Publishing
Journal of Physics: Conference Series	1445 (2020) 012006	doi:10.1088/1742	-6596/1445/1/012006

from l = 0 to lmax. The value of V_{eff} for each *l* is summed then multiplied by the number of electrons filling the skin (F [l] [n]), to get the total potential effective energy (V_0).

Then the third step is to solve the Schrödinger equation computationally. The values E and V_0 obtained from the first and second processes we use to get the wave function $(\psi_0(x))$. From the value of ψ_0 obtained in the fourth stage, we use it to determine ρ_N . The square of ψ_0 gives the electron distribution, the total electron density is the sum of the squares ψ_0 multiplied by the number of electrons that fill the F[[1] [n] orbitals. In mathematical form, electron density is written as follows





The process of B in this program flow is to determine the potential energy of the electron, such as Figure 3. There are three potential energy contributing here, Hartree potential energy (V_H) , potential electron energy with nucleus (V(r)), and *exchange-correlation* potential energy (V_{xc}) . So that the total potential energy is the sum of the three potential energies.



Figure 3. B process for finding potential energy

International Symposium on Nanoscience & N	anotechnology in Life Sci	ences 2017	IOP Publishing
Journal of Physics: Conference Series	1445 (2020) 012006	doi:10.1088/1742	2-6596/1445/1/012006

 V_H is potential energy from the interaction between electrons and electrons. This potential energy is obtained by solving the Poisson equation, with the form of the Poisson equation

$$\nabla^2 \phi(\vec{x}) = -4 \pi [kc] e \rho(\vec{x})$$

 $\phi(\vec{x})$ is electronic potential energy for one electron. So to find potential energy due to the interaction of electrons with other electrons, it needs to be multiplied by the electron density. The mathematical forms of V_H are as follows,

$$V_{H} = \frac{1}{2} \int \phi(\vec{x}) \rho(\vec{x}) \, dV$$

 V_{XC} is the energy obtained from interactions aside from V_H and V(r). The mathematical form of energy exchange-correlation $V_{XC} = f'_{XC} (\rho(\vec{x}))$ and $f_{XC} = \epsilon_{XC} \rho$. is the mathematical form of the chosen exchange-correlation. We chose the VWN exchange correlation, with a mathematical form

$$\varepsilon = A \left[\ln \frac{x^2}{X(x)} + \frac{2b}{Q} \tan^{-1} \left(\frac{Q}{2x+b} \right) - \frac{bx_0}{X(x_0)} \left(\ln \frac{(x-x_0)^2}{X(x)} + \frac{2(b+2x_0)}{Q} \tan^{-1} \left(\frac{Q}{2x+b} \right) \right) \right]$$

if V_{XC} is added to other potential energy it will get total potential energy value,

$$V = V(r) + V_H + V_{XC}$$

The C process in this program is to determine the E_N basic state energy, such as Figure 4 left. If you want to know the energy value, you can use the Schrödinger equation by knowing the value of the wave function and potential energy. Because we already know these two values, we can get the value of E_N , using the Schrödinger equation

$$\left(\frac{d^2}{dx^2} + V\right)\psi_0 = E_N\psi_0$$

The fourth process in Figure 4 right of the program flow is to determine the new electron density for the iteration process. The iteration process is carried out to correct the values previously obtained. Because this program is based on an approach, we do iterations as a step to get closer results.



Figure 4. Left: C Process, Finding ground-state energy. Right: D Process, routine to finding new density

3.2. Program Testing

Based on the program flow that we have designed, we have realized the flow into a DFT-based numerical calculation program. The program that we designed uses the Go programming language. The reason we chose the Go programming language was because according to Peiyi Tang's paper from the Department of Computer Science, the University of Arkansas stated that parallel programming using Go language was easier and more efficient.

We present the program calculation results with the Go language in Table 1. The parameters of the program calculation results that we have tabulated in the table consist of energy values, program calculation time, program number to achieve convergent values, and time taken for the program once

iteration. The value we get from the Go language is then displayed in graphical form to see the relationship between the energy value of the program's ability to iterate.

Table 1. The program calculation value					
	Atomio	Groundstate energy calculation			
Atom	Number	Value (Hartree)	Calcualtion	Ν	Iteration time
	Number	value (Halliee)	time (s)	Convergence	
Hydrogen (H)	1	-0.44567056136	11.76	25	0.10 - 0.23
Helium (He)	2	-2.83483568022	12.12	29	0.10 - 0.23
Carbon (C)	6	-37.4257485949	32.83	32	0.10 - 1.00
Oxygen (())	8	-74.4730768471	34.97	34	0.10 - 1.01
Chlorine (Cl)	17	-458.664179537	53.55	36	0.09 - 1.01
Iron (Fe)	26	-1261.09305585	15.58	36	0.09 - 1.01
Zink (Zn)	30	-1776.57384967	16.66	37	0.09 - 1.01

Figure 5 shows the graphical relationship between atomic number and iteration convergence. The atomic number axis and Y axis indicate the number of iterations needed by the program to reach convergence. It is seen that the larger the atomic number, the program will need more iterations to achieve convergent values. So the calculated time needed to calculate a large atomic number takes longer.



Figure 5. Relation between atomic number and confergence of the calculation

4. Conclution

The Go language program made gives H-atom energy results of -0.45 Hartree with a time of 11.76 seconds; He atom of -2.83 Hartree with a time of 12.12; C atom of -37.43 Hartree with a time of 32.83 seconds; O atom of -74.47 Hartree with 34.97 seconds; Cl at -458.66 Hartree with a time of 53.55 seconds; Fe at -1261.09 Hartree with a time of 15.58 seconds; Zn at -1776.57 Hartree with 16.66 seconds for Zn atoms.

References

- [1] Aisyah, N. D., Fadilla, R. N., Dipojono, H. K., & Rusydi, F. (2017). A Theoretical Study of Monodeuteriation Effect on the Rearrangement of Trans-HCOH to H 2 CO via Quantum Tunneling with DFT and WKB Approximation. Procedia Engineering, 170, 119-123.
- [2] Balbaert, I. (2012). The Way to Go a Thorough Introduction to the Go Programming Language. iUniverse.
- [3] ALFIANTO, E., et al. Implementation of density functional theory method on object-oriented programming (C++) to calculate energy band structure using the projector augmented wave (PAW). In: *Journal of Physics: Conference Series*. IOP Publishing, 2017. p. 012043.
- [4] Fadilla, R. N., Aisyah, N. D., Dipojono, H. K., & Rusydi, F. (2017). A Theoretical Study of the Rearranging Trans-HCOH to H 2 CO via Quantum Tunneling with DFT and WKB Approximation. Procedia Engineering, 170, 113-118.
- [5] Fadilla, R. N., Aisyah, N. D., Dipojono, H. K., & Rusydi, F. (2017). The first-principle study on the stability of trans-HCOH in various solvents. International Conference on Physical Instrumentation and Advanced Materials27 October 2016, Hotel Santika Premiere, Surabaya, Indonesia. 853. Surabaya: IOP Publishing Ltd.
- [6] I. Balbaert, The Way to Go a Thorough Introduction to the Go Programming Language, iUniverse, 2012
- [7] L. Prechelt, "An Empirical Comparison of C, C++, Java, Perl, Phyton, Rexx, and Tcl," 2000.
- [8] Prechelt, L. (2000). An Empirical Comparison of C, C++, Java, Perl, Phyton, Rexx, and Tcl.
- [9] Schreiner, P. R., Resienauer, H. P., Pickard IV, F. C., Simmonett, A. C., Allen, W. D., Matyus, E., et al. (2008). Capture of hydroxymethylene and its fast disappearance through tunneling. Nature, 453(12 June), 906.
- [10] Tang, P. (2010). Multi-core Parallel Programming in Go.