Design New Compound of Meisoindigo Derivative as Anti Breast Cancer Based on QSAR Approach

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https://doi.org/10.14710/jksa.23.9.305-311

1. Introduction

The discovery of new high-efficacy compounds requires a long experimental stage, including design, synthesis, identification, purification, and activity testing. One solution that can be offered to overcome this problem is the introduction of modeling using computers. Using modeling, a compound can be searched for a relationship model between structures, both electronic and geometric, from one or a group of molecules suspected of having certain activities [1].

In the field of health, the use of computer-aided modeling is constructive in research. For example, is the treatment of diseases that attack the brain is complicated because the delivery of drug molecules to the brain is blocked by the blood–brain barrier (BBB) molecules. One method to overcome this problem has been developed by using a new method using a computer molecular dynamics approach [2].

The scientific discipline in pharmacy, which is much helped by these developments, is medicinal chemistry, especially for the study of Quantitative Structure-Activity Relationship (QSAR). This is synergistic with the development of new drug discoveries that are increasingly expected to be more effective and efficient [3]. In computer experiments, calculations are performed using prescription algorithms written in programming languages, using theoretical experts’ models. This method allows the calculation of complex molecular properties with results that correlate significantly with laboratory experiments [4].

Drugs usually consist of complex molecules. Research and development of new drugs in the laboratory experiment are needed many times and high costs. Also, the results obtained are likely to be unsatisfactory, so that a series of laboratory work and experience becomes wasted and useless. Application of computational chemistry can be crucial in medicinal chemistry, mostly for drug design, chemical properties theory, and biological activity of a molecule [5].

This research attempts to design the new meisoindigo derivative compound structure as anti-breast cancer through the QSAR approach. Cancer is an abnormal growth of cells in the body’s tissues, which
gradually become malignant. These cells grow faster and cannot experience apoptosis. Based on data from GLOBOCAN, the International Agency for Research on Cancer (IARC), it was reported that in 2012, there were around 14 million new cases of cancer and 8.2 million deaths from cancer. The most cancer cases causing death were lung cancer (19.7%), breast cancer (12.9%), liver cancer (9.5%), and stomach cancer (8.9%). The number of cancer patients is expected to increase every year and is estimated to reach 23.6 million new cases per year by 2030 [6].

The type of cancer that most causes death in women, especially those aged 40 years and over, is breast cancer. Most cancers attack the left breast at the top near the arm. Based on data from the Indonesian Ministry of Health’s Data and Information Center (2015), the prevalence of breast cancer in West Nusa Tenggara Province was 0.2% [6].

One of the potential compounds for anticancer is the meisoindigo derivative compound. The in vitro test for breast cancer cells shows that meisoindigo derivative compounds have good activity against these cancer cells. Meisoindigo derivative compounds will bind to the Cyclin–Dependent Kinase 4 (CDK4) enzyme, which plays a role in the cell cycle, thereby preventing cell division [7].

A similar study was conducted by Lestari et al. [8]. This study used the PM3 semiempirical method. The conclusion obtained from this research is successfully composing the best QSAR equation model. Based on these equations, Lestari et al succeeded in finding the compound (E)-2- (1- ((3-ethylisoxazole-5-yl) methyl) -2-oxoindolin-3- ylidene) –N- (4-methoxyphenyl) acetamide with the value of IC50 5.31144 x 10^-5 (µM).

2. Methodology

2.1. Tool and Material

The research material used was meisoindigo derivative compound data, which amounted to 20, obtained from the literature that had inhibitory activity on CDK4. The structure of the parent compound of meisoindigo derivatives can be seen in Figure 1. Data on compounds with biological activities divided into two-part first is training set data and second is test set data. The data meisoindigo derivative compound can be seen in Table 1.

![Figure 1. The parent compound of meisoindigo derivative](image)

This study was designed by a computer device with specification Pentium core i5 6600 processor, 2 GB RAM, 1 TB Hard Drive. While the software is Hyperchem 8.0 for Windows to build a 3D model and SPSS 16.0 software for analysis of the QSAR equation.

| Compounds | R1 | R2 | R3 | Log1/IC50 |
|-----------|----|----|----|-----------|
| 1* H      | p-Cl-Ph | p-pyridinyl | 4.853 |
| 2 H       | p-Cl-Ph | 6-quinolinyl | 5.455 |
| 3 H       | 3-methylisoxazolyl | p-MeO-Ph | 5.397 |
| 4 H       | 3-methylisoxazolyl | p-pyridinyl | 5.443 |
| 5* H      | 3-methylisoxazolyl | 6-quinolinyl | 5.494 |
| 6 MeO     | 3-methylisoxazolyl | 6-quinolinyl | 5.638 |
| 7 H       | 2-naphthalenyl | 6-quinolinyl | 5.327 |
| 8 H       | p-CF3-Ph | 6-quinolinyl | 5.823 |
| 9 H       | p-Br-Ph | 6-quinolinyl | 5.602 |
| 10 H      | p-NO2-Ph | 6-quinolinyl | 5.619 |
| 11* Me    | p-Cl-Ph | 6-quinolinyl | 5.207 |
| 12 H      | 2,6-CI2-Ph | 6-quinolinyl | 5.387 |
| 13 H      | 3-methylisoxazolyl | 3-methylisoxazolyl | 5.408 |
| 14 H      | 3-methylisoxazolyl | m-pyridinyl | 4.872 |
| 15 MeO    | p-NH2-Ph | 6-quinolinyl | 4.554 |
| 16 MeO    | p-Br-Ph | 6-quinolinyl | 5.508 |
| 17 OH     | p-Br-Ph | 6-quinolinyl | 5.455 |
| 18 H      | H | Phenyl | 4.071 |
| 19* H     | H | 6-quinolinyl | 4.349 |
| 20 H      | Me | 6-quinolinyl | 4.191 |

Compounds with a sign (*) are compounds that are used as a test set.

2.2. Research Procedure

Molecular Modeling. Modeling the meisoindigo derivative compound's molecular structure was carried out using two parameters (PM3 and AM1 semiempirical method) with a Convergence Limit of 0.001 kcal/Å. Optimization was carried out based on the Polak-Ribiere algorithm to achieve the minimum energy with Root Mean Square (RMS) of 0.001 kcal/Å [9]. Descriptors that will be counted represent three main parameters, including hydrophobic, electronic, and steric parameters. The materials used were 20 types of meisoindigo derivative compounds obtained from the study [7], known to have inhibitory activity on CDK4. The structure of the parent compound from the meisoindigo derivative can be seen in Figure 1. Furthermore, the data of compounds with biological activities can be seen in Table 1.

QSAR Equation Analysis. QSAR equation analysis was performed by statistical analysis using SPSS with the backward and entered the combination method. This is done by entering all the variables and including a few selected independent variables (data descriptors). Several equation models are produced: the relationship between the physicochemical properties of the
independent variable and the anticancer activity (Log 1/IC50), which is the dependent variable. The models are then tested for validity [10]. The chosen QSAR model must meet all statistical criteria used and had specific descriptors representing hydrophobic, electronic, and steric parameters. Validity testing of several equation models was done by calculating the value of $r$ (correlation coefficient) and $r^2$ (coefficient of determination) close to 1, adjusted $r^2$ with the most significant value, smallest SE (standard error), $F_{count}/F_{table} > 1$, and PRESS (predicted residual sums of squares) with the smallest value [11].

**New Compound Design.** The design of the new compound was done by modifying the position and type of the substituent. The position of the substituent was focused on the active site of the compound. The active site was chosen because it has atoms responsible for anti-breast cancer activity from meisoindigo derivative compounds. The next step was to perform calculations using the best semiempirical method according to the method used in determining the QSAR compound against all new compounds. Theoretical anti-breast cancer activity, Log 1/IC50 from the yield compound, was calculated using the QSAR equation that had been selected. Compounds with a high Log 1/IC50 value indicate that these compounds have high anti-breast cancer activity, and these compounds can be proposed for synthesis [12].

The method used in this study is the semiempirical method. Researchers used two semiempirical methods (AM1 and PM3). The best method will be used as a basis for the design of a new compound. The selection of the best method is based on the effectiveness of using descriptors and statistical parameter aspects.}

**3. Results and Discussion**

The method commonly used to model organic compounds is using semiempirical methods, especially AM1 and PM3 methods. The meisoindigo compound is also an organic compound, so in this research also used the same method. One of the several semiempirical methods certainly has results that are closer to the actual molecule compared to other methods. Therefore, it is necessary to compare the results of molecular modeling between the two semiempirical methods.

The parent compounds used in this study were meisoindigo derivatives (Figure 1), which were optimized using a previously validated method (semiempirical PM3 and AM1 methods). These compounds are then described as model sticks equipped with an atomic net charge for each atom, as shown in Figure 2.

![Figure 2. The optimization results of meisoindigo compounds with model sticks are equipped with an atomic net charge, (a) using the AM1 method, (b) using the PM3 method](image)

The biological activity obtained from the literature [6] is the minimum inhibitory value or IC50. However, in this study, the researcher changed the IC50 data to form a logarithmic value assuming to facilitate data analysis, so that the distribution is not too far away. The logarithmic form used is Log 1/IC50, it is mean that the higher the log 1/IC50 value, the compound has better activity than compound whose lower of Log 1/IC50.

Descriptors used in this study are descriptors representing electronic parameters, hydrophobic, and steric. The electronic parameter consists of the atomic net charge (q), isolated atomic energy (Eat.is), dipole moment (μ), hydration energy (EH), binding energy (Eb). Descriptors representing hydrophobic parameters are the partition coefficient of n-octanol/water (Log P); meanwhile, descriptors representing steric parameters consist of the gradient (Gr) and surface area approx (SA). The atoms’ net charge is atoms C6, C8, C9, C10, N11, C13, O14, and C15.

The reason to choose of atomic net charge descriptor (q) is carried out considering that the atomic charge is critical in determining chemical reactions and physicochemical properties of a compound, and it is useful for measuring intermolecular interactions. The net charge of an atom can be a positive or negative value. The net charge of the atom depends on the groups that are bound to the atom. Positive atomic charges are caused by electron pulling groups’ presence so that the electron density becomes smaller. As for charges that have negative values due to methyl groups, alkyl groups, and halides, which are electron donor groups, the electron density becomes greater.

From the QSAR equation models in table 2 and table 3, the best candidate QSAR equation model can be chosen. The selection of the best QSAR equation model is made by calculating statistical parameters such as the value of $r^2$ (correlation coefficient), adjusted $r^2$, SD (Standard Deviation), $F_{count}/F_{table}$, and PRESS (Predicted Residual Sum of Square).
The selection of the best models, when viewed from the value of r and r², all models have values above 0.8. This means that all models have the requirements as a good model. The values of r and r² have the meaning of the strength of the correlation between the independent and dependent variables and how well the equation model’s linearity is to predict another equation model. Another parameter that can be used to determine the best model is the value of F_count/Table. The higher value of F_count/Table means that the level of significance is getting better. Another statistical parameter is the PRESS (Predicted Residual Sum of Square) value. A small PRESS value means that the model is better to be used as the best model. Based on table 2 and table 3, the AM1 method has the same good parameter value as the PM3 semiempirical method.

The QSAR equation formula can be obtained from various multivariate statistical methods that produce satisfactory results. The most widely used primary method is regression analysis. The method correlates several independent variables x (in the form of physical-chemical parameters in the Hansch method or the value of indicator variables in the Free-Wilson method) with non-independent or bound variables y (in the form of biological activity parameters of the compound). The AM1 method validation test was conducted by predicting breast anticancer activity using a model chosen according to the statistical parameters in table 2 (models 4, 6, 7, and model 9), then making a curve of the relationship between log 1/IC₅₀ experiments with log 1/IC₅₀ predictions. Table 4 shows the results of a comparison between the proposed models.

Table 2. QSAR equation model results from multilinear regression analysis with electronic, steric, and hydrophobic parameters using the AM1 method

| Model | Variable | R     | r²   | adjusted r² | SE   | F    | F_count/Table | PRESS |
|-------|----------|-------|------|-------------|------|------|---------------|--------|
| 1     | qC13, qC8, qC9, qO10, qN11, μ, Gr, Eat.is, Log P, EH, Eb, qC9, qO10 | 0.998 | 0.995 | 0.997 | 0.804 | 53.033 | 6.064 | 1.177 |
| 2     | qC13, qC8, μ, Gr, Eat.is, qN11, Log P, EH, Eb, qC9, qO10 | 0.995 | 0.99 | 0.962 | 0.1018 | 35.880 | 6.0441 | 128.6408 |
| 3     | qC13, qC8, μ, Eat.is, Log P, qN11, Eb, EH, qC9, qO10 | 0.969 | 0.938 | 0.815 | 0.1155 | 7.6296 | 1.611 | 371.487 |
| 4     | qC13, qC8, μ, Eat.is, Log P, qN11, EH, qC9, qO10 | 0.962 | 0.926 | 0.816 | 0.115 | 8.391 | 2.047 | 0.0005 |
| 5     | qC13, qC8, μ, Eat.is, Log P, qN11, qC9, qO10 | 0.959 | 0.919 | 0.827 | 0.218 | 9.969 | 2.675 | 0.002 |
| 6     | qC13, μ, Eat.is, qC9, Log P, qN11, qO10 | 0.938 | 0.88 | 0.774 | 0.249 | 8.346 | 2.384 | 0.0006 |
| 7     | qC13, μ, Eat.is, qC9, qO10 | 0.935 | 0.874 | 0.811 | 0.228 | 13.895 | 4.178 | 0.0001 |
| 8     | qC15, qC9, qO10, Log P, Eat.is, μ, qC13, EH, qN11 | 0.939 | 0.881 | 0.703 | 0.286 | 4.946 | 1.206 | 0.006 |
| 9     | qC15, qO14, qC6, qO10, Log P, Eat.is, μ, EH, qC9, qC13, qN11 | 0.955 | 0.913 | 0.672 | 0.301 | 3.792 | 0.638 | 0.00004 |

Table 3. The QSAR equation model results from multilinear regression analysis with electronic, steric, and hydrophobic parameters using the PM3 method

| Model | Variable | R     | r²   | adjusted r² | SE   | F    | F_count/Table | PRESS |
|-------|----------|-------|------|-------------|------|------|---------------|--------|
| 1     | qC13, Gr, qN11, Eat.is, μ, qC8, Log P, EH, Eb, qC9, qO10 | 0.925 | 0.856 | 0.28 | 0.445 | 1.485 | 0.169 | 2188.854 |
| 2     | qC13, qN11, Eat.is, μ, qC8, Log P, EH, Eb, qC9, qO10 | 0.923 | 0.852 | 0.446 | 0.391 | 2.098 | 0.353 | 655.080 |
| 3     | qC13, qN11, Eat.is, μ, qC8, Log P, EH, Eb, qC9, qO10 | 0.92 | 0.847 | 0.541 | 0.356 | 2.767 | 0.584 | 486.532 |
| 4     | EH, μ, qN11, Eat.is, qO10, qC9, qC15, Log P, qC13 | 0.927 | 0.859 | 0.647 | 0.311 | 4.054 | 0.989 | 0.0004 |
| 5     | qC6, qO14, qN11, μ, Eat.is, Log P, qC15, EH, qC9, qC13, qO10 | 0.968 | 0.937 | 0.764 | 0.254 | 5.420 | 0.913 | 0.004 |

Figure 3 shows the relationship between log 1/IC₅₀ experiments with log 1/IC₅₀ predictions for models 4, 6, 7, and 9.

![Graph showing the relationship between log 1/IC₅₀ experiments with log 1/IC₅₀ predictions for models 4, 6, 7, and 9.](image-url)
Figure 4 shows a curve of the relationship between log \( 1/IC_{50} \) experiments with log \( 1/IC_{50} \) predictions for models 4 and 5.

Based on Figure 5, the researcher concluded that the best model using the PM3 method is model 5 with the following formula:

\[
\log 1/IC_{50} = 44.316 + (-0.0000282 * \text{Eat.is}) + (-0.257 * \mu) \\
+ (3.495 * qC9) + (3.729 * qO10) + (0.862 * qC13)
\]

The best model obtained for the AM1 method uses isolated atomic energy descriptors, dipole moments, and atomic charges for C9, O10, and C13 atoms. The model does not involve oil/water partition coefficients and only involves three atomic charges. The partition coefficient, abbreviated P, is defined as the ratio of certain solute concentrations between the two solvents (dual-phase liquid), especially for non-ionized solutes. If one of the solvents is water and the other is a non-polar solvent, then the log P-value is the size of the lipophilicity or hydrophobicity. For this reason, theoretically, the log P parameter must be included in the calculation of the best model. Based on this reason, researchers conclude that the PM3 semiempirical method was chosen as the best model. On the other hand, this method is suitable for many organic molecules, and this method is also proven to be able to optimize the carbon structure [13].

Based on those reasons, the PM3 semiempirical method was chosen to be the best method. Model 5 with the PM3 method was used as a guiding model in designing and predicting new breast anticancer activities from melsoindigo derivative compounds. In designing
new compounds, selecting the main compound structure is based on the meisoindigo derivative compound, which has the best breast anticancer activity. The design of the compound begins by modifying the type and position of the substituent.

Descriptors of new compounds obtained from these calculations are then included in the best QSR equation model (model 5 PM3 Method). In designing a new compound, the parent compound's structure selection is based on the meisoindigo derivative compound, which has the best inhibitory activity. New meisoindigo derivative compounds that have been theoretically designed have relatively good inhibition values with insecticidal activity values from 5.120 to 6.992. The result of the calculation can be seen in table 6. The higher log 1/IC50 value of a compound, or the smaller IC50 value, means the compound’s activity is getting better. From several new compounds that have been designed, compounds with log 1/IC50 values can be selected, which are higher than the price of the synthesized log 1/IC50 compound. Based on this, the researchers concluded that the best compounds with higher activity than previous experiments were compounds (E) – 2- (2-oxo-1-(2- (trifluoromethyl) benzyl) indolin-3-ylidene) –N-(quinolin-7-yl)acetamide, the best compound image shown in Figure 5. The proposed compound has a log 1/IC50 value of 6.992 more than log 1/IC50 before.

![Figure 5. Compound (E)-2-(2-oxo-1-(2-(trifluoromethyl)benzyl)indolin-3-ylidene)-N-(quinolin-7-yl)acetamide](image-url)

**Table 6. Design of new compound of meisoindigo derivative**

| No. | Name                                                                 | Log 1/IC50 |
|-----|----------------------------------------------------------------------|------------|
| 21  | (E)-2-(1-(3-chlorobenzyl)-2-oxindolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 5.359      |
| 22  | (E)-2-(1-(2-chlorobenzyl)-2-oxindolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 5.452      |
| 23  | (E)-2-(1-(3-ethylisoaxo-5-yl)methyl)-2-oxindolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 5.619      |
| 24  | (E)-2-(1-(3-ethylisoaxo-5-yl)methyl)-2-oxindolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 5.306      |
| 25  | (E)-2-(1-(3-methylisoaxo-5-yl)methyl)-2-oxindolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 5.334      |
| 26  | (E)-2-(1-(3-ethylisoaxo-5-yl)methyl)-2-oxindolin-3-ylidene)-N-(quinolin-7-yl)acetamide | 5.426      |
| 27  | (E)-2-(1-(3-ethylisoaxo-5-yl)methyl)-5-methoxy-2-oxindolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 5.784      |
| 28  | (E)-2-(1-(naphthalen-1-yl)methyl)-2-oxindolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 5.377      |
| 29  | (E)-2-(2-oxo-1-(3-(trifluoromethyl)benzyl)indolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 6.067      |
| 30  | (E)-2-(2-oxo-1-(2-(trifluoromethyl)benzyl)indolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 6.455      |
| 31  | (E)-2-(2-oxo-1-(4-(trifluoromethyl)benzyl)indolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 6.562      |
| 32  | (E)-2-(2-oxo-1-(3-(trifluoromethyl)benzyl)indolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 6.262      |
| 33  | (E)-2-(2-oxo-1-(2-(trifluoromethyl)benzyl)indolin-3-ylidene)-N-(quinolin-7-yl)acetamide | 6.992      |
| 34  | (E)-2-(1-(3-bromobenzyl)-2-oxindolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 5.438      |
| 35  | (E)-2-(1-(2-bromobenzyl)-2-oxindolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 5.737      |
| 36  | (E)-2-(1-(3-nitrobenzyl)-2-oxindolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 5.676      |
| 37  | (E)-2-(1-(2-nitrobenzyl)-2-oxindolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 6.079      |
| 38  | (E)-2-(1-(2,5-dichlorobenzyl)-2-oxindolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 5.47       |
| 39  | (E)-2-(1-(2,6-dichlorobenzyl)-2-oxindolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 5.536      |
| 40  | (E)-2-(1-(3-bromobenzyl)-5-methoxy-2-oxindolin-3-ylidene)-N-(quinolin-6-yl)acetamide | 5.12       |
4. Conclusion

The best QSAR equation model of meisoindigo derivative as antitumor breast cancer using semiempirical PM3 method is as follows:

\[
\log I_{50} = 44.316 + (-0.0000282 \times \text{Eat.is}) + (-0.257 \times \mu) \\
+ (-0.054 \times \log P) + (0.014 \times EH) + (-7.441 \times qC6) + (-1.734 \times qC9) + (25.711 \times qO10) + (7.309 \times qN11) + (94.825 \times qC13) \\
+ (58.794 \times qO4) + (5.866 \times qC15)
\]

\( n = 20; r = 0.927; \text{adjusted } r^2 = 0.647; \text{SE} = 0.311; \)

\( \text{Fhit/Feasle} = 4.05; \text{PRESS}=0.00047 \)

Based on that model, new compounds which have predictive inhibition activity are better than existing ones is (E) -2- (2-oxo-1- (2- (trifluoromethyl) benzyl) indolin-3-ylidene) -N- (quolinin-7-yl) acetamide with log \(I_{50}\) prediction = 6.992

Acknowledgment

This work was supported by DIPA BLU (PNBP) Universitas Mataram with contract No. 1375/S/UN18.1/Li/PP/2018.

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