Comparing Generalized Linear Model of Count Data: Application towards Pteridophyta species

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Abstract

Pteridophyta is known as “paku-pakis” in Malay and it is one the flora species that exists in ecological system. Besides, Pteridophyta is one of the species that need to be preserved. Either flora or fauna, both are very important to preserve the ecosystem and control the pollution. In order to observe the species, the fundamental unit of all diversity metrics is a count of specific individuals. In some consequences, the uncorrected counts of observed species often used in measuring the diversity which ignore detection entirely and established methods to be used to account for missed species. Analysis of count data is widely used in engineering, public health, epidemiology, medical studies, ecology and many research of interest. Rarity increases the number of locations with zero detection in excess of those expected under simple models of abundance. The aims of this study are to compare the Generalized Linear Model (GLiM) in the application of Pteridophyta species of count data.

Keywords: Poisson model; Zero-Inflated Poisson; Negative Binomial; Akaike Information Criterion (AIC).

1. Introduction

Conservation of floral is important of floral is important in stabilizing the biodiversity of the earth. Consequently, ecology comprises the relation of the organisms to their environment as the plants are useful in reviewing the pattern of biotic response to their climate change and they are abundant and sensitive to the climate variables which are the temperature, seasonality and rainfall [4]. Besides, diversity estimates are central to the community and macro ecology and are frequently used in conversation ecology. The fundamental unit all of the diversity metrics is a count of species, individuals or both. Attempt to define the imperfect detection, the predictable consequences occur when the species are rare, missed individual’s results in false absences. Hence, uncorrected counts of observed species often used in the measure of diversity ignore detection together and established methods used to account for missed species do not disentangle detection from occurrence. Ecological data are normally described as the count observation with their non-negative values. In recent research, assessed the implication count data, the analysis of count data is widely used in medical studies, epidemiology, ecology and many research of interest. The rarity increases the number of locations with zero detection in excess of those expected under a simple model of abundances such as the Poisson regression analysis or the negative binomial regression analysis. Subsequently, in ecology, normally the count data appears as overdispersed and a common approach to deal with it is by the generalized linear model framework [3]. Besides, environmental factors or habitat condition are favorable to their species of interest.

Besides, policy statements of Malaysia’s National Policy on Biological Diversity said: “To conserve Malaysia’s biological diversity and to ensure that its components are utilized in a sustainable manner for the continued progress and socio-economic development of the nation”. Through that statement, as Malaysia covered by tropical floral, the humid tropics attempted to provide a climate that is suitable to support the rich and diverse life forms. Hence, it could sustain the population of flora and fauna on the natural ecological habitats.

The studies of floral kingdom include the non-seed bearing plants which indicate the lower plants and also the seed bearing plants which comprised the higher plants. Focused on the non-seed bearing plants, it covered on the algae, mosses and fern. Instead of that, the seed bearing plants consists of the angiosperms (flowering plants) and the gymnosperm (cone-bearing plants). Explored on the non-seed bearing plants, botanically, the fern is classified as Pteridophyta and they are common as it is edible to be eaten raw, cooked, preserved in brine or pickled. The ferns do not produce through seed but they produced by the spores. Besides, Pteridophyta has true roots and leaves with a short stem and sometimes called as rhizomes. Based on the previous study by [1], the classification of living ferns comprised four classes of order which is 11 orders and 37 families with living representatives of 11,500 species worldwide. Consequently, there are 1,165 species of pteridophytes in Malaysia; 647 species occur in Peninsular Malaysia and 750 in Sabah while 615 in Sarawak [2].

The generalized linear model has been used in analyzing the count observation as several standard statistical analyses of parametric models for non-normally distributed data covers the Poisson regression model, negative binomial regression model, zero-inflated models, and hurdle models [3]. These models have the power of parametric model and more flexible in handling repeated measures, multiple covariates and various structures of fixed and random effects as it has been assumed difference than the normal distribution. Poisson distribution defines as the number of events that occur in a fixed period of time and as the mean count increases, the distribution could be approximately normal. According to [8], to assess the relationship between the abundance of a species in...
for Social Science (SPSS) version 20 and Microsoft Excel 2010. The GLIM was applied as the dependent variable involved count data of discrete variables. In order to make the study more convincing, the relevant statistical analysis procedure has been included, so that the hypotheses could be achieved.

The logarithm of the response variable was linked to a linear function of explanatory variable such that:

$$\ln(\mu_i) = \beta' x_i$$ or $$\mu_i = \exp(\beta' x_i)$$

On the other hand, a standard Poisson regression model expressed the log outcome rate as a linear function of a set predictors.

2.2. Overdispersion checking

To further illuminate the Poisson regression analysis, the most important part was detecting the presence of overdispersion. If the equality of mean and variances was violated, an overdispersion feature of the problem could occur. There were three conditions which could describe the key of overdispersed model. The first condition indicates, if deviance/df > 1, then, overdispersion might be present. The second condition was, if deviance/df < 1, then underdispersion might occur. Lastly, $$\chi^2$$ with degrees of freedom equal to g (has a mean of g). Benefit for using the maximum likelihood method, the simple likelihood ratio test would be used to assess the adequacy of the negative binomial over the Poisson regression.

In order to overwhelm the problem of overdispersion, the generalized of Poisson which was the negative binomial regression model has been suggested by [10]. Therefore, the value of deviance which was used to calculate the overdispersion could be described as:

$$Deviance (D) = 2 \left[ \Sigma (y \ln \left( \frac{y}{\hat{y}} \right) - (y - \hat{y})) \right]$$

Besides, in ordinary least square regression analysis, similarly, the coefficient of determination, $$R^2$$, has been defining as the deviance in analyzing count data. This $$R^2$$ value was used to provide the descriptive information about the model fit.

$$R^2 = \frac{\Sigma (\hat{y} - \bar{y})^2}{\Sigma (y - \bar{y})^2}$$

where the observed value was y, the predicted value was $$\hat{y}$$ with the mean value of y. In addition, the model adequacy checking compares a fitted model to a saturated model. The difference between log-likelihood under two model:

$$D = 2 \ln \lambda = 2[\{l(b \text{ max}; y) - l(b; y)\}$$

where $$\{l(b \text{ max}; y)$$ the log likelihood function for the maximum model and $$\{l(b; y)$$ is the log likelihood function for testing the model. Thus, the hypothesis of the test is:

$$H_0:$$ The model is a good fit

$$H_1:$$ The model is not a good fit

The deviance based on $$\chi^2$$ with n-p degree of freedom. The null hypothesis is rejected if deviance, $$D > \chi^2_{n-p}$$. vice versa.

2.3. Negative Binomial regression analysis

Previously mentioned, if overdispersion exists, there is a possibility for the analysis to use negative binomial regression so that, the results could fit the data very well. Based on [6], if overdispersion exists, there are two properties normally associated
with the presence of overdispersion which is about the contagion and through excess zero. “Contagion” could be defined when the individuals are more grouped than expected as they happened individually while the excess zero is when the total observations happened to have more zero than what they would expect. In spite of that, both contagion and excess zeros may increase the variance relative to the mean which then contributes to the overdispersion. As overdispersion increased, in the statistical analysis of ecology, the result produced would be more accurate [5, 9]. In addition, negative binomial has been known as the generalized of Poisson regression as the assumption of Poisson which is the variance is not equal to the mean. The calculated value of negative binomial is:

$$\mu_i = \exp(X_i' \beta_i + \epsilon_i) = \exp(X_i' \beta_i) \exp(\epsilon_i)$$

where $$\exp(\epsilon_i) \sim \text{Gamma}(\alpha^{-1}, \alpha^{-1})$$ and the function of the density can be derived as:

$$f(y | X, \beta) = \frac{\Gamma(y_i + \alpha^{-1})}{\Gamma(y_i + 1) \Gamma(\alpha^{-1})} \left( \frac{\alpha^{-1}}{\alpha^{-1} + \mu} \right)^{y_i} \left( \frac{\mu}{\alpha^{-1} + \mu} \right)^{\alpha^{-1}}$$

where $$\Gamma$$ represent the gamma integral which specializes to a factorial integer argument. Besides, the estimation of the parameters is done by maximizing the log likelihood function;

$$\ln L(\beta) = \sum_{i=1}^{n} \left[ y_i \ln(\beta)' - \exp(\beta'X_i) - \ln y_i ! \right]$$

Therefore, the Maximum Likelihood Estimator is done by the numerical method generated using the computer-based iteration procedure in GenStat [10].

### 2.4. AIC of regression model

The smallest AIC values can be used to identify the appropriate or suitable method in modeling the observations of counts data. AIC was considered as GLIM models was part of nested models. Otherwise, the analysis must use Vuong test to compare the non-nested models such as the Poisson Hurdle versus negative binomial regression model.

### 2.5. Incidence rate ratio (IRR) value

Incidence rate ratio was another way in explaining the negative binomial regression model. From the final model which was the negative binomial, IRR indicates some explanation in statistical analysis which displays the best-selected variables and could accurately estimate the numbers of Pteridophyta grows that includes:

- IRR = 1 (numerator group and denominator group have same incidence rate)
- IRR > 1 (numerator group has higher incidence rate than denominator group)
- IRR < 1 (numerator group has a lower incidence rate than denominator group)

### 3. Results and Analysis

By using SAS statistical software, this study performed the Poisson regression analysis, followed by zero-inflated Poisson regression analysis and negative binomial regression analysis. These models are part of GLIM, which is used to analyse the count data as observation may indicate high zero values and having a non-normal observation. In order to analyse the count data, this study used PROC GENMOD, PROC UNIVARIATE and PROC IMPORT.

#### 3.1. Poisson Regression model

Based on Table 1, the results showed the summarization of the estimation score obtained by the Poisson regression model. Starting from the 95% of confidence limits, standard error and the P-value, the variable of Types (types of Pteridophyta species) and the variable of years (duration of Pteridophyta species grows) have been summarized as follow.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Estimate (95% CI)</th>
<th>Std. Error</th>
<th>P-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>4.8727 (4.7681, 4.9974)</td>
<td>0.0534</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Types: ALF</td>
<td>-0.0202 (-0.1595, 0.1191)</td>
<td>0.0711</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>Types: DDF</td>
<td>1.2669 (1.1560, 1.3779)</td>
<td>0.0566</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Types: GPF</td>
<td>0.6806 (0.5614, 0.5603)</td>
<td>0.0614</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Types: NBF</td>
<td>1.6606 (1.5537, 1.7675)</td>
<td>0.0545</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Types: VEF</td>
<td>0.0000 (0.0000, 0.0000)</td>
<td>0.0707</td>
<td>1.0000</td>
</tr>
<tr>
<td>Years: 4</td>
<td>-2.1894 (-2.3304, -2.0485)</td>
<td>0.0719</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Years: 8</td>
<td>-0.5082 (-0.5812, -0.4352)</td>
<td>0.0372</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Years: 12</td>
<td>0.2982 (0.2931, 0.3572)</td>
<td>0.0301</td>
<td>&gt;0.0001</td>
</tr>
<tr>
<td>Years: 20</td>
<td>0.0000 (0.0000, 0.0000)</td>
<td>0.0800</td>
<td>&gt;0.0001</td>
</tr>
</tbody>
</table>

Based on Table 1, when the probability values indicate less than α = 0.05, the study derived that the variable significantly affected the total numbers of Pteridophyta in that area. For the P-value of types of Pteridophyta, DDF, GPF and NDF were < 0.0001 while 4 years, 8 years and 12 years showed P-value < 0.0001. Hence, the variables significantly affected the total numbers of Pteridophyta in that area. However, the ALF, VEF, SFF have significantly not affected the species to grow.

$$\ln(\gamma) = 4.8727 -0.0202X1 + 1.2669X2 +0.6806X3 +1.6606X4 +0.0780X5 -2.1894X6 -0.5082X7 +0.2982X9 +0.2982X10$$

Therefore, the equation of Poisson regression model could be summarized as below.

$$\ln(\gamma) = 4.8727 + 1.2669DF + 0.6806GPF + 1.6606NBF - 2.1894Y4 + 0.0780Y5 - 2.1894Y6 + 0.2982Y7 + 0.2982Y8 + 0.2982Y9 + 0.2982Y10$$

The SAS procedure could be used as the followed.

**PROC GENMOD DATA=FIQ.PTERIDOPHYTA;**
**CLASS TYPES YEARS PARAM=GLM;**
**MODEL NUMBERS = TYPES YEARS/TYPE3**
**DIST=POISSON;**
**RUN;**

### 3.2. Goodness of fit Poisson regression model

By referring to the Table 2, since the P-value was 0.000 less that than α with the value of 0.05, the study rejects null hypothesis indicating that the Poisson regression model does not fit the data reasonably well.

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Deviance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chi sq</td>
<td>948.3119</td>
</tr>
<tr>
<td>P-value</td>
<td>0.000</td>
</tr>
</tbody>
</table>

The SAS procedure could be used as the followed.

**DATA FIQ.PTERIDOPHYTA;**
**DF=18; CHISQ=948.3119;**
**IPVALUE=1-PROBCHI(CHISQ,DF);**
**RUN;**
**PROC PRINT DATA=FIQ.PTERIDOPHYTA;**
**RUN;**
Since P-value < 0.05, the model does not fit the data. This study checks on the presence of overdispersion and excess zero before proceeding with the other models of GLiM.

3.3. Checking on the overdispersion

<table>
<thead>
<tr>
<th>Table 3: Overdispersion by deviance model</th>
</tr>
</thead>
<tbody>
<tr>
<td>Criterion</td>
</tr>
<tr>
<td>Value</td>
</tr>
<tr>
<td>Value/DF</td>
</tr>
</tbody>
</table>

Based on Table 3, overdispersed by deviance, the value of deviance was 948.3119 and the average value of deviance was 52.6840. Therefore, as the P-value equals to 52.6840 previously, which was greater than 1. This indicates that overdispersion exists. Checking the overdispersion was compulsory as it was one of the model adequacies checking for the GLiM. Besides, another method in checking the overdispersion was by comparing the mean and value of variance. If the value of variance was greater than the mean, thus, overdispersion exists.

3.4. Checking excess zero observation

By using the following SAS procedure;

```
PROC UNIVARIATE DATA = FIQ.PTERIDOPHYTA NOPRINT;
HISTOGRAM NUMBERS / MIDPOINTS = 0 TO 50 BY 10;
VSABLE = COUNT;
RUN;
```

Fig. 1: Excess zero values

Fig. 1 shows that zero value exists with four times of the observations. Since the zero values are not too many, this study needs to perform both zero-inflated Poisson and negative binomial to get accurate results for Pteridophyta species.

3.5. Zero-Inflated Poisson regression model

In order to perform zero-inflated Poisson regression model, this study uses the following SAS codes.

```
PROC GENMOD DATA=FIQ.PTERIDOPHYTA;
CLASS TYPES YEARS;
MODEL NUMBERS = TYPES YEARS/ DIST=ZIP;
ZERO MODEL NUMBERS /LINK=LOGIT;
RUN;
```

Table 4 shows the summarization of the estimation score obtained by the zero-inflated Poisson regression model.

### Table 4: Parameter estimate of zero-inflated Poisson regression model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Estimate (95% CI)</th>
<th>Std. Error</th>
<th>P-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>4.9100 (4.8035, 5.0145)</td>
<td>0.0333</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Types: ALF</td>
<td>-0.0202 (-0.1595, 0.1191)</td>
<td>0.0711</td>
<td>&gt;0.5000</td>
</tr>
<tr>
<td>Types: DDF</td>
<td>1.2158 (1.1047, 1.3270)</td>
<td>0.0567</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Types: GPF</td>
<td>0.6295 (0.5090, 0.7499)</td>
<td>0.0615</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Types: NBF</td>
<td>1.6065 (1.5024, 1.7166)</td>
<td>0.0547</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Types: SPF</td>
<td>-0.0779 (-0.2193, 0.0635)</td>
<td>0.0721</td>
<td>&gt;0.2000</td>
</tr>
<tr>
<td>Types: VEF</td>
<td>0.0000 (-0.1386, 0.1386)</td>
<td>0.0707</td>
<td>&gt;0.5000</td>
</tr>
<tr>
<td>Types: VLF</td>
<td>0.0000 (0.0000, 0.0000)</td>
<td>0.0000</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

Based on the following Table 5, variables DDF, GPF, NBF, 4 years, 8 years and 12 years are significant towards the observation of Pteridophyta.

Before that, by performing the goodness of fit, this study found that zero-inflated Poisson does not fit the data of Pteridophyta.

```
DATA FIQ.PTERIDOPHYTA; DDF=16; CHISQ=572.5962; PVALUE=1-PROBCHI(CHISQ,DF); RUN;
```

```
PROC GENMOD DATA=FIQ.PTERIDOPHYTA;
CLASS TYPES YEARS /
MODEL NUMBERS = TYPES YEARS/TYPE3 DIST=NB;
RUN;
```

### Table 5: Goodness of fit by zero-inflated Poisson model

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Deviance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Df</td>
<td>16</td>
</tr>
<tr>
<td>ChiSq</td>
<td>572.5962</td>
</tr>
<tr>
<td>P-value</td>
<td>0.0000</td>
</tr>
</tbody>
</table>

Since P-value < 0.05, the model does not fit the data. Then, this study proceeds with negative binomial GLiM.

3.6. Negative Binomial regression model

The SAS code below performs the negative binomial regression analysis.

```
PROC GENMOD DATA=FIQ.PTERIDOPHYTA;
CLASS TYPES YEARS /
PARAM=GLM;
MODEL NUMBERS = TYPES YEARS/TYPE3 DIST=NB;
RUN;
```

Based on Table 6, the results showed the summarization of the estimation score obtained by the negative binomial regression model.

### Table 6: Parameter estimate of Negative Binomial regression model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Estimate (95% CI)</th>
<th>Std. Error</th>
<th>P-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>4.7976 (4.0285, 5.5668)</td>
<td>0.3924</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Types: ALF</td>
<td>0.1207 (-0.7845, 0.1060)</td>
<td>0.4691</td>
<td>0.7938</td>
</tr>
<tr>
<td>Types: DDF</td>
<td>1.8649 (0.9204, 2.8094)</td>
<td>0.4819</td>
<td>0.0001</td>
</tr>
<tr>
<td>Types: GPF</td>
<td>0.0861 (0.1597, 0.20124)</td>
<td>0.4726</td>
<td>0.0216</td>
</tr>
<tr>
<td>Types: NBF</td>
<td>2.6087 (1.6442, 3.5732)</td>
<td>0.4921</td>
<td>&gt;0.0001</td>
</tr>
<tr>
<td>Types: SPF</td>
<td>0.1545 (-0.7751, 1.0841)</td>
<td>0.4743</td>
<td>0.7446</td>
</tr>
<tr>
<td>Types: VEF</td>
<td>0.0670 (-0.8449, 0.9788)</td>
<td>0.4652</td>
<td>0.8856</td>
</tr>
<tr>
<td>Types: VLF</td>
<td>0.0000 (0.0000, 0.0000)</td>
<td>0.0000</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Years: 4</td>
<td>-3.3066 (-4.0972, -2.5159)</td>
<td>0.4034</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Years: 8</td>
<td>-0.9391 (-1.6179, -0.2603)</td>
<td>0.3463</td>
<td>0.0067</td>
</tr>
<tr>
<td>Years: 12</td>
<td>0.0235 (-0.6628, 0.7098)</td>
<td>0.3502</td>
<td>0.9465</td>
</tr>
</tbody>
</table>

When the probability values indicate less than α = 0.05, the study derived that the variable significantly affected the total numbers of Pteridophyta in that time. For the P-value of types of Pteridophyta, DDF was 0.0001 less than 0.05 and NBF were less than 0.0001. While ALF equal to 0.7938, GPF equal to 0.0216, SPF was 0.7446 and VEF was 0.8856 were all greater than α, 0.05. Besides, 4 years has shown P-value less than 0.0001. The 8 years equal to 0.0067 less than 0.05, while for 12 years 0.9465 greater than 0.05.
Hence, those probability values were significant in predicting the total numbers of Pteridophyta if the variables were less than alpha value, 0.05. Overall, variables that were not significant comprised the ALF, SPF, VEF and 20 years. Besides, by checking the goodness of fit of negative binomial model.

**DATA** FIQ.PTERIDOPHYTA; DF=18; CHISQ=25.1724; PVAlue=1-PROBCHI(CHISQ,DF); RUN;

**PROC PRINT DATA=FIQ.PTERIDOPHYTA; RUN;**

| Table 7: Goodness of fit by negative binomial model |
|---------------------|---------------------|
| Criterion          | Deviance            |
| DF                 | 18                  |
| Chisq              | 25.1724             |
| P-value            | 0.12024             |

Based on Table 7, since P-value = 0.12024 > 0.05, the model does fit the data. Then, this study proved that *Pteridophyta* data fit by using the negative binomial GLIM. The formulated equation assessed by negative binomial regression model has been shown as below:

\[
\text{Numbers} = 121.2191 \exp (1.8649 \text{DDF} + 1.0861 \text{GPF} + 2.6087 \text{NBF} - 0.9391 \text{years}4 + 0.0235 \text{years}12)
\]

where \(\exp (\epsilon) \sim \Gamma(\alpha^{-1}, \alpha^{-1})\)

### 3.7. The Akaike Information Criterion (AIC)

**Table 8: AIC and BIC scores of both models**

<table>
<thead>
<tr>
<th>Models</th>
<th>Poisson Model</th>
<th>Zero-Inflated Poisson Model</th>
<th>Negative Binomial Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>AIC</td>
<td>1133.976</td>
<td>1010.2405</td>
<td>327.7790</td>
</tr>
</tbody>
</table>

Based on the results of goodness of fit previously by the negative binomial regression model and by observing the Akaike Information Criterion (AIC), the results obviously showed that negative binomial is the most suitable model that fit the *Pteridophyta* data. Based on the smallest values of the GLIM models, a negative binomial regression model fits with the smallest value of AIC and P-value greater than 0.05 (95% confidence interval).

### 3.8. Interpreting the incidence ratio rate (IRR) of Negative Binomial regression

**Table 9: Estimated IRR value based on negative binomial model**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Estimate</th>
<th>Standard Error</th>
<th>IRR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>4.7976</td>
<td>0.3924</td>
<td>121.223</td>
</tr>
<tr>
<td>Types ALF</td>
<td>0.1207</td>
<td>0.4691</td>
<td>1.128</td>
</tr>
<tr>
<td>Types DDF</td>
<td>1.8649</td>
<td>0.4819</td>
<td>6.355</td>
</tr>
<tr>
<td>Types GPF</td>
<td>1.0861</td>
<td>0.4726</td>
<td>2.963</td>
</tr>
<tr>
<td>Types NBF</td>
<td>2.6087</td>
<td>0.4921</td>
<td>13.581</td>
</tr>
<tr>
<td>Types SPF</td>
<td>0.1545</td>
<td>0.4743</td>
<td>1.167</td>
</tr>
<tr>
<td>Types VEF</td>
<td>0.06695</td>
<td>0.4652</td>
<td>1.069</td>
</tr>
<tr>
<td>Types VLF</td>
<td>0.0000</td>
<td>0.0000</td>
<td>1.000</td>
</tr>
<tr>
<td>Years 4</td>
<td>-3.3066</td>
<td>0.4034</td>
<td>0.037</td>
</tr>
<tr>
<td>Years 8</td>
<td>-0.9391</td>
<td>0.3463</td>
<td>0.391</td>
</tr>
<tr>
<td>Years 12</td>
<td>0.02349</td>
<td>0.3502</td>
<td>1.024</td>
</tr>
<tr>
<td>Years 20</td>
<td>0</td>
<td>0.0000</td>
<td>1.000</td>
</tr>
</tbody>
</table>

Based on Table 9, starting from ALF, the IRR = 1.128 (12.8%), the total number of Pteridophyta is expected to increase by 12.8% when involving ALF compared to VLF. Then, IRR = 6.455 (545.5%) for DDF, the total number of Pteridophyta is expected to increase by 545.5% when involving DDF compared to VLF. For GPF, IRR = 2.963 (196.3%) indicating that the total number of Pteridophyta is expected to increase by 196.3% when involving GPF compared to VLF. Besides, for NBF, the IRR = 13.581 (1258.1%) means that total number of Pteridophyta is expected to increase by 1258.1% when involving NBF compared to VLF. Besides, for SPF and VEF, the IRR values were 1.167 (16.7%) and 1.069 (6.9%) respectively. Thus, the total number of Pteridophyta is expected to be increased by 16.7% and 6.9% when involving SPF and VEF respectively compared to VLF. On the other hand, for the duration of plants grows, within 4 years, the IRR = 0.037 (-96.3%) which means that the total number of Pteridophyta is expected to be decreased by 96.3% when involving 4 years compared to 20 years. The IRR for 8 years was equal to 0.391 (-60.9%). That means the total number of Pteridophyta is expected to be decreased by 60.9% when involving 8 years compared to 20 years. For 12 years, the IRR = 1.024 (2.4%) indicates that total number of Pteridophyta is expected to be increased by 2.4% when involving 12 years compared to 20 years.

Hence, the results showed that all types of Pteridophyta seem to have influence in predicting the total numbers of Pteridophyta grows as all types showed increased values of IRR. While, the duration that is suitable for the Pteridophyta were within 12 years and 20 years.

### 4. Discussion

By approaching the objective of study consequently, the Poisson regression model and zero-inflated Poisson regression model does not fit the data really well compared to negative binomial regression analysis. Before that, by checking the model adequacy, the data showed the presence of overdispersion and this may lead the data to have not normally distributed due to the large extreme value of observations. In spite of that, the study proposed negative binomial regression analysis instead of Poisson regression model and zero-inflated Poisson. By proceeding the negative binomial, the study used AIC value and check the goodness of fit of the model. The outcome indicated that smaller value of AIC was in the negative binomial regression analysis and the model fit the data. Hence, a negative binomial regression model was proposed in analyzing the count data of the total number of Pteridophyta. Finally, the study used incidence rate ratio (IRR) of the negative binomial regression model in order to interpret the model and at the same time detecting the variable that eventually affects the total numbers of Pteridophyta to grow more efficiently. As a result, all types of Pteridophyta species seems to have influenced in predicting that species. While, 12 years and 20 years were a more suitable time in indicating the high number of Pteridophyta species.

### 5. Conclusion

The GLIM consists of several statistical analyses of parametric models for non-normally distributed data which includes the Poisson, negative binomial, zero-inflated and hurdle models. In this study, Poisson regression analysis, zero-inflated Poisson and negative binomial were used in predicting the total number of Pteridophyta species. Since the data was counted, it was difficult in accurately estimating the values. Besides, the data was limitedly taken on fixed duration of time which was within 4 years, 8 years, 12 years and 20 years. As the data was secondary data, it was difficult to estimate the certain time needed. In addition, in terms of the counting the total number of plants, this secondary data did not clearly explain the estimated values within what area, the width or how far the observation has been taken. Then, the data was only comprised the categorical data which was within the types of Pteridophyta and the duration of years the plants grow. The study recommended to use other types of the method in predicting the total number of Pteridophyta species and adding more suitable factors to increase the accuracy in predicting the Pteridophyta species.

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References