Detection of Interleukin 5 and 6 on Suspected Patients with Clinical Symptoms of Guttate Psoriasis

Anushiya Ahbiraami Letchumanan¹, Rosmani Ismail², Chin Chien Yee³
¹University of Selangor (UNISEL) Faculty of Engineering and Life Sciences, Jalan Zirkon A7/A, Seksyen 7, 40000 Shah Alam. Selangor, Malaysia.
²Tengku Ampuan Rahimah Hospital Ministry of Health Malaysia Jalan Langat, 41200 Klang. Selangor, Malaysia.
³Sungai Manila Klinik Kesihatan d/a Pejabat Kesihatan Kawasan Sandakan, Ministry of Health Malaysia Bangunan Persekutuan 90500 Sandakan Sabah, Malaysia.

Abstract:- Guttate Psoriasis is a second and third higher incident of skin disease influence on children and young adult with poor hygiene reported in Malaysia and the universe. In Malaysia, the diagnosis performed based on the clinical presentation and fewer studies are being conducted. In this study, we are investigating the presentations of Interleukin (IL) 5 and 6 levels, which act as a biomarker. The plasma samples collected from 46 hospitalised patients and two healthy controls with the age range of 1 to 12. The patients selected based on the clinical symptoms and readings of Differential Counts (DC) analysis. We analysed the plasma by using a quantitative Sandwich Enzyme Immunoassay method according to the manufacturer. Our findings show that levels of IL-5 and 6 classified into acute and chronic stages. Average readings of IL-5 and 6 were 0.121pg/mL and 0.061pg/mL. The IL-5 level increased by four folds than the controls; however, the IL-6 levels show infinite value on controls. In addition, data analysed by using T-Test and ANOVA. The study was significant in classifying the stages for Guttate Psoriasis. The DC analysis has a significant role in screening clinical symptoms of Guttate Psoriasis. It recommended for a diagnostic test since it has high sensitivity, specificity, and accuracy on detecting Guttate Psoriasis. They complimented it on the plasma in serology diagnosis when the clinical features and DC remain abnormal. Inconclusive measurement of cytokines was useful in diagnosis and it will perform further studies on a molecular level for the better prognosis.

Keywords:- Cytokines; Guttate Psoriasis; Paediatric; Differential Counts.

I. INTRODUCTION

Psoriasis is a skin disorder because of proliferation in the epidermis. It triggers by hereditary predisposition, environment, or immunological conditions. It encompasses about 1-3% of the entire community worldwide [1]. Guttate Psoriasis is a second and third higher scene of skin diseases that influence children and young adults with modest sanitation registered in Malaysia (3.2% in adult and 7.2% in paediatric) and universal (ranged 0.6-20%) [2], [3], [4]. The causative agents are Group A Beta-Haemolytic Streptococci (GABHS) species [5]. However, the clinical appearance of Guttate Psoriasis is producing red droplets on the face, ears, scalp, and upper extremities [6]. The relation between IL-5 and Psoriasis are an expansion, differentiate, and chemo-attractant factor for Eosinophils but it not involving in severity of the disease [7]. Eosinophils caused inflammation and tissue destruction in hypersensitivity by releasing toxic granule proteins, lipid mediators, cytokines, and chemokine [8]. IL-6 correlated with harshness of the condition by direct and indirect contacts as person-to-person approach, oral-faecal route, sharing of food and water and touching the surface of contaminant object [9]. At this stage, it extends to Plaque Psoriasis and other comorbidities as Osteoporosis, Obstructive Sleep Apnea (OSA), and Chronic Obstructive Pulmonary Diseases (COPD) [10], [11], [12]. In this investigation, we are enhancing on children assessment that suspected with clinical manifestations on Guttate Psoriasis.

A. Pathogenesis of Guttate Psoriasis

The bacteria have reversed action as it appears after excretion of not washing their urinary meatus [13], [14]. It causes to release an IL-1 passage in against the bacterial filtration. The IL-1 is an initial inflammatory cytokine as it took part in early cellular response to inflammatory stimuli [15]. The bacteria move inwards from the urethra, urinary bladder, ureter, and resides in the kidney [13]. In the kidney, it starts proliferate the bacteria, and it induced to the activation of IL-2. The kidney pump contaminated deoxygenated blood to the heart and delivered into the intrinsic flow [16]. T cell subgroup is effective in the incentive of keratinocyte growth that showed in the expression IL-2 [17]. In IL-3 passages, it affects the flow of blood within the body and induced destruction of tissues because of hypoxia. The bacterial proliferation in vessels stimulated white blood cells and destroying the bacteria [16]. Production of IL-4 could alter the homeostatic by suppressing the positive assessment of pro-inflammatory cytokines. In the interim, IL-4 down-regulates fusion of several cytokines that seen in Psoriasis [17]. Epidermal cells in Psoriasis have a strong linkage on IL-4 [18]. Skin eosinophilia and epidermal hyperplasia caused by enormous secretion of IL-5 owing to the exposure of pathogens [19]. We are expecting the afflicted children might experience from mild itchiness because of existence spot vesicles and discharge of fluid. Then IL-5 is serving as an acute phase of Guttate Psoriasis. We upheld the children should take appropriate diagnose and treatment as in the early stage than being treated in chronic. At instance we
cannot treat the concerned children with the same ointment and oral remedy as it prescribed for an adult. IL-6 generates new comorbidities, like multi-organ failure and enhances into other types of Psoriasis. 73% of patients with Psoriasis represent at least one persistent type of illness. The examples of conditions are inflammatory bowel syndrome, uveitis, and psychiatric disturbances are leading to Psoriasis. Inflammatory products produced by the skin lesion where it issues on systemic flow in worsening immune-mediated diseases. It proportions the same pathways in clinical manifestations of upper respiratory tract and urinary tract infections in developing the disease [20], [12]. We accomplish that IL-6 function as a chronic stage in Guttate Psoriasis where it develops into other difficulty in affected children. At this stage, it transmitted the disease from an infected individual to healthy people.

II. MATERIALS AND METHODS

- **Study Population**
  Plasma specimens from 46 local paediatric patients (24 boys and 22 girls), aged 1-12 years, hospitalised at HTAR from May to July 2017. The following norms applied in selection patients with no clinical record of an autoimmune syndrome, Psoriasis, and inflammation. They were suffering from Upper Respiratory Tract Infection (URTI), Urinary Tract Infection (UTI), Gastrointestinal Tract Infection (GITI), skin infection, or blood disease. Suspected Guttate Psoriasis specimens assigned based on high readings on DC. 2 healthy plasma specimens were available at UNISEL.

- **Measurement of Cytokines**
  Plasma IL-5 and 6 levels analysed by a quantitative Sandwich Enzyme-Linked Immunosorbent Assay (ELISA) method (R&D Systems a Biotechne Brand, USA & Canada). This method was effective in capturing and detecting the amount of antigen produces in between two layers of antibody. These assays detected only human cytokines and the Minimal Detectable Dose (MDD) in our lab were 0.06-1.08pg/mL for IL-5 and below 0.70pg/mL for IL-6.

- **Statistical Analysis**
  All data analysed applying the Statistical Package for Social Science (SPSS) 16.0 for Windows 7. Plasma cytokines analysed using T Test. Analysis of Variance (ANOVA) applied to analyse on DC, gender, and clinical manifestations. The statistical significance accepted as the p-value < 0.05.

III. RESULTS

A. Screening Test Analysis
  In our investigation, the DC specifications have a notable feature in screening on Guttate Psoriasis. The specimens preferred based on the high readings on Neutrophils, Monocytes, and Lymphocytes.

  The study illustrates significant differences in below specifications. It verifies that the DC plays essential aspects in the screening assessment for disease.

<table>
<thead>
<tr>
<th>Pairs</th>
<th>Tests</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>IL-5 and DC</td>
<td>t = 11.179</td>
<td>0.001</td>
</tr>
<tr>
<td>IL-6 and DC</td>
<td>t = 11.171</td>
<td>0.001</td>
</tr>
</tbody>
</table>

Table 1: T Test for DC IL-5 and 6

B. Clinical History on Guttate Psoriasis
  The analysis reveals no significant differences in the influence of clinical disorders in patients where p-value higher than 0.05. Then a diversity of health complications may promote to the progression Guttate Psoriasis.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>F</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>IL-5</td>
<td>1.650</td>
<td>0.151</td>
</tr>
<tr>
<td>IL-6</td>
<td>1.303</td>
<td>0.287</td>
</tr>
</tbody>
</table>

Table 2: ANOVA Analyses for Clinical Diagnoses, IL-5, and IL-6.

C. Analysis of Human IL-5 and 6
  It illustrates the health condition of IL-5 and 6 readings that correspond with Guttate Psoriasis. Two healthier paediatric have inadequate detection on IL-5 and 6 than the paediatric suspected for Guttate Psoriasis.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Stages</th>
<th>Number of Patients</th>
<th>Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td>IL-5</td>
<td>Acute</td>
<td>11</td>
<td>0.01</td>
</tr>
<tr>
<td>IL-5 and 6</td>
<td>Chronic</td>
<td>36</td>
<td>#</td>
</tr>
</tbody>
</table>

Table 3: Clinical Status of Guttate Psoriasis

  The analysis shows significant differences between IL-5 and 6 where the p-value fewer than 0.05. Therefore, it confirmed the roles of both cytokines in regulating the stages of Guttate Psoriasis.

<table>
<thead>
<tr>
<th>Pair</th>
<th>Test</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>IL-5 and 6</td>
<td>t = 8.995</td>
<td>0.001</td>
</tr>
</tbody>
</table>

Table 4: Paired Samples T-Test

D. Risk Factor Analysis
  The investigation determines that no significant differences in the aspect of genders in patients where the p-value were greater than 0.05. In outline, both genders were at correspond risks seen in the disease.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>F</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>IL-5 and Gender</td>
<td>0.770</td>
<td>0.385</td>
</tr>
<tr>
<td>IL-6 and Gender</td>
<td>0.049</td>
<td>0.826</td>
</tr>
</tbody>
</table>

Table 5: ANOVA analysis for gender, IL-5, and 6
E. Sensitivity, Specificity, and Accuracy

Keys:
TP: True Positive
TN: True Negative
FP: False Positive
FN: False Negative

\[
\begin{align*}
\text{Sensitivity} &= \frac{TP}{(TP + FN)} \\
\text{Specificity} &= \frac{TN}{(TN + FP)} \\
\text{Accuracy} &= \frac{(TP +TN)}{(TP + TN + FP + FN)}
\end{align*}
\]

[21]

The above study shows that the IL-5 and 6 were susceptible in identifying the infection among the suspected patients. However, the specificity of the assessments ruled out the patients without infection. The accuracy of the assessments was greater than specificity but lesser than the sensitivity.

Then these can enforce on a gold standard of diagnostic assessments in identifying Guttate Psoriasis.

<table>
<thead>
<tr>
<th></th>
<th>IL-5</th>
<th>IL-6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sensitivity</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Specificity</td>
<td>80</td>
<td>70</td>
</tr>
<tr>
<td>Accuracy</td>
<td>95</td>
<td>86</td>
</tr>
</tbody>
</table>

Table 1.7: Sensitivity, specificity, and accuracy of the tests

IV. DISCUSSION

People with intense Psoriasis were experiencing a large diversity of Neutrophil in contrasted with moderate and mild types [22]. In extreme presence of Neutrophil could contribute to an expeditious proliferation and differentiation on keratinocyte [23]. In pathological differences that observed in psoriatic skin bruises was the morphologic of T-cell and macrophage. The chemoattraction of Neutrophils within the epidermal compartments regulate the activation of Lymphocytes. They displayed Neutrophil in epidermis and papillary dermis at the early stages of illness [1]. Among Psoriasis patients have large circulating amount of Monocytes and that could figure out the harshness of Plaque Psoriasis. It further possesses an adhesive property in the pathogenesis of Psoriasis and Psoriatic Arthritis (PsA) [24]. T-Lymphocytes develop in a vulnerable environment among Psoriasis patients. Lymphocytes further facilitate in movement keratinocyte to epidermal cell in Psoriasis [14]. The URTI were correlates in preceding to Guttate Psoriasis but it not endorse in the investigation [5]. Psoriasis sufferers have a peculiar appearance on their digestive portion such as dystrophic modifications in epithelial cells and stromal infiltration on gastric and duodenal mucosa. A bacterial gut infection and eosinophilic gastroenteritis were causing malabsorption in Psoriasis patients [25]. UTI was another causative agent in causing Guttate Psoriasis and it justified in a research where the sufferer diagnosed for Guttate Psoriasis without a clinical record of URTI [20]. Cellulitis induced by Streptococcus species and it provokes Guttate Psoriasis [26]. Impetigo was a skin infection caused by Staphylococcus and GABHS species, and it was familiar among children [27]. If the condition persists, it would progress to a new disorder as Guttate Psoriasis, cellulitis, lymphangitis, septicaemia, scarlet fever, and poststreptococcal glomerulonephritis (PSGN) [28]. Iron Deficiency Anaemia (IDA) links with persistent inflammatory illnesses and not resemble in severing the Psoriasis condition [29]. Thrombocytopenia caused by drastic bacterial infection [30]. Thalassemia carries a significant prevalence in forming PsA [31]. The early researches emphasise the Guttate Psoriasis affected on children, young adult and it was common in both genders [2], [9], [20]. Our findings supported with published data on the elements of aged and genders with Guttate Psoriasis. An ample existence of IL-5 had in psoriatic lesions varied with controls by applying immunochemistry procedure [8]. IL-6 supports in T cell activation circulate a rapid cycle of keratinocyte during the initial progress of acute inflammation [32], [33]. However, the presence IL-5 appears not coincides with the severity of the disease [7].

The measured levels of serum IL-6 by ELISA method in Psoriasis patients it serves correlates with disease severity. There was a substantial rise on serum IL-6 on psoriatic abrasion than the healthy group [32]. Psoriasis can lead to systemic complication as cardiovascular defect, the metabolic disorder, and inflammatory bowel disease [20], [12], [34]. Sensitivity and specificity were proportional, meaning that as the sensitivity increases, the specificity decreases, and vice versa [35]. It can determine accuracy of a diagnostic assessment from sensitivity and specificity with existence pervasiveness. Sensitivity was the portion in true positives a diagnostic assessment identified. The analytical values of specificity suggest the possibility of a trial analyses a specific disease when the clinical aspects and DC remain anomalous. The measurement level cytokines was effective in analysis and more investigations will perform on a molecular level for a better prognosis.

V. CONCLUSION

They complemented it on the plasma serology diagnosis when the clinical aspects and DC remain anomalous. The measurement level cytokines was effective in analysis and more investigations will perform on a molecular level for a better prognosis.

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ETHICS APPROVED

This study approved by the Medical Research and Ethics Committee (MREC), Ministry of Health, and registered under the National Medical Research Registry (NMRR-17-760-35370).

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