SLE as Hematologic Disorder and Kozhikode Criteria

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ABSTRACT

Systemic Lupus Erythematosus is a multisystem connective tissue disease, characterized by a wide variety of clinical features and the presence of numerous auto-antibodies, circulating immune complexes and wide spread immunologically mediated tissue damage. The expression of the disease is greatly influenced by genetic, environmental, dietary, demographic and geographical factors. Most of the literatures on SLE are from the Western countries. Given the importance of diet, life style, genetic, environmental and racial factors, a change in the clinical profile of the disease should be anticipated in different parts of the world. This article combines the observations made in two well-conducted studies in a tertiary care center in North Kerala and describes SLE as a hematological disease in our sub set of population and on the criteria which we had developed for diagnosing SLE, the Kozhikode Criteria.

Keywords: SLE, Kozhikode criteria, Hematology

INTRODUCTION

Systemic Lupus Erythematosus is a multisystem connective tissue disease, characterized by a wide variety of clinical features, presence of numerous auto-antibodies, circulating immune complexes and wide spread immunologically mediated tissue damage [1]. The expression of the disease is greatly influenced by genetic, environmental, dietary, demographic and geographical factors. Considerable variation has been observed regarding the clinical manifestations of SLE among various geographical regions and ethnic groups. The first case of SLE in India was reported in 1955 [2]. Subsequent studies conducted in different parts of India described the various manifestations in Indian population with SLE.

Race has shown to be a major predictor in the clinical manifestations, serological pattern and morbidity in SLE [3,4]. Being the largest continent, both by area and population, Asia encompasses people of different diet, lifestyles, and socio cultural background with diverse ethnic groups. Diet and life style habits of the Indian population are never comparable to the western practices and standards. Importance of dietary factors is exemplified by an Indian study on Vitamin D levels and SLE [5]. Although a causal relationship could not be established, it was found that people with Vitamin D deficiency have severe disease. Interestingly Vitamin D deficiency is very high among Indians. High melanin concentration in the skin [6], avoidance of sunlight and poor food habits contribute to low levels of Vitamin D 3 in Indians [7], the same are the causes for other diseases as well [8].

Most of the literatures on SLE are from the West and the diagnostic guidelines are based on those observations only. Given the importance of diet, life style, genetic, environmental and racial factors, a change in the clinical profile of the disease should be anticipated in different parts of the world. Most of the western studies highlight SLE as a rheumatological and dermatological disease, the logic of which is questionable too. Blood and blood vessels contain more variety of antigens than that of other organs, and therefore naturally, in SLE where autoimmunity can develop against any antigen, the clinical manifestations should be more often hematological [9,10].

The criteria used for diagnosis of SLE till 2012 were the American College of Rheumatology (ACR) Criteria.

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However it should be understood that ACR criteria is not a diagnostic criteria, but a classification criteria. It can be employed only to identify and classify the disease in a person as SLE when there is a group of manifestations occurring sequentially or simultaneously, it does not serve the purpose of early diagnosis. Moreover, it requires 4 out of the 11 criteria to be present, before a pattern of symptom evolution could be classified as SLE. SLE is a disease, which has been proven to evolve over time with manifestations in various organs, which often happen sequentially over many years and not present simultaneously at presentation. It may take several years to satisfy the ACR criteria, which may result in delay in diagnosing and instituting treatment, which may have life threatening consequences. Moreover, the hematological manifestations of SLE are underrepresented in the ACR criteria.

Hematological manifestations of SLE are diverse and are often the presenting manifestation of the disease [9,11,12]. The major hematological manifestations are anemia due to hemolysis, MDS, B12 deficiency, Pure Red cell aplasia; leukopenia due to MDS or aplasia, B12 deficiency, hypersplenism; ITP like presentation and thrombocytopenia due to multiple other causes; Pancytopenia and anti phospholipid antibody syndrome with its varied presentation like cerebral vein thrombosis, deep vein thrombosis, Budd Chiari syndrome, Pulmonary thromboembolism with its varied presentations (including primary pulmonary hypertension-like presentation).Other manifestations seen are lymphadenopathy and hepatosplenomegaly mimicking lymphoma or leukemia, myelofibrosis, macrophage activation syndrome (HLH), TTP, pure red cell aplasia, MDS, etc., are not uncommon. It has been observed in our studies that SLE presents with hematological manifestations alone, without features of musculoskeletal, dermatological or other system involvement [9]. In almost all these cases presenting with hematological manifestations, it was only because a high index of suspicion that, they were diagnosed and treated early, to the advantage of the patient. The diagnosis in these cases was primarily by good clinical evaluation, looking for evidence of an autoimmune phenomenon and after ruling out other causes. Many such cases, which presented initially with manifestations of autoimmunity in one tissue or organ alone, were ANA negative and did not satisfy the ACR criteria, but did so on follow up [9]. A study from central rural India also showed that the manifestations in SLE were more Hematological and immunological [11]. Two separate studies from South India have shown that mucocutaneous manifestations are less prevalent [12,13]. Taking into account the aforementioned factors, we proposed the need for a practical guideline to diagnose SLE and framed a new criterion named as the Kozhikode criteria [9,10].

KOHIKODE CRITERIA

Major/essential criteria
1. Presence of an unresolved autoimmune disorder, which is known to occur with SLE (Chronic ITP, Immune hemolytic anemia, autoimmune hypothyroidism and autoimmune hepatitis).
2. No other causes identified, other than autoimmunity, for the said problem, by clinical reasoning and investigations.

Minor criteria
1. Another co existing autoimmune disorder/any other clinical or laboratory evidence of autoimmunity.
2. Positive ANA.
3. Positive anti ds DNA.
4. Sustained and definitive response to steroids and immunosuppressant even after 6 months of follow up.

If the patient has 2 essential and 2 or more minor criteria, they can be diagnosed as SLE.

Our first study was to establish the fact that SLE presents more often with hematological manifestations, and the second study was to establish the validation of the Kozhikode Criteria, both were done in a tertiary care center in North Kerala sequentially [10,14].

MATERIALS AND METHODS

Both the studies were conducted in Government Medical College, Kozhikode, after getting approval from the institutional research committee. The first study was conducted in the year 2009-2010, by Sasidharan et al. [10] and the second one in the year 2013, by Arathi et al. [14]. Both the studies included diagnosed cases of Systemic Lupus Erythematosus, in the Departments of Internal Medicine, Rheumatology, Hematology and Nephrology Departments of Government Medical College, Kozhikode. The studies included already diagnosed cases of SLE and the newly diagnosed cases during their respective study periods. The cases were followed up for a period of one year. There was a considerable overlap in patients in both the studies, since patients already under follow up were included.

Clinical diagnosis of SLE was considered in those presenting with unsettled clinical problems and conditions that were autoimmune in nature, like chronic ITP, hypothyroidism, autoimmune hemolytic anemia, vitiligo, alopecia and all other autoimmune disorders with some laboratory evidences to suggest autoimmunity.

Basic investigations constituting complete blood count, including red cell indices, ESR, renal and liver function tests, urine routine, peripheral smear, ANA, anti-ds DNA were done for all cases. Further investigations like reticulocyte count, Coombs test, ANA profile, radiological
tests, tissue biopsy or cytology, including bone marrow, were done in relevant cases. Data was collected using structured personal interview and clinical examination. The first study included patients who satisfied either the ACR criteria or the Kozhikode Criteria. The new criteria was essential, to include those patients who did not satisfy the ACR criteria, but were strongly suggestive of having an autoimmune etiology.

The second study was aimed at validating the Kozhikode Criteria [14]. For this purpose both the ACR and the Kozhikode Criteria were applied to all diagnosed cases of SLE. All newly diagnosed cases that were diagnosed as SLE using the Kozhikode Criteria were also subjected to the ACR criteria. We found that large number of them did not satisfy the ACR criteria, but all of them who satisfied the ACR criteria also satisfied the Kozhikode Criteria. Those who did not satisfy the ACR criteria initially were reviewed after a period of 6 months and the ACR criteria was applied again to see if they satisfied it and thereby to prove whether the Kozhikode Criteria helps in early diagnosis of SLE. In addition, those who were already under follow up were studied by reviewing their clinical features and treatment details and noted whether or not they satisfied the ACR criteria at the beginning and by calculating the average time required to satisfy the ACR criteria.

For assessing the influence of diet and life style, we recruited the caretakers of patients who matched with the cases in age and sex, utilizing a case control design. Dietary information was estimated using the semi quantitative food frequency questionnaire derived from the Integrated Disease Surveillance Project, Non Communicable disease Survey Questionnaire. There were six possible responses, regarding how frequent a particular food item was consumed. The six responses were, nil/less than once in a month, 1-3 times in a month, 1-2 times in a week, 3-4 times in a week, 1-2 times in a day, 3-4 times in a day.

RESULTS

The first study [10] included 108 patients and the second study [14] included 71 patients. Male to female ratio in the first study was 1:10 and in the second study was 1:9. The most frequent clinical manifestations in both the studies were hematological (Figure 1).

Hematological manifestations were the most common in the study conducted by Arathi et al. too [14] (Figure 2).

![Figure 1. Most common clinical manifestation in SLE observed by Sasidharan et al. [10].](image1)

![Figure 2. Clinical manifestations of SLE as observed by Arathi et al. [14].](image2)
It was seen in both the studies that, a large proportion of cases presented with hematological manifestation as the first symptom (42.69%). There was an inverse relationship between musculoskeletal and hematological manifestations (p value<0.001). There were 10 cases of hypothyroidism and 6 cases of autoimmune hepatitis in the study by Sasidharan et al. [10]. These diseases are not given any representation in the ACR criteria. The most common hematological manifestation was Immune Thrombocytopenic purpura (Figure 3).

**Figure 3.** Spectrum of hematological manifestations as observed by Sasidharan et al. [10].

Anemia was seen in 62.9% cases. It was multifactorial. It was due to nutritional, autoimmune and anemia of chronic disease.

The commonest manifestation of APLA was Cerebral Venous Thrombosis (CVT) and abortions.

Thrombocytopenia was seen in 7 cases of APLA. All the patients had only mild thrombocytopenia that did not warrant any intervention (Figure 4).

**Figure 4.** Clinical manifestation of anti-phospholipid antibody syndrome, observed by Sasidharan et al. [10].

In the study conducted by Arathi et al. [14], for validation of the Kozhikode criteria, there were a total of 71 cases. Out of this 30 were new cases and 41 were previously diagnosed cases of SLE, already under follow up. Of the 71 cases, 45 satisfied Kozhikode Criteria alone, but only 22 satisfied both ACR and Kozhikode Criteria and 4 did not satisfy either criterion. Altogether, 67 patients satisfied the Kozhikode Criteria at the beginning of the study. There was no such group of patients who satisfied the ACR criteria alone but not the Kozhikode criteria, validating the new criterion. Both these observations are highlighting the utility of the new criterion. Out of the 30 new cases, 26 individuals satisfied the Kozhikode criteria, whereas only 6 were satisfying the ACR criteria. The 26 newly diagnosed cases, which satisfied the Kozhikode criteria alone, were followed up for a period of 6 months and it was observed that only 2 of them satisfied the AR criteria at the end of 6 months (Figure 5).
Figure 5. Distribution of cases based on whether or not they satisfied the Kozhikode criteria [14].

Of the 4 cases which did not satisfy either of the criteria, 2 of them satisfied the Kozhikode criteria at the end of 6 months. One of them had refractory oral ulcers, which subsequently became ANA positive and responded well to steroids. The second patient was a case of alopecia and ITP who responded well to steroids and later became ANA positive. None of the 4 satisfied the ACR criteria even at the end of 6 months. Of the 2 patients who did not satisfy either of the criteria, one patient had refractory oral ulcers and the other patient had lost for follow up.

Among the patients who were already under follow up, it was found that 16 of them who all had satisfied the Kozhikode criteria in the initial diagnosis itself, had satisfied the ACR criteria only during variable periods of follow up as given in Table 1.

Table 1. Average time required to satisfy the ACR criteria among the cases [12].

<table>
<thead>
<tr>
<th>Serial Number</th>
<th>Diagnosis</th>
<th>Average time from diagnosis to satisfying the ACR criteria (in months)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>ITP</td>
<td>32</td>
</tr>
<tr>
<td>2</td>
<td>ITP</td>
<td>72</td>
</tr>
<tr>
<td>3</td>
<td>ITP, Hypothyroidism</td>
<td>40</td>
</tr>
<tr>
<td>4</td>
<td>AIHA, Polyarthritis</td>
<td>7</td>
</tr>
<tr>
<td>5</td>
<td>AIHA, Nephrotic Syndrome</td>
<td>64</td>
</tr>
<tr>
<td>6</td>
<td>Polyarthritis</td>
<td>4</td>
</tr>
<tr>
<td>7</td>
<td>APLA, raised ESR</td>
<td>14</td>
</tr>
<tr>
<td>8</td>
<td>CVA, ITP, Raised ESR</td>
<td>144</td>
</tr>
<tr>
<td>9</td>
<td>Anemia, Splenomegaly</td>
<td>6</td>
</tr>
<tr>
<td>10</td>
<td>Secondary Sjogrens syndrome</td>
<td>4</td>
</tr>
<tr>
<td>11</td>
<td>Thyrotoxicosis</td>
<td>11</td>
</tr>
<tr>
<td>12</td>
<td>Polyarthritis</td>
<td>8</td>
</tr>
<tr>
<td>13</td>
<td>Polyarthritis</td>
<td>4</td>
</tr>
<tr>
<td>14</td>
<td>Anemia, High ESR</td>
<td>12</td>
</tr>
<tr>
<td>15</td>
<td>Pancytopenia</td>
<td>12</td>
</tr>
<tr>
<td>16</td>
<td>MDS</td>
<td>60</td>
</tr>
</tbody>
</table>

Abbreviations: ACR: American College of Rheumatology; ITP: Immune Thrombocytopenic Purpura; AIHA: Autoimmune Hemolytic Anemia; ESR: Erythrocyte Sedimentation Rate; MDS: Myelodysplastic Syndrome; CVA: Cerebro Vascular Accident
The most common clinical manifestation was hematological and it was unusual to see any joint symptom in this subset. Arthritis was seen in combination with other symptoms but not alone as the initial manifestation. Among the new cases, only two were ANA positive at the time of initial clinical presentation. They were diagnosed and treated as SLE due to strong index of suspicion. Those who were ANA positive were also positive for double stranded DNA. Of the ANA negative patients among new cases, only 2 became ANA positive at 6 months of follow up.

Majority of the patients belonged to moderate socio economic status. Among those diagnosed as SLE, most of them, had a sedentary life style (p<0.05) (Figure 6).

Comparison of diet among cases and controls showed statistically significant decrease in intake of pulses, green leafy vegetables, legumes and fruits (p=0.000). The intake of junk food and canned food was statistically significant among cases than in controls (p=0.000).

**DISCUSSION**

In both the studies, it was found that hematological manifestations were the commonest and the presenting symptom. This was contrary to the description of the disease in most western books [15,16]. Our findings emphasize the fact that hematological manifestations are a common presenting symptom and may be missed if the index of suspicion is low. Some studies in different parts of the world have evaluated the hematological findings in SLE and the prevalence rates obtained in them are comparable to our observations [17-20]. But hematological manifestations as the initial presentation and its proper inclusion in the criteria for diagnosing SLE were not addressed in any of these studies.

There was a significant inverse relation between the presence of musculoskeletal symptoms and hematological manifestations. The commonly co existing abnormality of autoimmune hypothyroidism in patients presenting with hematological manifestations is under represented in the ACR criteria. No such studies are found in the literature.

Twelve patients in the first study were ANA negative and did not satisfy the ACR criteria and 28 cases in the second study were ANA negative and 24 cases in the second study did not satisfy the ACR criteria. Thus ANA negativity does not rule out SLE in early stages. They all had the clinical diagnosis of possible SLE or evolving SLE at the time of initial presentation itself. In 1982, Mc Hardy and Rennie [21] investigating a cohort of SLE patients in Aberdeen, had suggested a prevalence of 8.9% for ANA negative SLE.

Gladman et al. [22] and Ferrerio et al. [23], in two separate studies found that the prevalence of ANA negative SLE was approximately 5%. This emphasizes the importance of relying on individualized clinical judgment rather than on the existing criteria alone for treating the disease and the need for rigorous follow up of all suspected cases, even if they are ANA negative. Based on these observations, Sasidharan et al. [10] proposed the new criteria- The Kozhikode Criteria.

The second study was undertaken to validate the Kozhikode criteria as an alternative to the ACR criteria. The observations of the study has shown the utility of the Kozhikode criteria for early diagnosis of SLE and the observations mentioned are supporting the superiority of the Kozhikode criteria over the ACR criteria for the early diagnosis of SLE. The study has thus validated the Kozhikode Criteria as a useful tool for early diagnosis of SLE.

It was also observed that several of the patients were ANA negative at the time of presentation and they become ANA positive during follow up only. All these patients were clinically diagnosed as SLE or evolving SLE and all of them had satisfied the Kozhikode criteria and not the ACR criteria.

None of the patients were doing exercise on a regular basis. The level of physical activity in the newly diagnosed cases when compared to the controls showed statistically

**Figure 6.** Level of physical activity in SLE patients [14].
significant decrease in physical activity among the cases. Although there were studies establishing the role of lack of regular exercise and sedentary lifestyle increasing the mortality in SLE, there were no studies relating them to the etiology of SLE [24,25].

The comparison between diets of cases and controls showed statistically significant difference, showing poor intake of pulses, green leafy vegetables and fruits. The intake of junk food, fried meat and canned food was higher among cases than controls, with a significant p value.

Table 2. Comparison between ACR and Kozhikode criteria [14].

<table>
<thead>
<tr>
<th>ACR Criteria</th>
<th>Kozhikode Criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Classification criteria. Cannot be used for diagnosing SLE</td>
<td>Diagnostic criteria</td>
</tr>
<tr>
<td>Requires at least 4 out of 11 criteria, most of which are not present at the time of initial presentation and could be present at any time in a patient's history- delays diagnosis</td>
<td>Requires 4 of the 6 and are easily available features occurring early on, as the diagnosis of SLE is based on a clinical suspicion. Promotes early diagnosis</td>
</tr>
<tr>
<td>Under representation of hematological manifestations, which is the commonest at least in our sub set of population</td>
<td>Adequate representation of hematological manifestations</td>
</tr>
<tr>
<td>Presence of anti-nuclear antibodies given equal weightage as other criteria</td>
<td>Presence of antinuclear antibody is only a minor criteria and helps to identify ANA negative SLE</td>
</tr>
</tbody>
</table>

REFERENCES


