A Study on Biochemical Parameters in Patients with Rickettsial Infection

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Abstract

Background: The traditional views of tick-borne rickettsioses as endemic diseases with largely focal distributions and limited host and geographic ranges, predetermined seasonality and defined tick associations became obsolete or at least very incomplete. This expansion of awareness about the existence of other rickettsial agents with varied clinical and epidemiological attributes has been thoroughly reviewed but it has presented new challenges to the medical and public health communities. Subjects and Methods: The clinical presentation and multiple organ dysfunctions in these patients were evaluated with a special focus on the renal manifestation and hepatic manifestations. The study population included all the patients presenting fever, rash and were diagnosed with rickettsial disease by clinical examination. A total of 60 subjects, satisfying the inclusion and exclusion criteria were included in the final analysis. The sample size was calculated assuming the expected proportion of rickettsial infection as 11% among fever cases as per previously published studies, with a precision of 8% and 95% confidence level. Results: The mean AST in the 80 titre was 140.43 ± 79.92, it was 285.89 ± 184.95 in 160 titre, 364.92 ± 89.69 in 320 titre group and 579.29 ± 106.26 in 640 titre group. The mean difference of AST 145.47 in 160 titre group was statistically significant (P value<0.001), 224.50 in 320 titre group was statistically significant (p value<0.001), and in 640 titre group 438.86 was statistically significant. (P- Value <0.001). Conclusion: The study has highlighted the need to have a high index of suspicion to enhance the diagnosis of ricketsial diseases and also the strong association between weifelixititre and liver and renal dysfunction.

Keywords: Rickettsioses, Weil Felix Titre, Biochemical Parameters.

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Introduction

Right from the very early time in the history, there were many insults caused by the Rickettsial infections. The most evident was that of the epidemic typhus upshot in Russia during the late 1910s and early 1920s where in about three million lives were spared. There is convincing evidence of this disease to be considered a public health threat with its emergence characteristics on at-risk population. In the 1940s the United States (US) reported an alarming rise of murine typhus cases, which was then followed by RMSF surge in 1970s and by the ehrlichioses (human) ascent later in the 1990s.[1] The increased burden of the disease in a specific geographical region has added insight into the transmission cycle happening in vectors, that envisages trans-ovarian and trans-stadial route of transmission of Rickettsial pathogens occurring in vectors.[2] This further added the epidemiological values to this disease.

The incidence of tick-borne rickettsial diseases is peaking. It is currently going through its second pronounced increase in the last 40 years. Since the 1970s, four endemic rickettsioses Rocky Mountain Spotted fever (RMSF), Mediterranean spotted fever (MSF), North Asian tick typhus (NATT) and Queensland tick typhus (QTT), have been on a continuous increase. The incidence of Japanese spotted fever has also increased steadily since its discovery in the mid-1980s. It is possible that other tick-borne rickettsial infections have shown similar increases but only these more common and severe diseases have much useful, albeit based on the limits imposed by contemporary views, disease surveillance information. The incidence of RMSF has raised its toll to the maximum US during 2004. The ticks (Rhipicephalussanguineus) were noted in the brown dogs of the residents which showed positivity of R. rickettsii organisms in it.[3]

Blanton L. S. et. Al.[4] have reviewed the understanding of this disease by the countries and the precautions taken by the travelers to prevent this from occurring. The reports were daunting as the study found increasing prevalence of disease in tropics (travelers most preferred location) in addition to its re-emergence in Bulgaria following poor vector control measures. Furthermore, the sequel of the disease (Rickettsia rickettsii) sustains the same level in Central and South America. There was also evidence suggesting the declining efficacy of fluoroquinolones that were used for the treatment of milder conditions. This exposes the weakened preparation of the countries to control this disease.
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Sagin D. D. et. Al.\textsuperscript{[5]} have identified and described the Rickettsial infection in remote Orang Ulu villages in upper Rejung River, Sarawak, Malaysia. People in 5 Orang Ulu villages in Sarawak, Malaysia, there were 9.6% who were positive for typhus out of which, 3.8% were positive for tick typhus (7/11), while scrub typhus was positive in (4/11) or endemic typhus positivity was found in (1/11). The incidence of typhus was found to be higher among semi-nomadic Penans than that of the settled Kayans. 

Mittal V. et. Al.\textsuperscript{[6]} in their quest to find the profile of Rickettsial infections in Delhi have detected scrub typhus (48.2%) as the commonest Rickettsial infections followed by spotted fever group (27.5%) and typhus group (6.8%) between 2005 and 2009. They have also found Leptotomibiumdeliensie mite on the rodents during the entomological survey. 

Rahi M. et. Al.\textsuperscript{[7]} have postulated that the scrub typhus, murine flea-borne typhus, Indian tick typhus and Q fever are the major groups of rickettsioses, commonly reported diseases in India. Rickettsial infections are generally incapacitating and difficult to diagnose; untreated cases have case fatality rates as high as 30-45 percent with multiple organ dysfunctions, when not diagnosed promptly and appropriately treated. 

Ahmad S. et. Al.\textsuperscript{[8]} have reported nine cases of scrub typhus from the Garhwal region of the newly created north Indian state of Uttarakhand, a region not previously known to harbor the vector. It stressed the significance of entomological studies that are needed to find the density of the vector and for institution of vector control measures in order to prevent this relatively benign, yet potentially fatal, clinical entity from spiraling into a major public health issue. Being an emerging and re-emerging disease this attribute is critical for a disease of public health importance. “The traditional views of tick-borne rickettsioses as endemic diseases with largely focal distributions and limited host and geographic ranges, predetermined seasonality and defined tick associations became obsolete or at least very incomplete. This expansion of awareness about the existence of other rickettsial agents with varied clinical and epidemiological attributes has been thoroughly reviewed but it has presented new challenges to the medical and public health communities. The paradigm shift is due to the fact that numerous rickettsiae of unknown to variable degrees of pathogenicity for humans co-circulate in overlapping geographic regions and may even be found in the same tick species.”

**Subjects and Methods**

The study was a prospective observational study. Cases of fever admitted in the study period in our hospital were followed up on the basis of weilfelix positivity. The clinical presentation and multiple organ dysfunctions in these patients were evaluated with a special focus on the renal manifestation and hepatic manifestations. The study population included all the patients presenting fever, rash and were diagnosed with rickettsial disease by clinical examination. 

A total of 60 subjects, satisfying the inclusion and exclusion criteria were included in the final analysis. The sample size was calculated assuming the expected proportion of rickettsial infection as 11% among fever cases as per previously published studies, with a precision of 8% and 95% confidence level.

The required sample size, as per the above calculation was 59. To account for a 5% nonparticipation rate it was decided to sample 63 subjects, so that final analysis can include not less than 59 subjects. The final analysis has included 60 subjects at the end of data collection period.

All the eligible study subjects, satisfying inclusion and exclusion criteria were recruited into the study by convenient sampling. 

The association between quantitative blood pressure variables and any positive weilfelix was assessed by comparing the mean values. The mean differences along with their 95% CI were presented. ANOVA was used to assess statistical significance. The association between all symptoms of the disease and weilfelix titre was assessed by cross tabulation and comparison of percentages. Chi square test was used to test statistical significance.

**Results**

**Table 1:** Comparison of mean hemoglobin across study groups (N=60)

| Overall Weil-Felix titre | Haemoglobin Mean ± SD | Mean Difference | 95% Confidence Interval for Mean | P Value |
|--------------------------|-----------------------|----------------|-------------------------------|---------|
|                          | Lower Bond            | Upper Bond     |                               |         |
| 80                       | 11.88 ± 0.67          |                |                               |         |
| 160                      | 10.17 ± 1.77          | 1.71           | 0.58 - 2.83                  | 0.004   |
| 320                      | 9.78 ± 2.59           | 2.10           | 0.84 - 3.35                  | 0.001   |
| 640                      | 8.66 ± 2.25           | 3.22           | 1.67 - 4.78                  | <0.001  |

The mean hemoglobin in the 80 titre was 11.88 ± 0.67, it was 10.17 ± 1.77 in 160 titre, 9.78 ± 2.59 in 320 titre group and 8.66 ± 2.25 in 640 titre group. The mean difference of hemoglobin 1.71 in 160 titre group was statistically significant (p value<0.001), 2.10 in 320 titre group was statistically significant (p value<0.001), and in 640 titre group 3.22 was statically significant. (P- Value<0.001).

**Table 2:** Comparison of mean total count across study groups (N=60)

| Overall Weil-Felix titre | Total Count Mean ± SD | Mean Difference | 95% Confidence Interval for Mean | P Value |
|--------------------------|-----------------------|----------------|-------------------------------|---------|
|                          | Lower Bond            | Upper Bond     |                               |         |
| 80                       | 7978.57 ± 2895.8      |                |                               |         |
| 160                      | 7094.74 ± 2285.33     | 883.83         | -759.06 - 2526.73             | 0.286   |
| 320                      | 7465.38 ± 2693.46     | 513.19         | 2344.34                      | 0.577   |
| 640                      | 5864.29 ± 2118.71     | 2114.29        | -150.29 - 4378.86             | 0.067   |

The mean total count in the 80 titre was 7978.57 ± 2895.8, it was 7094.74 ± 2285.33 in 160 titre, 7465.38 ± 2693.46 in 320 titre group and 5864.29 ± 2118.71 in 640 titre group. The mean difference of total count 883.83 in 160 titre group was statistically not significant (p value0.286), 513.19 in 320 titre group was statistically not significant (p value0.577), and in 640 titre group 2114.29 was statically not significant. (P- value0.067).
The mean platelet in the 80 titre was 52333.98 ± 47613.68, it was 60631.71 ± 35052.44 in 160 titre, 58153.85 ± 18365.94 in 320 titre group and 21285.71 ± 10209.71 in 640 titre group. The mean difference of platelet 8297.74 in 160 titre group was statistically not significant (p value 0.468), and in 640 titre group was statistically significant (p value <0.001), 224.50 in 320 titre group was statistically not significant (p value 0.146), 2.87 in 320 titre group was statistically not significant (p value <0.001), and in 640 titre group 31048.26 was statistically not significant. (P value 0.052).

The mean total bilirubin in the 80 titre was 1.54 ± 0.51, it was 2 ± 1.11 in 160 titre, 4.41 ± 1.42 in 320 titre group and 6.63 ± 0.76 in 640 titre group. The mean difference of total bilirubin 0.46 in 160 titre group was statistically not significant (p value 0.146), 2.87 in 320 titre group was statistically significant (p value <0.001) and in 640 titre group 5.09 was statistically significant. (P value <0.001).

The mean AST in the 80 titre was 140.43 ± 79.92, it was 285.89 ± 184.95 in 160 titre, 364.92 ± 89.69 in 320 titre group and 579.29 ± 106.26 in 640 titre group. The mean difference of AST 145.47 in 160 titre group was statistically significant (p value <0.001), 224.50 in 320 titre group was statistically significant (p value <0.001), and in 640 titre group 438.86 was statistically significant. (P value <0.001).

The mean ALT in the 80 titre was 154.52 ± 112.16, it was 295.11 ± 200.03 in 160 titre, 375.23 ± 113.79 in 320 titre group and 484.14 ± 67.93 in 640 titre group. The mean difference of ALT 140.58 in 160 titre group was statistically significant (p value 0.003), 220.71 in 320 titre group was statistically significant (p value <0.001), and in 640 titre group 329.62 was statistically significant. (P value <0.001).

The mean ALP in the 80 titre was 149.74 ± 64.42, it was 209.74 ± 85.09 in 160 titre, 254.54 ± 67.09 in 320 titre group and 356.43 ± 94.55 in 640 titre group. The mean difference of ALP 59.97 in 160 titre group was statistically significant (p value 0.015), 104.78 in 320 titre group was statistically significant (p value <0.001), and in 640 titre group 206.67 was statistically significant.

The mean urea in the 80 titre was 40.86 ± 7.92, it was 46.53 ± 14.59 in 160 titre, 60.85 ± 15.77 in 320 titre group and 72.29 ± 14.67 in 640 titre group. The mean difference of urea 5.67 in 160 titre group was statistically not significant (p value 0.172), 19.99 in 320 titre group was statistically significant (p value <0.001) and in 640 titre group 31.43 was statistically significant (P value <0.001).

The mean sodium in the 80 titre was 129.76 ± 7.49, it was 119.74 ± 10.28 in 160 titre, 107.54 ± 7.45 in 320 titre group and 100.29 ± 7.7 in 640 titre group. The mean difference of sodium 10.03 in 160 titre group was statistically significant (p value 0.172), 19.99 in 320 titre group was statistically significant (p value <0.001) and in 640 titre group 29.48 was statistically significant. (P value <0.001).
The mean serum creatinine in the 80 titre was 1.32 ± 0.38, it was 1.67 ± 0.69 in 160 titre, 2.43 ± 0.64 in 320 titre group and 3.33 ± 0.87 in 640 titre group. The mean difference of serum creatinine 0.35 in 160 titre group was statistically not significant (p value 0.077), 1.11 in 320 titre group was statistically significant (p value<0.001) and in 640 titre group 2.01 was statically significant. (P-Value<0.001).

**Discussion**

The mean hemoglobin in the 80 titre was 11.88 ± 0.67, it was 10.17 ± 1.77 in 160 titre, 9.78 ± 2.59 in 320 titre group and 8.66 ± 2.25 in 640 titre group. The mean difference of hemoglobin 1.71 in 160 titre group was statistically significant (p value= 0.004), 2.10 in 320 titre group was statistically significant (p value<0.001), and in 640titre group 3.22 was statically significant. (P-Value<0.001).

The mean total count in the 80 titre was 7978.57 ± 2895.8, it was 7094.74 ± 2285.33 in 160 titre, 7465.38 ± 2693.46 in 320 titre group and 5864.29 ± 2118.71 in 640 titre group. The mean difference of total count 883.83 in 160 titre group was statistically not significant (p value <0.286), 513.19 in 320 titre group was statistically not significant (p value=0.577) and in 640 titre group 2114.29 was statically not significant. (P-value 0.067) The mean platelet in the 80 titre was 52333.98 ± 47613.68, it was 60631.71 ± 35052.44 in 160 titre, 58153.85 ± 18365.94 in 320 titre group and 21285.71 ± 10209.71 in 640 titre group.

The mean difference of platelet 8297.74 in 160 titre group was statistically not significant (p value= 0.468), 5819.87 in 320 titre group was statistically not significant (p value=0.648) and in 640 titre group 31048.29 was statistically not significant. (P-value=0.052) Chang K. et. al., in a study on 81 cases have shown only 10 (12.3%) had leukocytosis (leukocyte count ≥ 12.0 × 10⁶ cells/L) and 8 (13.6%) had leukopenia (≤ 4.0 × 10⁶ cells/L). Nearly 75% of the 81 cases had normal leukocyte counts at initial presentation. Thrombocytopenia, (platelet count < 100 × 10⁶ cells/L), was present in 30 (37.0%) cases. In addition, prolongation of activated partial thromboplastin time (36 of 53, 67.9%) was noted more commonly than prolongation of prothrombin time (16 of 51, 31.4%).[9] Among people with icterus, 10 (33.33%) people had 160 Overall Weil-Felix titre. The number of people with 320 and 640 Overall Weil Felix titre was 13 (43.33%) and 7 (23.33%) in people with icterus. Among people without icterus higher proportion of people had 80 and 160 titre. None of them had 320 and 640 titre. The mean total bilirubin in the 80 titre was 1.54 ± 0.51, it was 2 ± 1.11 in 160 titre, 4.41 ± 1.42 in 320 titre group and 6.63 ± 0.76 in 640 titre group. The mean difference of total bilirubin 0.46 in 160 titre group was statistically not significant (p value0.146), 2.87 in 320 titre group was statistically significant (p value<0.001) and in 640 titre group 5.09 was statistically significant. (P- Value<0.001).

In one study, of the 49 cases with available serum levels of total bilirubin, hyperbilirubinemia was found in 17 (34.7%) cases and 10 (20.4%) cases had total bilirubin levels ≥ 34.2 μmol/L.[9] The mean AST in the 80 titre was 140.43 ± 79.92, it was 285.89 ± 184.95 in 160 titre, 364.92 ± 89.69 in 320 titre group and 579.29 ± 106.26 in 640 titre group. The mean difference of AST 145.47 in 160 titre group was statistically significant (p value<0.001), 224.50 in 320 titre group was statistically significant (p value<0.001) and in 640 titre group 438.86 was statistically significant. (P- Value <0.001) The mean ALT in the 80 titre was 154.52 ± 112.16, it was 295.11 ± 200.03 in 160 titre, 375.23 ± 113.79 in 320 titre group and 484.14 ± 67.93 in 640 titre group.

The mean difference of ALT 140.58 in 160 titre group was statistically significant (p value 0.003), 220.71 in 320 titre group was statistically significant (p value<0.001) and in 640 titre group 329.62 was statically significant. (P- Value<0.001) The mean ALP in the 80 titre was 149.76 ± 64.42, it was 209.74 ± 85.09 in 160 titre, 254.54 ± 67.09 in 320 titre group and 356.43 ± 94.55 in 640 titre group. The mean difference of ALP 59.97 in 160 titre group was statistically significant (p value 0.015), 104.78 in 320 titre group was statistically significant (p value<0.001) and in 640 titre group 206.67 was statistically significant. (P- Value <0.001) In a study by Chang K. et. al., 19 (25.7%) had normal serum levels of AST or ALT (≤ 50 U/L) whereas serum ALT levels were five times greater than normal levels in 9 (12.2%) of 74 cases. Of 51 cases, 36 (70.6%) had elevated serum levels of alkaline phosphatase (> 126 U/L) and 35 (87.5%) of 40 cases had elevated levels of lactate dehydrogenase (> 250 U/L).[9]

The mean urea in the 80 titre was 40.86 ± 7.92, it was 46.53 ± 14.59 in 160 titre, 60.85 ± 15.77 in 320 titre group and 72.29 ± 14.67 in 640 titre group. The mean difference of urea 5.67 in 160 titre group was statistically not significant (p value 0.172), 19.99 in 320 titre group was statistically significant (p value<0.001) and in 640titre group 31.43 was statistically significant. (P- Value<0.001) The mean serum sodium in the 80 titre was 129.76 ± 7.49, it was 119.74 ± 10.28 in 160 titre, 107.54 ± 7.45 in320 titre group and 100.29 ± 7.7 in 640 titre group. The mean difference of serum sodium 10.03 in 160 titre group was statistically significant (p value<0.001), 22.22 in 320 titre group was statistically significant (p value<0.001) and in 640 titre group 29.48 was statistically significant. (P-Value<0.001).

The mean serum creatinine in the 80 titre was 1.32 ± 0.38, it was 1.67 ± 0.69 in 160 titre, 2.43 ± 0.64 in 320 titre group and 3.33 ± 0.87 in 640 titre group. The mean difference of serum creatinine 0.35 in 160 titre group was statistically not significant (p value=0.052), 2.10 in 320 titre group was statistically significant (p value<0.001) and in 640 titre group 2.01 was statically significant. (P-Value<0.001).

**Conclusion**

The mean hemoglobin values were statistically significantly lower in people with higher weilfelixtitre. All the liver function parameters and renal function parameters were...
significantly higher in people with weifeliixtire of 320 and 640, when compared to people with 80 weifeliixtire (p value < 0.05).

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