CLINICAL VERSUS LIGHT’S CRITERIA VERSUS SERUM PLEURAL FLUID ALBUMIN GRADIENT IN DIFFERENTIATING BETWEEN TRANSUDATIVE AND EXUDATIVE PLEURAL EFFUSIONS

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ABSTRACT

BACKGROUND
The first and important step is to classify pleural effusion into transudates and exudates. Traditionally, Light’s criteria are used to separate transudative from exudative pleural effusions. Light’s criteria misidentify about 20% of transudates as exudates, particularly in patients with heart failure on diuretics. In such cases serum pleural fluid albumin gradient is used to correctly identify transudates. Apart from Light’s criteria, the nature of the effusion can also be assessed by clinical examination. Very few studies have been done to evaluate the efficacy of clinical judgement with respect to Light’s criteria in determining the transudative or exudative nature of the pleural effusion.¹² Few studies have been done to evaluate the efficacy of Light’s criteria and serum pleural fluid albumin in determining transudative and exudative pleural effusions.¹³ But no studies were done to evaluate the efficacy of clinical judgement, Light’s criteria and serum pleural fluid albumin gradient in differentiating transudative and exudative pleural effusions.

MATERIALS AND METHODS
The observational, descriptive, cross-sectional study was conducted on 60 patients coming to the outpatient department as well as those admitted to the wards in Department of Pulmonary Medicine, Maharajah Institute of Medical Sciences during the period November 2013 to October 2015 with evidence of pleural effusion. Pleural fluid analysis is done to differentiate between transudative and exudative effusions, additionally Light’s criteria and serum pleural fluid albumin gradient is used for all samples.

RESULTS
Of the 60 patients, 50 were males and 10 were females. The age of the patients ranged between 15 - 85 years. Of the 60 effusions, 20 (33.3%) are transudates and 40 (66.6%) are exudates. Among the transudates CHF was the commonest disease and among exudates tuberculosis was the leading disease followed by synpneumonic effusion. Transudates were common in older age group, whereas exudates were common in younger and middle age group. Of 20 transudative effusions, clinical presumption could rightly classify all of them as transudates. Light’s criteria could classify only 12 of them as transudates and it misclassified 8 cases, of which (6 cases of CHF, 1 case of CKD, 1 case of Cirrhosis) on diuretic as exudates. SAPA could rightly classify all of them as transudates. Out of the 40 exudative effusions, clinical presumption could rightly classify 39 effusions, but misclassified 1 case of CHF as transudate. Light’s criteria could rightly classify all the 40 exudative effusions as exudates. SAPA could rightly classify 39 effusions, but misclassified 1 case of synpneumonic effusion as transudate.

CONCLUSION
The present study shows that the clinical criteria and SAPA are superior to Light’s criteria in identifying the transudative effusions (100% vs 60% vs 100%). Light’s criteria identified exudative effusions better than clinical criteria and SAPA (100% vs 97.5% vs 97.5%). So, in primary health centres where biochemical analysis is not available, clinical criteria can be used to separate transudates and exudates.

KEY WORDS
Transudates, Exudates, Clinical Diagnosis, Light’s Criteria, Serum Pleural Fluid Albumin Gradient, CHF, Cirrhosis, TB, Synpneumonic Effusion.

Apart from Light’s criteria, the nature of the effusion can also be assessed by clinical examination symptomatology and all available information including Chest X-ray, ECG, 2D-Echo, USG abdomen, blood and serum biochemistry etc. If clinical presumption can accurately identify the transudates and exudates, the cost and morbidity associated with diagnostic thoracentesis can be avoided.

Very few studies were done to evaluate the efficacy of clinical judgement with respect to Light’s criteria in determining the transudative or exudative nature of the pleural effusion.1,2 Few studies were done to evaluate the efficacy of Light’s criteria and Serum pleural fluid albumin in determining transudative and exudative pleural effusions.3,4 But no studies were done to evaluate the efficacy of clinical judgement, Light’s criteria and serum pleural fluid albumin gradient in differentiating transudative and exudative pleural effusions.

**Objectives**

To assess the ability of clinical versus Light’s criteria versus serum pleural fluid albumin gradient in differentiating between transudative and exudative pleural effusions.

**MATERIALS AND METHODS**

The observational, descriptive, cross-sectional study was conducted on 60 patients coming to the outpatient department as well as those admitted to the wards in Department of Pulmonary Medicine, Maharajah Institute of Medical Sciences during the period from November 2013 to October 2015 with evidence of pleural effusion. In this study, 60 consecutive patients with evidence of pleural effusion on chest radiographs were screened.

The patients who are seropositive for HIV and having any bleeding diathesis are excluded from the study.

All the 60 patients in the study met the inclusion criteria. Written informed consent was obtained from each study subject. Detailed history was taken, thorough clinical examination was done including CXR and clinical diagnosis was formed and then lab data like CBP, Blood and Urine biochemistry were obtained. After these each patient was subjected to diagnostic thoracentesis. ECG, 2D-Echo, USG abdomen and LFT were done in relevant cases.

**All the Patients Selected were Subjected to the following Investigations**

1. Sputum examination for M. Tuberculosis by direct smear for AFB on 2 consecutive days.
2. X-ray chest, PA view.
4. Pleural fluid analysed for total count, differential count, smear and culture for AFB, total protein, glucose and albumin levels, ADA levels, Gram staining and culture and sensitivity, malignant cytology by smear and cell block.
5. Pleural biopsy using Abrams pleural biopsy needle.
6. Serum protein and albumin were also sent.
7. Bronchial washings for malignant cytology and transbronchial biopsy, CECT thorax, FNAC of Lymph node (selected cases).
8. ECG, 2D-echo, USG-Abdomen, renal profile, LFT (in selected cases).
9. The following biochemical parameters were estimated and calculated: (1) The criteria of Light et al (namely pleural fluid/ serum protein ratio, pleural fluid/ serum LDH ratio, pleural fluid LDH concentration). In this study pleural fluid/ serum protein ratio alone is considered as according to Light’s criteria exudates can meet any one of the criteria. (2) Albumin gradient (Serum albumin concentration minus pleural effusion albumin concentration). The clinical presumption of the nature of the effusion (Transudate or Exudate) was based on all available information obtained before performing thoracentesis and was compared with that obtained from biochemical criteria.

**Diagnostic Thoracentesis**

A diagnostic thoracentesis is performed on every patient of pleural effusion with the fluid thickness of more than 10 mm on decubitus chest radiograph.

**Technique**

Once the site for the thoracentesis is identified, the skin surrounding the site is cleaned thoroughly with an antiseptic solution. Then local anaesthesia is given with 2% xylocaine to skin, subcutaneous tissue, muscles and parietal pleura. Then 20 cc syringe with 22-G needle is introduced through the intercostal space at the upper border of lower rib and 10 - 20 cc of pleural fluid is aspirated.

**Complications**

Vasovagal shock, pneumothorax, infections of the pleural space, haemothorax, chest pain and cough.

**Statistical Methods**

As the study is a descriptive study here, categorical data is represented as proportions and quantitative data as mean and standard deviations and data will be analysed using excel sheets. This data is used to calculate sensitivity, PPV and accuracy.

**RESULTS**

A total of 60 patients were evaluated. After necessary diagnostic workup, the effusions were definitively classified as transudates in 20 patients (33.3%) and as exudates in 40 patients (66.6%).

<table>
<thead>
<tr>
<th>Transudates</th>
<th>Exudates</th>
</tr>
</thead>
<tbody>
<tr>
<td>CHF</td>
<td>14</td>
</tr>
<tr>
<td>Cirrhosis</td>
<td>01</td>
</tr>
<tr>
<td>Anaemia and Hypoproteinaemia</td>
<td>01</td>
</tr>
<tr>
<td>CKD</td>
<td>04</td>
</tr>
<tr>
<td>Rheumatoid Effusion</td>
<td>01</td>
</tr>
<tr>
<td>CHF</td>
<td>01</td>
</tr>
</tbody>
</table>

| Total | 20 (33.3%) | Total | 40 (66.6%) |

**Table 1**

**Transudative Effusion**

CHF is by far the commonest transudative effusion (70%) in our study. TB (45%) and synpneumonic effusion (35%) together accounted 80% of exudative effusion in our study of 20 transudative effusions, clinical presumption could rightly classify all of them as transudates. Light’s criteria could classify only 12 of them as transudates and it misclassified 8 cases, of which (6 cases of CHF, 1 case of CKD, one case of Cirrhosis) on diuretic as exudates. Serum pleural fluid albumin gradient could rightly classify all of them as transudates.
**Exudative Effusion**

Out of the 40 exudative effusions clinical presumption could rightly classify 39 effusions, but misclassified 1 case of CHF as transudate. Light’s criteria could rightly classify all the 40 exudative effusions as exudates. Serum pleural fluid albumin gradient could rightly classify 39 effusions, but misclassified 1 case of synpneumonic effusion as transudate.

**Table 2**

<table>
<thead>
<tr>
<th>Final Diagnosis</th>
<th>Clinical Presumption</th>
<th>Light’s Criteria</th>
<th>SAPA</th>
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</thead>
<tbody>
<tr>
<td>CHF</td>
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<td>CHF</td>
<td>08</td>
</tr>
<tr>
<td>Cirrhosis</td>
<td>01</td>
<td>Cirrhosis</td>
<td>00</td>
</tr>
<tr>
<td>Anaemia and</td>
<td>01</td>
<td>Anaemia and</td>
<td>01</td>
</tr>
<tr>
<td>Hypoproteinaemia</td>
<td></td>
<td>Hypoproteinaemia</td>
<td></td>
</tr>
<tr>
<td>CKD</td>
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</tr>
<tr>
<td>Total</td>
<td>20</td>
<td>Total</td>
<td>12</td>
</tr>
</tbody>
</table>

**Table 3**

<table>
<thead>
<tr>
<th>Final Diagnosis</th>
<th>Clinical Presumption</th>
<th>Light’s Criteria</th>
<th>SAPA</th>
</tr>
</thead>
<tbody>
<tr>
<td>TB</td>
<td>18</td>
<td>TB</td>
<td>18</td>
</tr>
<tr>
<td>Malignancy</td>
<td>04</td>
<td>Malignancy</td>
<td>04</td>
</tr>
<tr>
<td>Synpneumonic Effusion</td>
<td>14</td>
<td>Synpneumonic Effusion</td>
<td>14</td>
</tr>
<tr>
<td>Pancreatic Effusion</td>
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<td>Rheumatoid Effusion</td>
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</tr>
<tr>
<td>CHF</td>
<td>01</td>
<td>CHF</td>
<td>07</td>
</tr>
<tr>
<td>Total</td>
<td>40</td>
<td>Total</td>
<td>48</td>
</tr>
</tbody>
</table>

**Table 4. Sensitivity, PPV and Accuracy of Clinical Criteria**

- Sensitivity for Transudates: 20/20 * 100 = 100%
- Sensitivity for Exudates: 39/40 * 100 = 97.5%
- PPV (Positive Predictive Value) for transudates: 20/21 * 100 = 95.2%
- PPV (Positive Predictive Value) for exudates: 39/39 * 100 = 100%
- Accuracy of Clinical Criteria: 59/60 * 100 = 98.3%

**Table 5. Sensitivity, PPV and Accuracy of Light’s Criteria**

- Sensitivity for Transudates: 12/20 * 100 = 60%
- Sensitivity for Exudates: 40/40 * 100 = 100%
- PPV (Positive Predictive Value) for transudates: 12/12 * 100 = 100%
- PPV (Positive Predictive Value) for exudates: 40/48 * 100 = 83.30%
- Accuracy of Light’s Criteria: 52/60 * 100 = 86.6%

**Table 6. Sensitivity, PPV and Accuracy of SAPA**

- Sensitivity for Transudates: 20/20 * 100 = 100%
- Sensitivity for Exudates: 39/40 * 100 = 97.5%
- PPV (Positive Predictive Value) for Transudates: 20/21 * 100 = 95.2%
- PPV (Positive Predictive Value) for Exudates: 39/39 * 100 = 100%
Accuracy of SAPA: 59/60 * 100 = 98.3%

**DISCUSSION**

The first step in the diagnosis of pleural effusion is classifying them into transudates and exudates. Light’s criteria are the standard for differentiating transudative and exudative pleural effusions, but main disadvantage is misclassification of 20% - 30% of transudates which were related to diuretic therapy as exudates.

In a study done by [Romero et al1, Burgess et al2, Vives et al3, Gazquez I et al] used a number of alternative criteria such as pleural fluid cholesterol levels, serum and pleural fluid albumin gradient, serum and pleural fluid protein gradient, pleural fluid and serum bilirubin ratio, cholinesterase ratio etc., which showed that Light’s criteria correctly classified exudates but misclassified transudates which were on diuretic therapy as exudates which were correctly identified by SAPA. All the criteria invariably require diagnostic thoracentesis, which is associated with certain risk and expenditure.

The nature of the pleural effusion can also be assessed by clinical examination based on the symptomatology and lab data such as X-ray Chest, ECG, Blood and Urine Biochemistry, Complete Blood Picture (CBP) with the added help of USG abdomen and 2D Echo. This study is aimed at studying the utility of clinical judgement, Light’s criteria and SAPA in determining the nature of pleural effusion and to determine its superiority or inferiority with respect to each other. A total of 60 patients were evaluated, of which 50 were males and 10 were females. The age of the patients ranged between 15 - 85 years. After necessary diagnostic workup, the effusions were definitively classified as transudates in 20 patients (33.3%) and as exudates in 40 patients (66.6%). Among the transudates, CHF was the commonest disease and among exudates tuberculosis was the leading disease followed by sympneumonic effusion. Transudates were common in older age group, whereas exudates were common in younger and middle age group.

**Analysis of Transudative Effusion**

Out of the 20 confirmed transudative effusions, clinical criteria could rightly classify all of them as transudates. Whereas Light’s criteria could classify only 12 of them as transudates. It misclassified 8 (6 had CHF, 1 had CKD and 1 had cirrhosis) of them as exudates, which were on diuretic therapy. These 8 cases were proved as transudates by SAPA (Serum and pleural fluid albumin gradient). The statistical analysis of the results showed that the sensitivity for transudative pleural effusion was 60% by Light’s criteria, 100% by clinical criteria and 100% by SAPA. It is because 8 out of 20 cases of transudates were misclassified as exudates, Light’s criteria lost accuracy. The sensitivity levels for transudates in the present study are comparable to the 77% sensitivity of Santiago Romero and Alfredro et al study, where a large group of 297 patients were studied between 1986-89.1

In the present study 8/20 (40%) of transudates were misclassified, which is close to 23% misclassified as transudates in the Santiago Romero et al study2 and 29% misclassified as transudates in Bielsa S, Porcel JM and Castellote J et al study.5 Another study by Peterman and Speicher who evaluated 495 patients with pleural effusions, 33% of CHF cases were wrongly classified as exudates. Bernard J Roth, MD; Thomas F O’Meara, MD studied 59 patients with effusions, of which 41 were exudates and 18 were transudates. Light’s criteria misclassified 5 transudative effusions as exudates, whereas serum pleural fluid albumin gradient correctly classified as having transudates.3

MC Dhar, S Chaudhuri, K Basu, TJ Sau, D Paland, K Mitra studied 50 patients of which Light’s criteria diagnosed all the 35 cases of exudates, but 2 cases of heart failure. 68 (Transudate) were misclassified as exudate. Using serum-effusion albumin gradient, all heart failure patients were correctly classified as having transudate.4

In the study of Burgess et al3 the gradient had a sensitivity and specificity of 87% and 92%, respectively. Dr. Padmasree Dantu and Dr. Srinivas Pusuluri5 studied 50 patients with pleural effusions, of which Light’s criteria correctly identified all the exudates but misdiagnosed 2 cases out of 6 transudates (cases of cardiac failure). By using albumin gradient of 1.2 g/dL or less to indicate exudate and values more than 1.2 g/dL to indicate transudate, all the patients (30 exudates and 20 transudates) were correctly diagnosed. Light’s criteria are accurate for identifying exudates, but not so much in the case of transudates. The serum-effusion albumin gradient is accurate equally for both exudates and transudates.

In the present study 20/20 (100%) of transudates were correctly identified as transudates by SAPA, which is close to Romero-Candeira and colleagues2 who studied 64 patients with transudative pleural effusions and reported that the Light’s criteria identified 75% correctly and the serum-pleural fluid albumin gradient identified 86% correctly. In Bielsa S, Porcel JM and Castellote J et al study5: 107 of 364 transudates (29%) caused by Congestive Heart Failure (CHF) were misclassified as exudates. In these 107 instances, a serum-pleural fluid albumin gradient greater than 1.2 gm/dl, which was only performed in 36 patients identified 83% correctly.

**Analysis of the Exudative Effusion**

In the present study, Light’s criteria proved to have 100% sensitivity for exudates with a PPV for exudates reaching 88.30%. On the other hand, clinical criteria and SAPA proved to have less sensitivity, i.e. 97.5% for exudates. In the Santiago Romero et al study [1986-1989], the sensitivity for exudates by Light’s criteria is 98%. It is the only study which examined the usefulness of clinical judgement alone and its added value to biochemical criteria. In their study, clinical criteria had a sensitivity of 94% for exudates. It is because 4 exudative effusions, due to trapped lung caused by TB and atelectasis due to lung cancer were wrongly classified as transudates. The exudative effusions in their study were caused by Pneumonia and Breast cancer. But in the present study, none of the exudates was misclassified by Light’s criteria.

Since the size of the study group is small future studies with a large group are required, so that better clinical criteria can be evolved for more perfect separation of transudates and exudates, so that unnecessary expenditure and morbidity
associated with thoracentesis can be avoided. It is also suggested that thoracentesis is not indicated in patients with CHF, Cirrhosis etc., on routine clinical practice and treatment of the systemic condition will resolve the effusion in most cases.

CONCLUSION
The results in the present study suggest that using clinical criteria, pleural effusions can be classified satisfactorily and in case of transudative effusions, there is no need of thoracentesis. So, in primary health centres where biochemical analysis is not available, clinical criteria can be used to identify transudates and exudates. The management can be directed towards underlying systemic condition and cost and morbidity associated with thoracentesis can be avoided as well.

REFERENCES