Lipedematous Scalp: A Case Report

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Introduction

Lipedematous scalp (LS) and Lipedematous Alopecia (LA) are two uncommon disorders, which have been clinically characterized by thickening of the subcutaneous layer of the scalp and a boggy scalp on palpation. While there is no hair abnormality in LS, shortening of hair and hair loss are observed in the thickened part of the scalp in LA. Nevertheless, the etiology and pathogenesis of both disorders are unknown and no approved association with any other diseases is reported.

LS was first described by Cornbleet in 1935. The patient was a 44-year-old black woman who had complaints about thickening and softening of her scalp which felt like a cotton. A few years later, Lee et al. reported the second case in a Korean woman. Since then, ten cases, all in females, have been reported (Table 1).

In 1965 and for the first time, Coskey et al. introduce LA as a new term to describe the disorders observed in two black females aged 28 and 75 who were examined for the shortening of hair and thickening of the subcutaneous layer of the scalp. So far, the total reported number of LA cases is 18, out of whom 14 cases were females and 4 cases were males. Considering the fact that the main pathology observed in both disorders are obvious increase in the diameter of the lipid tissue, it is most probable that LS and LA are the same or two manifestations of one disease.

Case Report

The patient was a 42-year-old woman who was visited for the softening of scalp and a continuous headache. The problem had started two years ago with a headache which became worse and continued for a longer period each time. Gradually, the vertex became soft and thickened. Even though she had been examined by different physicians, her problem was gradually aggravated and she had complaints about headaches all over her head as well as a feeling of pain and stiffness in her neck when visited. She had no history of taking medications, underlying systemic diseases or head trauma (Figure 1).

Considering the patient's age, hair density was normal and no apparent abnormality was seen. The appearance of the scalp skin seemed normal. When pressing the scalp down, it felt spongy and soft and became depressed but returned to the same level after the pressure was removed. The patient had tenderness and a severe pain during the examination. The spongy feeling was observed in the whole scalp but in the margins of the hair growth. The patient had no family history for such diseases and her children were normal. Blood biochemical analysis was normal.
Biopsy of the scalp showed normal epidermis, the diameter of dermis was obviously reduced while the diameter of lipid layer had extremely increased and as a result had reached the limit of superficial dermis. Mucin deposition was not observed by Alcian blue staining. Hair follicles and Adnexes were normal (Figure 2).

Considering the patient’s complaints i.e. headaches and neck stiffness and pain, she was referred to consult a neurologist and a neurosurgeon. CT scan and MRI were normal. Obstruction and CSF leakage, as possible causes of headaches and scalp swelling, were ruled out.

The maximum thickness of scalp measured by MRI was 17 mm which was observed in the vertex area. The thickness of other parts was slightly less (Figure 3, 4).

**Discussion**

LS is an uncommon disorder which is mainly characterized by spongy swelling and increase of the scalp thickness without any noticeable hair loss. When hair loss is included, the term LA is applied. Most of the cases reported so far are black women, although reports of men and white patients are increasing. As a consequence, the previous opinion about these disorders being limited to a specific race is now questionable. Nevertheless, the higher number of female patients could be related to the role of female hormones in the pathogenesis of the disease.

The thickness of scalp is naturally 5-6 mm². For the cases reported so far, the thickness is ranging between 10 to 18 mm measured with different methods including a needle introduced into the scalp, ultrasonography, MRI and CT scan which is about 2-3 times more than the normal range. For the case reported in this article, the scalp thickness measured by MRI was 17 mm which is 3 times more than normal.

To date, it is not known if the movement of the lipid tissue toward the skin surface is compensatory.

### Table 1: Reported cases of Lipedematous scalp (LS) and Lipedematous Alopecia (LA)

<table>
<thead>
<tr>
<th>Author, year</th>
<th>Age, sex, race</th>
<th>Duration</th>
<th>Association</th>
<th>Scalp thickness</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Lipedematous scalp</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cornbleet, 1935</td>
<td>44,F,B</td>
<td>6 years</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Lee, 1994</td>
<td>32,F, A</td>
<td>3 years</td>
<td>-</td>
<td>10.7</td>
</tr>
<tr>
<td>Scheufler, 2003</td>
<td>51,F,W</td>
<td>5 years</td>
<td>Fever, headache, Syncopeal attacks</td>
<td>15 mm</td>
</tr>
<tr>
<td>Bukhari, 2004</td>
<td>51,F,W</td>
<td>6 months</td>
<td>Hypercholesterolemia, Supraventricular tachycardia</td>
<td>19.2 mm</td>
</tr>
<tr>
<td>Martin, 2005</td>
<td>48,F,W</td>
<td>2 months</td>
<td>-</td>
<td>10.8 mm</td>
</tr>
<tr>
<td>Ifihi, 2005</td>
<td>55,F,B</td>
<td>?</td>
<td>DLE</td>
<td>10.15</td>
</tr>
<tr>
<td>Darlin, 2006</td>
<td>10,F,A</td>
<td>6 months</td>
<td>Asthma, headache</td>
<td>9.8 mm</td>
</tr>
<tr>
<td>Yasar, 2007</td>
<td>62,F,W</td>
<td>50 years</td>
<td>positive family history</td>
<td>18 mm</td>
</tr>
<tr>
<td>Mansur, 2007</td>
<td>46,F,W</td>
<td>16mm</td>
<td>Nevus lipomatosus</td>
<td>13 mm</td>
</tr>
<tr>
<td>Kazak, 2008</td>
<td>50,F,W</td>
<td>6 months</td>
<td>Hypertension</td>
<td>10.8 mm</td>
</tr>
<tr>
<td>Shabanzadeh, 2008</td>
<td>42,F,A</td>
<td>2 years</td>
<td>Headache, scalp tenderness, Neck stiffness</td>
<td>17 mm</td>
</tr>
</tbody>
</table>

| **Lipedematous alopecia** | | | | |
| Coskey, 1961 | 28,F,B | 2 years | Diabetes mellitus | 15 mm |
| Curtis, 1964 | 62,F,B | 15 years | Skin and joint hyperelasticity | 15 mm |
| Kane, 1998 | 49,F,B | 6 months | - | 12.6 mm |
| Foi, 2000 | 18,F,B | 7 years | Renal insufficiency | 9 mm |
| Rejiama, 2000 | 30,M,A | 7 years | 16 mm |
| Tiscornia, 2002 | 69,F,W | 6 months | 10 mm |
| Mohsin, 2004 | 8,F,W | 1 year | 11 mm |
| Martin, 2005 | 77,F,W | 1 year | Sjogren's syndrome | 9.2 mm |
| High, 2005 | 57,F,B | 10 years | DLE | 12-15 mm |
| Piraccini, 2006 | 48,M,W | 10 years | Androgenetic alopecia | 11 mm |
| Yasar, 2007 | 45,F,W | 4 years | - | 12 mm |
| Farmanje, 2007 | 45,F,A | 5 years | Freckles, café-au-lait spots | 10.7 mm |
| Tipp, 2008 | 67,F,W | 1 year | - | ? |

A: Asian  B: Black  F: Female  M: Male  W: White
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Figure 1: Clinical manifestations of lipedematous alopecia: boggy scalp without hair loss

Figure 2: Histopathology view; the diameter of dermis was obviously reduced while the diameter of lipid layer had extremely increased (H&E *10)

Figure 3: MRI imaging demonstrated no abnormality in brain and thickness of scalp measured as 17 mm

or reactive. The important pathology in a report by Fair et al. is demonstrated to be lipid tissue edema together with disruption and degeneration of lipid tissue without hypertrophy. Martin, Moravvej and Tiscornia observed dilated lymphatic vessels. Since lymphangiectasia is only reported in patients with LA it could be the cause of hair loss seen in such patients. The thickening of lipid tissue and its hyperplasia have also been reported in some cases. Coskey reported free-floating lipid droplets in subcutaneous layer. Mucin deposition in superficial dermis was observed in one case. Linear telangiectasia is also described through dermoscopy.

In some cases, scalp thickening is localized while it is widespread in others. In our patient, all hair-bearing areas of the scalp were asymmetrically thickened.

So far, four male patients have been reported. This indicates the effect of X chromosome, or
female hormones on the etiology of the disease. Meanwhile, the number of patients having different races; 9 black, 14 white and 6 Asian patients, shows that in spite of previous views, these diseases are prevalent in all races and there is no limitation to a specific race.

Concomitant diseases include diabetes mellitus, asthma, sjogren’s syndrome, freckles and café-au-lait on the trunk, androgenetic alopecia, skin hyperelasticity and joint laxity, renal insufficiency, nevus lipomatosus, hyperlipidemia, hypertension. In one case, LA occurred in a patient with DLE after several years.

It seems that these associations are accidental and to date, there is no evidence of any specific disease associated with LS and LA. In addition, Lipedema of the lower legs can occur in the shin, leg and buttocks (but not foot) and is painful with a histology similar to LS and LA but no concomitance has been reported yet.

The progression of LS and LA in most patients is gradually continued for a period of few years and is stabilized afterwards. It is also interesting to know that so far, in none of the reported cases of LS, hair loss has been observed in the lipedematous areas even several years after the diagnosis which shows that LS is not a precursor or an introduction to LA.

Despite the use of systemic and intralesional steroid for the treatment of these disorders, a permanent and satisfactory result has not been obtained so far and the treatment is symptomatic.

References
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