Lichen sclerosus et atrophicus and sexual abuse

Shirley A Warrington, Camille de San Lazaro

Abstract

Aims—The aetiology of lichen sclerosus et atrophicus (LSA) is unknown. A series of 42 cases of this uncommon condition is reported. The aim of this study was to identify associations of LSA and document the association with sexual abuse.

Methods—Information about the patients was obtained by retrospective case note review and some patients were contacted by telephone for further information.

Results—In 12 cases there was evidence of sexual abuse. The abused group were slightly older than the non-abused group but were similar in all other respects. All three patients who presented over the age of 12 years had evidence of sexual abuse. Genital trauma was recalled by the patient or found at examination in 17 cases. Evidence of autoimmunity was present in five cases. Positive microbiological isolates were obtained in 18 cases. In only 11 cases were there no associated factors. The symptoms of LSA started between the ages of 3 and 7 years in most patients. The usual symptoms were related to genital skin involvement, and symptoms related to bladder and bowel function were common (50%).

Conclusion—In this large series of paediatric LSA, associations with trauma, autoimmunity, and infection were noted. There was a high rate of coexisting sexual abuse with LSA, possibly due to genital trauma.

(Arch Dis Child 1996;75:512–516)

Keywords: lichen sclerosus et atrophicus, sexual abuse.

Lichen sclerosus et atrophicus (LSA) is a chronic condition of the epithelium and dermis which is characterised by ivory or white shiny macules and papules that form homogeneous hypopigmented areas. The affected skin shows a tendency to fine wrinkling. A characteristic 'figure of eight' pattern is seen when both the vulva and the perianal areas are involved. Involved skin is sharply demarcated from surrounding normal skin. Hymenal tissue is uninvolved and should appear smooth and symmetrical.

The condition may affect all areas of the body, at all ages, and in both sexes. It is most frequently found in the fifth and sixth decades, but also affects prepubertal girls and premenopausal women. An increased risk of malignancy (squamous cell carcinoma) has been found in adult women and long term surveillance is suggested. Boys as well as men may be affected. In boys LSA may cause phimosis. Adult men are also affected on the penis or elsewhere, and in older men it is the commonest cause of balanitis xerotica obliterans (a possibly premalignant condition). In young girls LSA may be asymptomatic or the lesions may itch, become infected, or cause pain and bleeding. Common presenting symptoms are vulvovaginitis, pruritus, bleeding, blistering, pain on defecation and urinary symptoms. LSA is managed with good hygiene, bland emollients, and mild corticosteroids. The aim of treatment is generally to control symptoms, and the condition often improves at puberty. The aetiology of LSA is unknown. In some patients there is evidence to implicate genetic, hormonal, and autoimmune related factors. It is well recognised that lichen sclerosus may occur at sites of trauma.

LSA may be mistaken for acute trauma because the involved skin is friable and prone to bleeding from trivial trauma. Unexplained or unexpected bleeding or purpura may suggest that a significant injury has occurred. This has resulted in patients with LSA being the subject of child sexual abuse investigations. The differential diagnosis of abuse and LSA is an important issue. We therefore evaluated the case records of all our patients with LSA for evidence of abuse and other associated factors.

Patients and methods

Forty two patients with the established diagnosis of LSA were identified by retrospective review of 1617 readily accessible case notes. These notes are held securely by one of us (CSL) for a variety of reasons. This population represented a substantial proportion of the patients referred to the paediatric forensic team over a seven year period (1989–95) and all patients had undergone a genital examination. These patients were referred to one of the authors (CSL) for assessment of suspected child sexual abuse, advice on management of LSA, or investigation of genital symptoms.

Figure 1 Age distribution at diagnosis.
The assessment for children referred to the clinic usually involves a long consultation with a consultant paediatrician (up to 90 minutes) and includes a clinical history, general physical examination, and a genital examination. Some patients were also seen at a special paediatric infectious diseases clinic where appropriate infection screening was carried out as previously described.

The case notes of the 42 cases of LSA were carefully reviewed for demographic details, source of referral, presenting symptoms, further details of the history, and family history. The main reasons for referral were divided into three categories: management of LSA, advice on genital symptoms, and suspected child sexual abuse. Evidence was sought for associated factors, particularly infection, autoimmune disease, and trauma, including sexual trauma from abuse. Details of examination findings and the person making the initial diagnosis were documented. In those cases in which a child sexual abuse inquiry had occurred, case conference minutes were studied and outcome data sought. Additional information about past medical history and family history was obtained from some parents when this was practicable.

**Results**

Forty two girls with LSA were identified by case note review. All but two were white. Age of symptom onset ranged from a few months to 13 years. Age at diagnosis ranged from 3.9 to 14.0 years with a mean age of 7.5 years (fig 1).

**REFERRAL SOURCES**

General practitioners referred 16 cases, paediatricians 17 cases, and child protection agencies five cases. The remaining four were referred by other specialists, including one by a cardia surgeon, when lesions suggesting sexual trauma were noted at urethral catheterisation before surgery.

Diagnosis before referral had been made by three general practitioners and by 10 of the referring paediatricians. In one case a retrospective diagnosis was made from photographs taken in a previous child abuse inquiry. This child also showed gross hymenal damage (fig 2).

The reasons for referral differed, general practitioners referring for symptom assessment, and paediatricians and child protection agencies referring because of suspected child sexual abuse.

**ASSOCIATED FACTORS (fig 3)**

**Injury**

A history of accidental trauma or findings of abusive genital trauma were associated with the development of LSA in 17 children. The history of accidental trauma, obtained in six cases, took the form of a variety of memorable falls recalled by the child or parent, preceding the development of LSA. One of these children was also sexually abused. Details of the 12 children in whom evidence of sexual abuse was found are shown in table 1. Of these, six made allegations resulting in convictions in two. The other six showed damaged hymens and other physical and behavioural signs. Concern about three other children, one with a short tear to

### Table 1  Details of cases with evidence of sexual abuse

<table>
<thead>
<tr>
<th>Case No</th>
<th>Age (years)</th>
<th>Allegation</th>
<th>Findings</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>6</td>
<td>Yes</td>
<td>Thickened scar across hymen. Disrupted hymen</td>
<td>Conviction obtained</td>
</tr>
<tr>
<td>2</td>
<td>7</td>
<td>Yes</td>
<td>Healed posterior transection and scarred attenuated left free edge of hymen. Shown in fig 2</td>
<td>Conviction obtained. Father convicted of indecent assault on child. Grandfather—offences of indecent exposure</td>
</tr>
<tr>
<td>3</td>
<td>13</td>
<td>Yes</td>
<td>Bruising inner thighs—no adequate explanation</td>
<td>Violence in home. Brother physically abused and showing sexualised behaviour</td>
</tr>
<tr>
<td>4</td>
<td>7</td>
<td>Yes</td>
<td>Intact hymen. Anal laxity with distortion</td>
<td>Father arrested for allegations of sexual abuse against uncle, aunt, and another boy</td>
</tr>
<tr>
<td>5</td>
<td>4</td>
<td>No</td>
<td>Attenuated hymen with rolled edge. Disruption and nodular scarring</td>
<td>Proven in court. Withdrawn behaviour. Father had record of indecent exposure</td>
</tr>
<tr>
<td>6</td>
<td>6</td>
<td>No</td>
<td>Hymen tethered to labia minora? Post-traumatic adhesions</td>
<td>Highly sexualised behaviour. Nightmares and sleepwalking</td>
</tr>
<tr>
<td>7</td>
<td>7</td>
<td>No</td>
<td>Thin adhesion from inferior free edge of minora to the hymen</td>
<td>Spent long periods unsupervised with known abuser (stepgrandfather). Mother also sexually abused by him. Police investigation</td>
</tr>
<tr>
<td>8</td>
<td>7</td>
<td>Yes</td>
<td>Nodular hymen with cleft</td>
<td>Exposed to inappropriate sexual activity</td>
</tr>
<tr>
<td>9</td>
<td>13</td>
<td>No</td>
<td>Transsected hymen with scarring</td>
<td>Possible sexual abuse in stepfather's previous family</td>
</tr>
<tr>
<td>10</td>
<td>7</td>
<td>No</td>
<td>Disrupted hymen</td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>9</td>
<td>No</td>
<td>Scar at midline of hymen with deep cleft and attenuation</td>
<td></td>
</tr>
<tr>
<td>12</td>
<td>14</td>
<td>Yes</td>
<td>Widely disrupted hymen</td>
<td></td>
</tr>
</tbody>
</table>

![Figure 2](http://adc.bmj.com/)

*Figure 2  Case 2: 7 year old girl alleging penetration by two family members. A guilty plea was obtained from one and conviction of the other. Healed posterior transection and scarred attenuated left free edge of hymen. Pallor, friability, and bleeding from lesions of LSA.*
the hymen, one exposed to a known offender, and one with genital herpes simplex, has not so far developed into a finding of sexual abuse.

**Autoimmune disease**

Thirty one case notes yielded information about autoimmune disease. One child had extensive morphea and two had arthritis. Three other children were serum positive for autoantibodies (one Rh factor, one antimitochondrial, and one thyroid antibodies). Of the 33 cases where a full family history had been taken, nine were positive for a first or second degree relative (27% of those evaluated).

**Infection screening**

Results were available in 32 cases of which 21 isolates were seen in 18 children. Warts were not seen in any child. Six children had more than one infectious agent identified. Streptococcus, staphylococcus, Escherichia coli, and bacteroides were isolated from both groups of children. Candida and gardnerella were only isolated in the group of children where abuse had occurred (table 2).

**SYMPTOM HISTORY**

The frequency of presenting symptoms differed between the 12 children who were known to have been abused and the 30 children where abuse was not found.

**Table 3 Frequency of presenting symptoms in non-abused and abused cases**

<table>
<thead>
<tr>
<th>Presenting symptoms</th>
<th>Incidence (%)</th>
<th>Non-abused (n=30)</th>
<th>Abused (n=12)</th>
<th>All (n=42)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vulvovaginitis</td>
<td>22 (73)</td>
<td>6 (50)</td>
<td>28 (67)</td>
<td></td>
</tr>
<tr>
<td>Bleeding</td>
<td>13 (43)</td>
<td>3 (25)</td>
<td>16 (38)</td>
<td></td>
</tr>
<tr>
<td>Pruritis</td>
<td>9 (30)</td>
<td>1 (8)</td>
<td>10 (24)</td>
<td></td>
</tr>
<tr>
<td>Urinary</td>
<td>8 (27)</td>
<td>4 (33)</td>
<td>12 (29)</td>
<td></td>
</tr>
<tr>
<td>Bowel symptoms</td>
<td>7 (23)</td>
<td>4 (33)</td>
<td>11 (26)</td>
<td></td>
</tr>
<tr>
<td>Purpura/blisters</td>
<td>6 (20)</td>
<td>1 (8)</td>
<td>7 (17)</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>1 (3)</td>
<td>1 (8)</td>
<td>2 (5)</td>
<td></td>
</tr>
<tr>
<td>None</td>
<td>2 (7)</td>
<td>2 (17)</td>
<td>4 (10)</td>
<td></td>
</tr>
</tbody>
</table>

Abused children with LSA presented less often with genitourinary symptoms and also presented later (abused children with vulvovaginitis 50%, others 73%; abused children with bleeding 25%, others 43%). Some 33% of the abused group presented first with bowel symptoms (pain on defecation and constipation) compared with 23% of the others. Only 8% of abused girls were diagnosed with LSA before 6 years of age, compared with 23% in the non-abused group. About 25% of the abused girls presented with LSA for the first time after the age of 12 years. All the non-abused girls presented before this age (fig 4).

Table 3 shows the frequency of presenting symptoms. The majority of patients presented with symptoms related to genital skin involvement, soreness, bleeding, discharge, and pruritis. Symptoms relating to bladder and bowel function were seen in 50% of cases. These symptoms, which included dysuria, dribbling, and stool withholding, appeared to relate to the presence of open tender areas of fissuring and resolved rapidly with treatment. Some patients were asymptomatic, and LSA was discovered incidentally in four cases.

The reasons for referral were compared with outcome diagnosis. Of 12 children referred for symptoms alone, one was found to have LSA coexisting with sexual abuse. Of 12 girls referred for suspected sexual abuse, three were found to have LSA alone and nine were confirmed to have been abused. Eighteen children were referred with the established diagnosis of LSA, and in two of these coexisting sexual abuse was found (fig 5).

**Discussion**

The incidence of LSA remains unknown, even in adults. Wallace suggested that the frequency varies between one in 300 to less than one in 1000 of all new patients in a general hospital, depending on the liaison between the skin and gynaecological departments. The disease is much commoner in adults than in children: in a series of 290 cases only 7% occurred below the age of 16. No surveys have been carried out to establish how commonly the condition occurs. In Newcastle, the Lindisfarne Centre is a tertiary referral centre with a specialist population. Seventeen of our cases came from Newcastle with a population of 280 000. These did not represent all the LSA in Newcastle.
Figure 5  Reason for referral compared with final diagnosis.

<table>
<thead>
<tr>
<th>Reason for referral:</th>
<th>Management of LSA</th>
<th>Assessment of symptoms</th>
<th>Suspected abuse</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>18</td>
<td>12</td>
<td>12</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Final diagnosis:</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>LSA only</td>
<td>16</td>
<td>11</td>
<td>3</td>
</tr>
<tr>
<td>LSA and abuse</td>
<td>2</td>
<td>1</td>
<td>9</td>
</tr>
<tr>
<td>Abuse only</td>
<td>0</td>
<td>0</td>
<td>N/A</td>
</tr>
</tbody>
</table>

LSA and abuse only for referral compared to Berth-Jones in anitis often atrophic vulvitis, for patients diagnosed may influence of maternal are than rather have referred with population attempted have found to be combined and is likely to relate to local topical steroids to treat the LSA. The presence of candida and gardnerella has been associated with sexual abuse and our findings support this suggestion. No warts were found in any of our cases, in spite of the finding of Kiene et al of four of 18 cases of LSA being infected with human papillomavirus type 16. More specific testing for wart virus in children with LSA may provide further evidence of this possible association. Early literature notes the occurrence of the Koebner phenomenon in patients with LSA. Chronic frictional injury from waist bands and bra straps is well known to precipitate the appearance of lichen sclerosus lesions. Harrington reports extragenital lesions occurring at sites of acute trauma—for example, oven burns. Six of our patients could recall a definite accidental injury to the genital area. Some patients diagnosed then. This is the peak age for atrophic vulvitis, when oestrogen concentrations are at their nadir. No cases of LSA were diagnosed below the age of 3 years. The influence of maternal oestrogens lasts until the age of 2 years and this may be a protective factor explaining the rarity of LSA below this age.

The peak age for diagnosis of LSA in our series was found to be 6 to 8 years with 64% of patients diagnosed then. This is the peak age for atrophic vulvitis, when oestrogen concentrations are at their nadir. No cases of LSA were diagnosed below the age of 3 years. The influence of maternal oestrogens lasts until the age of 2 years and this may be a protective factor explaining the rarity of LSA below this age.

All the 42 patients in this series were female. The condition is known to be much commoner in girls than boys. Histological evidence of balanitis xerotica obliterans (as the condition is often termed in the male) can frequently be found after circumcision for scarring phimosis. Thus boys with LSA are more likely to be referred to urologists or surgeons than to paediatricians. Berth-Jones et al reported that two thirds of their patients presented with soreness, a figure similar to our numbers presenting with vulvovaginitis (a syndrome including soreness and discharge). The Leicester group found that urinary symptoms occurred in 66% of their patients and bowel symptoms in 83% as compared with half that incidence of symptoms reported by our patients. Chernosky et al found that pruritus was described in 49% of cases, whereas we found that only half that number of patients reported this symptom. The differing presentation of genital symptoms and bowel symptoms in children with LSA and abuse raises the possibility that symptoms, particularly relating to the genital area, may be unreported by children or families involved in sexual abuse, whereas bowel symptoms may be perceived as less threatening and also tend more to mandate clinical attention.

Associated factors were commonly found with LSA in our patients. It has been suggested that genetic factors are involved in the aetiology of LSA. We found two siblings with the condition. Details of the family history of autoimmune disease showed that the affected family members were generally the patient’s grandparents reflecting the younger age of the patient’s parents compared with the usual age of onset of these diseases. This is similar to the incidence of autoimmune disease in relatives previously reported and adult studies have found a high rate of autoantibodies in patients with LSA. One girl had patchy morphea. The frequency of occurrence of morphea in combination with LSA has been described as statistically significant.

Positive microbiological isolates have not previously been reported in LSA. In this series positive swabs were obtained in 18 cases (56% of those tested) and negative ones in 14 cases. In a comparable normal population the rate of carriage of isolates is unknown. However, these figures are considerably higher than those found in a previous study carried out by this department where only 4% of children had positive isolates. The high rate of positive isolates is likely to relate to secondary colonisation of compromised damaged skin in LSA. We have no evidence that these positive isolates are causative, but all positive isolates were treated, as potentially the children required the application of local topical steroids to treat the LSA.

Six of our patients could recall a definite accidental injury to the genital area. Some
authors have suggested that LSA is secondary to sexual abuse26-28 while others have disputed this.29 In 12 of our patients there was evidence of such abuse. In one of the children who had been abused there was also a history of accidental injury preceding by months the development of the disease. The changes seen in LSA in the abused group may be triggered by intermittent frictional trauma to the genital area. However, we believe that the recurrent presence of foreign substances, like semen, saliva, or lubricants, should be also considered a possible trigger.

There are numerous reports of LSA being mistaken for childhood sexual abuse.14-18 In our study this occurred in three cases. The issue raised by these cases remains, as always, that of the role of physical findings in formulating an opinion about abuse. The sexually abused victim commonly demonstrates no physical signs, and such findings as there are rarely reflect the degree of trauma that children report. Most detected physical abnormalities at the genital area are neither amenable to dating nor to understanding of chronicity. A single event of digital penetration in a small child could be reflected in gross abnormality years later; frequent, painful friction between the labia may result in minimal, if any, physical abnormality. These limitations underpin difficulties practitioners experience when an incidental finding of physical abnormality in an apparently well adjusted symptom free child is made. In practice it is rarely easy to design a meaningful child protection process in these situations. Our clinical experience of over 2500 sexually abused children suggests that it is unusual to see superficial skin tears, bleeding, and bruising of external structures in acute sexual assault with no disclosure or signs of hynenal trauma. This is an unusual constella-

1 Wallace HJ. Lichen sclerosus et atrophicus. Transac
tions of St
2 Meyrick Thomas RH, Ridley CM, McGilligan DH, Black
MM. Lichen sclerosus et atrophicus and autoimmunity—a
3 Ridley CM. Lichen sclerosus et atrophicus. BMJ 1987;294:
1295-6.
4 Tremaine RDL, Miller RAW. Lichen sclerosus et atrophicus.
5 Clark JA, Muller SA. Lichen sclerosus et atrophicus in children:
6 Berti-Jones J, Graham Brown RAC, Burns DA. Lichen
sclerosus et atrophicus—a review of 15 cases in young girls.
7 Newbrun ES, Rubenstein ML, Kennedy CTS. The developmen-
ton of lichen sclerosus et atrophicus in monogynous twin
8 Davidson DC, Clarke MDB, Keen HB. Lichen sclerosus et
atrophicus in children misdiagnosed as sexual abuse
9 Shrier JA, Ray MC. Familial occurrence of lichen sclerosus
10 Friedrich EG, Kalra PS. Serum levels of sex hormones in
vulvar lichen sclerosus, and the effect of topical testoster-
11 Goolamali SK, Barnes EW, Irvine WJ, Shuster S. Organ-
specific antibodies in patients with lichen sclerosus. BMJ
12 Harrington CI, Dunsmore IR. An investigation into the
incidence of autoimmune disorders in patients with lichen
13 Meyrick Thomas RH, Ridley CM, Black MM. The associ-
ation of lichen sclerosus et atrophicus and autoimmune
14 Lavery HA, Pinkerton JHM, Callender M. The associ-
ation of lichen sclerosus et atrophieus and primary herpe-
15 Chernosky ME, Derbes VJ, Burks JW. Lichen sclerosus et
16 Handfield-Jones SE, Hindle FRJ, Kennedy CTC. Lichen
sclerosus et atrophicus in children misdiagnosed as sexual
17 Priestley BL, Bleehen SS. Lichen sclerosus et atrophicus in
children misdiagnosed as sexual abuse [letter]. BMJ 1987;
295:211.
18 Jowsey C, Kirby P, Fuqasy DM. Genital lichen sclerosus
19 Bays J, Carol J. Genital and anal conditions confused with
20 de San Lazaro C. Making paediatric assessment in
suspected sexual abuse: a therapeutic experience. Arch
21 Steele AM, de San Lazaro C. Transcultural care for
sexually transmissible organisms. Arch Dis Child 1994;71:
425-7.
22 Chalmers RJG, Burton PA, Bennett RF, Goring CC, Smith
PJB. Lichen sclerosus et atrophicus. Arch Dermatol
1984;120:1025-7.
23 Ridley CM. Genital lichen sclerosus (lichen sclerosus et
atrophicus) in childhood and adolescence. J R Soc Med
1993;86:69-75.
24 Argenti AC, Lachman PJ, Hanso D, Bass D. Sexually trans-
mittted diseases in children and evidence of sexual abuse.
25 Kene P, Milde-Langosch K, Löng T. Human papillomavi-
rus infection in vulvar lesions of lichen sclerosus et atrophie-
26 Harrington CI. Lichen sclerosus [letter]. Arch Dis Child
1990;65:335.
27 Priestley BL, Bleehen SS. Lichen sclerosus and sexual abuse
28 de San Lazaro C. Lichen sclerosus [letter]. Arch Dis Child
1990;65:1184.
29 Berti-Jones J, Graham Brown RAC, Burns DA. Lichen
Lichen sclerosus et atrophicus and sexual abuse.

S A Warrington and C de San Lazaro

*Arch Dis Child* 1996 75: 512-516
doi: 10.1136/adc.75.6.512

Updated information and services can be found at:
http://adc.bmj.com/content/75/6/512

**Email alerting service**

Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

**Notes**

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/