Clinicopathological Analysis of de Quervain’s Disease

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Abstract

Excised extensor retinacula of the first compartment and tenosynovium from 35 patients (6 men and 29 women) with de Quervain’s disease were examined by light and electron microscopy to investigate the pathogenic mechanism. The patients, aged from 22-78 years, averaging 50 years, comprised the study group. Two hundred and thirty-two specimens from cadavers of 95 men and 75 women were macroscopically examined as the control. In the study group, the extensor retinaculum and tenosynovium were macroscopically thickened, and were histologically classified into 4 groups based on presence or absence of septum, and the location of retinacular thickening. Morphologically, the thickening of the tenosynovium and retinaculum was due to fibrosis in every layer, although fibroses were seen mainly in the middle layer. The ratios of proliferation of fibroblasts, myxoid changes and/or hyaline degeneration, and vascular proliferation were varied between layers. Minimal round cell infiltration was found in the retinaculum as well as in the tenosynovium. The results also indicate that the Iwahara-Nozue test can be used to accurately predict relatively greater thickening of the retinaculum on the extensor pollicis brevis side. Based on clinicopathological analyses, it appears that de Quervain’s disease is induced not only by extrinsic factors such as superficial friction but also by intrinsic factors.

KEYWORDS: de Quervain’s disease, stenosing tenosynovitis, ultrastructural study, histopathology

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Clinicopathological Analysis of de Quervain’s Disease

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Excised extensor retinacula of the first compartment and tenosynovium from 35 patients (6 men and 29 women) with de Quervain’s disease were examined by light and electron microscopy to investigate the pathogenic mechanism. The patients, aged from 22–78 years, averaging 50 years, comprised the study group. Two hundred and thirty-two specimens from cadavers of 95 men and 75 women were macroscopically examined as the control. In the study group, the extensor retinaculum and tenosynovium were macroscopically thickened, and were histologically classified into 4 groups based on presence or absence of septum, and the location of retinacular thickening. Morphologically, the thickening of the tenosynovium and retinaculum was due to fibrosis in every layer, although fibroses were seen mainly in the middle layer. The ratios of proliferation of fibroblasts, myxoid changes and/or hyaline degeneration, and vascular proliferation were varied between layers. Minimal round cell infiltration was found in the retinaculum as well as in the tenosynovium. The results also indicate that the Iwashara-Nozue test can be used to accurately predict relatively greater thickening of the retinaculum on the extensor pollicis brevis side. Based on clinicopathological analyses, it appears that de Quervain’s disease is induced not only by extrinsic factors such as superficial friction but also by intrinsic factors.

Key words: de Quervain’s disease, stenosing tenosynovitis, ultrastructural study, histopathology

In 1895, de Quervain described a previously unrecognized condition affecting the tendon sheath and tendons of the extensor pollicis brevis (EPB), and abductor pollicis longus (APL) that occupy the first compartment on the dorsal surface of the radius near the styloid process (1). The condition became known as de Quervain’s disease, or stenosing tenovaginitis at the first compartment of the extensor retinaculum. Although the disorder is relatively common, there are few histopathological studies in the literature (2, 3). The early reports focused on inflammation and identified three etiologic factors: repetitive movement, and anatomical and hormonal factors. Repetitive movement was considered to be an extrinsic factor, and anatomical and hormonal factors were intrinsic. The presence or absence of a septum and number of tendons have been discussed in relation to the onset mechanism of de Quervain’s disease (4, 5). To better understand the onset mechanism, we examined tissue structures of the first compartment of the extensor retinaculum of patients diagnosed as having de Quervain’s disease, and compared the results with a cadaveric control group. In the early reports, chronic inflammation has been cited as the main cause of this disorder. However, Ippolito et al. (6) observed degenerative changes in collagen fibers by electron microscopy. De Quervain declared that the synovial surface was uninvolved, and the adjacent tissue was thickened but not due to inflammatory reactions or round cell infiltration. He never found any gross pathological changes in the tendons (1).

In this study, surgical specimens from the patients with de Quervain’s disease were examined by light and electron microscopy, and the correlation between the morphology and functional disturbances in clinical presentation are discussed.

Materials and Methods

Thirty-five patients diagnosed as having de Quervain’s disease who were treated in our department from 1985 to 1992 comprise the study group. Our diagnostic criteria of de Quervain’s disease were: a) pain on the radial side of

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Table 1  
Age and sex distribution of patients and cadavers examined

<table>
<thead>
<tr>
<th>Gender</th>
<th>Number of patients and cadavers classified by age (year)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0~19</td>
</tr>
<tr>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Men</td>
<td>0</td>
</tr>
<tr>
<td>Women</td>
<td>0</td>
</tr>
</tbody>
</table>

The specimens from the study group for examination by scanning electron microscopy (SEM) were fixed with 2.5% glutaraldehyde (buffered to pH 7.4 with 0.1M PBS), freeze-dried and cracked in liquid nitrogen after washing in 0.2M cold cacodylic acid buffer. Some were frozen and cracked to examine the central portion. Each sample was dried, evaporated and examined by SEM (JSM-T20 type, JEOL Ltd.). The specimens for examination by transmission electron microscopy (TEM) were fixed after washing by the same method as SEM, and post-fixed with 1% osmium tetroxide. Double staining was done with uranyl acetate and ruthenium red. They were then examined by TEM (JEM-100S type, JEOL Ltd.).

The specimens were classified into four groups based on anatomical criteria. The results of statistical analyses were subjected to chi-square test (with Yates’ correction in the case of degree of freedom). A p value of <0.05 was considered significant for all tests.

Results

Anatomical and histopathological variations in the first compartment of extensor retinaculum are summarized in Table 2. A septum was observed more frequently in the study group than in controls. EPB and APL were more numerous in patients with de Quervain’s disease than in cadavers. The percentage of specimens containing septa was 74% in the study group and 49% in the control group, and the difference was statistically significant (p < 0.01). The numbers of EPB and APL tendons in the de Quervain’s disease were also higher than in the cadavers, and this difference was also statistically significant (p < 0.001) (Tables 1, 2). In the control group, there was no significant difference by gender in the proportion of septa found
Table 2  Variations in the first compartment of extensor retinaculum

<table>
<thead>
<tr>
<th>Gender</th>
<th>Group</th>
<th>Number of EPB</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>0</td>
</tr>
<tr>
<td>Patients&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td>1</td>
</tr>
<tr>
<td>Men</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Women</td>
<td>10</td>
<td>9</td>
</tr>
<tr>
<td>Total</td>
<td>12</td>
<td>9</td>
</tr>
<tr>
<td>Cadaveric controls&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
<td>(+)&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Men</td>
<td>61</td>
<td>60</td>
</tr>
<tr>
<td>Women</td>
<td>53</td>
<td>58</td>
</tr>
<tr>
<td>Total</td>
<td>114</td>
<td>118</td>
</tr>
</tbody>
</table>

<sup>a</sup> 35 samples from 35 patients  
<sup>b</sup> 232 samples from 170 cadavers  
<sup>c</sup> presence of septum  
<sup>d</sup> absence of septum  

or in the number of EPB and APL tendons. In the study group, both the retinaculum and tenosynovium were thickened in 30 patients, and only the retinaculum was thickened in 5 patients (Fig. 1). The diameters of retinacula for both groups are illustrating in Fig. 2. The number of women was higher than men with de Quervain’s disease.

The tenosynovium and retinaculum consisted of the synovial, inner, intermediate, and the outer layers of the retinaculum (Fig. 3). The histological findings in the study group were classified into 4 groups according to the presence or absence of septum and the retinaculum diameter (Fig. 3): Group I, a septum was present and the retinaculum of both EPB and APL sides were equally thickened (n = 12); Group II, a septum was present and the retinaculum was thickened on both sides, although the EPB side was thicker (n = 9); Group III, a septum was present and retinaculum was thickened on both sides, and the APL side was thicker (n = 5); Group IV, the retinaculum was thickened and the septum was absent (n = 9). EPB and APL tendons were not enlarged in any of these groups.

All of the patients with de Quervain’s disease were diagnosed according to their Finkelstein test score. Twenty-six patients also tested positive in the Iwahara-Nozue test before surgery. This test was positive in 3 out of the 4 groups; 12 out of 12 patients in Group I, 9 out of 9 patients in Group II and 5 out of 9 patients in Group IV. The 5 patients in Group IV had greater thickening on the EPB side. Iwahara-Nozue test results were negative in all 5 patients in Group III. Retinaculum was thickened in all 4 groups. Although the Iwahara-Nozue test originally was not designed to indicate EPB thickening, our
results showed a statistically significant correlation between positive test results and EPB thickening. The test results for Groups I and IV indicated that there was some difference other than the presence or absence of the

Fig. 2  De Quervain’s disease and normal control. A: de Quervain’s disease, B: normal control.

Fig. 3  Histology of classified types. A: Group I (I: superficial synovial layer, II: inner layer of the retinaculum, III: intermediate layer, IV: outer layer) (HE, X17); B: Group II (HE, X17); C: Group III (HE, X17); D: Group IV (HE, X17).
The total percentages of histological findings in the retinacula of all 4 groups are shown in Table 3. Histologically, the thickening of the tenosynovium and retinaculum was caused by fibrosis in every layer (Fig. 4 A–D). There were also chronic inflammatory changes in all layers. In the superficial tenosynovium, the number of layers of synovial cells and capillaries increased (Fig. 4A).

**Table 3** Histological findings in 35 patients

<table>
<thead>
<tr>
<th>Grade</th>
<th>Capillary proliferation</th>
<th>Round cell infiltration</th>
<th>Fibroblast proliferation</th>
<th>Fibrosis</th>
<th>Hyaline degeneration</th>
<th>Myxoid degeneration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Severe</td>
<td>11</td>
<td>0</td>
<td>20</td>
<td>34</td>
<td>0</td>
<td>26</td>
</tr>
<tr>
<td>Moderate</td>
<td>31</td>
<td>12</td>
<td>46</td>
<td>29</td>
<td>14</td>
<td>34</td>
</tr>
<tr>
<td>Minimal</td>
<td>26</td>
<td>17</td>
<td>20</td>
<td>20</td>
<td>9</td>
<td>11</td>
</tr>
<tr>
<td>Absent</td>
<td>26</td>
<td>71</td>
<td>14</td>
<td>17</td>
<td>77</td>
<td>29</td>
</tr>
<tr>
<td>Total</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
</tbody>
</table>

**Fig. 4** Light microscopic findings of the retinaculum. A: Vascular proliferation in the first layer (HE, X120); B: Myxoid lesion in the second layer (HE, X120); C: Regularly arranged collagen fibrils and irregular fibrils around small vessels in the third layer (HE, X120); D: Round cell infiltration around the small vessels and edema in the fourth layer (HE, X120).
Round-cells had minimally infiltrated the edematous stroma (Fig. 4C). Fibroblast proliferation and an increase of collagen fibrils were observed in the inner and outermost layers (Fig. 4B, C). Capillary proliferation with stromal myxoid and/or hyaline degeneration was found in the intermediate layer (Fig. 4C). An increase of the dense fibrous connective tissue, containing numerous capillaries, and lymphocytic infiltration was also found in this layer (Fig. 4D). The fibril network of these layers was easily identified under the polarized light microscope (Fig. 5). The tendons in situ appeared to be normal in these patients.

Under SEM, the cracked surface disclosed the proliferation of synovial cells. Columnar cells just beneath the synovial cells were arranged in regular parallel rows (Fig.

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**Fig. 5**  Fibrillar structure of the 4 layers under polarized light microscope in Group 1 (HE, X17).

**Fig. 6**  Scanning electron micrographs of the retinaculum in de Quervain's disease. A: Proliferation of superficial cells; B: The surface of the first layer; C: Dense collagen bundles in the fourth layer; D: Irregular collagen fibrils around capillaries (*) in the third layer.
6A). The surface of the synovial layer was elevated by 15 μm, showing an almost normal appearance (Fig. 6B). Dense collagen bundles of 15 μm wide were found in the outer layer of the retinaculum (Fig. 6C). Scattered and irregularly shrunken collagen fibrils, suggesting newly formed collagen fibrils, were mainly observed around the capillaries in the intermediate layer of the retinaculum (Fig. 6D). Under TEM, the bundles of collagen fibrils in the synovial layer were found around the fibroblasts containing rough endoplasmic reticula in their cytoplasm (Fig. 7). In the intermediate layer, histiocytes with many electron-dense particles in the extracellular space were observed.

Discussion

According to Muckart (9), the first reference on stenosing tendovaginitis of EPB and APL at the radial styloid was found in the 13th edition of Gray’s Anatomy (1893) under the title “Washer-Woman Sprain”. De Quervain published his original paper that described the disorder in 1895 (1). Stenosing tenosynovitis was defined as the disturbance of tendon movement due to chronic inflammation resulting in pain on the radial side of the wrist. However, there have been a few reports of histopathology or etiology (2, 9).

Finkelstein claimed that, in mild cases, the synovial membrane was thickened, and that the loose connective-tissue layer was considerably thickened and vascularized. In severe cases, the synovial layer was completely destroyed and was totally replaced by hyalinized collagenous tissues, and the loose connective-tissue layer was compressed and thinned by the thickened hyalinized tissue. The ligamentous layer was markedly thickened and undergoes hyaline degeneration and or cartilaginous transformation. Finkelstein concluded that stenosing tendovaginitis could be produced by thermal, chemical and mechanical stimulation, and proposed that an inflammation was caused by friction between tendon and tenosynovium (2). Repetitive superficial friction produced inflammatory reactions of round cell infiltration and edema in the synovial layer. Our findings suggest that inflammation was not induced by superficial friction, because additional minimal-to-severe inflammatory changes were found in all four layers (Fig. 4). We also noted a high incidence of capillary and round cell proliferation in cases that ranged from minimal to severe, which indicated a chronic inflammatory condition. In the future, we will clarify the inflammatory reaction by the analyses of fibrosis and type of collagen produced in each layer.

According to Muckart (9), the chronic inflammatory reaction found in both fibrous retinaculum and synovium and the presence of hemosiderin in the synovium supported the contention that de Quervain’s disease was an acute episode superimposed on a chronic condition. Previous histologic examination of de Quervain’s disease (2, 3, 9-12) showed an inflammatory reaction in the constricted fibrous sheath. Ippolito et al. (6) observed an increased amount of collagen but no round cells in the thickened sheath in their ultrastructural study. In our study, the patient’s specimens clearly showed round cell infiltration (12 %) and fibrosis (63 %) (Table 3).

Although there was a clear relationship between the histological changes and the degree of disturbance of movement, it seemed that the inflammatory reaction reflected either different stages of the disease or different parts of the involved retinaculum and tenosynovium. This may reflect a selective grouping by the health care industry of patients that require different kinds of treatment. Each researcher may be investigating different cases that represent different stages and involve the different layers of the retinaculum. This could explain some of the apparent contradictions in the reports that we have cited.

The Iwahara-Nozue test helps to identify the cases where the focus of pain is on the EPB rather than on the APL side. This test also informs the surgeon about the probability of the presence of a septum and the need to
look carefully for supernumerary tendons behind the septum. The persistence of pain after surgery may indicate that supernumerary tendons had been overlooked because they were obscured by the septum. Hopefully, in the future, precise identification of a focus of pain in the EPB side may allow preservation of the APL in patients who respond to anti-inflammatory agents. Because corticosteroids inhibit the inflammatory reaction and the increased synthesis of collagen, or other extracellular matrix (6), these hormones may be the treatment of first choice for de Quervain's disease. Orthopaedic surgeons must consider the high probability of thickening of the retinaculum in this disease when they make precise diagnoses and recommending treatment.

Occupational activities have been frequently cited in discussion of etiology (2, 9–11, 13, 14). Finkelstein (2) stated that cumulative trauma was the most common cause. There were many case reports where the causes were playing the piano, using a typewriter, handwriting, washing, wringing clothes, chopping wood, carrying heavy objects, farm labor, cutting cloth with heavy scissors, etc. However, Thompson et al. (15) reviewed 419 cases of peritendinitis crepitsans occurring in factory workers after a period of intense effort where only two patients developed de Quervain’s disease. These results suggest that friction does not play an important role in the development of stenosing tenovaginitis.

De Quervain’s disease was, in the past, thought to consist of anatomic variations of the retinacula and tendons (4, 5, 9, 13, 14, 16). In our study, the incidence of the presence of a septum was also higher in patients with de Quervain’s disease than in normal individuals (Table 2). However, there was no gender difference in the occurrence of a septum and the number of EPB and APL tendons in the cadavers (7). We agree with the anatomic factor explanation that the existence of a septum may be a predisposing etiologic factor in complaints of limitation of movement (Fig. 8). However, the high incidence in both groups of women and the differential distribution by gender and age group (Table 1) indicate that some hormonal factors may be associated.

We, therefore, conclude that the onset mechanism of de Quervain’s disease includes intrinsic factors such as hormonal and anatomical variation (Table 1, 2), and extrinsic factors such as sports and occupational activities (Fig. 8). The high incidence of this disorder in women aged 40–60 years, may be mainly due to intrinsic factors, especially hormonal. The target organs seemed to be both the tenosynovium and the extensor retinaculum where thickening results in the disturbance of tendon gliding. Iwahara-Nozue test results indicated that impairment is directly related to the degree of thickening of the retinaculum in the EPB side. Complaints of limited movement with pain associated with extrinsic factors such
as sports and occupational activities are relatively acute and transient. Therefore, the extrinsic factors are probably not a primary cause in the chronic cases of this disease.

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