CLINICAL AND BIOPSY FINDINGS IN TEMPORAL ARTERITIS

BY

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Hutchinson (1890) described a man aged 80 years who was prevented from wearing his hat by painful, inflamed temporal arteries. Horton, Magath, and Brown (1932) described two more patients and called the syndrome temporal arteritis. Originally thought to be a localized, self-limiting, and benign disorder, inflammation of the temporal arteries is often part of a widespread arteritis (Cooke, Cloake, Govan, and Colbeck, 1946) which can cause blindness (Jennings, 1938) and death (Harrison, 1948).

The varied presentation of the disorder was emphasized by Paulley and Hughes (1960) and often the diagnosis is not made with certainty without histological examination of an affected artery. The histological appearances are also variable. The media is usually the site of collections of inflammatory cells and giant cells apparently related to fragmentation of the internal elastic lamina. Giant cells are not always found (Harrison, 1948; Paulley and Hughes, 1960) and in some cases intimal fibrosis predominates, with little evidence of cellular infiltration. In the face of a variable clinical picture and variable histology, we have studied 37 biopsied cases of temporal arteritis in an attempt to relate the clinical and histological features.

Methods

The medical records of all patients in the Radcliffe Infirmary thought to have been suffering from temporal arteritis in the period from 1945 to 1965 were reviewed. Those in whom biopsy of a superficial temporal artery had been carried out were selected for further study. This yielded 37 cases. Details of the symptoms, length of history, the clinical state of the arteries, and the laboratory investigations were extracted.

Independently one of us (M.S.D.), without knowledge of the clinical information, reviewed the biopsy sections.

Results

I. Clinical Examination of the Temporal Arteries

These were recorded as normal in appearance and to palpation in nine cases. In twenty cases the arteries were tender and, in addition, in five of these they were noted to be nodular or "beaded". In the remaining eight patients, although the vessels were not tender, there was reduction or absence of the normal pulsation.

II. Histological Appearances

Sections were taken at several different levels from each biopsy. They were stained with haematoxylin and eosin, by Masson's trichrome method, and by the Orcein method for demonstrating elastic tissue. It was possible to classify the biopsies into three broad groups: those showing giant cell arteritis, those showing arteritis but without giant cells, and those showing intimal fibrosis only.

(a) Giant cell arteritis (Fig. 1, overleaf).—Present in 23 cases. In these vessels the lumen of the artery frequently showed obliteration or gross distortion because of the presence of fibrous tissue. Thrombosis was not present. The striking features were the presence of very plump fibroblasts and the large quantity of metachromatic material, seen in Azure A preparations, surrounding them. Evidence of re-canalization with the formation of numerous new vascular channels, occasionally possessing smooth muscle in their walls, was also present. In some of the sections haemosiderin could be seen. In this portion of the artery the inflammatory cellular infiltrate was concentrated near the internal elastic lamina or its remains. It consisted most commonly of lymphocytes but occasionally included neutrophil polymorphonuclear leucocytes and eosinophils. The giant cells were invariably of the Langhans variety with peripherally-placed nuclei, and they were always present in the region of a break in the
internal elastic lamina. They were most frequently situated in the outermost part of the fibrosed intima. Their derivation is uncertain but there is some slight tinctorial evidence, seen particularly in the trichrome stains, that they may possibly be derived from smooth muscle cells. The media often showed rather striking preservation and where smooth muscle was seen there was little or no cellular infiltration. Focal breaches occurred in the media adjacent to a break in the internal elastic lamina, and at this point giant cells and lymphocytes could be seen. The adventitia occasionally showed a giant cell infiltration but most commonly exhibited a generalized infiltration with lymphocytes.
(b) Arteritis without giant cells (Fig. 2).—Because of the focal nature of the disease, considerable caution must be observed in claiming that giant cells are absent in any case of temporal arteritis. However, in five cases they were not seen in sections taken at several different levels and in general these showed a rather more acute infiltration with more neutrophil polymorphonuclear leucocytes and eosinophils than in the first group. In one of these cases thrombus was seen in the lumen.

Fig. 2(a).—Temporal artery, showing severe arteritis with no giant cells.

Fig. 2(b).—Arteritis with no giant cells. (Haematoxylin and eosin preparation.)
(c) **Intimal fibrosis** (Fig. 3).—In nine arteries the predominant finding was the presence of intimal fibrosis. The fibrous tissue was in most cases rather acellular and there did not appear to be an excess of metachromatic material.

The internal elastic lamina in these vessels was for the most part intact, though an occasional small breach could be seen. The media was essentially normal but the adventitia occasionally showed focal lymphocytic aggregates.
III. Relationship between Clinical and Pathological Features

Table I sets out the clinical and laboratory features of the cases according to their subdivision into three groups on the basis of histological appearances. There is no evidence that length of history, erythrocyte sedimentation rate, or other feature is strongly linked with any particular histological picture. The biopsies showing a "burnt-out" arteritis with marked intimal fibrosis did not prove to have come from patients with longer histories, less systemic upset, or lower erythrocyte sedimentation rates.

<table>
<thead>
<tr>
<th>Table I</th>
<th>CORRELATION OF CLINICAL AND BIOPIX FEATURES OF 37 CASES OF TEMPORAL ARTERITIS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical Features</td>
<td>Histological Type</td>
</tr>
<tr>
<td></td>
<td>Arteritis</td>
</tr>
<tr>
<td>No. of Cases</td>
<td>5 23 9</td>
</tr>
<tr>
<td>Mean Age (yrs)</td>
<td>72 70 69</td>
</tr>
<tr>
<td>Sex</td>
<td>Female</td>
</tr>
<tr>
<td></td>
<td>Male</td>
</tr>
<tr>
<td>Length of History (mths)</td>
<td>9 5 9</td>
</tr>
<tr>
<td>Signs</td>
<td>Headache</td>
</tr>
<tr>
<td></td>
<td>Malaise</td>
</tr>
<tr>
<td></td>
<td>Weight loss</td>
</tr>
<tr>
<td></td>
<td>Anorexia</td>
</tr>
<tr>
<td></td>
<td>Visual loss</td>
</tr>
<tr>
<td></td>
<td>Fever</td>
</tr>
<tr>
<td>Mean Erythrocyte Sedimentation Rate (mm./hr)</td>
<td>94 78 95</td>
</tr>
<tr>
<td>White Blood Cell Count</td>
<td>10,000 10,000 8,000</td>
</tr>
</tbody>
</table>

Table II relates the clinical state of the temporal arteries at the time of biopsy with the histological appearance of the specimen. The important finding is that a vessel which is clinically normal in all respects may nevertheless yield evidence of an arteritis (5/9) and may contain giant cells (4/9).

<table>
<thead>
<tr>
<th>Table II</th>
<th>COMPARISON BETWEEN THE CLINICAL STATE OF THE TEMPORAL ARTERY AND THE BIOPIX FINDINGS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical State of the Artery</td>
<td>No. of Cases</td>
</tr>
<tr>
<td></td>
<td>Arteritis</td>
</tr>
<tr>
<td>Normal</td>
<td>9</td>
</tr>
<tr>
<td>Tender</td>
<td>20</td>
</tr>
<tr>
<td>Loss of Pulsation</td>
<td>8</td>
</tr>
</tbody>
</table>

One patient was re-biopsied on two further occasions. In all three instances the artery was tender when biopsied. In one of the biopsy specimens there was evidence of an arteritis. The other specimens showed intimal fibrosis.

Deterioration of vision was noted by eleven patients and in Table III the clinical and pathological information on these patients is compared with that of those who had no such complications. No clear differences emerge which might be related to the pathogenesis of the ocular involvement or used to predict it. It was noteworthy that two of the patients with ocular involvement had clinically normal temporal arteries.

<table>
<thead>
<tr>
<th>Table III</th>
<th>COMPARISON OF CASES WITH AND WITHOUT OCULAR INVOLVEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical Features</td>
<td>No Eye Involvement</td>
</tr>
<tr>
<td>No. of Cases</td>
<td>26</td>
</tr>
<tr>
<td>Mean Age (yrs)</td>
<td>74</td>
</tr>
<tr>
<td>Sex</td>
<td>Male</td>
</tr>
<tr>
<td></td>
<td>Female</td>
</tr>
<tr>
<td>Length of History (mths)</td>
<td>6</td>
</tr>
<tr>
<td>Signs and Symptoms (per cent.)</td>
<td></td>
</tr>
<tr>
<td>Headache</td>
<td>100</td>
</tr>
<tr>
<td>Malaise</td>
<td>50</td>
</tr>
<tr>
<td>Weight loss</td>
<td>50</td>
</tr>
<tr>
<td>Anorexia</td>
<td>50</td>
</tr>
<tr>
<td>Muscle pain</td>
<td>46</td>
</tr>
<tr>
<td>Joint pain</td>
<td>15</td>
</tr>
<tr>
<td>Jaw pain</td>
<td>23</td>
</tr>
<tr>
<td>Fever</td>
<td>54</td>
</tr>
<tr>
<td>Mean Hb (g./per cent.)</td>
<td></td>
</tr>
<tr>
<td>Erythrocyte Sedimentation Rate (mm./hr)</td>
<td>90</td>
</tr>
<tr>
<td>White Blood Cell Count</td>
<td>10,000</td>
</tr>
<tr>
<td>Temporal Artery (per cent.)</td>
<td></td>
</tr>
<tr>
<td>Tender</td>
<td>54</td>
</tr>
<tr>
<td>Loss of Pulse</td>
<td>20</td>
</tr>
<tr>
<td>Normal</td>
<td>27</td>
</tr>
<tr>
<td>Biopsy (per cent.)</td>
<td></td>
</tr>
<tr>
<td>Arteritis</td>
<td>15</td>
</tr>
<tr>
<td>Giant cells</td>
<td>65</td>
</tr>
<tr>
<td>Intimal fibrosis</td>
<td>20</td>
</tr>
</tbody>
</table>

Discussion

In all of the present cases the biopsy sections were thought to confirm the clinical diagnosis of temporal (giant cell) arteritis. Much has been written about the difficulty of distinguishing temporal arteritis from polyarteritis nodosa on histological grounds (Jennings, 1938; Harrison, 1948; Ross Russell, 1959). There is usually little difficulty if the clinical picture is considered together with the histological evidence (Harrison, 1948). Although giant cell arteritis has been seen at autopsy to involve nearly all medium and large sized arteries (Cooke and others, 1946; Harrison, 1948; Ross Russell, 1959), clinical evidence of the involvement of visceral vessels is rare (Ross Russell, 1959). Renal involvement has never been described clinically although the renal artery has been found to be affected at autopsy (Sproul, 1942). The clinical predominance of symptoms related to cranial vessels despite the widespread nature of the disease remains unexplained.

Nine of the 37 patients had no detectable clinical abnormality of the superficial temporal arteries despite having headache and biopsy involvement of these vessels. Miller Fisher (1961) has claimed that
minor abnormalities of the vessels can always be
detected by palpation in such circumstances, and
that furthermore careful palpation is as reliable as
arterial biopsy in making a confident early diagnosis.
Most physicians do not find this so and one in four
of the cases biopsied at this hospital have had
apparently normal temporal arteries. It is of great
interest that these vessels were nevertheless produc-
tive of the same histological pattern as were tender,
nodular, occluded vessels. Positive biopsies have
also been obtained from clinically normal vessels in
cases of polymyalgia rheumatica (Alestig and Barr,
1963), which itself may develop into temporal
arteritis (Harrison and Bevan, 1967). Hollenhorst,
Brown, Wagener, and Shick (1960) mentioned that
they obtained diagnostic biopsies from apparently
normal vessels but they did not say how often this
situation arose and the point has received little
attention. As early diagnosis is so vital in the
anticipation of ocular complications, it is important
to realise a clinically normal artery may yield a
positive biopsy.

The biopsy findings reported here fell into three
groups, one showing the features of an arteritis with
marked cellular infiltration, the second and largest
group containing evidence of an arteritis together
with multi-nucleate giant cells, and the third showing
little cellular infiltration but striking intimal fibrosis.
It was thought that these three subgroups might
have represented acute, subacute, and chronic
stages of the arteritis, but this was not supported by
the study of the clinical features of the cases. Intimal fibrosis was the predominating feature of
the biopsy of several acute florid cases. Harrison,
Harrison, and Kopelman (1955) re-biopsied two
cases of temporal arteritis after starting steroid
treatment. The earlier biopsies had shown giant
cells and marked lymphocytic infiltration; after
treatment both showed intimal fibrosis. Harrison
and others concluded that intimal fibrosis represented
an inactive phase of the disease and that its presence
in a biopsy could be used to assess progress and
even to guide the dosage of steroid. Our findings do
not support that contention. Rather the finding of
intimal fibrosis can be considered to carry the same
significance as the finding of giant cells. It is not
known why the histological picture is so variable.
The vessel is often involved in a focal manner
(Harrison, 1948) and it is possible that in some cases
an insufficient length of the artery was biopsied,
though in fact 10 to 15 mm. were taken in each
case. The failure to find giant cells in each case
may be due to the patchy involvement of the vessel
and this view would receive indirect support from
our patient who had three biopsies. On each occa-
sion the vessel was tender and the erythrocyte
sedimentation rate elevated. On two occasions the
sections showed intimal fibrosis and on the other a
cellular "arteritis".

Visual failure continues to occur in 30 to 50 per
cent. of cases (Harrison, 1948; Roux, 1954; Ross
Russell, 1959; Hollenhorst and others, 1960;
present series). This that suggests our ability to
anticipate its appearance has not improved greatly.
The eleven cases with partial or complete loss of
vision in this series were studied from both the
clinical and the pathological standpoint. Nothing
appeared to demarcate them from those who did not
suffer this complication. There is thus as yet no
way of predicting which cases may develop visual
failure. It is important to note that in two of the
eleven cases the temporal arteries appeared entirely
normal on clinical examination.

Summary

The clinical features of 37 cases of temporal
arteritis were compared with the histological
appearance of their arterial biopsies. One in four
of the cases had clinically normal temporal arteries,
but these yielded the same microscopical evidence of
a giant cell arteritis as did tender, nodular, occluded
vessels. Eleven patients with visual failure did not
appear to differ clinically or histologically from
those without ocular involvement. The different
histological appearances obtained on arterial biopsy
are discussed. It is stressed that the finding of
clinically normal temporal arteries does not rule out
the diagnosis of temporal arteritis.

REFERENCES

Alestig, K., and Barr, J. (1963). Lancet, 1, 1228 (Giant-cell arteritis: A biopsy study of polymyalgia
rheumatica, including one case of Takayasu's disease).
47 (Temporal arteritis: a generalized vascular disease).
re-emphasis of the value of careful palpation).
Clinical and biopsy findings in temporal arteritis


Résultats de l’examen clinique et des biopsies dans 37 cas d’artérite temporaire

RÉSUMÉ

On a comparé les signes cliniques de 37 cas d’artérite temporaire avec l’aspect histologique de leurs biopsies artérielles. Dans un cas sur quatre les artères temporales étaient cliniquement normales mais à l’examen histologique on y trouva les mêmes signes d’artérite à cellules géantes que dans les vaisseaux douloureux, noduleux et obstrués. Onze malades ayant des troubles visuels ne paraissaient pas différer cliniquement ou histologiquement de ceux qui n’avaient pas d’atteinte oculaire. On discute les différents aspects histologiques observés dans les biopsies artérielles. On souligne que le fait de trouver des artères temporales cliniquement normales ne permet pas d’exclure l’artérite temporaire.

Hallazgos clínicos y de biopsia en 37 casos de arteritis temporal

SUMARIO

Se compararon los rasgos clínicos de 37 casos de arteritis temporal con el aspecto histológico de sus biopsias arteriales. En una cuarta parte de los casos las arterias temporales fueron clínicamente normales, pero histológicamente presentaron el cuadro de arteritis con células gigantes similar al encontrado en los vasos dolores, nodulares y ocluidos. Once enfermos con distúrbios visuales no parecieron diferir clínicamente o histológicamente de los sin afectación ocular. Se discuten las diferentes apariencias histológicas observadas en biopsias arteriales. Se destaca el hecho de que la normalidad clínica de arterias temporales no excluye la existencia de una arteritis temporal.