The past decade has witnessed a practical re-orientation to the problem of so-called lymphadenoid goitre or struma lymphomatosa of Hashimoto. No longer is controversy so deeply centred as to whether this condition follows the epithelial hyperplasia of Graves or precedes the fibrosis of Riedel. Stress must now be laid on its practical aspects, the clinical recognition, treatment and, if possible, the final cure.

It is now becoming increasingly recognized that this thyroidal state is part of a general systemic dysfunction affecting not only the other endocrine glands, the pituitary, hypothalamus and cerebral cortex, but also the vital internal organs, such as the liver, spleen and kidneys.

The results of my work, published in 1951, 1952 and 1953, will be referred to here in the briefest outline. I came to the conclusion that, following unremitting and continued stress, the thyroid gland underwent successive pathological changes which were mirrored by the clinical condition of the patient. The epithelial tissue increases by steady hyperplasia. At the height of epithelial hyperplasia lymphoid tissue appears and increases as perivascular lymphorrhages and as focal lymphoid follicles; the epithelial tissue becomes hypoplastic. Following the greatest and diffuse hyperplasia of lymphoid tissue the latter in turn becomes hypoplastic. Fibrous tissue then appears on the scene, at first focally, increasing pari passu with diminishing lymphoid tissue, to end in the terminal diffuse fibrosis.

In 1951 I named the progressive phases of the degenerating thyroid gland (Fig. 1):
LEVITT: Lymphoid Goitres

1. Epithelial hyperplasia.
2. Lympho-epithelial hyperplasia (Graves’s constitution).
3. Focal lymphoid hyperplasia (lymphoid thyrotoxicosis).
4. Diffuse lymphoid hyperplasia (lymphadenoid goitre).
5. Fibrolymphoid hyperplasia (Hashimoto-like disease).
6. Fibrosis (Riedel’s fibrosis in sharp contrast to Riedel’s inflammatory chronic thyroiditis).

Table 1.—The Microscopical Appearance of the Thyroid in the Lymphoid Phases

<table>
<thead>
<tr>
<th></th>
<th>1 Epithelial</th>
<th>2 Lympho-epithelial</th>
<th>3 Focal Lymphoid</th>
<th>4 Diffuse Lymphoid</th>
<th>5 Fibrolymphoid</th>
<th>6 Fibrous</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Epithelium</td>
<td>++++++++ Very hyperplastic</td>
<td>++ Less Still less hyperplastic</td>
<td>++++ Hypoplastic</td>
<td>++ More hypoplastic</td>
<td>+</td>
<td>Atrophied</td>
</tr>
<tr>
<td>2. Lymphoid</td>
<td>—</td>
<td>Small lymphorrhages Focal with germinal centres</td>
<td>—</td>
<td>—</td>
<td>++</td>
<td>Diminished</td>
</tr>
<tr>
<td>3. Fibrous</td>
<td>—</td>
<td>—</td>
<td>Diminished</td>
<td>—</td>
<td>—</td>
<td>++</td>
</tr>
<tr>
<td>4. Colloid</td>
<td>Maximal</td>
<td>Less</td>
<td>Focal</td>
<td>—</td>
<td>—</td>
<td>++</td>
</tr>
<tr>
<td>5. ‘Giant’ cells</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>++</td>
</tr>
<tr>
<td>6. Oxyphilic cells</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>++</td>
</tr>
</tbody>
</table>

Table 2.—The Macroscopical Appearance of the Thyroid in the Lymphoid Phases

<table>
<thead>
<tr>
<th></th>
<th>1 Epithelial</th>
<th>2 Lympho-epithelial</th>
<th>3 Focal Lymphoid</th>
<th>4 Diffuse Lymphoid</th>
<th>5 Fibrolymphoid</th>
<th>6 Fibrous</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Size</td>
<td>+ Smooth</td>
<td>+++ Pebbly</td>
<td>+++ More opaque</td>
<td>+++ Lobular</td>
<td>++ Nodular</td>
<td>+</td>
</tr>
<tr>
<td>2. Surface Capsule</td>
<td>Translucent</td>
<td>Still shiny</td>
<td>Densely opaque</td>
<td>Densely opaque</td>
<td>Fine extra-capillary adhesions</td>
<td>Hard lobulated by trabeculae</td>
</tr>
<tr>
<td>3. Consistency</td>
<td>Soft</td>
<td>Less soft</td>
<td>Firmer</td>
<td>Rubbery homogeneous</td>
<td>Hard lobulated by trabeculae</td>
<td>Very hard; cuts with difficulty; grates on knife</td>
</tr>
<tr>
<td>4. Colour</td>
<td>Beef-red</td>
<td>Lighter red</td>
<td>Pink</td>
<td>Pinkish yellow</td>
<td>Yellowish white</td>
<td>White</td>
</tr>
<tr>
<td>5. Vascularity</td>
<td>+++++</td>
<td>++++</td>
<td>+++</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>6. Pressure</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

Table 3.—The Clinical Appearance of the Patient in the Lymphoid Phases

<table>
<thead>
<tr>
<th></th>
<th>1 Epithelial</th>
<th>2 Lympho-epithelial</th>
<th>3 Focal Lymphoid</th>
<th>4 Diffuse Lymphoid</th>
<th>5 Fibrolymphoid</th>
<th>6 Fibrous</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Thyrotoxicity</td>
<td>maximal 40</td>
<td>30</td>
<td>moderate 28</td>
<td>minimal</td>
<td>absent 7</td>
<td>0</td>
</tr>
<tr>
<td>2. Exophthalmos</td>
<td>rising 32</td>
<td>maximal 57</td>
<td>falling 50</td>
<td>minimal</td>
<td>absent 20</td>
<td>0</td>
</tr>
<tr>
<td>3. Lid-retraction</td>
<td>rising 15</td>
<td>still rising 20</td>
<td>maximal 22</td>
<td>falling 13</td>
<td>minimal 7</td>
<td>0</td>
</tr>
<tr>
<td>4. Pre-operative loss in weight</td>
<td>maximal 81</td>
<td>high 78</td>
<td>falling 74</td>
<td>moderate 55</td>
<td>minimal 40</td>
<td>lowest 20</td>
</tr>
<tr>
<td>5. Post-operative gain in weight</td>
<td>minimal 44</td>
<td>increasing 69</td>
<td>moderate 73</td>
<td>still high 71</td>
<td>rising 90</td>
<td>maximal 100</td>
</tr>
<tr>
<td>6. Post-operative hypothyroidism</td>
<td>minimal 2</td>
<td>low 8</td>
<td>moderate 14</td>
<td>rising 27</td>
<td>fast rising 67</td>
<td>maximal 80</td>
</tr>
</tbody>
</table>

* expressed as a percentage of patients in each phase.
I further suggested that this sequence might progress to any phase and persist as such or may occasionally revert to a previous phase. The appearance of fibrous tissue would, however, ensure irreversibility of the affected part of the gland.

The epithelial element appears to be chiefly concerned with the growth and metabolism of the thyroid gland. The lymphoid deposition is in part reflected by the antibody-antigen reaction and the production of immunity against disease, as stressed in 'The Thyroid' (1953). The fibrous element appears to be part of the healing process, replacing the degenerated and atrophied epithelial and lymphoid tissues, but taking little or no part in the physiological and biochemical aspects of this phased process. Its complications are chiefly clinical.

As the phases advance the ages of the patients increase in concord with the gradual obliteration of the lumina of the blood vessels.

The microscopical features of these six phases are carefully defined in Table 1 and the macroscopical appearances of the gland are detailed in Table 2. The clinical criteria of the phases are shown in Table 3 and are expressed as a percentage of the 1,410 epithelio-lymphoid goitres which I have studied (Fig. 2). The truly epithelial goitres in this special series total 706, and the lymphoid goitres constitute 704 patients.

The pre-operative clinical, the operative macroscopical and post-operative clinical and microscopical criteria are postulated for each of these phases of the lymphoid goitre. Thus the clinician, the surgeon, the pathologist and, ultimately, the biochemist dealing with these lymphoid goitre patients for the first time become able to pinpoint the exact state of progression of this generalized systemic condition. Treatment becomes equally definitive for each phase.

It is this wide 'spectrum' of 704 lymphoid goitre patients who have been variously described by the terms Graves's constitution, lymphoid thyrotoxicosis, lymphadenoid goitre, Hashimoto's disease and Riedel's disease. More frequently, however, they have been included under the general designation 'Hashimoto's disease.'

Riedel, in all, reported three patients (Fig. 3), the first of whom appeared to suffer from fibrosis of the thyroid (my phase 6), a non-inflammatory condition, the termination of this phased chain-process. In sharp contradistinction, his last two patients manifested chronic thyroiditis, a truly inflammatory condition which has no bearing on this spectrum of the six phases and will not be discussed here.

Hashimoto described four patients in whom he suspected and suggested a gradational progressive change. The first patient presented diffuse lymphoid hyperplasia (phase 4); the intermediate, second and third, that of fibrolymphoid hyperplasia (phase 5); the final patient suffered from fibrosis (phase 6) which very remarkably resembled the first of Riedel's three patients; but, as I emphasized in 1952, it differed from Riedel's inflammatory condition of chronic thyroiditis seen in his last two patients.
The spectrum of so-called struma lymphomatosa of Hashimoto was so very wide that advance in the true scientific understanding of this process had become seriously hampered. Many different and apparently conflicting features had been attributed to the condition during the past half-century from 1900 to 1950. I found that the separation of this chain-process into six phases as described, and defining the attributes of each, very greatly simplified its understanding. Physiological, biochemical, radioiodine, clinical, prognostic and therapeutic features and correlations have become simplified and the stage of the patient’s progression can be easily ‘pinpointed.’

Biochemical Tests

The extensive range of reported biochemical findings reflects the wide spectrum of Hashimoto’s struma lymphomatosa generally assessed by different workers, not only in separate centres, but also in the same institution.

Biochemical evidence of thyroidal failure may be associated with (a) an apparently euthyroid individual without symptoms or signs of thyroidal failure or (b) a hypothyroid patient with definite symptoms or signs of diminished thyroidal function. The former has been designated by Skillern and his colleagues at the Cleveland Clinic (1956) as compensated thyroidal failure and the latter group as uncompensated thyroidal failure. Any degree of thyroidal maladjustment intermediate between these may occur. The complexity of this problem becomes intensified if the whole spectrum is lumped under the all-comprehensive term ‘struma lymphomatosa or Hashimoto’s disease’ instead of unravelling the whole into its parts. This analysis I attempted to do in 1951 by defining and correlating the progressive phases of the lymphoid goitre.

The more advanced phases of the lymphoid goitre, especially diffuse lymphoid hyperplasia and fibrolymphoid hyperplasia, show the following features:

- The basal metabolic rate (B.M.R.) is usually low and forms a valuable index.
- The serum cholesterol is elevated above 260 mg. per 100 ml. in the hypothyroid range and being occasionally above the myxoedematous range of 400 mg. per cent. It is a more valuable index in the later myxoedematous lymphoid phases.
- The erythrocyte sedimentation rate (E.S.R.) is raised.
- The plasma proteins. Electrophoretic studies show very characteristic changes which not only serve to diagnose the condition, but also appear to differentiate it from malignant tumours of the thyroid. The superposition of malignant change on lymphoid goitres, as reported by an increasing number of observers, notably Lindsay (1955), makes this problem more difficult, however, to assess.

The albumens are usually low or fall to a low-normal figure.
- The beta globulins may be high or reach the higher levels of normality.
- The gamma globulins are very characteristically raised.

In a separate study of 40 cancers of the thyroid gland (1957) I have not found the gamma globulin greatly raised, although such an isolated occurrence has been reported by Robbins, Rall and Rawson (1956) in a functional thyroid carcinoma.

Liver function tests. These are useful indices for diagnosis, prognosis and treatment of this systemic disease. The non-specific nature of these measurements, however, emphasize the need for viewing the biochemical picture as part of a whole and correlating it with the overall survey.
The *serum colloidal gold test*, as observed by Cooke and Wilder (1954), forms a most valuable index in the evaluation of the earlier lymphoid phases where thyroidal failure is less severe. Its raised value may be due to the precipitation of the colloidal gold by the greatly increased content of the gamma globulin in the plasma protein.

The *turbidity levels* of thymol, zinc sulphate and ammonium sulphate are all raised with the greater deposition of lymphoid tissue.

The *thymol flocculation test* becomes positive in the later lymphoid phases and is a more useful index than the *cephalin flocculation test*.

### Radioiodine Tests

The I¹³¹ tests for so-called Hashimoto’s struma lymphomatosa, as recorded in the literature, show a wide range. They measure the change in the epithelial element and give no direct indication of the status of the lymphoid or fibrous tissues.

The *thyroid uptake* shows an extraordinarily wide variation. The 24-hour uptake has been reported as varying from below 10 per cent. to above 95 per cent. This confirms the suspicion that the histological diagnosis, not only in different centres, but also in the same institution, covers a wide spectrum of lymphoid goitre, ranging probably from lympho-epithelial hyperplasia (phase 2) to fibrosis (phase 6).

This scatter of recorded results for ‘Hashimoto’s disease’ is seen in the range of reported biological half-life, varying from 2 to 10 days in the same institution.

The *thyrotropin stimulation test* shows no increased uptake, indicating that the endogenous thyrotropin is already producing maximal stimulation of the epithelial cells of the thyroid.

The *potassium perchlorate test* has been described by Morgans and Trotter (1957). The administration of potassium perchlorate after the exhibition of I¹³¹ was often followed by considerable loss of the radiiodine previously taken up by the so-called lymphadenoid goitre of Hashimoto. The extent of this resultant discharge of absorbed radiiodine from the thyroid is an index of the organic binding of iodine by the thyroid gland.

The *total protein-bound iodine* (P.B.I.) is low or attains the lower level of normality.

The *serum butanol extractable iodine* (B.E.I.) gives a similar result, indicating a low thyroid hormone level.

The *topographical survey*, as seen by the automatic scanner, shows an even, diffuse distribution of uptake delineating the size and shape of the affected thyroid gland unlike malignant disease or cystic change.

**Autoradiography** confirms this even distribution in lymphoid goitres.

### Immunological Tests

In 1953 I recorded in ‘The Thyroid’ (p. 149, 357) that hypoalbuminaemia, associated with an absolute increase of beta and gamma globulins, accompanied the degeneration of the thyroid during the later lymphoid phases when the tendency to myxoedema was greater. I then suggested that these lymphoid stores in the thyroid were concerned with antibody formation and that the extensive removal of an enlarged thyroid in the phase of diffuse lymphoid hyperplasia or its radical destruction by radiotherapy was unwise and diminished its ability to sustain an immunological action.

In order to detect and at the same time evaluate the strength of antigen-antibody reactions in lymphoid goitres, the following tests are proving of value:

- **Precipitin test.** This usually gives a reliable reading and is readily performed. The precipitate is produced in vitro by the interaction of antigen derived from exogenous thyroglobulin with that of the antibody already present in the patient’s serum.

- **The complement fixation test**, on the other hand, is much more sensitive in man and therefore requires greater care in assessment. Cross reactions with heterologous species may occur at an early stage.

- **Tanned cell haemagglutination test.** This has proved very valuable in the hands of the Witebsky team; it also gives considerable cross reaction. The erythrocytes (modified by dilute tannic acid) are coated with the appropriate dilution of antigenic thyroid extracts. These tanned red cells become agglutinated on the addition of serum from the lymphoid goitre patient which contains the antibody. Great sensitivity is obtained even in high dilution.

- **The skin reactivity test**, also used by Witebsky, gives a positive reaction measured by the diameter of the erythema and intensity of the dermal induration.

In 1955 Witebsky and colleagues found that the degree of lymphoid change in rabbits immunized with extracts of their own thyroids appeared to be proportional to their antibody titre. Precipitins obtained in the sera of these animals were found to be organ-specific against the rabbit thyroid.

The precipitin test in man, as used by Roitt and his co-workers (1956) at the Middlesex Hospital, London, marked an important advance. They found that patients with ‘Hashimoto’s disease’ gave a positive precipitin test, unlike normal subjects and patients with other thyroid
diseases. Furthermore, the antibodies were found to be organ-specific to the human thyroid gland. It was suggested that patients with lymphadenoid goitre were immunized against human thyroglobulin and that the destruction of their thyroid gland resulted from the progressive interaction of the thyroglobulin in the gland with the autoantibody present in the patient's serum.

Fluorescein stain test, as evolved by White (1957) of the London Hospital, marks a further recent advance. Preparations of sections of human thyroid are treated with fluorescein conjugates of the globulin fraction of the same patient's serum which contains the antibodies. Localization of the conjugate occurs in the intracytoplasmic colloid. The same conjugate will localize to other human and monkey thyroid sections. These facts support the hypothesis that the antigen concerned is present in other thyroids than those pathologically involved. Normal thyroids show antigen in the acini and within the colloid. The thyroids of lymphoid goitres show the antigen free among the lymphoid cells surrounding the acini.

The possible role of periacinar colloid extrusion in the production of lymphoid goitres was reported by Hellwig (1951) and by Levitt (1952 and 1953). The lymphoid hyperplasias of the other target endocrine glands such as the adrenal cortices, gonads and secondary sexual organs may most probably be part of a similar systemic immunological process.

**Treatment**

Adequate treatment by means of desiccated thyroid produces a diminution in size of the thyroid and a decrease in the hypothyroid stigmata. The radioiodine and biochemical reactive levels may thus be made to revert to normal or near-normal levels.

The danger of malignant change in a lymphoid goitre cannot be stressed too strongly. If any suspicion of such neoplastic degeneration is awakened, adequate incisional biopsy is necessary. Needle biopsies are easy to do, but often provide misleading information.

The chief features of treatment are summarized in Table 4. As the phases advance from 1 to 6 the extent of the thyroidectomy must of necessity become smaller because the possibility of post-operative hypothyroidism increases steadily. The size of the residue left behind must therefore become larger in order to minimize this myxoeedematous change. The necessity for ligation of the arterial pedicles becomes minimal after the phase of focal lymphoid hyperplasia. The need for post-operative thyroid medication increases steadily from phase 3 to phase 6.

**Focal lymphoid hyperplasia** is adequately treated by means of desiccated thyroid if there is no danger of malignant change.

**Diffuse lymphoid hyperplasia** is still reversible, reacting well to thyroid therapy.

** Fibrolymphoid hyperplasia,** with its increasing fibrosis, may produce severe pressure phenomena. The enlarged lymphoid element reacts well to thyroid substitution therapy. Its steadily augmenting fibrosis indicates the necessity for relief of pressure by isthmectomy or wedge excision. If severe constriction ensues, limited partial thyroidectomy is required, followed by adequate post-operative thyroid medication. This ensures a maintained and balanced euthyroid state. The operative intervention affords the added opportunity for excluding any malignant change.

The **rare terminal fibrosis** always requires surgical intervention and treatment by wedge excision, as the differentiation from malignant disease is very difficult and constictive processes are always present.
The Future

In the same manner as I have defined the different criteria for the phases of the degenerating thyroid, I am completing the overall biochemical picture for each of these six phases. This data, especially for the most recently evolved tests, is not yet statistically significant. Similar radioiodine trends correlating the microscopic with the macroscopic and clinical features have become generally obvious.

It would, for instance, be important to assess definitively when and how the titre of the precipitin test changes in each of these phases, the different types and contours of the gold colloidal test, the changing intensity of the turbidity tests and to plot the accurate levels of the plasma proteins, especially the gamma globulin in each phase. Then, and then only, will we have an accurate and scientific yardstick to evaluate the validity of this concept of phased progression.

Finer assessment of subjective and clinical change in thyroidal dysfunction, with its periodic or rhythmic fluxes subclinical manifestations and the variations of measurement by differently trained observers, is notoriously difficult. Basically, we require an accurate pathological picture of the predominant phase of the whole thyroid gland. In addition, we have to accumulate an adequate number of objective biochemical and radioiodine and immunological criteria for assessing the validity of this concept.

A large, virtually unexplored, field lies wide open.

The apparent wide range of the recorded biochemical and radioiodine levels in so-called struma lymphomatosa or 'Hashimoto's disease' may fall into place when studied from these analysed phases.

I would reiterate the plea I made in 1952, that the many eponymous designations associated with the 'dark ages' of thyroid disease be replaced by more precisely defined terminology and nomenclature.

Conclusion

A representative international conference would constitute an admirable forum for finally bringing order into the contentious problem of lymphoid goitre.

T. Levitt,
88 Harley Street, London, W.1

BIBLIOGRAPHY

COOKE, R. T., and WILDER, E. (1956), Lancet i, 984.
LEVITT, T. (1953), 'The Thyroid: a Physiological, Pathological, Clinical and Surgical Study', Edinburgh, E. S. Livingstone.
LEVITT, T. (1957), in process of publication.
WHITE, R. G. (1957), personal communication.

H. K. LEWIS & Co. Ltd.
Medical Publishers and Booksellers
136 GOWER STREET
LONDON, W.C.1
(Adjoining University College and Hospital)

Medical Lending Library

ANNUAL SUBSCRIPTION from £1 17s. 6d.

Prospectus post free on application

Bi-monthly List of New Books and New Editions added to the Library sent post free on request

The Library Catalogue revised to December, 1949, containing a classified index of authors and subjects.

To subscribers 10/- net; To non-subscribers 17/6 net. Postage 1/9
Supplement 1950 to 1952. To subscribers 1/6 net; to non-
subscribers 3/- net; postage 8d.

NEW BOOKS ADDED IMMEDIATELY UPON PUBLICATION

July 1957
Lymphoid Goitres

T. Levitt

Postgrad Med J 1957 33: 352-358
doi: 10.1136/pgmj.33.381.352

Updated information and services can be found at:
http://pmj.bmj.com/content/33/381/352.citation

These include:

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/