Demonstration of Treponema pallidum in tissues
By a modification of the rapid method of silver impregnation of Ito, Ohtani, and Haba (1968)

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A new method for the demonstration of Treponema pallidum in tissues was described by Ito, Ohtani, and Haba (1968). Silver impregnation of treponemes could be carried out in a simple and rapid way, using their new method, which was based on the same property of silver nitrate as the older methods. Treponemes have been demonstrated by this method, subsequently modified by Ohtani (1969a), in papular lesions of secondary syphilis and in experimental syphilis in the rabbit.

We have used essentially the same technique in five cases of recent syphilis, using experimentally-induced syphilis in the rabbit testis as a control.

Materials and methods
The cases investigated may be classified as follows:
(1) Primary syphilis, lower lip;
(2) Primary syphilis, inguinal lymph node;
(3) and (4) Secondary syphilis, papular lesions of the trunk;
(5) Secondary syphilis, corymb-like lesions.

Rabbit testis, inoculated with Nichols strain treponemes, was used as a control.

Biopsy specimens were fixed in alcohol and/or in Bouin liquid, and embedded in paraffin. Sections 5–10μ thick and frozen sections were used for the staining procedure.

Technique
The method described by Ohtani was used first; with experience we made slight modifications to the original procedure. The following solutions remained active for months, stored under refrigeration:

1. Uranyl nitrate 1·0 g.
2. Silver nitrate 1·0 g.†
3. Sodium bisulphite 6·0 g.
4. Sodium sulphite (anhydrous) 1·0 g.
5. Hydroquinone 5·0 g
6. Phenidone (Geigy) 0·2 g.
7. Sodium lauryl-benzene-sulphonate 0·3 g.
8. Pyridine 20 ml.
10. Distilled water 300 ml.

Procedure
At room temperature:
(1) after deparaffinization, bring sections down to water;
(2) Solution (1) 5 min.;
(3) wash in two changes of distilled water;
(4) Solution (2) 20–40 min.;
(5) wash in two changes of distilled water;
(6) Solution (3) 10 min.;
(7) wash in distilled water; dehydrate, clear in xylene and mount in Canada balsam. Harris haematoxylin and eosin were used on some occasions as a counterstain.

Results
Treponema pallidum was identified both in the primary and in the secondary syphilitic lesions; only the so-called ‘corymb-like’ secondary syphilis was completely devoid of treponemes. The typical spiral

*Ito and other (1968) and Ohtani (1969a), added laurylpyridinium chloride, 0·3 ml. as a cationic surface active agent. In a first attempt, we used 0·3 ml. of a 10 per cent. solution, but we later discarded it, observing no substantial differences in the staining properties. On the other hand, Ohtani (1969b) maintained that laurylpyridinium chloride is ‘not very active’ in this phase of the procedure.
†Ito and others used 2 g.; Ohtani (1969a) used 0·2 g. silver nitrate.
formation was recognizable in the majority of the sections. However, the structure of the treponeme was not fully identifiable in sections from the epidermis of the lower lip in a case of primary syphilis (Fig. 1); we believe that this could be due to the fact that the patient had previously treated the lesion with an antibiotic ointment.

In the inguinal lymph node of primary syphilis, treponemes were typically arranged near to or within the walls of small vessels; some of them were detected entering the blood stream of large vessels (Fig. 2).

In secondary syphilitic papular lesions, treponemes were usually encountered among the cells infiltrating

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**FIG. 1** Treponemes, altered by topical antibiotic treatment, among the epidermal cells of the lower lip, in a case of primary syphilis. × 540

**FIG. 2** Treponemes in the walls of a large blood vessel, some of which are entering the blood stream. Inguinal lymph node from a case of primary syphilis. × 540
the lesion or in the proximity of small blood vessels (Fig. 3).
Many treponemes were present in the interstitial tissues of the rabbit testis, thus providing good control material for the various modifications of the procedure which we tried (Fig. 4).
Surprisingly, no treponemes were found in the corymb-like lesions of secondary syphilis. We were

**FIG. 3** Treponemes lying in the cellular infiltrate of the dermis in a case of secondary syphilis. × 600

**FIG. 4** Abundance of treponemes in the interstitial tissue of the rabbit testis, not counterstained. × 800
unable to explain this by our own experience or by reference to the literature.

Comment and conclusions

Walter, Smith, Isaac, and Gager (1969) described a new modification of a silver stain impregnation method for *Treponema pallidum*. Their entire procedure, though demonstrably valuable, seems to be over complicated compared with the method presented here. It involves more than twenty steps, and six solutions which cannot be stored; moreover, the method works at temperatures ranging from 60° to 70-75°C. At all events it indicates the measure of interest in this matter and justifies attempts to evolve new procedures. The opportunity of confirming data obtained in different laboratories seemed to us another good reason to have undertaken the present work.

In substantiating all the conclusions of the Japanese workers who first described this method, we agree that the advantages can be summarized as follows:

(1) This staining method is more rapid and much simpler than those described in the past.

(2) The entire procedure can be performed on ordinary histological sections and not on the whole specimen as in other methods, thus obtaining a better homogeneous staining.

(3) It offers the prospect of employing counterstains.

(4) The staining procedure was successful using either alcohol or formalin or Bouin liquid for fixing; the method also worked on frozen sections.

(5) Both the silver nitrate and the reducing solutions can be used at room temperature, and all solutions can be preserved for months without loss of activity.

In conclusion, we believe that the method described above represents a significant improvement on the older techniques and deserves wider recognition. The slight modifications to the original technique proposed here, though not essential, gave an equally distinct and clear impregnation of *Treponema pallidum*.

Summary

The authors describe the results and advantages offered by their modification of a method of demonstrating treponemes in tissues by silver impregnation. The reliability and simplicity of this method has been proved in cases of human syphilis and in experimental syphilis of the rabbit testis.

References

OHTANI, M. (1969a) Ibid., no. 82, p. 1
—— (1969b) Ibid., no. 70, p. 1

Mise en évidence de *T. pallidum* dans les tissus par une modification de la méthode rapide d'impregnation à l'argent d'Ito, Otaha, et Haba (1968)

SOMMAIRE

Les auteurs exposent les résultats et les avantages que présente la modification qu'ils ont apporté à une méthode de mise en évidence de tréponèmes dans les tissus par une impregnation argentique. Le degré de confiance et la simplicité de cette méthode ont été démontrés dans des cas de syphilis humaine et sur les testicules de lapin au cours de la syphilis expérimentale.