STEVENS-JOHNSON SYNDROME TREATED WITH CORTICOSTEROIDS


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The Stevens-Johnson Syndrome, otherwise known as erythema multiforme exudativum, has lately become more widely recognised as being liable to present as an acute medical emergency. The case described here is of special interest for several reasons: it would seem to be the most severe example of the disease ever recorded, which did not end fatally; it was complicated by acute dysphagia necessitating treatment with intravenous fluids, by acute retention of urine, and by bilateral corneal ulceration which finally healed without scarring; and it responded well to treatment with steroid hormones, both systemic and topical.

Case Report

Mary X, a laundress, unmarried, aged 26 years, was admitted to hospital on 12th November, 1956.

Present History

Some three weeks previously she had consulted her general practitioner because of nausea, occasional vomiting, and intermittent colicky pain, radiating from the right iliac fossa upwards and retrosternally. The pain had no connection with meals, micturition or defaecation. There was no story of menstrual upset; her menses were usually heavy with a good deal of pain. She was diagnosed as suffering from nervous dyspepsia, and treated with aluminium hydroxide and phenobarbitone tablets; she took the latter for 14 days in a dose of ½ gr. thrice daily. On 10th November she became listless; her eyes were inflamed and itchy, her throat sore, and a frontal headache developed. She was notified to us as probably having an acute respiratory infection, more especially as there was an epidemic of acute laryngo-tracheo-bronchitis in the area at the time.

Past History

Measles, mumps and varicella in childhood. Menarche at 13 years of age. General health excellent. No known contact with any infectious disease; no contact with dead birds or sick animals, or with the new detergents. No previous allergy of any kind.

Clinical Findings on Admission

The patient was a slim, listless girl, slightly drowsy, but not dyspnoeic; temperature 99.4° F., pulse 96/min., respirations 28/min. She had a generalised mild, but itchy macular erythema on her face, trunk and limbs, but not on her scalp, with additional facial oedema, and small ulcers on the lips and buccal mucosa. Her eyes were swollen and red. The lung fields were clinically clear, the blood pressure 130/70 mm. Hg., and the cardiovascular system normal except for systolic murmurs at the mitral and pulmonic areas. In the abdomen there was localised direct tenderness and slight guarding in the right iliac fossa and slight left-sided subcostal tenderness. No masses were palpable, and bowel sounds were normal. Posteriorly there was moderate tenderness of the spinal processes of the upper thoracic vertebrae (especially T4), and slight tenderness in both loins. Per rectum slight ballooning was noted, together with tenderness forwards and to the right.

Immediate Investigations

The urine was chemically and microscopically normal, and sterile on culture. Blood examination showed photohaemoglobin 14.4 G/100 ml., P.C.V. 44 per cent., M.C.H.C. 32 g. per cent., white cells 9,900/cu. mm. (neutrophils 82 per cent., lymphocytes 11 per cent., monocytes 7 per cent., eosinophils nil, basophils nil), films normal. Chest skigram showed no abnormality.

Diagnosis and Treatment

It seemed that the picture of coryza with rash was most like the Stevens-Johnson Syndrome, although atypical measles could not be entirely excluded. As the patient was not dangerously ill and seemed likely to have some abdominal surgical condition present as well, it was decided not to use steroid hormones meantime. Treatment
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FIGURE 1.

DISEASE
STEVEN'S-JOHNSON
SYNDROME

Notes of Case
MARY X.

Age 26

STEREIDS
Intravenous Hydrocortisone.
Free Alcohol.
Intramuscular Cortisone Acetate.

ANTIBIOTICS
Oral Phenoxy-Methylpenicillin.
Intramuscular Benzylpenicillin.
Intramuscular Streptomycin Sulphate.

DATE OF ADMISSION
12.11.56.

Pulse Resp.

Entered at Stationers Hall Printed and Published by Wollaston & CO.® Gould's Clinical Chart

was started with oral diphenhydramine (50 mg. q.i.d.) and oral phenoxymethyl penicillin (250 mg. q.i.d.).

Progress

November 13. Patient drowsy; temperature 103° F.; rash widespread and intense. Dysphagia and dysphonia apparent. Antibiotic changed to intramuscular benzylpenicillin (500,000 U. b.d.).

November 14. Condition much worse. Tongue ulcerated, swollen and dry; dysphagia complete; eyelids ulcerated and conjunctivae markedly reddened; micturition very painful. The abdominal symptoms and signs had nearly disappeared, and it was decided that treatment with steroid hormones and intravenous fluid was now imperative.

Accordingly, diphenhydramine was stopped, and a slow intravenous (stick-in) drip set up of 5 per cent. aqueous glucose; 75 mg. hydrocortisone free alcohol were administered in this medium during the next 8 hours, and thereafter cortisone acetate was given intramuscularly every 8 hours (see Fig. 2). Benzylpenicillin was continued intramuscularly and streptomycin sulphate (0.5 G. b.d.) begun by the same route. Sedation was provided by intravenous morphine in doses of 1/12 gr.

FIGURE 2

Mary X

Stevens-Johnson Syndrome

DOSAGE OF STEROID HORMONES
A—Hydrocortisone-Free Alcohol

<table>
<thead>
<tr>
<th>Date</th>
<th>Route</th>
<th>Total Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>14/11</td>
<td>I.V.</td>
<td>75 mg. in 750 ml. 5 per cent. aqueous glucose</td>
</tr>
</tbody>
</table>

B—Cortisone Acetate and Potassium Chloride

<table>
<thead>
<tr>
<th>Date</th>
<th>Total Dose I.M. Cortisone Acetate</th>
<th>Total Dose Oral KCl</th>
</tr>
</thead>
<tbody>
<tr>
<td>15/11</td>
<td>300 mg.</td>
<td>4 g.</td>
</tr>
<tr>
<td>16/11</td>
<td>100 mg.</td>
<td>4 g.</td>
</tr>
<tr>
<td>17/11</td>
<td>100 mg.</td>
<td>4 g.</td>
</tr>
<tr>
<td>18/11</td>
<td>50 mg.</td>
<td>4 g.</td>
</tr>
<tr>
<td>19/11</td>
<td>50 mg.</td>
<td>4 g.</td>
</tr>
<tr>
<td>20/11</td>
<td>50 mg.</td>
<td>4 g.</td>
</tr>
<tr>
<td>21/11</td>
<td>50 mg.</td>
<td>4 g.</td>
</tr>
<tr>
<td>22/11</td>
<td>50 mg.</td>
<td>4 g.</td>
</tr>
<tr>
<td>23/11</td>
<td>50 mg.</td>
<td>4 g.</td>
</tr>
<tr>
<td>24/11</td>
<td>37.5 mg.</td>
<td>4 g.</td>
</tr>
<tr>
<td>25/11</td>
<td>12.5 mg.</td>
<td>4 g.</td>
</tr>
<tr>
<td>26/11</td>
<td>—</td>
<td>2 g.</td>
</tr>
<tr>
<td>Total</td>
<td>925 mg.</td>
<td>46 g.</td>
</tr>
</tbody>
</table>
November 15. Slight general improvement apparent, but dysphagia still severe, and acute retention of urine now present. Gross oedema of the vulva completely obliterated the normal contours of the labia majora and minora, and there was ulceration of the skin around the vaginal introitus. Catheterization was accomplished with great difficulty and 30 fl. oz. of clear urine obtained. Treatment continued with antibiotics and intramuscular cortisone acetate, and oral potassium chloride (in solution) begun in doses of 1 g. q.i.d., which the patient managed to swallow. Silicone cream applied routinely to all pressure points.

November 16. Dysphagia had decreased sufficiently for oral fluids to be taken without great difficulty, and the intravenous glucose was therefore stopped; altogether 7.5 pints had been given. Spontaneous voluntary micturition returned. The condition of the eyes began to give cause for anxiety. Until now they had been only moderately inflamed, and no local therapy had been given. However, this evening, the lids became rapidly oedematous and heavily crusted along the margins. Frequent swabbing with normal saline was started, and penicillin drops instilled. Elsewhere the skin eruption was getting no worse; on the face most of the original papules had progressed through the vesicular stage to crusting. On the trunk and limbs most of the lesions remained as papules, slowly diminishing in size from about 1/4 in. in diameter, but becoming increasingly red. Pruritus had nearly disappeared. This evening, for the first time since admission, the temperature decreased to 98°F.

November 17. Acute urinary retention recurred which necessitated catheterisation; vulval erosion was very marked. Fluids could be swallowed fairly easily, but not solids. Both eyes were fully closed by oedema, and sloughing of the lids was gross; tenderness was severe enough to preclude the possibility of proper examination except under a general anaesthetic. She was taken to theatre, anaesthetised with cyclopropane and oxygen, and examination and débridement carried out. Both corneae were found, on testing with fluorescein, to be widely ulcerated. Local treatment was begun as follows: Saline swabbing of lids every 2 hours, followed by the instillation of 1 per cent. hydrocortisone acetate drops and 1 per cent. chloramphenicol ointment, and the instillation of 1 per cent. atropine sulphate drops every 4 hours.

November 18. Slow improvement maintained. She could now sit up and speak faintly. Both eyes were partly open, with no crusting, vision being, of course, completely obscured by the local treatment. Anuria was still present, and catheterisation again performed. This time vulval oedema had decreased sufficiently to make the location of the urethral meatus fairly easy. Ketonuria was moderate. Penicillin therapy was changed back to phenoxymethyl penicillin orally; streptomycin dosage changed to 1 g. once daily.

November 19. Taking oral fluids well. Crusts separating from face and lips; eyes satisfactory. After a small saline enema voluntary micturition became established.


November 22. Very miserable and tired, but no signs of psychosis developing. This state seemed due to her being continually disturbed for treatment of some kind, and to the effect of prolonged starvation. Sedation with intramuscular morphine (½ gr.) and chlorpromazine (25 mg.) being employed.

November 23. Irritable and obtuse. All sedatives stopped and deliberate bullying used to induce her to take nourishment, including 'Complan' (Glaxo).

November 24. Marked improvement; now cooperative and more cheerful. Facial crusts clearing; diction restricted mainly by crusts on the lips. Vulval oedema minimal. Antibiotics stopped.

November 25. Eating semi-solids. Face be-
gining to assume normal contours. Systemic cortisone therapy stopped. Corneal ulceration minimal; local treatment altered so that atropine henceforth applied only 12-hourly and hydrocortisone and chloramphenicol 4-hourly.

November 26. Pharyngitis with cervical lymphadenopathy. Her mouth could now open sufficiently to permit proper inspection; the mucosa of the cheeks, tongue, hard and soft palate, and uvula presented a crazy-paving or 'geographic' appearance with plaques of regenerating epithelium; 1.25 Mega U. of fortified benethamine penicillinil ('Triplopen': Glaxo) given intramuscularly. Potassium chloride stopped.

November 29. Left cornea completely healed. Treatment continued to both eyes as before.

December 2. Full diet taken for the first time. Attitude cheerful and co-operative; sleeping well. Skin of labia minora still slightly ulcerated, but no dysuria. Recrudescence of acute pharyngitis; 1.25 Mega U. intramuscular 'Triplopen' again given.

December 3. Eye treatment altered as follows: 6 per cent. sulphacetamide drops applied to both eyes four times daily (but not at night) in place of the chloramphenicol ointment, hydrocortisone drops applied four times daily likewise, and atropine drops instilled 12-hourly as before. Allowed up in a chair.

December 7. Patient happy and laughing, but not euphoric; her starting to wear sunglasses to offset her photophobia enabled her cubicle, for the first time, to be unscreened to admit full daylight. All facial crusts had now separated; general desquamation continued over the rest of the body, with faint brownish staining. Hydrocortisone eye drops stopped, sulphacetamide continued, and atropine applied henceforth only once daily.

December 9. Patient up walking about.

December 14. Both corneas completely healed. All local eye treatment stopped.

December 22. Patient allowed home. Slight photophobia present; pupils not fully mobile. Scaling heavy on the soles of the feet, but only slightly elsewhere. Five per cent. aqueous witch hazel was proving an excellent astringent to minimise any pruritus and tenderness of skin and nail roots. Facial appearance nearly normal, except for the total absence of eyelashes; pigmentation minimal, being mostly restricted to a faint malar redness. No trace remained of the original tenderness in her abdomen and back.

Special Investigations

The following were all done within a few days of admission and were all negative: Blood cultures, cultures from catheter specimens of urine, serum Wasserman and Meinicke Reactions, serum Widal Reactions (B. Typhosus H. & O., B. Paratyphosus A. H. & O., B. Paratyphosus B. H. & O., non-specific salmonella, Brucella abortus and Brucella melitensis), and pregnancy tests for urinary choriomic gonadotrophins.

Serum potassium, sodium, chloride and urea estimations were made in case starvation plus the effect of steroid therapy should produce a biochemical imbalance, but values remained normal, and serial readings of blood pressure were also all normal. It was noted that at the onset of hormone therapy the cloting time shortened to about 15 sec., but this effect lasted only for about two days.

Blood examinations on November 23, November 29 and December 20 showed red cell and white cell values within normal limits, and never an eosinophilia of more than 1 per cent.

Throat swabs taken at the attack of pharyngitis on November 26 were negative on culture for Strep. haem., B. diphtheriae and Vincent's organisms.

Chest skiagram on December 17 showed no abnormality. E.S.R. on December 18 was 28 mm. (Westergren) in the first hour. Electrocardiogram on December 20 showed no abnormality.

During convalescence two serum specimens were tested for evidence of recent virus infection. Complement-fixation tests for the following were all negative:

Influenza soluble antigen (types A, B and C), newborn pneumonitis virus antigen, adenovirus group antigen, psittacosis-L.G.V. group antigen, and Q fever (R. burnetii antigen).

Further Progress

When seen as an out-patient again on 14.1.57, Mary's general condition was excellent.

Skin. Rash completely healed in all areas. Minimal brown pigmentation in the malar areas, and slight telangiectatic mottling on the nose.

Nails: Shedding of nails progressing in both hands and feet. The old nails detached at the roots and new ones growing below them.

Abdomen. Occasional left hypochondriac pain, especially on laughing; and occasional mild retrosternal pain (heartburn). Rib cage mobile at left 6th costal cartilage; pressure there reproduced this pain. Most suggestive of a slipping rib syndrome.


Genito-urinary system. No dysuria. Menstruation not yet established.
Discussion

The eponym Stevens-Johnson Syndrome dates from the original paper by Stevens and Johnson in 1922, but cases of a similar nature had been described previously in the 19th century; Rosenberg and Rosenberg (1940) mention a case recorded by Alibert and Bazin in 1822. However, most authorities give credit to von Hebra for the first description of the disease in 1860. It was then known as erythema multiforme exudativum and consisted of skin erythema, ulceration at the orifices, and general constitutional upset. In 1916, Reiter described a syndrome affecting German soldiers, and consisting of urethritis, conjunctivitis and arthritis. This is currently regarded as being closely related to the so-called non-bacterial or non-specific urethritis which has displaced gonorrhoea as the most prevalent venereal disease in U.K. Epstein (1947) compared Reiter’s Disease with keratitis blenorrhagica and psoriasis arthropathica, and Robinson and McCrumb (1950) reviewed a variety of cases of erythema multiforme exudativum previously described and regarded them as being all varieties of the so-called ‘muco-cutaneous syndrome.’ The matter is still highly controversial and a wealth of literature exists dealing with the arguments. The Commission on Acute Respiratory Diseases (1946) declared that the Stevens-Johnson Syndrome was an eponym for which there was no justification.

Some writers consider von Hebra’s cases to have been so mild that they would have an erythema multiforme exudativum (Hebra) described in contradistinction to the other acute erythema multiforme exudativum (Stevens-Johnson). Cooper (1955) considers the term ‘muco-cutaneous—ocular syndrome’ sufficient to include all cases of the von Hebra and Stevens-Johnson Syndromes and Reiter’s Disease.

The diagnosis in the typical case of Stevens-Johnson Syndrome rests on the presence of ulcerative (or ‘exudative’) erythema multiforme, which may involve, besides the skin, the mucous membrane of the alimentary canal, the respiratory system, the genito-urinary tract, and the eyes. Haemorrhagic neuroretinitis (Cooper 1955) and meningitis (Todd 1953) have, on occasion, been described as features; pneumonia as shown by skiagram may be much more extensive than is thought from clinical findings (Cooper, 1955). Shedding of nails is a fairly common sequel (Romer, 1953, and Humphrey, 1955). Dressner (1949) reported a case with ECG changes suggestive of pericarditis; joint effusions were also present.

Mauriello (1954) reviewed a series of 14 patients; four out of 13 were suspected initially of having aphthous ulcers and one of 10 with eye signs was seen by an ophthalmologist for one week before any other signs developed.

Ashby and Lazar (1951) have conducted the most extensive review to date, comprising 81 cases. The sex incidence was approximately 9:2 in favour of males; maximal incidence was in December, January and February, with a secondary rise in May. The skin of the scalp was always free from lesions. Urethritis was common, but retention (and then ‘transient’) was present in one case only. Conjunctivitis was present in 74 patients, being bilateral in all but four. Seven had keratitis, and of these five went totally blind, one had residual scarring, and only one made a complete recovery without scarring. The death rate in the 81 cases was eight, of which seven had pneumonia present. The actual cause of death was non-specific: it has been attributed on occasions (Librach, 1955) to circulatory collapse.

The etiology of the Stevens-Johnson Syndrome remains obscure. Some patients have recurrent attacks (Finland, 1948). Numerous viruses have been incriminated, e.g. vaccinia (Grant, 1953), atypical pneumonia (Mauriello, 1954), psittacosis (Finland, 1948) and mumps, foot and mouth disease, and herpes simplex (Cooper, 1955), but most writers consider the condition to be a non-specific allergic reaction, notwithstanding the fact that even mild eosinophilia is seldom noted (excepting Goldfarb (1946), Finland (1948) and Vandermeer et al. (1953)) and that the ‘incubation period’ is extremely variable. Foods, bacterial toxins, aspirin, barbiturates and sulphonamides are commonly mentioned (e.g. Cooper, 1955), less commonly phenolphthalein (Mauriello, 1954), hydantoinates (Librach, 1955), and phenylbutazone (Conway, 1956) have been blamed. Barbiturates and sulphonamides are the two groups of drugs most commonly labelled as causative, and here again controversy arises, for some authorities (Sommerville, 1957) would consider my case to be one of barbiturate intoxication rather than the Stevens-Johnson Syndrome.

Treatment is no less controversial. Some authorities treat cases conservatively; others use antihistamines, although no one has claimed that they are of much use. Jones (1951) treated three cases with chlorotricycline, and in one of them, being used as his own control, relapse occurred with a placebo; Grant (1953) recorded an immediate response to chloramphenicol; whilst the Lancet (1951) in an editorial on the subject pointed out the considerable risk of inducing moniliasis by broad-spectrum antibiotic therapy.

Lately, steroid hormones have been more widely used. Librach (1955) considers they are good if employed early enough, and Fishman (1951) has used them topically on skin lesions. Corticotro-
phen is variable in response (Campbell 1955),
perhaps due to variations in the potency of dif-
ferent preparations, and perhaps due to the
adrenals not responding to stimulation because of
toxic depression. There seems to be no record to
date of intravenous hydrocortisone or topical
hydrocortisone having been used to arrest the
progress of an acute case of the disease.

The severest case previously recorded in the
literature appears to be that of Dresner (1949)
already mentioned. Dysphagia was severe, but
never sufficient to prevent oral fluids being taken;
melena was marked. There was no corneal
ulceration, but retinal haemorrhages were present.
In spite of severe proteinuria and dysuria, acute
urinary retention did not occur. Rectal saline was
administered early and blood transfusion employed
later.

My own patient, Mary, was more critically ill, in
the early stages, than any case I have seen re-
corded, but for reasons different from Dresner’s
patient. Her acute dysphagia made intravenous
nourishment a necessity, which is in itself a
difficult problem without being complicated by
cortisone therapy and urinary retention. The
latter, indeed, seemed at one stage as if it would
require suprapubic cystostomy to relieve it, be-
cause the external genitalia were swollen and
ulcerated almost beyond recognition. The em-
ployment of continuous drainage by indwelling
urethral catheter was thought inadvisable in view
of the risk of producing pressure necrosis of the
mucosa. In addition to all this she developed a
fulminating bilateral keratitis. In retrospect, there
seems no doubt that Mary owes her life and her
vision to corticosteroid therapy; it was quite re-
markable how quickly she began to improve after
the intravenous hydrocortisone, and how quickly
the ocular oedema and exudation subsided during
treatment with hydrocortisone drops.

Probably had bolder use of intravenous hydro-
cortisone been made, clinical response would have
been more dramatic. A total of 75 mg. was given
in about eight hours; 200 mg. in eight hours is
recommended by some writers (Pfizer, 1956).
Mary’s chest X-ray was clear, but the possibility
of an intra-abdominal lesion being present, such
as peptic ulcer, which had caused her original
symptoms, was a constant worry, and even al-
though she now presents with symptoms and
signs suggestive of a left slipping rib, her original
vomitting and epigastric pain remain unexplained.

In the presence of acute dysphagia the only
alternative to intravenous hydrocortisone was an
intramuscular preparation. Absorption here is
slow compared to the oral route and may take
from three to six days for one dose (Cope, 1956);
accordingly, Mary was kept on the intramuscular
preparation even after she could swallow, to
avoid the possibility of marked fluctuations in the
blood hydrocortisone level. One need be less
concerned about the gradual cessation of steroid
administration by the intramuscular route than by
the oral route because of the depot effect already
mentioned (Cope, 1956).

It has been assumed that Mary’s condition was
due to allergy to phenobarbitone. Now that she
has recovered, the obvious way to prove this
hypothesis would be to administer oral phenol-
barbitone again in small doses to see if her con-
dition recurred, but I think that such a procedure
would be quite indefensible in this particular
instance. Mary has therefore been told that she is
highly allergic to all barbiturate preparations and
that she is on no account to accept tablets or in-
jections at any time without so informing the
practitioner treating her.

As has been already noted, there was no past
history of drug administration or contact with
sick animals or dead birds, or of previous allergic
conditions.

Now that chemical analogues of cortisone are
becoming more widely used, prednisolone would
probably, by virtue of its lesser effect on serum
electrolytes, be a more convenient drug to use in
treating a similar case in the future.

Summary

A case is described of Stevens-Johnson Syn-
drome, featuring acute dysphagia, acute retention
of urine and bilateral corneal ulceration.

It is suggested that this case may be the most
severe example of the condition ever recorded.

The successful employment of steroid therapy,
both parenteral and topical, is described in detail.

The clinical features and etiology of the Stevens-
Johnson and allied Syndromes are discussed
briefly.

It is suggested that prednisolone rather than
cortisone or hydrocortisone would be more suit-
able for the treatment of similar cases in the future.

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mission to publish this case; Dr. J. Boyd Adams,
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assistance.

Bibliography continued on page 392.
An end-to-side bypass homograft was inserted from the aorta to the femoral artery, but shortly after releasing the clamp it was evident that thrombosis had occurred in the graft. Efforts to obtain a flow were unsuccessful and the graft was therefore removed.

He made a straightforward post-operative recovery and his symptoms were unaltered.

**Case 2** (an R.A.F. flight-sergeant, age 46)

This patient had noticed increasing weakness of the left leg for one year. He was unable to walk further than 150 yards because of this weakness, but had no actual pain. He recovered after resting for a short period and was able to walk a further 150 yards.

On examination the colour and nutrition of the left leg were good and there was no obvious wasting. All pulses were present in this leg, but they were very weak. The pulses in the right leg were of normal volume. His blood pressure was 120/80 mm. There were no other abnormalities on physical examination or laboratory investigation.

An aortogram performed at the R.A.F. Hospital, Ely, showed normal appearances except for an abnormally narrow but smooth external iliac artery.

At laparotomy the aorta, right iliac and left common iliac vessels were of normal calibre, soft and thin-walled with no evidence of atheroma. The left external iliac artery was extremely small, but also soft and thin-walled. The internal iliac artery was present, but also small.

An end-to-side bypass homograft was inserted from the common iliac to the femoral artery, which was normal in every way, the graft being passed under the inguinal ligament. After completing the anastomosis the clamps at both ends of the external iliac were not removed until a good flow had been established through the graft for some minutes. It was thought that this measure might reduce the likelihood of clotting in the graft.

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FIG. 2.—Bypass around a hypoplastic left external iliac artery (case No. 2).

Post-operatively he made a straightforward recovery and had full ankle pulses. An aortogram (Fig. 2) demonstrated the graft to be functioning well. The narrow external iliac vessel can also be seen in this picture.

When seen two months after operation he was able to walk an unlimited distance.

In both these patients the arteriographic appearances and the operation findings suggested that the iliac arteries were congenitally hypoplastic. It was not felt justifiable to remove a section of the vessel for histology.
Stevens-Johnson Syndrome Treated with Corticosteroids

J. H. Mitchell

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