TREATMENT OF GONORRHOEA WITH NALIDIXIC ACID*

BY

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Nalidixic acid (Negram), a naphthyridine derivative, is a new chemotherapeutic agent, which is highly active in vitro against a wide range of Gram-negative organisms and has been reported to be particularly effective in vivo in urinary tract infections (Barlow, 1963; Lishman and Swinney, 1963; Ward-McQuaid, Jichlinski, and Macis, 1963). Although in laboratory tests the gonococcus was found to be very sensitive to nalidixic acid, early clinical trials with doses usually recommended for urinary tract infections had been disappointing†. The only positive results were reported from Mexico by Aparicio (1962), who treated thirty cases of gonorrhoea, 23 in the male and 7 in the female, with 1 g. four times a day over 5 to 8 days and attained apparent cure in 24. However, he followed up the patients for not more than a week and the majority for less than 5 days.

Present Study

Preliminary Investigations

(1) Cultures of Neisseria gonorrhoeae obtained from eight unselected male and female patients attending the clinic at St. Luke's Hospital in Bradford were tested for sensitivity to nalidixic acid with 30 μg. disks and all were found to be sensitive.

(2) Two male Pakistanis with acute uncomplicated urethral gonorrhoea, aged 21 and 27 years respectively, were admitted to hospital for treatment with a single dose of 4 g. (8 tablets) of Negram to assess the clinical effect and tolerance of this dose and to estimate the blood levels of the drug over 8 hours.

The patients were examined 6, 24, and 48 hours after treatment. The immediate response to nalidixic acid was comparable with that observed in penicillin-treated cases. After 6 hours there was a considerable relief from symptoms and the previously purulent urethral discharge changed to a more scanty muco-purulent one in which extracellular pairs of distorted gonococci, larger and more deeply red-stained than usual, were found; the organisms could not be cultured. After 24 hours there was a further subjective improvement and the gonococcus disappeared from smears and cultures. After 48 hours there was a clinical relapse and the smears and cultures became positive again.

In both cases the gonococcus was sensitive in vitro to nalidixic acid and penicillin disks, both before treatment and after its failure. There were no side-effects. The patients were successfully re-treated with 600,000 units procaine penicillin fortified by 200,000 units potassium benzyl-penicillin.

Blood samples for spectrophotofluorometric assay of nalidixic acid plasma levels were collected from the patients immediately before and 1, 2, 4, 6, and 8 hours after the drug was taken (Fig. 1, opposite). In one the peak was higher and the rise and decline steeper than in the other, but in both cases the level after 6 hours was still above 25 μg. per ml.

These preliminary investigations confirmed that nalidixic acid was active against the gonococcus both in vitro and in vivo, and that after a single dose of 4 g. a high plasma level was attained within 1 or 2 hours and was maintained for 4 to 5 hours. It appeared that further trials to arrive at an effective dosage schedule were worth while.

Clinical Material and Method

Over a period of 8 weeks (from March 9 to May 4, 1964) seventy-four more male patients with acute uncomplicated urethral gonorrhoea were treated with nalidixic acid by various dosage schedules. Eight were treated in hospital.

Table I (opposite) shows the age and race distribution of the patients, including the two preliminary cases (making a total of 76); it is fairly representative of a cross-section of the male population attending the Bradford Clinic with gonorrhoea.

The Asians were mostly Pakistanis; of the Europeans, thirteen were English, one Irish, and one Hungarian.

Treatment was started immediately after the diagnosis was made by Gram-stained smear, but in all but four

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* Paper read at the MSSVD Meeting in Dublin on May 29, 1964.
† Unpublished results in the files of the Bayer Products Company.

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RESULTS

The results in relation to dosage are shown in Table II.

Table II
Dosage and Results

<table>
<thead>
<tr>
<th>Dosage (g.)</th>
<th>Results</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>No further dose</td>
<td>4</td>
<td>—</td>
</tr>
<tr>
<td>One further dose</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>Two further doses</td>
<td>5</td>
<td>3</td>
</tr>
<tr>
<td>Three further doses</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td>Four further doses</td>
<td>10</td>
<td>14</td>
</tr>
<tr>
<td>Five further doses</td>
<td>3</td>
<td>10</td>
</tr>
</tbody>
</table>

The group treated with a single dose includes the two preliminary patients and two more who vomited after the first dose and discontinued treatment. In the following five groups, treatment consisted of an initial dose of 4 g. and from one to five further doses of 2 g. at 6-hrly intervals. In general the success rate increased with the number of doses given; in the group of nineteen patients treated with 14 g. over 30 hrs, there were only three definite failures. The highest failure rate, excluding the single dose group, was in the group of eight patients who, after a loading dose of 4 g., received from two to six subsequent doses of less than 2 g. (1 or 1.5 g.); here there were six failures in the seven patients who were followed up. Of the three patients who were treated with five equal doses of 2 g. over 24 hrs, one was cured and two failed, and of the four who were treated with an initial dose of 3 g. and three doses of 2 g. over 12 hrs two were cured and two failed.

All the re-infections occurred within the first fortnight and sexual re-exposure was admitted by the patients.

Of the defaulters*, five ceased to attend within 10 days of the completion of treatment, one received some tablets (presumably sulphonamides) from his own doctor during the follow-up period, and one was found to have syphilis for which treatment was started at the clinic; the last case is later described in detail.

* Since the report was read two of the defaulters re-appeared at the clinic with a re-infection: one 4 weeks after treatment with 12 g. over 24 hrs, the other 6 weeks after treatment with 14 g. over 30 hrs.

cases, the diagnosis was confirmed by culture and the organisms were tested for disk-sensitivity to nalidixic acid and penicillin. In the first forty cases the commercially available 30 μg. nalidixic acid disks were used, but later the tests were performed with disks of 5 μg. potency.

Although all the patients, except those who have defaulted, are still kept under review, for the purpose of this report the tests for cure were performed 3 weeks after treatment, when the fluid expressed by prostatic massage was examined by smear and culture. Before that the patients were examined at least twice during the first week and the majority also twice during the second week.

As a rule cases which failed to respond to nalidixic acid were treated with penicillin, but seven patients were treated twice with nalidixic acid, viz. five after failure of penicillin, all again without success, and two after re-infection, both successfully. This gives a total of 83 treatments performed at nine different dosage schedules.

# FIG.

1. Plasma levels of nalidixic acid in two cases after a single 4 g. oral dose.

**TABLE I**

Age and Race Distribution

<table>
<thead>
<tr>
<th>Age Group (yrs)</th>
<th>Racial Group</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Asian</td>
<td>West Indian</td>
</tr>
<tr>
<td>20-29</td>
<td>30</td>
<td>3</td>
</tr>
<tr>
<td>30-39</td>
<td>16</td>
<td>4</td>
</tr>
<tr>
<td>40-49</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>50+</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>49</td>
<td>9</td>
</tr>
</tbody>
</table>
Treatment failures (Table III) all became apparent within the first week. In three cases, of which two were treated with 12 g. over 24 hrs and one with 14 g. over 30 hrs, the gonococcus was resistant to nalidixic acid in vitro; in two of them the urethral discharge remained purulent from the first and the gonococcus was found in smears and cultures after 24 hrs; in the third case the gonococcus was sensitive before treatment and resistant after the relapse which occurred on the third day.

Penicillin Resistant Cases (Table IV).—In thirteen cases the gonococcus was resistant in vitro to penicillin disks of potencies ranging from 1/2 to 1/4 of a unit. Six were successfully treated with nalidixic acid and seven failed, including the five in which the drug was given a second time. The failures were retreated with large doses of penicillin. In all the seven cases in which the gonococcus was tested for sensitivity to streptomycin, it was found to be resistant.

**TABLE IV**

<table>
<thead>
<tr>
<th>Dosage of Negram</th>
<th>Result Failed</th>
<th>Result Succeeded</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>4 g. stat</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ 2 g. in 6 hrs</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>+ Two doses of 2 g. 6-hrly</td>
<td>2</td>
<td>2</td>
<td>4</td>
</tr>
<tr>
<td>+ Four doses of 2 g. 6-hrly</td>
<td>1</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>+ Five doses of 2 g. 6-hrly</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>+ Four doses of 1 g. 6-hrly</td>
<td>2</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td>3 g. stat</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ Three doses of 2 g. 4-hrly</td>
<td>2</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td>Total</td>
<td>7</td>
<td>6</td>
<td>13</td>
</tr>
</tbody>
</table>

Patients Sensitized to Penicillin.—Two patients who were known to be sensitized to penicillin were successfully treated with nalidixic acid, 12 g. over 24 hrs and 14 g. over 30 hrs respectively.

Plasma Levels (Fig. 2).—Plasma levels of the drug were estimated by spectrophotofluorometric assay at various times during and after treatment in the ten patients admitted to hospital and single estimations were made in nine of the patients treated at the clinic. Blood samples from the latter were collected between 20 and 24 hrs and in two cases 30 hrs after

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**TABLE III**

<table>
<thead>
<tr>
<th>Dosage</th>
<th>Second Positive Culture after Less than 2 days</th>
<th>3 days</th>
<th>4 days</th>
<th>5 days</th>
<th>6 days</th>
<th>7 days</th>
</tr>
</thead>
<tbody>
<tr>
<td>4 g. stat</td>
<td>+ No further treatment</td>
<td>4</td>
<td>4</td>
<td>4</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>+ 2 g. in 6 hrs</td>
<td>2</td>
<td>1</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>+ Two doses of 2 g. 6-hrly</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>+ Three doses of 2 g. 6-hrly</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>+ Four doses of 2 g. 6-hrly</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>+ Five doses of 2 g. 6-hrly</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>+ Variable Dosage 1 to 2 days</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>2 g. 6-hrly for 24 hrs</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 g. stat + 2 g. 4-hrly for 12 hrs</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
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the first dose and between 5 and 8 hrs after the last dose of the drug. Plasma levels varied considerably irrespective of the dosage, but there was an obvious correlation between the plasma levels and the clinical response. Excluding the early cases in which the plasma level readings were limited to 8 hrs, all the successes but one are seen in the upper half and all the failures except one in the lower half of the graph, the level of 25 µg./ml. being more or less the dividing line. The single high plasma level associated with failure is explicable, as in this case the gonococcus was resistant to nalidixic acid in vitro.

Associated Disorders (Table V).—There were two cases of primary syphilis (briefly described later), nine of non-gonorrhoeal urethritis of which three were due to trichomonas infection, six of asymptomatic non-specific prostatitis which were detected on routine examination for cure of gonorrhoea 3 weeks after treatment, and four cases of scabies. Positive serological tests for syphilis were detected in two West Indians who had suffered from yaws in childhood and the gonococcal complement-fixation test was positive in a Pakistani with a history of two previous gonorrhoeal infections.

<table>
<thead>
<tr>
<th>TABLE V ASSOCIATED DISORDERS</th>
<th>No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary syphilis</td>
<td>2</td>
</tr>
<tr>
<td>Non-gonococcal urethritis</td>
<td>9</td>
</tr>
<tr>
<td>Prostatitis</td>
<td>6</td>
</tr>
<tr>
<td>Positive standard tests for syphilis (yaws)</td>
<td>2</td>
</tr>
<tr>
<td>Positive gonococcal complement-fixation test</td>
<td>1</td>
</tr>
<tr>
<td>Scabies</td>
<td>4</td>
</tr>
</tbody>
</table>

Of the two patients with primary syphilis one, a Pakistani aged 29, already had an ulcer on the glans penis when he was first examined and was found to be suffering from acute gonorrhoea. The ulcer was secondarily infected with pyogenic organisms and had been treated with an antiseptic ointment and the search for Treponema pallidum was unavailing. He was treated with nalidixic acid (4 g. initially and two doses of 2 g.) and when he attended after 48 hours numerous Treponema pallida were detected in a clear exudate from the ulcer. Penicillin treatment was started and for convenience this patient was included in the group of defaulters. The serological tests in this case were positive.

The second patient, an unmarried Englishman aged 30 years, attended the clinic with gonorrhoea 11 days after sexual exposure. He was treated with 12 g. of nalidixic acid over 24 hrs and was seen on the 2nd, 3rd, and 7th day after treatment. There was no urethral discharge from the 3rd day onwards. When he attended on the 10th day a papule about 8 mm. in diameter with a slightly eroded surface and well defined outlines was noted on the inner aspect of the prepuce and Treponema pallidum was demonstrated in the serum from the lesion. Before treatment with penicillin was started prostatic smears and cultures were taken; as they were negative, this patient was included in the group of nalidixic acid successes. The serological tests in this case were negative.

Side-effects (Table VI).—These occurred in twenty patients, but were not severe and did not prevent the patients from working. The most frequent complaint was dizziness which in seven cases was associated either with headache or with nausea. One patient had transient diplopia after each dose of the drug. The four patients who vomited were all Pakistanis working at night and all of them, it was later discovered, had received the first dose on an empty stomach. All of them were treatment failures. There were no side-effects in any of the patients who were treated in hospital.

| TABLE VI SIDE-EFFECTS IN 83 COURSES OF TREATMENT |
|-----------------------------------------------|--|
| Side-Effects                                   | No. of Cases | Percentage |
| Dizziness                                     | 15           | 18         |
| Headache                                      | 4            | 5          |
| Vomiting                                      | 4            | 5          |
| Nausea                                        | 3            | 4          |
| Diplopia                                      | 1            | 1          |

Conclusion

The results of this trial indicate that nalidixic acid has a definite antigenorrhoeal action. In spite, however, of the high sensitivity of the gonococcus to the drug in vitro, in clinical practice large doses are needed to attain a significant proportion of cures. The drug should be taken at intervals of not less than 6 hrs over at least 30 hrs; individual doses should not be less than 2 g. and an initial loading dose of 4 g. is indicated. Even then in some patients an adequate plasma level, which appears to be in the region of 25 µg./ml. may not be reached or maintained owing probably either to poor absorption or to rapid elimination of the drug.

Obviously nalidixic acid has all the disadvantages of any drug requiring multiple doses by mouth and cannot be recommended for the routine treatment of gonorrhoea. The fact, demonstrated by two cases in...
this series, that it has no effect whatever upon
Treponema pallidum makes it a probable drug of
choice in cases of gonorrhoea in which a con-
comitant early syphilis is suspected. It may also be
useful for treatment of gonorrhoea in patients
sensitized to penicillin.

Summary

76 male patients with acute uncomplicated
urethral gonorrhoea were treated with nalidixic acid
using nine different dosage schedules.
In general the success rate increased with the
increase in dosage and a significant proportion of
cures was attained with an initial dose of 4 g. and
five further doses of 2 g. given at 6-hrly intervals.
Plasma levels of the drug were assayed in nineteen
patients and the results suggest that 25 μg./ml. is
probably the lowest effective level.
In two cases of concomitant primary syphilis,
nalidixic acid showed no effect upon Treponema
pallidum.

The unusually large doses of the drug which were
used in this trial were well tolerated and the side-
effects (which occurred in twenty patients) were on
the whole not severe.
It is concluded that nalidixic acid cannot be
recommended for routine treatment of gonorrhoea,
but that it may be useful in cases in which con-
comitant early syphilis is suspected and in cases of
sensitization to penicillin.

I am indebted to Dr G. B. Hill and Mr H. W. Hurst of
the Clinical Evaluation Unit of the Bayer Products
Company for the generous supply of Negram tablets and
disks and for practical help; to Mr A. Robertson of
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imations of the drug plasma levels; and to Dr H. G. Smith,
Director of the Public Health Laboratory in Bradford, for
sensitivity tests.

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Traitement de la gonorrhée par l’acide nalidixique

RÉSUMÉ

76 patients atteints de gonorrhée urétrale aigüe non
complicée furent traités par l’acide nalidixique en
utilisant neuf méthodes différentes. En général les succès
augmentèrent avec la dose et une proportion notable de
guérison fut obtenue en traitant par une dose initiale de
4 g. suivie de 5 doses de 2 g. données à 6 heures d’intervalle.
Le taux plasmatique des médicaments fut déterminé
chez 19 patients et les résultats suggérèrent que 25μg./ml.
est probablement le taux le plus bas d’efficacité.
Dans deux cas de syphilis primaire surajoutée l’acide
nalidixique n’eut aucun effet sur le tréponème pâle.
Les doses, plus importantes que d’habitude, utilisées
dans cette expérience furent bien tolérées et les effets
secondaires qui survinrent chez vingt malades ne furent
pas sévères dans l’ensemble.
En conclusion l’acide nalidixique ne peut être recom-
mandé pour le traitement de routine de la gonorrhée, mais
il peut être utile en cas de syphilis précoce surajoutée et en
cas de sensibilisation à la pénicilline.
Treatment of Gonorrhoea with Nalidixic Acid

Leonard Z. Oller

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doi: 10.1136/sti.40.4.256

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