Rauwolfia and Guanethidine: Pharmacology and Clinical Use in Treatment of Hypertension*

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BY COINCIDENCE, reserpine, which is the oldest and weakest, and guanethidine, which is the latest and most potent of the modern antihypertensive drugs, share a common basic pharmacologic action. Both agents lead to depletion of the body stores of norepinephrine and thus, to varying degrees, interference with the function of the sympathetic nervous system.1-4 The precise mechanism by which they prevent the storage of norepinephrine is unknown, but evidence for its immediate release is seen in the transient rise in blood pressure and cardiac output after intravenous injection of guanethidine.4 Reserpine produces no such acute changes in man, but a similar release has been observed in animal preparations.5,6

Since both these agents lead to depletion of the sympathetic transmitter substance, the vascular smooth muscle retains its responsiveness to norepinephrine; indeed, an increased sensitivity is observed. The pressor effect of certain vasoconstrictor drugs, however, is diminished. These drugs—such as tyramine, amphetamine and ephedrine—depend for their action upon the presence of stored catecholamines in the tissue.7-9

While guanethidine and reserpine share a similar mode of action, some important differences in their pharmacologic properties must be recognized. Guanethidine appears to produce actual blockade of adrenergic neurones in addition to depleting catecholamines. It does not cross the blood-brain barrier and depletes cerebral stores of norepinephrine11 as well as those of hydroxytryptamine (serotonin).12 To this may be attributed its sedative and psychologic and certain endocrine effects.13

Therefore it was originally believed that the site of action of reserpine was a central one, probably in the hypothalamus.14 However, the observation that at lower doses of reserpine the peripheral stores of norepinephrine are more depleted than the central ones15 has led to the conclusion that, as far as its hypotensive action is concerned, the main site of action of reserpine might be a peripheral one.

While the pharmacologic mode of action of reserpine and guanethidine tend to be similar, their clinical effects are conspicuous by contrasts rather than by comparisons.

**Reserpine: Clinical Effects**

Reserpine is a mild antihypertensive agent. In oral doses of 0.8 to 1.5 mg./day a reduction of approximately 17/11 mm. Hg in blood pressure has been found in two controlled double-blind studies.16,17 Both the recumbent and standing pressures are affected without reduction in cardiac output,19 renal blood flow, glomerular filtration rate19 or cerebral blood flow.20 The reduction in pressure progresses slowly over a period of weeks and persists for a similar period after the drug has been discontinued.

The side effects of reserpine (Table 1), although few and comparatively infrequent, are nonetheless serious since their gradual onset may delay detection; mental depression has been severe enough to result in suicide. Consequently, it is recommended that the maximal oral dose for long-term therapy should be limited to 0.5

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TABLE I
Side Effects of Reserpine

Nasal stuffiness
Increased appetite and weight gain
Fluid retention
Increased gastric acidity with possible aggravation of peptic ulcer
Diarrhea
Vascular collapse under anesthesia
Nightmares
Depression
Drowsiness
Tremor

mg./day. The reduced antihypertensive effect of the smaller dose can to some extent be offset by simultaneous use of a diuretic agent.

Clinical Application: Reserpine therefore can be recommended for the milder cases of hypertension in which the diastolic pressure does not exceed 105 mm. Hg. Oddly enough, parenteral reserpine (1–3 mg. intramuscularly) can be recommended in emergencies. The gradual fall in pressure over a period of 3 to 4 hours is rarely excessive in degree, and the minimal effect on renal function is ideal for the severely hypertensive subject. The accompanying sedation, however, may make the neurologic assessment of the patient difficult. Therefore, in the presence of hypertensive encephalopathy or cerebrovascular disease it is wiser to use a short-acting ganglion-blocking agent like camphodium (Arfonad) or parenteral pentolinium.

GUANETHIDINE: CLINICAL EFFECTS

In contrast to the gentle if insidious effects of reserpine, guanethidine is rough hewn, producing extensive sympathetic blockade and consequently more obtrusive side effects.

The onset of action is quite slow. It takes a few hours to achieve maximum blockade after parenteral injection. Intestinal absorption is incomplete, and the excretion of its metabolites is prolonged, only 24 per cent appearing in the urine 24 hours after a single oral dose. Therefore, two to three days are required to achieve a steady response with oral dosage. The effect on the blood pressure, like that of other autonomc blocking agents, is evident mainly in the standing position.

Guanethidine produces its hypotensive action by reducing both cardiac output and total peripheral resistance in recumbent position. There is also a small reduction in inulin and PAH clearances. On physical exertion the blood pressure may fall further in patients taking large doses of guanethidine, almost certainly because normal vasoconstriction fails in inactive vascular areas while dilation is occurring in the active muscles. The fall in pressure occurs in spite of an increase in cardiac output during exercise.

Hypotensive Effects: The fall in blood pressure on standing is a frequent cause of difficulty. The standing blood pressure is lowest in the morning on arising at which time patients

TABLE II
Therapeutic Effect of Guanethidine in 11 Hypertensive Patients

<table>
<thead>
<tr>
<th>Blood Pressures (mm. Hg)</th>
<th>Mean Dose (mg./day)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Recumbent</td>
<td>Standing</td>
</tr>
<tr>
<td>A.M.</td>
<td>P.M.</td>
</tr>
<tr>
<td>Before treatment</td>
<td>190/114</td>
</tr>
<tr>
<td>After treatment</td>
<td>166/104</td>
</tr>
<tr>
<td>Effect</td>
<td>-24/-10</td>
</tr>
</tbody>
</table>
frequently complain of fatigue, lassitude, and then may faint quite suddenly. Since the difference between the standing and recumbent pressures decreases as the day progresses, this effect may be missed when the patient attends the physician's office. It has further been shown that hypotensive effects of exercise are greater in the morning than later in the day. The magnitude of the effect both on the recumbent and standing pressures varies considerably from one patient to another as does the dosage required. A mean dose of 60 mg/day in a group of 11 hypertensives recording their blood pressures at home produced a mean reduction of 34/10 mm Hg recumbent and 68/17 mm Hg standing in morning readings. By the evening there was a mean increase of 15/8 mm Hg recumbent and 25/5 mm Hg standing pressure (Table II). The cause of this loss of therapeutic effect during the day is unknown.

Although they have similar effects on the blood pressure, there is no relationship between the sensitivity of the blood pressure to ganglion-blocking drugs and guanethidine, but little evidence has been seen for the development of tolerance with the latter.

The chief disadvantage of guanethidine, therefore, is that in many patients it is impossible to maintain an adequate reduction in recumbent pressure without unbecoming postural symptoms in the morning. Furthermore, if the pressure rises much during the day, the expected benefit from the drug will be limited to a very short period of time.

Side Effects: In addition to its hypotensive effect, guanethidine produces certain nonspecific effects (Table III); diarrhea has been the most prominent. From our own experience and from a review of the literature, diarrhea appeared to occur whenever the dosage of the drug was high or was increased rapidly. Weight gain or edema has been observed, and mental depression has been reported, as have parotid pain, nasal stuffiness, failure of ejaculation and muscle tremors. The cause of these side effects is not understood, but they are readily reversible by reducing the dosage or withdrawing the drug.

Clinical Application: At present the most powerful antihypertensive agents are the autonomic blocking agents. Of these guanethidine is undoubtedly the most effective in relation to its nonspecific side effects. This drug is, therefore, required by most patients with severe hypertension. The gradual onset of action and prolonged effect, however, make the drug unsuitable for use in emergency cases.

Side effects can be minimized by increasing the dosage gradually: 12.5 mg./day is the best initial daily dose increased to 25, 50, 75, 100 or 150 mg./day in weekly steps until the required effect is reached. If this gradual program is thought to be dangerously slow for the patient with severe disease, it is preferable to supplement it temporarily with a ganglion-blocking agent (mecamylamine) than to accelerate the regimen. The action of all blocking agents is variable; if an effective dose of the drug is to be used, inevitably occasions will arise when symptoms of the excessive blockade are experienced, e.g., after an infection, unusual exercise, in hot humid weather or even after a large meal. To help the patient maintain an optimal therapeutic response to the drug, he should be made aware of the symptoms he may encounter and how to deal with them.

In view of the inevitable fluctuations in pressure with autonomic blocking agents, they should be used with care in patients with advanced coronary or cerebrovascular diseases. However, they are not necessarily contraindicated in these conditions when severe hypertension is present; in fact, angina may be relieved as effective control of blood pressure is achieved.

SUMMARY

Both reserpine and guanethidine appear to utilize the same fundamental pharmacological mode of action which leads to depletion of tissue catecholamines.

Guanethidine produces effects on the blood pressure similar to those of the ganglion-blocking agents without the troublesome side effects. It is an extremely potent agent, however, and is likely to produce symptoms attributable to postural hypotension.

REFERENCES


