PAROXYSMAL TACHYCARDIA; AETIOLOGY AND PROGNOSIS OF ONE HUNDRED CASES

BY

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The early papers on paroxysmal tachycardia gave its essential features with great accuracy, and the modern reader will enjoy their lucid descriptions. After the classic account by Cotton (1867) with pulse tracings by the great Burdon-Sanderson, many cases were reported, and Bristowe (1887) and Bouveret (1889) collected these and added more of their own. Bristowe's paper was "On recurrent palpitations of extreme rapidity in persons otherwise apparently healthy" and Bouveret first introduced the modern name in "De la Tachycardie essentielle paroxystique"; he stressed that it did not produce organic heart disease, but thought it a serious malady, and as he recorded a fatal outcome in many of the longer attacks he may be responsible for the guarded prognosis often given in books. A few of these attacks were perhaps paroxysmal flutter, but with this exception the papers just quoted might still be read as accurate accounts of much of our clinical knowledge. Herringham (1897) emphasized that paroxysms often started in childhood and that the patient lived to a good age. Hoffmann (1900) wrote a book analysing 135 reported cases.

For some time little more was added, till the electrocardiogram made our knowledge more precise and enabled us to distinguish paroxysmal auricular flutter and to differentiate the various types of paroxysmal tachycardia. Hume (1930) has reviewed the whole subject, giving a full list of references, and for this reason most papers are only referred to here in connection with special points as they arise. Many of the large number of recent papers deal with the electrocardiographic features of single cases and pay less attention to the clinical picture of paroxysmal tachycardia as a whole.

This paper is based on observations on one hundred cases seen during the years 1925–32. A few with no features of special interest may have escaped indexing, but they were not many, and otherwise it is an unselected series. Cardiograms * of the paroxysms were obtained in 42; 30 others were seen by

* A good short word has not yet been coined for electrocardiogram, which is wasteful of time and space now it is in such constant use; no confusion is likely to arise by the substitution of the word cardiogram.
a doctor (occasionally by one of us) in the attack, though no graphic record was obtained; and in 28 the diagnosis was made on the history alone. The occurrence of paroxysms in five students at Guy’s Hospital during the years 1928–30 suggested that the condition was not uncommon, but only two more have been seen in the subsequent seven years.

Many of these patients have been followed over a long period, and as the immediate prognosis is nearly always good, this is essential for any complete picture. We have followed 16 until their death, not always from heart disease; 53 have been seen or have replied fully to letters in 1937–38 and most of these and another 13 were seen in 1934–35, and we have no reason to expect any great change in their condition. The remaining 18 were last seen during 1925–32 and generally attacks had already occurred for some years. The series is therefore likely to give a fair view of those cases severe enough to come under medical observation, and has been followed long enough to give some approximation to a correct prognosis.

DIAGNOSIS

Often the attack is not seen and there is only the history; this applied to 28 of our patients. Reliance must then be placed on the abrupt onset, and as a rule on the equally abrupt end. The rapid beating of the heart is nearly always the first symptom felt, whatever others may follow. If the patient is able to define the number of attacks and their duration it is almost diagnostic, since ordinary palpitation is much more vague and indefinite in its character. Enquiry must then be made as to whether the heart was regular or irregular, and, perhaps surprisingly, the patient’s statement usually proves to be reliable, and only one had to be excluded from this series when a subsequent record showed it to be paroxysmal fibrillation. This suddenness of onset is the most reliable single symptom for diagnosis—“As if the pendulum of a clock had suddenly lost its weight” in the words of one patient. During the attack the heart action is a constant steady fluttering, and this word is often used.

The end of the attack is usually as sudden as the onset, but when the heart is diseased the patient may often be incapable of saying exactly when the end came. “I think it ended a few minutes ago, but it is still going very fast” was one description. This is specially so when the heart muscle is becoming worse, and in one such observation the rate appeared to fall gradually from 180 to 100 in the course of three hours, simple tachycardia having followed the paroxysm so that the exact ending was difficult to detect clinically: the offset is therefore less reliable than the onset. Most other symptoms are too variable to be reliable, and extrasystoles before or after the attack may obscure the clear picture.

When the attack is seen the diagnosis rarely remains doubtful, though it may not be thought of if the heart rate is 140 or less. It is sometimes difficult to distinguish paroxysms of tachycardia from those of flutter if a constant degree of heart block makes the ventricular rate regular, and impossible if the diagnosis has to be made on the history alone. The auricular rate in flutter is
generally 240–360; 4:1 heart block is rare in an untreated patient, and regular 3:1 heart block extremely rare; but 2:1 heart block is common and gives a rate of 120–180, i.e. within the usual limits of paroxysmal tachycardia. Again, flutter with the auricle at a slow rate without heart block (220–240) may be difficult or impossible to distinguish from paroxysmal tachycardia even on the cardiogram, and the decision may have to be made on clinical grounds. Lastly, a paroxysm of tachycardia may fall above its usual limits of 240 (Fig. 1).

FIG. 1.—Ventricular tachycardia, auricle 128, ventricle 274; from a boy whose heart was otherwise normal (Case 69). The auricular waves have been marked with a • in lead II. The time marker in this and subsequent records indicates fifths and twenty-fifths of a second.

Apart from the rate, the following points may help to distinguish between paroxysmal flutter and tachycardia. If the regular rhythm is occasionally interrupted by a slower or irregular rhythm; if the rate can be halved by pressure on the neck; or if in the longer attacks the rate can be halved or made irregular with digitalis, the attack is flutter. Flutter, like fibrillation, is more often established than paroxysmal, but Parkinson and Bedford (1927) found that nearly one quarter of their cases of flutter had the paroxysmal form. Paroxysmal tachycardia very rarely lasts more than ten days and generally not more than four days: therefore the longer an attack of unknown origin lasts, the more likely it is to be flutter, and after ten days this becomes more certain. If the patient has high blood pressure or arteriosclerosis or if congestive failure develops rapidly, flutter must be thought of but cannot be diagnosed merely on these grounds. Occasionally even when a cardiogram is obtained there may still be difficulties, e.g. slow flutter without heart block, or a faster rate with 2:1 heart block where the P and T waves cannot be clearly distinguished.
Every effort should be made to obtain a cardiogram of an attack; not only to differentiate flutter, but to decide the variety of paroxysmal tachycardia, which cannot be determined in any other way and is of value in prognosis. It may still be difficult or impossible to subdivide the supraventricular into the various types of auricular and nodal, but fortunately this seems of less practical importance.

Of the 42 paroxysms recorded, 8 were ventricular. In three it was possible to identify P, occurring regularly at a slower rate than the ventricular complexes and quite unrelated to them, and so to prove that the paroxysms were ventricular; the auricular rates were 150, 130, and 100, and the ventricular 220, 272, and 166 respectively (Fig. 1). In the remaining five ventricular records it was not possible to identify P, or there was retrograde conduction from the ventricle (Fig. 2). The type shown in Fig. 2 of very short paroxysms, recurring frequently, may be found with a normal heart and seems to be of no serious significance even though it is ventricular.

If the QRS wave in the paroxysm was not widened or grossly aberrant, the focus was counted supraventricular. If QRS was widened and aberrant, it was compared with the QRS in the free intervals, and if different in conformation we considered the focus ventricular. Fig. 3 is an example we have called ventricular (though the P waves could not be seen) because the ventricular complexes of the attack differ from those of normal rhythm. This evidence is not conclusive, for the paroxysm may sometimes be auricular with aberrant
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conduction to the ventricle. If, on the other hand, QRS in the free interval was similar to that in the attack, we considered the focus as supraventricular. Such a paroxysm is shown in Fig. 4; it appears ventricular, but the QRS complexes in the free intervals and in the attacks are identical; further, ventricular extrasystoles during normal rhythm did not resemble the ventricular waves of the paroxysm.

Fig. 3.—(A) Normal rhythm, and, (B) two days later, ventricular tachycardia, rate 207; there is no clear evidence of auricular activity but a suggestion of possible P waves, low on each second ventricular downstroke in lead III (Case 5).

Fig. 4.—Supraventricular tachycardia, rate 175, simulating ventricular tachycardia because of the permanent defect of conduction; and normal rhythm; both from an elderly man with myocardial disease (Case 38).
Levine (1927–28) has drawn attention to the slight irregularity in the rate and sounds in some ventricular paroxysms which may suggest the diagnosis on clinical grounds alone. On measuring the cardiograms this irregularity was present in two of our cases, but too slight to be appreciable to the finger or ear. Any gross irregularity is likely to indicate paroxysmal flutter with changing degrees of block.

Paroxysms of nodal tachycardia can usually be differentiated, but in many supraventricular paroxysms the exact site of origin cannot be defined. An example of auricular paroxysmal tachycardia is shown in Fig. 5. On the picture alone it could not be distinguished from simple tachycardia; if the rate were a little faster P would be hidden in T, and one could not decide if it was an auricular attack with P hidden in T, or a nodal attack with P hidden in QRS. This seems to be a common difficulty when the beginning or ending of the attack is not seen (Figs. 9, 10).

Inversion of P indicates an abnormal focus in or near the A-V node, and its relation to QRS—whether before it with a short P-R interval, incorporated with it, or following it—indicates the level in the node of such focus. In two patients where the position and conformation of the P wave showed that the attack was nodal, it was inverted and preceded the QRS with a P-R interval that was normal (Fig. 6); this is sometimes described as superior or high nodal. In three more it was just before or with the QRS and was located by its large size (Fig. 7) or by recording a complete attack (Fig. 15a). This nodal group is relatively large in some published series and evidently includes many that we have left unclassified. The P wave was inverted and followed QRS in eight (Fig. 8), and these have been grouped as inferior or low nodal. In 13 of our 34 supraventricular cases there was no doubt of their nodal origin, two being high nodal, three nodal, and eight low nodal, and another three were
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Fig. 6.—Nodal tachycardia of the “high nodal” type, rate 169, with the rhythm constantly changing; from a boy whose heart was otherwise normal (Case 2).

Fig. 7.—Nodal tachycardia of “intermediate” type, rate 125; and normal rhythm; both from a woman with mitral stenosis (Case 87). The large P wave is incorporated with QRS on the downstroke of R.

probably nodal. This left 21 other supraventricular; three were auricular but in the others there were no clear indications of the type (Figs. 9, 10).

Ritchie (1926) reviewed 138 cases, adding 14 of his own, and found one-third were ventricular. Hume (1930) collected the classification from various series as follows: auricular 40; superior nodal 8; inferior nodal 18; nodal 100; unclassified supraventricular 32; ventricular 55. But this is merely a
FIG. 8.—Nodal tachycardia of “low nodal” type, lead II only: (A) from a woman with mitral stenosis, rate 171 (Case 52); (B) from a woman with rheumatic aortic incompetence, rate 234 (Case 70).

FIG. 9.—Supraventricular tachycardia, from two women with mitral stenosis: (A) rate 199 (Case 36); (B) rate 185 (Case 60).

FIG. 10.—Supraventricular tachycardia, rate 172: and normal rhythm; both from an elderly man whose heart was otherwise normal (Case 58).
summary of some very discordant series, and we think that many classed as
auricular or nodal should be left as unclassified supraventricular. The propor-
tion of ventricular attacks in these series of Hume and Ritchie is in our
opinion much higher than is an unselected series. Ventricular paroxysms
occur most often in patients admitted to hospital for heart disease or even for
failure, and so tend to be observed and reported; many of these patients
have been treated with digitalis and have had advanced heart disease; and the
paroxysms have occurred under these conditions on a few occasions, rather than
periodically over a long time as with ordinary paroxysmal tachycardia. As
instances, three of the five attacks reported by Gilchrist (1925–26) were almost
terminal, and of the 65 patients reported by Strauss (1930) more than half
were under treatment with digitalis for their heart failure and four-fifths had
died within three months.

<table>
<thead>
<tr>
<th>Type of Extrasystoles recorded in the free intervals</th>
</tr>
</thead>
<tbody>
<tr>
<td>Supraventricular only</td>
</tr>
<tr>
<td>-----------------------</td>
</tr>
<tr>
<td>Supraventricular (34 cases)</td>
</tr>
<tr>
<td>Ventricular (8 cases)</td>
</tr>
</tbody>
</table>

It is said that the type of extrasystole observed in the free intervals will
indicate the type of paroxysm and that extrasystoles are frequently observed
in most cases of paroxysmal tachycardia. Often, however, this is not so, and as an example, in one patient seen each few weeks for ten years no extrasystoles
have been noted. They were recorded in 16 of the 42 with records of their
paroxysms (see Table I). Where supraventricular extrasystoles were observed
(whether alone or with ventricular extrasystoles) the paroxysms were always
of the supraventricular type, though auricular or nodal extrasystoles did not
always indicate that the paroxysms were from the same focus. But where ventricular extrasystoles were observed the converse did not hold true: if
they were the only type observed the paroxysms might be ventricular or
supraventricular, though more likely to be the former; but if they were of
both types the paroxysms were always supraventricular.

ÆTIOLOGY

The first question to decide in each patient with paroxysmal tachycardia is
whether there is any organic heart disease. This is essential for prognosis,
and even slight signs may be very important. The ætiology of our hundred
cases is shown in Table II, and rather more than half had no evidence of heart
disease. The inclusion of those where there was no graphic record of the
attack does not invalidate these conclusions about the ætiology; the only
difference was that fewer records were obtained of the normals because they
had less reason for attending hospital regularly.

TABLE II
AETIOLOGY OF 100 CASES OF PAROXYSMAL TACHYCARDIA

<table>
<thead>
<tr>
<th>Aetiological Group</th>
<th>Number of Cases</th>
<th>Number with Cardiogram during Attack</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Males</td>
<td>Females</td>
</tr>
<tr>
<td>Normal</td>
<td>34</td>
<td>21</td>
</tr>
<tr>
<td>Rheumatic</td>
<td>4</td>
<td>15</td>
</tr>
<tr>
<td>Syphilitic</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Hyperpietic</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>Thyro-toxic</td>
<td>0</td>
<td>4</td>
</tr>
<tr>
<td>Myocardial</td>
<td>9</td>
<td>3</td>
</tr>
<tr>
<td>Total</td>
<td>52</td>
<td>48</td>
</tr>
</tbody>
</table>

Men and women were equally affected, as the excess of women in the
rheumatic group was balanced by the men in the myocardial and normal
groups. The age incidence—the age at which the attacks started—is shown
in Table III and Figure 11. In 11 per cent. they started before the patient was

![Figure 11](http://heart.bmj.com/)

**Fig. 11.**—Diagram showing the age incidence at onset of paroxysmal tachycardia.

Above: 1. Hyperpietic and myocardial groups combined; the hyperpietic are shaded
and are mostly between 35 and 65, the myocardial are more evenly spread at all ages.
Below: 2. Normal group, at all ages with the main incidence between 15 and 40.
3. Rheumatic group, with the main incidence between 20 and 45 years of age.
fifteen, in 65 per cent. between fifteen and thirty-nine, and in another 12 per cent. before fifty, making 88 per cent. where the attacks started before this age.

**TABLE III**

**AGE INCIDENCE AT ONSET OF PAROXYSMS**

<table>
<thead>
<tr>
<th>Age—in years</th>
<th>Number of Cases in each Group</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Normal</td>
<td>Rheumatic</td>
</tr>
<tr>
<td>0-4</td>
<td>2</td>
<td>—</td>
</tr>
<tr>
<td>5-9</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>10-14</td>
<td>3</td>
<td>—</td>
</tr>
<tr>
<td>15-19</td>
<td>11</td>
<td>—</td>
</tr>
<tr>
<td>20-24</td>
<td>10</td>
<td>6</td>
</tr>
<tr>
<td>25-29</td>
<td>7</td>
<td>4</td>
</tr>
<tr>
<td>30-34</td>
<td>7</td>
<td>3</td>
</tr>
<tr>
<td>35-39</td>
<td>7</td>
<td>2</td>
</tr>
<tr>
<td>40-44</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>45-49</td>
<td>3</td>
<td>—</td>
</tr>
<tr>
<td>50-54</td>
<td>2</td>
<td>—</td>
</tr>
<tr>
<td>55-59</td>
<td>1</td>
<td>—</td>
</tr>
<tr>
<td>60-</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>All ages</td>
<td>59*</td>
<td>19</td>
</tr>
</tbody>
</table>

* Includes 4 with goitre.

Most were hospital patients, and in those seen in practice since completing this series there have been fewer in the rheumatic group and more elderly patients with myocardial and coronary disease. They have been classified in the following six groups.

1. **Rheumatic Group**: 19 cases.—In all but one of these the attacks started between 20 and 44 years of age. There was valvular disease in 17, 10 with mitral, 5 with mitral and aortic, and 2 with aortic lesions; the other 2 had a rheumatic history only.

2. **Syphilitic Group**: 2 cases.—Both had aortic incompetence; and the paroxysms did not seem to influence the natural course of their disease. As there were no special features they have been included in the myocardial group for the tables, etc.

3. **Hyperpietic Group**: 8 cases.—Five of these, with an average age of 62 and an average blood pressure 180/105, may be taken as typical of the group. Their attacks were of relatively recent onset. Two elderly men were included, but probably the raised pressure was incidental, as their attacks had been present for 15 and 20 years; two were not included, as paroxysms had occurred for 40 years. The eighth was unusual, as her attacks and high blood pressure both started after the removal of cystic ovaries. A less strict standard might have added one or two to this group, but among the normals there were very few who were even on the borderline.

4. **Myocardial Group**: 12 cases.—In many ways this was the most important group, though rather mixed. It comprised all those with heart disease who
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could not be included elsewhere. Generally the signs were unmistakable, but occasionally even in young people the course of events showed serious heart disease with the paroxysms as the only early symptom, with a slight cardiac enlargement and/or an abnormal cardiogram as the only sign. They are discussed in more detail in the section on prognosis.

5. Thyrotoxic Group: 4 cases.—All were women with a goitre, but none had clinical hyperthyroidism. All have done well, and as they had no heart disease are included for subsequent discussion with the normal group. It is surprising that toxic goitre should be so much more frequently the cause of paroxysmal fibrillation (15 per cent.; Parkinson and Campbell, 1930) than of paroxysmal tachycardia.

6. Normal Group: 55 cases.—The presence of paroxysms is not evidence that there is organic heart disease, and this group includes all those with hearts that were thought to be normal except for the paroxysms. Examples are given in the section on prognosis (see p. 149) of paroxysms for twenty or even sixty years without demonstrable heart disease at the end. Possibly a few have been wrongly included, but it seems certain that this group is much the largest. Details of the age when the attacks started are given in Table III. Naturally in this group the patients were younger, and nineteen were under 20, fifteen from 20–29, and twelve from 30–39, leaving only seven over this age and only three over 50; 72 per cent. were between 15 and 39 years of age at the onset of their attacks.

The aetiology of the 42 with records of the paroxysms are shown in Table IV. The only points of interest are the rarity of ventricular attacks in the normal and rheumatic groups, and that just over half of the nodal attacks were found in the rheumatic group.

**TABLE IV**

**TYPE OF PAROXYSMS IN DIFFERENT AETIOLOGICAL GROUPS**

<table>
<thead>
<tr>
<th>Type of Paroxysms</th>
<th>Aetiological Group</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Normal</td>
<td>Rheumatic</td>
</tr>
<tr>
<td>Supraventricular</td>
<td>16</td>
<td>13*</td>
</tr>
<tr>
<td>Ventricular</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Number with cardiograms</td>
<td>19</td>
<td>13</td>
</tr>
<tr>
<td>Total number with paroxysms</td>
<td>59</td>
<td>19</td>
</tr>
</tbody>
</table>

* Of these 13, 7 were nodal.

**CAUSATION OF PAROXYSMS**

Although it is important from the practical point of view to decide whether there is organic heart disease or not, this does not explain the initiation of a
paroxysm. There must be some instability of the normal pacemaker, either absolute or relative to other parts of the heart muscle, and this may sometimes be shown by the occurrence of other abnormal rhythms. There was a respiratory sinus arrhythmia of sufficient magnitude to be unusual in 7; and extrasystoles were observed in 26, indicating that parts of the heart other than the S-A node were over-ready to initiate impulses. Two others showed nodal rhythm and sino-aurious block.

Peculiar changes were seen in the cardiogram of one patient. When lying on her right side there was a short P-R interval of 0.08 sec., a large wide R, and an S-T interval below the iso-electric level (Fig. 12). On her left side

![Fig. 12.—Changes which could generally be induced in one patient: (A) lying on the right side; (B) lying on the left side (Case 30).](image)

lead III changed, the P-R interval 0.14 sec., a smaller R, and larger S with upright T waves; the QRS waves varied with respiration more in this position than on her right side. These two types were obtained frequently, and sometimes occurred apart from changes of position, suddenly from beat to beat (Fig. 13). On one occasion the type with a large wide R was obtained by holding her

![Fig. 13.—Spontaneous changes, illustrating the short P-R interval with complexes of bundle branch block type (Case 30).](image)
breath after a deep inspiration, the S type being present at the beginning and again after expiration.

Wolff, Parkinson, and White (1930) published 11 cases prone to paroxysms with somewhat similar changes, for in the free intervals there was a short P-R interval and a bundle branch lesion, both being abolished by exercise. Evidently the S-A node is in some cases unduly over susceptible to outside influences, such as respiration or changes of position; and in others the relative excitability of the S-A node and the extranodal tissues is altered, extrasystoles resulting.

In addition to this underlying instability, some exciting factor must initiate the paroxysms from an ectopic focus. Many attacks, as is well known, start for no apparent reason and without any warning, but sometimes the patient knows the usual exciting factor, and more rarely may be able to initiate an attack voluntarily.

The beginning or end of a paroxysm are not often recorded except in those studied intensively in hospital, but we were fortunate in recording them in six patients. An unusual type where the onset and offset of short attacks were constantly obtained in many records is shown in Fig. 6; at one time it was hard to obtain a record without these changes, but now seven years later they are less frequent. The end of two attacks and another beginning soon after are shown in Fig. 14; deep inspiration seemed to cause the short returns of normal rhythm. Generally her attacks lasted for many hours, but that morning her condition was specially unstable. Short paroxysms from two other patients are shown in Fig. 15; the first, probably nodal, occurred spontaneously; the second could be produced almost regularly by deep inspiration. Another patient had her first attack when her doctor, examining her lungs after pneumonia, asked her to take a deep breath. The beginning of a paroxysm and a

Fig. 14.—Onset and offset of paroxysms, rates 180 and 190, induced by deep breathing; lead II only (Case 82).
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Fig. 15.—(A) Short nodal paroxysm, rate 174, occurring spontaneously in a woman with mitral stenosis (Case 46); (B) Short paroxysm, rate 158, induced by deep inspiration in a woman whose heart was normal (Lead I only).

complete short one are shown in Fig. 16; here again the short attack was exceptional, and there was no obvious exciting cause.

Fig. 16.—Onset of paroxysm, rate 171, in lead I, and complete short paroxysm in lead III; occurring spontaneously in a woman with rheumatic heart disease (Case 60).

A paroxysm interrupting auricular fibrillation is shown in Fig. 17. This was the only time we were able to induce an attack by means of the stimulus

Fig. 17.—Paroxysm at a rate of 141, induced by a loud noise; in a man with established auricular fibrillation and myocardial disease (Case 56). In a long paper tracing the third beat of the paroxysm (shown above the word ‘noise’) was the only one that differed from the others.
which the patient thought was generally responsible for his attacks—in this instance the unexpected noise of dropping tins on a concrete floor. Another patient attributed several attacks to sudden unexpected noises, such as banging a door.

Sudden movements, such as stooping, turning on one side, jumping, or running upstairs, often start an attack, and where there is an ascertainable cause it is most often of this type. Such movements nearly always provided the stimulus in nine of our cases, for the most part with normal hearts; but similar movements stopped attacks in six others, and this may be useful in advising treatment. One patient was particularly liable to an attack when playing tennis, especially at the moment he raised his arm to serve; he was able to play if he took quinidine before a game. Another had his first attack when he leant out of bed to reach a urinal, and his most recent when he dropped his newspaper in the train and reached down to pick it up.

Such stories are far too common to be due to chance, though often no cause can be detected. Apart from these movements there is a somewhat heterogeneous group of conditions which seem to increase the instability of the S-A node so that any minor cause may excite an attack. Such conditions are anxiety, overwork, over-exertion, pregnancy, obesity, indigestion, smoking, and infection, and these will be considered shortly.

Anxiety.—The patient’s state of mind is important. Emotional upsets, excitement, worry due to financial or domestic causes, or anxiety about the supposed gravity of the cardiac state may induce more frequent attacks; in one the first and only attack occurred whilst he was engrossed in a game of chess, and this often happened in another patient. In a bank clerk the heavy work during the period of the War Loan conversion changed infrequent paroxysms into daily events. In many patients the attacks became less frequent and less important when they knew that the heart was otherwise normal. We are speaking of the effect of anxiety on normal people; there is no special association with pathological anxiety states.

Infection.—There is little evidence that infection, acute or chronic, plays any important part. In two patients the first attacks started during convalescence from pneumonia and continued for many years. This might suggest a focus of myocardial damage, but there were no other signs of heart disease, and no such history of infections could be obtained in others. First attacks followed influenza, acute frontal sinusitis, rheumatic fever, tonsillectomy, etc., but six such cases in a hundred is not very convincing.

Chronic focal infections can be accepted as the cause of paroxysms only if their effective treatment produces an improvement. In this series results have been disappointing and no case has shown much improvement. At present there seems no real evidence for an infective group of cases. On general principles focal sepsis should be treated when it is present, but one should never assume that this will cure paroxysmal tachycardia.

Smoking.—There was some improvement in three cases when cigarettes were reduced. Reduction of smoking even down to complete abstinence failed to influence the frequency or severity of attacks in others. There are
people who are unable to smoke because of the frequent extrasystoles induced, and paroxysms of tachycardia occur in some of them; but tobacco is not often the cause of paroxysms though over-indulgence may increase their frequency.

Exercise.—In six patients attacks were increased by exertion greater than that to which they were accustomed, e.g. working overtime, cross country running, and football when out of training; and reduction of exercise led to improvement. In two others the first attacks had followed on exercise, in one a game of hockey and in the other long distance running, but they continued without such causes. In two exertion had the opposite effect; one of them could generally stop an attack by walking or running; the other, who was leading a sedentary life, was improved by daily exercise. Most of these had no heart disease, but especially in those with rheumatic hearts more frequent paroxysms were often due to their doing more than was advisable for the condition of their hearts.

Indigestion.—No organic disease of the stomach or intestine was found in any of these patients. In two, attacks usually occurred at a fixed time after a meal, and in three a period of indigestion led to an increase in attacks. In two others their frequency was greatly diminished by treatment of their indigestion, but two doctors were disappointed that effective treatment of their indigestion failed to reduce the attacks. Sometimes the disturbance of digestive function produced by the paroxysm is mistaken for its cause. One patient attributed his unusually long attack to indigestion, though he did not usually suffer in this way with shorter ones; there was, however, congestion of the liver owing to a seven-day attack at a rapid rate. There were severe gastric symptoms, nausea, retching, and vomiting, in eight others. Spontaneous vomiting heralded the termination in three, and induced vomiting was the best method of ending an attack in others. Such symptoms were only observed in a minority, and flatulent dyspepsia is only rarely the cause. Nevertheless enquiries should always be made along these lines, and may reveal an effective method of treatment.

Allergic Conditions.—We have found no association between paroxysmal tachycardia and asthma, urticaria, migraine, or any conditions which are thought to be associated with allergy.

Pregnancy.—Five of these patients and several we have seen since were unfavourably influenced by pregnancy. In three the attacks were more frequent, in two they started at that time. On the other hand, two improved, one for the time and the other after as well as during her pregnancy. Anderson (1932) also has reported a case where paroxysms stopped during pregnancy and (1933) another where they occurred only at this time. In no case in our series except the one described (see p. 156) was there any reason to suppose that pregnancy should have been terminated on account of paroxysms. Meyer, Lackner, and Schochet (1930) reporting two cases stated they had only been able to find four or five examples of paroxysmal tachycardia in pregnancy; we think they are by no means uncommon and are often mistaken for simple palpitation.
Obesity.—This is mentioned because in two cases, both comparatively young but over-stout, reduction in weight was followed by improvement.

Intrathoracic Disease.—This may cause paroxysms, and two patients, seen since this series, had their first attacks as an early symptom of a mediastinal neoplasm. In two others they occurred during the course of pulmonary tuberculosis. Graber (1925–26) has reported a case where lymphadenomatous glands were pressing on the vagus and we have recently seen an attack in a young man with a Riedel’s goitre where the subsequent attacks of pain showed that there was pressure on the vagus. Paroxysmal auricular fibrillation is a more common sequel of thoracic neoplasms, especially when there is a pleural effusion.

SYMPTOMATOLOGY OF PAROXYSMS

Besides the sensation of palpitation complaint may be made of a variety of symptoms, any of which may be more or less prominent and exceptionally may mask the true nature of the paroxysm, especially in children. Rarely the patient may be unaware he is experiencing an attack, when, of course, the diagnosis depends on chance observation. The general symptoms, palpitation, weakness, faintness, dizziness, anxiety, or breathlessness, will not be considered in detail as they are of little special significance in themselves. Any degree of incapacity up to complete prostration may be met with, and may be attributed to any of these sensations. On the whole our experience corresponds with that of other observers—that the incapacity depends mainly on the state of the heart muscle, on the length and to a less extent on the rate of the attack, and on the mentality of the patient.

If the heart is healthy the symptoms are generally discomfort rather than real disability. One man thought it was sufficient if he came out of the scrum and took an easier position in the field when an attack started during a game of Rugby football. Another was able to carry on his routine work at the hospital and lecture, though he certainly found it more of an effort. At the other extreme some patients developed congestive failure. Many of those with rheumatic or other heart disease had little more disability than most of the normals, yet with severe heart disease any attack may be very grave. The importance of the duration and of the rate is discussed later. Shortly one can say that if an attack lasts for one or two days the symptoms and disability generally become important, even if the heart is healthy; and that a healthy heart will support short attacks even at such a rapid rate as 200 surprisingly well.

Loss of Consciousness.—Unconsciousness during the attack is very rare. Two patients, both with heart disease, said that occasionally they had lost consciousness just after the onset. As the hundred cases represent several thousands of attacks, the rarity of loss of consciousness is obvious.

Anginal Pain.—In addition to discomfort or tightness in the chest, present in most attacks and of no special significance, twelve patients had more severe pain of an anginal type during attacks, some occasionally, others when the
the attack had lasted a definite number of hours. In ten it was felt beneath the sternum or just to one side and in two under the left breast; sometimes it spread down the left arm (five times), down both arms (once), or up into the angle of the jaw (twice). Only in two was the pain really severe, and one of these had angina on exertion apart from his paroxysms. In four where nitrates were used the pain was eased. Seven of these twelve had hearts which appeared normal except for their attacks, and two had mitral stenosis only. The presence of pain, even of a severe type simulating angina, does not indicate an unfavourable prognosis; this must be judged mainly by the condition of the heart during the free periods. Only one of these twelve has died, six years after the onset of the paroxysms and angina.

White and Camp (1931–32) have drawn attention to the occasional occurrence of anginal pain in paroxysmal tachycardia or fibrillation and the need for distinguishing it from angina of effort, but they do not stress how benign it may be under these conditions. The relationship of paroxysmal tachycardia to coronary thrombosis is discussed later (see p. 148).

**Congestive Failure.**—This was observed during the attack in nine patients. One, who so far as we could judge had no other evidence of heart disease, had experienced several attacks without failure, but was seen on the eighth day of a ten-day attack with commencing failure, when the rate was 196 and the blood pressure had fallen from 140 to 105. Two children with hearts that were otherwise normal developed failure in attacks lasting two or three days, but the rates were unusually fast. The remaining six had diseased hearts—four myocardial, one rheumatic, and one hyperpietic. In seven the failure became apparent on about the third day, in one with rheumatic heart disease within twenty-four hours. Not many attacks lasting three days were observed in normals, as they were less likely to seek medical advice, but the figures quoted indicate that the development of congestive failure is dependent as a rule on heart disease. Its occurrence does not seem so serious as might be expected, especially where the heart is normal or even well compensated in the free intervals.

**The Rate in Paroxysms**

The rate of one or more attacks was observed in 70 cases. It is stated by Lewis (1920) to lie between 160 and 200 usually, with a range from 110 to 220. These upper limits are not high enough, and it is not very rare for the rate to be up to 240 or even faster. In our cases the rate was between 160 and 200 in 45 per cent. and between 140 and 240 in 90 per cent.

In three the rate was less than 140, viz. 136, 126, and 124. The last, who had frequent nodal attacks, is illustrated in Fig. 7. At such a slow rate the diagnosis may be difficult, and here it was not made until she was seen in an attack.

In two children the rate was over 250 (272 and 300, see Fig. 1). They were exceptional in many ways and have been described elsewhere (Campbell, 1937); the second may have been paroxysmal flutter. In four others (all supra-ventricular), the recorded rates were 248, 230, 240, and 234 (see Figs. 8b
and 18). Two had no evidence of heart disease and had already had attacks for fifteen and seventeen years. The other two had rheumatic lesions.

### TABLE V

**Heart Rate in Paroxysms**

<table>
<thead>
<tr>
<th>Type of Paroxysmal Tachycardia</th>
<th>Number of cases recorded at each Rate</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nodal</td>
<td>2</td>
<td>5</td>
</tr>
<tr>
<td>Other Supra-ventricular</td>
<td>1</td>
<td>6</td>
</tr>
<tr>
<td>Ventricular</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Paroxysms without a cardio-gram</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>Total</td>
<td>3</td>
<td>15</td>
</tr>
</tbody>
</table>

Table V summarizes the findings and shows how the rates were distributed among the different types of paroxysms. There were cardiograms of 55 attacks from 42 patients (two from the same patient being counted because the rate was sometimes different), and of 30 others observed clinically. Slow and fast rates were found in each type. Analysis in various ways shows few conclusions except that nodal attacks tend to be a little slower and ventricular attacks a little faster; and that sometimes it is a bad sign when the rate of the attacks in any patient falls during the course of months or years.

In the nodal attacks the average rate was 170 and the commonest range was 140 to 180; one fallacy may be that at the slower rates it is easier to distinguish the P wave and so to diagnose nodal tachycardia. In the other supra-ventricular attacks the average rate was 187 and the usual range was 150–230 with more below 190. In the ventricular the average was 198, the usual range was the same, but more were above 190.

The presence or absence of disease did not appear to influence the rate, the average for the normal cases being 190, for the rheumatic 178, for the myocardial 180, and for the hyperpietic 185. The range in all the larger groups was much the same and included fast and slow attacks.

The rate in different attacks is said to keep about constant. This is our general experience and cardiograms obtained in different attacks from several cases were practically identical after an interval of years. In many patients, however, they varied by as much as 10 to 15 per cent. In short attacks the rate may vary considerably within a few minutes. Here it may represent a different ectopic focus.

Where rates were accurately observed in several paroxysms one third showed a difference of between twelve and thirty beats per minute; most of these had heart disease. The period over which paroxysms have occurred did not in itself cause a change in rate. The changes were not more common
in any one type of attack or in different diseases of the heart. There was some evidence that as the efficiency of a diseased heart lessens, the rate of the paroxysms may fall; but others observed up to the time of death maintained a constant rate to the end. For example, in two a fall of 12 and 30 beats per minute preceded the onset of fibrillation, and in two others the rate slowed as they became worse—26 and 22 beats per minute. An increase in rate as the myocardium fails is less common.

Comparing the rates recorded graphically with those counted personally, there is no reason to think that the "clinical" rates are not accurate, except that there were fewer under 160, perhaps because paroxysms at a slow rate are less likely to be diagnosed without a graphic record. The rate should always be taken for half a minute. Generally when the rate is said to be uncountable it is because an attempt has been made to count the pulse by the finger instead of the heart by the ear.

**THE LENGTH OF ATTACKS**

Paroxysms last for hours much more commonly than for days. Though variable, they tend to have a more or less customary length in any one subject, who can generally report that the attacks last between, say, two and eight hours, or less than half an hour, or more than a day, and is surprised if his past experience does not prove a useful guide of what to expect. In two patients, for example, the length of many of their attacks were recorded over a series of years. In one (Case 30) there were 122 paroxysms in eleven years (1927-38) and the length of 89 of these known; the number of paroxysms of different lengths was as follows:

<table>
<thead>
<tr>
<th>Length</th>
<th>Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-4 hours</td>
<td>8</td>
</tr>
<tr>
<td>5-8</td>
<td>19</td>
</tr>
<tr>
<td>9-12</td>
<td>24</td>
</tr>
<tr>
<td>13-16</td>
<td>14</td>
</tr>
<tr>
<td>17-20</td>
<td>5</td>
</tr>
<tr>
<td>21-24</td>
<td>10</td>
</tr>
<tr>
<td>25-48</td>
<td>7</td>
</tr>
<tr>
<td>17 and 29 days</td>
<td>2</td>
</tr>
</tbody>
</table>

In the other (Case 70) the length of 88 paroxysms in the years 1924-32 was recorded as follows:

<table>
<thead>
<tr>
<th>Length</th>
<th>Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Less than ½ hour</td>
<td>55</td>
</tr>
<tr>
<td>½-1½ hours</td>
<td>15</td>
</tr>
<tr>
<td>2-3</td>
<td>9</td>
</tr>
<tr>
<td>4-5</td>
<td>8</td>
</tr>
<tr>
<td>8</td>
<td>1</td>
</tr>
</tbody>
</table>

In these two, though there is variation round the customary length in each case, it is easy to give an average range and to say that what would have been a long attack for one would have been a short attack for the other.

More than half of these patients generally had paroxysms of less than two
hours, and in seven only did they habitually last for a day or more, even though those with longer attacks are more likely to come under observation. Table VI shows their usual duration in this series, though often it was difficult to decide such precise limits. The longer attacks occurred a little more often in those with heart disease, but the difference was not great.

TABLE VI

<table>
<thead>
<tr>
<th>The Customary Length of Paroxysms</th>
<th>Patients with no Heart Disease except the Paroxysms</th>
<th>Patients with Heart Disease</th>
<th>Total Number of Patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Up to 30 minutes</td>
<td>23</td>
<td>12</td>
<td>35</td>
</tr>
<tr>
<td>, 2 hours</td>
<td>15</td>
<td>11</td>
<td>26</td>
</tr>
<tr>
<td>, 6 hours</td>
<td>12</td>
<td>6</td>
<td>18</td>
</tr>
<tr>
<td>, 12 hours</td>
<td>5</td>
<td>5</td>
<td>10</td>
</tr>
<tr>
<td>, 24 hours</td>
<td>1</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>Over 24 hours</td>
<td>3</td>
<td>4</td>
<td>7</td>
</tr>
<tr>
<td>Total</td>
<td>59</td>
<td>41</td>
<td>100</td>
</tr>
</tbody>
</table>

The longer paroxysms are of more importance, for the distress and disability and even the likelihood of congestive failure are increased. In addition to the 7 that habitually had attacks lasting more than twenty-four hours, 22 others had some of this length, making a total of 29 with some or all of their attacks lasting more than a day. The attacks were not often this length and never greatly exceeded it in ten of these, six with heart disease and four without, and did not produce serious symptoms, or any change in the general condition.

The remaining 19 with long attacks are illustrated in Table VII. There were six (including two small children) with hearts that were normal except for the paroxysms and four with rheumatic heart disease, most of the remainder falling into the myocardial group. The long attacks were least frequent in those without other evidence of heart disease—11 per cent. of the normal, 21 per cent. of the rheumatic, and 33 per cent. of the other groups having these long attacks.

The rate in these longer attacks was generally faster than the average for all the paroxysms; in only three was it under 180 and in ten of the nineteen it was 200 or over, while among the other cases there were more under 180 than at 200 or over. The average rate in these long attacks was 208 (or 198 if the two exceptionally fast ones be excluded), instead of 186 for the series as a whole. All types of attacks were represented, but there was a much higher proportion of ventricular paroxysms than of the other types, nearly half the former lasting for more than twenty-four hours.

Even among these longer attacks relatively few lasted for more than two or three days, and they were of this length in eight patients. In two they lasted 4 days; in two up to 7 days; and in five, 10 days; only in two did they last longer than 10 days. One of these (Case 15) is described in the
### TABLE VII

**Paroxysms of Unusual Length**

<table>
<thead>
<tr>
<th>Case Number</th>
<th>Age (years)</th>
<th>Etiological Group</th>
<th>Paroxysms</th>
<th>Length of Paroxysms</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Rate (beats/min)</td>
<td>Type*</td>
<td>Usual—Less Usual in hours</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>21</td>
<td>Myocardial</td>
<td>210 V.</td>
<td>Three attacks only of 5 to 10 days.</td>
</tr>
<tr>
<td>15</td>
<td>24</td>
<td>Myocardial</td>
<td>190 —</td>
<td>8-48 7-8 days, three 14 and 17 days and 4 weeks.</td>
</tr>
<tr>
<td>69</td>
<td>1</td>
<td>Normal</td>
<td>272 V.</td>
<td>48 Several days.</td>
</tr>
<tr>
<td>78</td>
<td>1</td>
<td>Normal</td>
<td>300 S.</td>
<td>Two attacks only of 3 days.</td>
</tr>
<tr>
<td>84</td>
<td>27</td>
<td>Rheumatic</td>
<td>175 N.</td>
<td>6-48 Up to 7 days.</td>
</tr>
<tr>
<td>85</td>
<td>20</td>
<td>Rheumatic</td>
<td>200 S.</td>
<td>12-36 Up to 4 days.</td>
</tr>
<tr>
<td>88</td>
<td>40</td>
<td>Normal</td>
<td>— —</td>
<td>10-30 Up to 3 days.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>30</td>
<td>Myocardial</td>
<td>205 V.</td>
<td>Several of 10 days.</td>
</tr>
<tr>
<td>16</td>
<td>53</td>
<td>Myocardial</td>
<td>160 —</td>
<td>12-24 One of 7 days.</td>
</tr>
<tr>
<td>30</td>
<td>17</td>
<td>Normal</td>
<td>220 S.</td>
<td>10-18 Two 29 and 17 days; several 36-50 hours.</td>
</tr>
<tr>
<td>36</td>
<td>44</td>
<td>Rheumatic</td>
<td>240 S.</td>
<td>1-1 One of 2 days.</td>
</tr>
<tr>
<td>47</td>
<td>48</td>
<td>Syphilitic</td>
<td>— —</td>
<td>2 One of 2 days.</td>
</tr>
<tr>
<td>43</td>
<td>48</td>
<td>Normal</td>
<td>190 S.</td>
<td>1-6 One of 10 days; several 1-2 days.</td>
</tr>
<tr>
<td>55</td>
<td>41</td>
<td>Myocardial</td>
<td>200 —</td>
<td>4-12 Several of 3 days.</td>
</tr>
<tr>
<td>61</td>
<td>50</td>
<td>Hyperpietic to</td>
<td>172 —</td>
<td>1-48 Several over 2 days.</td>
</tr>
<tr>
<td>68</td>
<td>25</td>
<td>Myocardial</td>
<td>200 —</td>
<td>Variable Four attacks 7-10 days.</td>
</tr>
<tr>
<td>75</td>
<td>20</td>
<td>Normal</td>
<td>180 —</td>
<td>1-3 Sometimes up to 3 days.</td>
</tr>
<tr>
<td>80</td>
<td>41</td>
<td>Rheumatic</td>
<td>217 N.</td>
<td>1-1 Sometimes up to 4 days.</td>
</tr>
<tr>
<td>82</td>
<td>12</td>
<td>Normal</td>
<td>197 S.</td>
<td>2-12 Some up to 7 and 10 days.</td>
</tr>
</tbody>
</table>

* N, nodal; S, supraventricular; V, ventricular paroxysms.

section on prognosis; an attack in which congestive failure developed lasted 17 days. There had been shorter ones for sixteen years, but longer attacks recurred and he died five years later in one that lasted more than 28 days. The other, aged 38, had suffered from attacks for twenty years and during five years’ observation had about ten a year lasting generally for 10–18 hours and never more than 48 hours. She was admitted to another hospital when one attack had lasted 24 days and the heart rate remained over 200 in spite of treatment with digitalis. As it still continued after 29 days she was transferred to Guy’s Hospital, but it stopped during her journey. She soon felt as well as usual, though for a few days the T waves were rather flatter. The frequency and duration of her paroxysms were not permanently increased (see Table IX), and in other respects her heart could be passed as normal.

After another seven years she had a second long attack of 17 days treated with digitalis in her local hospital. A cardiogram of a subsequent attack showed no evidence of progressive myocardial disease as it was almost identical with the first of 1927 (Fig. 18), except for some left axis deviation.

These attacks of exceptional length are very unusual and naturally more serious, but it is possible for them to occur without there being any disease of
the heart and for complete recovery to take place. The great majority of paroxysms last for hours rather than for days, but 30 per cent. of our patients had one or more attacks lasting for a day, 10 per cent. of these very rarely, 10 per cent. occasionally, and 10 per cent. habitually. Only two had attacks lasting longer than ten days, though Gallavardin (1930) in a paper on paroxysmal tachycardia of long duration found that 11 out of 160 cases had some attacks as long as this.

**TREATMENT**

The treatment of the patient during a paroxysm and his management during the free intervals must be distinguished.

During an attack rest should generally be advised, but rest in bed is not needed unless special symptoms or the condition of the heart at other times make it advisable. Pressure on the vagus or on the carotid sinus is often effective, and also many tricks of sudden movement (e.g. bending down to open a low drawer); perhaps a forced expiration with a closed glottis after a deep inspiration is the most generally successful. If an attack persists towards evening, bed with a sedative or hypnotic will generally lead to its arrest.

Quinidine, 5 grains, repeated if necessary in two hours, is probably the most useful medicinal treatment. The cessation of attacks after digitalis has often been reported, but it is doubtful if it is effective, and generally it should not be given unless indicated by the state of the heart apart from the paroxysms or by the development of failure in a long attack. If there is any likelihood that the attack may be paroxysmal auricular flutter, full digitalization is indicated, both from the point of view of diagnosis and treatment.
Of newer remedies meckolin, 15 mg. intramuscularly, has often been successful in reported cases, but has been unpleasant and not very successful in our experience. So far prostigmin, 0.5 to 1.0 mg. subcutaneously, has seemed to us more effective and harmless to use; but it is early to decide its special indications and limitations. Morphia is not desirable in ordinary recurrent attacks, but may be used and may be effective if an attack is causing serious anxiety to the medical attendant.

After the paroxysm the most important question is to decide the condition of the heart. If thorough examination justifies it, reassurance about the nature of the attacks is often all that is called for, and this alone will sometimes greatly diminish their incidence. The general regulation of the patients' life as regards food, work, and exercise must next be attended to, and in many cases a sedative such as bromides will be all else that is needed. If in spite of this the attacks are frequent and troublesome regular administration of quinidine for a time, 3 to 5 grains, t.i.d., will often succeed. Regular digitalization is rarely indicated and has not been useful in our experience. Several cases where these methods have been successful and some where they have not are given in this paper, especially in the section on course and prognosis.

**COURSE AND PROGNOSIS**

*The Immediate Outlook at the time of a Paroxysm.*

This is excellent, and however long a paroxysm has lasted, whatever the rate, and however ill the patient may seem, he may be expected to recover. The only exceptions are if the paroxysms are of the rarer ventricular type, or occur after the heart has begun to fail, or in the course of coronary thrombosis; even so, though the risk is greater, he will generally recover from the immediate attack.

It is often stated that a patient may die in a paroxysm; this is rare, and it creates just the impression it is necessary to avoid in the large majority, where there is no danger. Only five in this series of a hundred have died suddenly or during a paroxysm, including one from an overdose of morphia and one after a miscarriage; and four of them had been liable to paroxysms for long periods (two for over 20 years and two for nearly 10 years) and had had hundreds of attacks before their deaths. Only one died soon after the onset of his paroxysms. Two at least, including the last, had ventricular paroxysmal tachycardia, and almost certainly died with ventricular fibrillation (Cases 5 and 79). A boy of seventeen under the care of Dr. Parkinson, who had suffered for five years from long and frequent ventricular paroxysms with gross congestive failure, provides more definite evidence. After an ordinary day with a short attack in the morning, he suddenly said that his heart had stopped and fell down dead, apparently after only a few breaths. From his words he must have known this was something quite different from the usual attacks and it was probably ventricular fibrillation. Because of this possibility the outlook in ventricular paroxysms is more serious, and this is one reason for trying to obtain an electrocardiogram of the paroxysm; fortunately they are
much less common. With supraventricular paroxysms the outlook is excellent, apart from other unfavourable findings in the heart.

It is equally important when the attacks are only of recent onset to decide if they may indicate progressive heart disease. The age of the patient is some help, and of those whose first attack came after 40, only 25 per cent. had normal hearts; while of those under 40, more than 60 per cent. had normal hearts. The following is an instance where paroxysms were the first symptom of commencing disease of the coronary artery. A man of 54 had a few paroxysms lasting two hours; he had no pain and no cardiac symptoms previously. After a year he began to get angina of effort, and during the next two years this was induced more easily by diminishing amounts of exertion. There have been no further paroxysms. This is an important and not uncommon variety; if paroxysms are of recent onset they are much more likely to be associated with organic heart disease in the elderly than in the young.

When a paroxysm occurs after coronary occlusion the immediate prognosis is not as bad as might appear, though it must add to the gravity. It generally stops spontaneously or after treatment, and the outlook depends more on the extent of the infarct. By chance none of these cases were examples of paroxysmal tachycardia secondary to coronary thrombosis, but one seen since illustrates the association. A man of 52 had a severe cardiac infarct, with pain for several hours in spite of morphia. His progress was satisfactory for ten days, until ventricular paroxysmal tachycardia started at a rate of 148. Quinidine, 30 grains daily, failed to arrest it, and he died after it had persisted for sixteen days. If, therefore, the patient is seen for the first time during a paroxysm, it is wiser not to make a final diagnosis of the heart condition at once. Murmurs may be difficult to hear, and if it has lasted long the blood pressure may have fallen and the heart may be dilated. The cardiogram may be abnormal for a few days after a long attack, so the complete examination should take place later than this. If at this stage the condition of the heart does not warrant a bad prognosis, the paroxysms may be looked on as an inconvenience rather than a serious matter.

The Ultimate Prognosis

This depends mainly on the condition of the heart in the free intervals, and to a lesser extent on the future frequency of the attacks and the effect, if any, they may have on the heart. Over 80 per cent. of these patients have already had attacks for more than five years and 47 per cent. for more than ten years, so that there has been ample opportunity to observe any unfavourable changes. The period which has elapsed since the onset of their paroxysms is shown in Table VIII. In the second and third columns they are divided into those who were alive when last heard of and those who have died, most of the deaths being due to the general course of their heart disease. In the last three columns they have been divided into three groups, those with no heart disease (including four with goitre), those with rheumatic heart disease, and those with hyperpyetic
or myocardial disease; as would be expected, the results are very different in these three groups.

### TABLE VIII

**Length of Life After the Onset of Paroxysms**

<table>
<thead>
<tr>
<th>Number of Years since onset of Paroxysms</th>
<th>Number of Cases</th>
<th>Number of Cases</th>
<th>Number of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Still living</td>
<td>Dead</td>
<td>Total</td>
</tr>
<tr>
<td>Up to 5</td>
<td>16</td>
<td>2</td>
<td>18</td>
</tr>
<tr>
<td>5-9</td>
<td>28</td>
<td>7</td>
<td>35</td>
</tr>
<tr>
<td>10-14</td>
<td>13</td>
<td>3</td>
<td>16</td>
</tr>
<tr>
<td>15-19</td>
<td>10</td>
<td>0</td>
<td>10</td>
</tr>
<tr>
<td>20-24</td>
<td>4</td>
<td>1</td>
<td>5</td>
</tr>
<tr>
<td>25-29</td>
<td>9</td>
<td>2</td>
<td>11</td>
</tr>
<tr>
<td>30-39</td>
<td>2</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>40-49</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>50 and more</td>
<td>2</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td>Total</td>
<td>84</td>
<td>16</td>
<td>100</td>
</tr>
</tbody>
</table>

* The numbers in brackets indicate those who have died.

1. **Normal Group.**—Where the heart was otherwise normal, serious complications were rare and a decrease in the frequency of the attacks from some change in the patient’s life was quite as likely as an increase. The follow-up of these patients and close contact with many of them for years has convinced us that as a rule paroxysms do not cause progressive deterioration and still less disease of the heart, though some are very resistant to treatment and continue to have paroxysms with the same frequency over many years. Some examples may be given.

In one patient paroxysms started when he was 9 and continued through life. A few months before his death at 69, although he had developed lymphatic leukaemia which was held in check by deep X-ray therapy, he had no signs of heart disease, a normal electrocardiogram with good T waves, and was able to lead an active life (Case 1).

Another started having paroxysms when 50; at 75, though they were rather frequent and he had suffered from an enlarged prostate for seven years, he was still in business and did not notice any dyspnea. There were no signs of cardiac or arterial disease, and his cardiogram was normal except for ventricular extrasystoles. He died three years later a few days after prostatectomy (Case 51).

A third was 68 when he was first seen and paroxysms had started at 23; they lasted up to two hours and he was rarely two weeks without. He was treated with bromides and luminal, and at 75, though his blood pressure was 180/110 and his heart a little enlarged, he had no complaint except of “a dulcimer constantly playing all the old tunes that had run through his life”
this had started two years before and was associated with increasing nerve deafness. His paroxysms were shorter and less severe (Case 66).

The cardiograms of these three cases are shown in Fig. 19.

![Cardiograms](image)

Fig. 19.—Normal electrocardiograms from three men who have had paroxysms for (A) 60 years (Case 1, aged 69); (B) 55 years (Case 66, aged 74); and (C) 28 years (Case 51, aged 75).

A fourth started having attacks when he was 7, and has rarely been three months without and sometimes has two or three in a month, lasting three to four hours or less commonly a day. He was seen when 56 with slight dyspnoea, with his aorta a little diluted and with broad QRS complexes in the cardiogram—findings which are not rare at his age. After another 10 years he has not aged much and there has been no substantial change in the condition of his heart or of his paroxysms, and he still leads an active life. He may die of heart disease, but in view of his long story this appears incidental (Case 89).

Table IX shows the frequency and length of paroxysms in a patient on

<table>
<thead>
<tr>
<th>Year</th>
<th>Number of Attacks</th>
<th>Length of Attack in Hours</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Average</td>
</tr>
<tr>
<td>1927</td>
<td>6</td>
<td>13</td>
</tr>
<tr>
<td>1928</td>
<td>7</td>
<td>14</td>
</tr>
<tr>
<td>1929</td>
<td>10</td>
<td>17</td>
</tr>
<tr>
<td>1930</td>
<td>7</td>
<td>16</td>
</tr>
<tr>
<td>1931</td>
<td>16</td>
<td>20</td>
</tr>
<tr>
<td>1932</td>
<td>7</td>
<td>21</td>
</tr>
<tr>
<td>1933</td>
<td>10</td>
<td>17</td>
</tr>
<tr>
<td>1934</td>
<td>19</td>
<td>9</td>
</tr>
<tr>
<td>1935</td>
<td>12</td>
<td>13</td>
</tr>
<tr>
<td>1936</td>
<td>11</td>
<td>9</td>
</tr>
<tr>
<td>1937</td>
<td>5</td>
<td>20</td>
</tr>
<tr>
<td>1938</td>
<td>4</td>
<td>25</td>
</tr>
</tbody>
</table>

* Excluding attack of 29 days.† Excluding attack of 17 days.
PAROXYSMAL TACHYCARDIA

whom no treatment, medicinal or general, has had any influence. But after 29 years of paroxysms her health has not been impaired and during 12 years' observation there has been no progressive increase in their frequency.

All these were free from organic heart disease. Three others are still without any signs of heart disease or of impaired health after more than 20 years of attacks.

2. Rheumatic Group.—Here paroxysmal tachycardia might be expected to be more serious, but generally this was not so. Most of the rheumatic patients have been satisfactory (see Table VIII), and they have done as well as could have been expected had they been free from paroxysms. Details of three follow: the first was one of the few instances where her progress was downhill from about the time the paroxysms started; the second was successfully treated with quinidine; the third was not influenced by treatment, but got on well for eleven years with paroxysms and for another nine with fibrillation.

A woman of 35 with mitral stenosis had her first paroxysm while hop-picking, and subsequently short attacks occurred frequently up to three or four daily. Three years later fibrillation became established and she responded well to treatment with digitalis, but after another two years abandoned treatment for months and was admitted with congestive failure and a cerebral embolism, and died after three weeks. Autopsy showed extreme mitral stenosis, left intra-auricular thrombosis, and pulmonary infarcts (Case 46).

The second had rheumatic fever at 21 and paroxysms started at 33, after her third pregnancy. At 46, when first seen, the heart was slightly enlarged, with aortic incompetence. For 8 years most of her attacks were reported and their frequency and duration, shown in Table X, were greatly reduced with quinidine; when this was stopped they became more frequent, after which she again resumed quinidine with success. Several attacks were observed, always at about the same rate, those recorded

### Table X

**Frequency and Length of Paroxysms in Case 70, Showing the Influence of Quinidine**

<table>
<thead>
<tr>
<th>Date</th>
<th>Period of Time in Months</th>
<th>Number of Paroxysms</th>
<th>Average Length in Hours (approx.)</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>1924-25</td>
<td>10</td>
<td>39</td>
<td>3.9</td>
<td>1.5</td>
</tr>
<tr>
<td>1925-26</td>
<td>15</td>
<td>20</td>
<td>1.3</td>
<td>1.0</td>
</tr>
<tr>
<td>1926</td>
<td>2</td>
<td>10</td>
<td>5.0</td>
<td>0.5</td>
</tr>
<tr>
<td>1927</td>
<td>4</td>
<td>6</td>
<td>1.5</td>
<td>0.5</td>
</tr>
<tr>
<td>1927</td>
<td>2</td>
<td>20</td>
<td>10.0</td>
<td>0.5</td>
</tr>
<tr>
<td>1927</td>
<td>7</td>
<td>16</td>
<td>2.3</td>
<td>1.5</td>
</tr>
<tr>
<td>1928</td>
<td>12</td>
<td>13*</td>
<td>1.1</td>
<td>0.5</td>
</tr>
<tr>
<td>1929</td>
<td>12</td>
<td>4†</td>
<td>0.3</td>
<td>2.0</td>
</tr>
<tr>
<td>1930</td>
<td>12</td>
<td>6</td>
<td>0.5</td>
<td>3.0</td>
</tr>
<tr>
<td>1931</td>
<td>12</td>
<td>4</td>
<td>0.3</td>
<td>1.5</td>
</tr>
<tr>
<td>1932</td>
<td>12</td>
<td>3</td>
<td>0.2</td>
<td>1.5</td>
</tr>
<tr>
<td>1933–35</td>
<td>36</td>
<td>15</td>
<td>0.4</td>
<td></td>
</tr>
<tr>
<td>122</td>
<td>85</td>
<td>0.7</td>
<td>1.5</td>
<td>With Quinidine.</td>
</tr>
<tr>
<td>14</td>
<td>69</td>
<td>3.0</td>
<td>1.0</td>
<td>Without Quinidine.</td>
</tr>
</tbody>
</table>

* Including 6 in two weeks with influenza.
† Excluding several short attacks in two weeks with influenza.
MAURICE CAMPBELL AND G. A. ELLIOTT

clinically at about 240, those recorded graphically at 232, 225, and 233 (Fig. 8b). Her condition 14 years after she was first seen had not changed for the worse, and with ten grains of quinidine daily her attacks were still well controlled, about one each three months. Her blood pressure was almost the same as 14 years before and the heart seemed little larger. A paroxysm was recorded at a rate of 210, rather slower than before (Case 70).

The third had rheumatic fever when 5 years old, after which she was slightly short of breath. At 20 she had sudden palpitation when walking upstairs, and following this attacks lasting from one to twenty-four hours. She was first seen when 29 with a rather large heart and with mitral and aortic lesions. Paroxysms occurred frequently (Fig. 8a) and were not reduced in frequency by strophanthin, digitalis, or quinidine; perhaps they were least frequent when she was taking bromides. Generally she had one or two attacks a week, and there was little change for two years. Fibrillation then became established when she was 31; the paroxysm before this lasting four days and seeming different because her heart had been irregular (paroxysmal fibrillation). Nine years later, after marriage and three years in India, fibrillation was still well controlled with digitalis (Case 52).

**Paroxysmal Tachycardia and Auricular Fibrillation.**—The relationship between these two may be considered here as it concerns rheumatic patients mainly. There is some connection, though it is not very close. Paroxysms interrupted fibrillation in one patient (Fig. 17). There were 7 others, 5 with rheumatic heart disease and 2 without, where fibrillation eventually became established and details are shown in Table XI. Another after having paroxysms for twenty-four years began to have these interspaced with paroxysmal auricular fibrillation (both recorded); she still maintains normal rhythm for a time only.
PAROXYSMAL TACHYCARDIA

3. Hyperpietic and Myocardial Groups. These patients have done less well than those with normal or rheumatic hearts, as nearly half of them have died. Two had syphilitic heart disease and eight had high blood pressure; in none of these did the occurrence of paroxysms seem of special significance, as the four who have died had serious heart disease which appeared to run its usual course. The other six in spite of their paroxysms have continued in moderate health—one after 26 years and another after 15 years, but possibly these two ought to be transferred to the normal group as the blood pressure may have developed later than and independently of the paroxysms. In all the prognosis was that to be expected from the condition of the heart quite apart from the paroxysms.

The myocardial group must be considered in rather more detail, and in view of the more serious outlook it is noteworthy that four of the eight with ventricular paroxysmal tachycardia belonged to this group. Four of the twelve in the myocardial group were elderly patients and would probably have come under observation for their heart condition quite apart from the paroxysms, which were unimportant compared with the other symptoms and findings; three of these have died.

The fifth was 58 and would not have been seen but for his paroxysms. Although he had no other symptoms his heart was enlarged and the cardiogram showed biphasic T waves in lead I and widened QRS complexes. Even so, although one paroxysm lasted nine days and led to congestive failure, they have not proved serious, since after six
years he is still well, with no progressive changes in the cardiogram or in the size of the heart (Case 16).

Three others would have passed as normal except for the cardiograms in the free intervals; as all were between 26 and 40 the abnormality is the more remarkable. The changes that may be found in the cardiogram for hours or days after a long attack were not regarded as proof of heart disease, though they mean that this possibility has to be excluded with care.

One had a tuberculous hip excised when he was 11 and at 30 had his first paroxysm. Generally they lasted for an hour only, but on five occasions for seven to ten days so that he was in hospital. The abnormal cardiogram was the only evidence of heart disease, but now after 15 years he is becoming short of breath on exertion. He has some calcified areas in the mediastinum and at the apices of the lungs, so possibly some tuberculous disease spreading to the pericardium may be responsible for his paroxysms and for the abnormal cardiogram (Case 11).

One started having paroxysms when she was 12 years old. She was seen when 27 because although they were not frequent and only lasted five or six hours she felt ill for some days after them. Nothing was found except scoliosis and a cardiogram with flat T waves and rather wide QRS complexes. Five years later she was better, her attacks shorter, and her cardiogram more normal (Case 6).

The third has more frequent attacks which have ceased to trouble him since he found that they can be stopped easily by bending down to open a low drawer; he has no other symptoms, but his cardiogram is still not normal (Case 68).

In the last two the abnormal signs in the cardiograms have become less, and now after six years they are in good health and without other symptoms. Possibly the paroxysms and abnormal cardiograms were the only signs of a temporary myocarditis from which there has been good recovery. Another patient, included as normal because nothing can now be detected in his heart (Case 43), had his first paroxysm when he leant out of bed during his convalescence from pneumonia; he may have had a pneumococcal myocarditis and recovered completely except for the tendency to paroxysms, but there is no proof of this.

The remaining four (Cases 5, 15, 79, and 81) had abnormal cardiograms and some enlargement of the heart without obvious cause. Their ages when first seen were 21, 39, 37, and 18. They form an important sub-division as they include three of the five patients who have died suddenly or during attacks. All are fully described in the next section and in two the fatal result was probably due to ventricular fibrillation associated with ventricular paroxysms. The third is more disturbing because he provides the only instance where death was due to a paroxysm without the certainty of preceding heart disease, and where such changes as were found seem to have been caused by the paroxysms rather than to have caused them—a process which might be expected on a priori grounds, but is in our view most exceptional. Paroxysms had occurred for 21 years, and where so little could be found at the autopsy it is unlikely the enlargement of the heart was present at the start or that there could then have been any coronary disease; the simplest explanation seems to be that attacks which were inconvenient rather than serious for 15 years began to last longer and to produce congestive failure, and that one lasted long
PAROXYSMAL TACHYCARDIA

enough to cause his death. Without this case, we should have said more
dogmatically that paroxysmal tachycardia alone never proved fatal. Even
with myocardial disease, slight or serious, the prognosis is generally that of
the heart disease and is not much affected by the paroxysms.

Patients who have Died

Some details about the 16 deaths during the period of observation (1925–38)
will be given. At first sight this number suggests a bad prognosis, but in 4
the cause of death was nothing to do with the heart and in 7 others their
length of life was quite up to the natural expectation of their heart disease, had
there been no paroxysms. In 5 of the 16, however, death took place during
a paroxysm and these must be considered fully, as even 5 out of 100 would be
a serious proportion, were there no other reason for the deaths.

Two of the 4 patients whose deaths were not due to their hearts had suffered
from paroxysms for 60 and 28 years and died from leukaemia and after prosta-
tectomy respectively; neither had any signs of heart disease even near the end
of life. (Cases 1 and 51, see p. 149.) The third died two years after an
operation for cancer of the breast with recurrences; paroxysms are not
uncommon when there are secondary deposits in the thorax, but here they
occurred for some years before the neoplasm. The fourth died two years
later from cerebral hemorhage.

In 7 patients death was caused by heart disease, but its course did not
seem to be influenced by the paroxysms. One had a blood pressure of 200/110
and pulsus alternans; a second, whose paroxysms began after fibrillation had
been arrested by quinidine, died six years later with congestive failure; a
third had severe angina for six years before his death from coronary occlusion;
and a fourth had syphilitic aortic incompetence and an aneurysm. Two had
rheumatic heart disease: one, with mitral stenosis and paroxysms for three
years, died two years after the onset of fibrillation; and the other, with a
large heart and mitral and aortic lesions, lived eleven years after the onset of
frequent paroxysms; the seventh (Forman, 1931) had a calcified pericardium
and died five years after the onset of his attacks.

So far as we know none of these 11 died unexpectedly or during paroxysms
and we have no reason to think the length of life was shortened by them,
though sometimes they were the first sign that the condition of the heart was
starting to go downhill. This may be so, but is by no means the rule, and
many patients with heart disease have done well without any progressive loss
of reserve.

The remaining 5 are more significant, since they died suddenly or during
paroxysms. The only one of these five that was thought to have a normal
heart almost certainly died from an overdose of morphia. Of the other four
two died suddenly and two with congestive failure.

The first was seen when his attacks, though infrequent, had already been present
for 15 years; and after another 11 years his wife wrote to report his death in a
paroxysm. To further enquiries his doctor replied that as on previous occasions when
called in he had given half a grain of morphia, saying it was sufficient for the night; but the patient had sent for a second doctor, who, unaware of the previous treatment, had given another injection of morphia, which probably caused his death (Case 28).

The second had mitral stenosis and moderate enlargement of the heart, and her supraventricular paroxysms began when she was 27; at 36 they had not greatly changed her condition and recurred about each six weeks and lasted for six to forty-eight hours (once seven days). She was admitted to hospital on the fourth day of an attack in the fourth month of pregnancy with anaemia and commencing failure, and as all efforts to stop it were unsuccessful her failure became worse. She lost a good deal of blood after a miscarriage and died on the seventeenth day. Here the attack was the immediate cause of the failure which led to her death, but the anaemia and the miscarriage were complicating factors (Case 84).

The third died suddenly when he was 45. He was refused for the army at 18 because of his heart, and again during the War at 26; he had some dyspnea and palpitation, and when 37 lost consciousness after a paroxysm. Seen two years later, after a similar attack, he had a large heart mainly to the left, a loud systolic murmur, some widening of the aorta without high blood pressure, and a partial bundle branch block, and it was difficult to decide the exact nature of his heart disease. Several short paroxysms of ventricular tachycardia were observed at a rate of 186. Six years later we learnt that he had died suddenly while carrying an electric cooker downstairs. The coroner kindly supplied this report—"Heart 540 grams, hypertrophy and dilatation, chiefly of ventricles; muscle thin and flabby with fibroid degeneration; mitral and tricuspid valves themselves normal but incompetent; coronary arteries moderate atheroma, orifices narrowed; no rheumatic or syphilitic disease." Here it seemed probable that disease of the coronary arteries started at an early age and progressed till it led to his death. He was never an example of the common type, where the paroxysms were the main presenting symptom, because the physical signs and dyspnea which led to his rejection from the army were present several years before the paroxysms (Case 79).

The fourth also died suddenly. His health had always been good except for diphtheria at 9, when he was in bed five months with paralysis of the soft palate and dilatation of his heart. Though some extrasystoles persisted he became free from symptoms, and when 19 took up heavy work, lifting crates that weighed 72 lb.; he was able to carry on with this and with boxing and football without dyspnea. When he was 21 he woke up in a paroxysm, which lasted ten days at a rate of about 200. After three weeks he felt fit and returned to work, but had shorter attacks of one and five days. When first seen five months later, his heart rate was 72 with frequent extrasystoles. The blood pressure was 120/85, and there were no physical signs except slight enlargement of the heart. His cardiogram showed poor T waves and ventricular extrasystoles of two types. He was admitted to hospital next day, with a ventricular paroxysm at a rate of 220 (Fig. 3); the blood pressure was under 100, and already there was some enlargement of the liver and slight edema of the feet. Pressure on the carotid sinus and digitalis and strophanthide failed to arrest the attack, which stopped on the fifth day after twenty grains of quinidine, the rate falling gradually from 220 to 194 as a result of quinidine, and then suddenly to 80. The signs of congestive failure quickly disappeared and within two days he felt quite well, but the changes in the cardiogram (Fig. 21) led us to keep him in hospital for three weeks, by which time the inversion of T I had disappeared. Because of these changes and because the paroxysms were ventricular and quickly led to congestive failure, his doctor was given a guarded prognosis. He kept well for two weeks and perhaps unwisely took tint. dig., min. xx daily. His doctor wrote that as he was starting for hospital for his next routine visit he suddenly fell down dead; there was no autopsy (Case 5).

The fifth patient started attacks when he was 24 and for the next 15 years they recurred each six or nine months, lasting from a few hours up to one or even two days. They were merely an inconvenience and he continued work as an engine driver.
In 1931, when he was 39, two attacks lasted 7 and 8 days, and for the first time he was ill enough to go to bed. Two months later he was admitted with congestive failure after 14 days of an attack which stopped when he had been given twenty minims of tincture of digitalis t.i.d. for three days. A cardiogram taken the next day when he seemed almost well showed some inversion of T in leads II and III, but ten days later these had gone, to be followed after a week by changes in the S-T intervals (Fig. 22).

Probably the inversion of T was due to the congestive failure and not to the digitalis or to coronary disease; the findings at the autopsy described later and somewhat similar changes after other long paroxysms seem to justify this view.

Fig. 21.—Changes simulating those of cardiac infarction, produced by a 10-day ventricular paroxysm; (A) 2 days after, (B) 4 days after, and (C) 3 weeks after (Case 5).

Fig. 22.—Changes after a 17-day paroxysm, simulating the changes after cardiac infarction; (A) one day after, (B) eleven days after, (C) eighteen days after, and (D) four years after (Case 15).
Though he returned to work after some months he was never really well. In 1932 he had attacks of 8 and 14 days, and in 1933 nearly every month lasting about 4 days. In 1934 they generally lasted about 7 days and he was often away from work. Although there had been no more congestive failure he was able to do progressively less, and in 1935 had a pulmonary infarct during an attack. After this he gave up work and the attacks rarely lasted more than 4 or 5 hours. In 1936 his heart sounds were normal, his arteries soft, and his blood pressure 130/80; but an orthodiagram showed slight enlargement and the cardiogram (Fig. 22D) showed more depression of S-T in lead I than in 1933. Nothing else abnormal could be found; he had no attacks in hospital and was advised to increase his activity slowly and to take quinidine regularly; two months later he wrote that he had been better.

In 1937 he had another attack and after resting in bed three weeks he called in his doctor, but the paroxysm persisted, and there was progressive failure with jaundice and orthoponera. On admission in the fifth week under the care of a colleague the heart rate was between 180 and 190, and he was dangerously ill. He was given 60 minims of tincture of digitalis, and during the night 1/2 gr. morphia, but died next morning. At autopsy the heart weighed 450 grams (it seems unlikely that it was this weight the year before). There was no valvular disease, no signs of past syphilis, and very little atheroma of the aorta. The myocardium showed no scarring, and the coronary arteries were patent. There were 400 c.c. of slightly turbid fluid in the pericardium and the right pleural cavity contained a litre of blood-stained fluid. The lungs showed one very large and two smaller infarcts, as well as old scars. There was a large granular liver with nutmeg changes, and a firm dark spleen about twice the normal size (Case 15).

**Summary and Conclusions**

One hundred unselected cases of paroxysmal tachycardia have been studied and followed for some years. In 42 the diagnosis was confirmed by the electrocardiograph, in 30 by observation of an attack, and in 28 by the history alone. The criteria of diagnosis, when this has to be made on history, have been described; the sudden onset of palpitation being the most reliable single symptom. There may sometimes be difficulty in distinguishing paroxysmal flutter and tachycardia.

Of the 42 attacks with graphic records, 8 were ventricular and 34 supraventricular; 11 of the latter were nodal, but in many of the other 23 the site of origin could not be defined more precisely. Extrasystoles were observed between attacks in twenty-one, but were of minor assistance in predicting the type of the attack. Ventricular paroxysms were very uncommon without serious heart disease.

There were 41 of these cases with heart disease, 19 rheumatic, 2 syphilitic, 8 hyperpietic, and 12 myocardial. There was no heart disease other than the arrhythmia in 59, though 4 of these had a goitre.

The rate was between 160 and 200 in nearly half; it was between 140 and 240 in 90 per cent., but occasionally outside this wide range. There was no great difference between the various aetiological groups. Nodal attacks tended to be a little slower and ventricular attacks were rather more often above 190, but even above this rate ventricular attacks formed a small minority.

Paroxysms are generally of short duration, lasting for hours rather than for days. In 61 the customary duration was less than two hours, and in
another 28 twelve hours or less. There were 4 where it was about twenty-
four hours and only 7 where it was longer than this. But 18 others, making
29 in all, sometimes had attacks lasting more than one day, viz. over twenty-
four hours, 10 cases; two or three days, 8 cases; up to seven days, 4 cases;
up to ten days, 5 cases; and two to four weeks, 2 cases. One third, therefore,
of our patients had some attacks lasting more than a day, 10 per cent. rarely,
10 per cent. often, and 10 per cent. habitually. Long attacks included an
undue proportion of the ventricular paroxysms and were more common in
those with myocardial disease.

Paroxysmal tachycardia is a symptom rather than a disease. In a minority
of patients it accompanies serious heart disease, when, of course, the prognosis
is grave. Such cases are nearly always under observation for their heart
disease before the onset of paroxysms. Ventricular paroxysms form a fairly
large proportion of this group and are rare otherwise. In most patients
paroxysmal tachycardia is not in itself of any grave significance. It is due to
reflex causes more often than to any primary change in the heart muscle.
This applies not only to the majority whose hearts are otherwise normal but
also to most of those with rheumatic heart disease and to some of those with
other myocardial disease.

There is no close association between paroxysmal tachycardia and
paroxysmal auricular fibrillation. In some of the rheumatic cases and less
often in others paroxysms of fibrillation may alternate with or replace paroxysms
of tachycardia. In the rheumatic cases established fibrillation then becomes
a possibility in the near future.

The prognosis of paroxysmal tachycardia as regards life is therefore excel-
alent, unless it is of the rare ventricular type, unless appearing relatively late in
life it is the first indication of disease of the coronary arteries, or unless before
the paroxysms have started there is already serious heart disease. Three of
these patients have lived fifty years after the onset of their paroxysms, another
18 for more than twenty years, and another 26, making 47 per cent., for more
than ten years, and most of these are still in good health. Paroxysmal tachy-
cardia does not produce heart disease, even when it continues throughout
life, though one possible exception to this statement has been quoted. In
general the prognosis depends on the condition of the heart muscle and should
be decided without reference to the paroxysms. There is no constant tendency
for the paroxysms to get worse as life advances, and usually some form of
treatment can be found which will reduce the frequency and the discomfort
produced by the attacks.

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REFERENCES

—— (1933). Ibid., 1, 224.