Determinants of the Duration of the Refractory Period of the Atrioventricular Nodal System in Man *

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One of the fundamental characteristics of nervous tissue, skeletal muscle, and myocardium is their refractoriness for short periods following depolarization (1). This period, during which a propagated action potential cannot be evoked by a stimulus, has been designated the "effective refractory period" (2), and the effective refractory period thus limits the maximal rate at which depolarizations can occur. When normal atrioventricular (AV) conduction takes place, the ventricular contraction rate is limited by the AV nodal system, which has been shown in experimental animals to have a longer effective refractory period than atrial or ventricular tissue (3). In spite of the fundamental importance of the refractory period of the AV nodal conduction system, this interval has heretofore been estimated only from electrocardiograms obtained from patients with spontaneous cardiac arrhythmias (4). The presence of arrhythmias may be associated with abnormalities of conduction, and the analysis of such electrocardiograms does not allow systematic investigation of the factors that influence this period in normal subjects. In the present investigation a practical method for the measurement of the effective refractory period of the AV nodal system in conscious human subjects is described. With this method normal basal values for this period were established, and the effects of tachycardia, exercise, atropine, and of various sympathomimetic drugs on its duration are presented.

Methods

A bipolar electrode catheter ¹ was passed into the right atrium and positioned against the mid-portion of the lateral wall of the right atrium. A slight bend was

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plex is evoked. Figure 2A shows an electrocardiogram in which the premature stimulus, introduced between S _1_ and S _2_, initiates a P wave and a QRS complex (although the latter is delayed, and as a consequence S _2_ is not conducted to the ventricles). In contrast, in the tracing shown in Figure 2B, the premature stimulus (between S _1_ and S _2_) is delivered 30 msec earlier, the atrial depolarization wave reaches the AV conduction system during the AVRP, the impulse is not transmitted to the ventricles, and no QRS complex occurs. Since the intervals were shortened by steps of 10 msec, alterations in the AVRP of less than 10 msec could not be detected by this technique.

After determination of the AVRP by the extra-stimulus method described above, the effects of one or more interventions on this measurement were studied. Whenever the effects of more than one intervention were determined in the same patient, a control measurement was repeated between the interventions. The effects of induced tachycardia on the AVRP were determined in nine patients by first making the measurement at a rate just greater than the sinus rate, and by repeating it 5 minutes after the rate had been elevated by the electrical pacemaker. The effects of the following drugs infused intravenously were studied: isoproterenol, 1.0 to 1.5 μg per minute (seven patients); methoxamine, 0.95 to 1.40 mg per minute (two patients); phenylephrine, 60 to 240 μg per minute (six patients); norepinephrine, 1.6 to 3.2 μg per minute (three patients); and atropine, 0.5 mg to 1.0 mg as a single injection (11 patients). In two patients the effects of phenylephrine on AVRP were studied both before and after atropine. In seven patients the effects on the AVRP of leg exercise, consisting of pedaling a bicycle ergometer in the supine position at 40 rpm at an external work load of approximately 500 ft-lbs per minute for 5 minutes, were determined.

In 15 patients the AVRP was also measured by a second technique, termed the "increasing rate" method, described in detail previously (6), and diagrammatically illustrated in Figure 1C. The basic rate of stimulation was gradually increased until an interval between stimuli was reached which was so short that a stimulus resulted in atrial depolarization but was not transmitted to the ventricles.

A total of 33 patients ranging in age from 11 to 59 years were studied in the postabsorptive state, 1 hour after the administration of 100 mg sodium pentobarbital. None of the patients had atrioventricular or intraventricular conduction abnormalities, the P-R intervals and QRS durations in each patient being normal in the basal state. No patients had experienced congestive heart failure or had marked limitation of cardiac reserve at the time of the study. None of the patients were receiving digitalis, quinidine, or procaine amide, and the serum electrolytes were always within normal limits. Eleven of the patients had functional heart murmurs, six had mitral stenosis, five had atrial septal defects, and three had mild aortic stenosis. In one patient each the diagnosis was ventricular septal defect, mild aortic regurgitation, idiopathic hypertrophic subaortic stenosis, right ventricular myxoma, patent ductus arteriosus, cardiomyopathy, status postclosure of a ventricular septal defect, and status postaortic valve replacement for aortic stenosis. In every instance the measurements of AVRP followed the diagnostic cardiac catheterization.

**Results**

I. **Control values.** The control values of the AVRP in 20 patients studied by the extra-stimulus method averaged 350 ± 53 msec (SD), a value which was significantly shorter (p < 0.01) than that obtained by the increasing rate method in 15 patients, which averaged 413 ± 52 msec.

II. **Tachycardia (Figure 3A).** In nine patients the AVRP was first measured at a rate just above the spontaneous sinus rate, which ranged from 68 to 102 and averaged 84 per minute; at this rate the AVRP ranged between 250 and 430 and averaged 344 msec. The atrial rate was then increased to between 109 and 131 (average = 120 per minute), and the AVRP rose significantly (p < 0.02) to between 340 and 450 msec (average = 390 msec). Thus, abbreviation of the R-R interval by an average of 215 msec (30% of control) was associated with a prolongation of the AVRP by an average of 55 msec (16% of control).

III. **Atropine (Figure 3B).** In 11 patients the heart rate before atropine ranged between 68 and 92 and averaged 77 per minute, while the AVRP ranged between 280 and 430 and averaged 344 msec. Atropine resulted in an elevation of heart rate to an average of 103 per minute. In contrast to the finding when heart rate was increased by
electrical stimulation, AVRP declined significantly ($p < 0.01$), falling to between 160 and 330 msec (average = 246 msec). Thus, an average fall of the R-R interval of 195 msec (25% of control) was associated with a decline of the AVRP averaging 98 msec (29% of control).

IV. Isoproterenol (Figure 3C). In seven patients the control heart rate averaged 82 per minute and rose to an average level of 95 per minute during the isoproterenol infusion. AVRP ranged between 300 and 500 and averaged 405 msec during the control period, falling in each patient to an average value of 300 msec, which was significantly lower ($p < 0.01$) than that noted during the control period. Thus, an average decrease in the R-R interval of 90 msec (12% of control) was

![ECG and arterial pressure recordings](image)

**Fig. 2**. **Simultaneous recordings of electrocardiogram and arterial pressure pulse during determination of AVRP.** $S_1$ to $S_6$ represent electrical stimuli. In tracing A (top) the extra stimulus, delivered between $S_2$ and $S_3$, was transmitted to the ventricles, but in tracing B (bottom) the extra stimulus, delivered between $S_3$ and $S_4$, was not transmitted.
associated with an abbreviation of the AVRP, which averaged 105 msec (26% of control).

V. Exercise (Figure 3D). In seven patients the heart rate at rest ranged between 60 and 95 and averaged 72 per minute, while the AVRP ranged between 320 and 520, averaging 390 msec. Muscular exercise raised heart rate to a mean value of 117 per minute, while AVRP fell significantly (p < 0.01) to an average value of 280 msec. Thus, an average decline of the R-R interval of 320 msec (28% of control) was associated with an abbreviation of the AVRP, which averaged 110 msec (39% of control).

VI. Phenylephrine and methoxamine (Figure
The administration of phenylephrine and methoxamine, in doses sufficient to elevate the systolic pressure by an average of 56 mm Hg and the diastolic pressure by 28 mm Hg, slowed heart rate, from an average value of 75 per minute during the control period to one of 56 per minute. This was associated with a significant prolongation of the AVRP from an average of 326 to 578 msec (p < 0.01). Thus, an average increase of the R-R interval of 274 msec (34% of control) was associated with a lengthening of the AVRP, which averaged 252 msec (77% of control).

VII. Phenylephrine and atropine (Figure 4B). In two patients the effects of phenylephrine were first studied, as described above, and after a second control measurement atropine was administered. After redetermination of the AVRP phenylephrine was reinfused to produce an identical elevation of arterial pressure. In these two patients the individual effects of phenylephrine and atropine on both the AVRP and the R-R interval were similar to those already described (Figure 4A and Figure 3B). However, when phenylephrine was reinfused after atropine, neither the AVRP nor the R-R interval was altered.

VIII. Norepinephrine (Figure 4C). The administration of norepinephrine to three patients slowed heart rate from an average of 78 per minute to 70 per minute while AVRP increased from an average value of 330 to 410 msec.

Discussion

Previous studies on the refractory period of the AV nodal conduction system have been carried out in a variety of experimental animal preparations. Krayenbuehl and collaborators showed that epinephrine shortens the functional refractory period of the AV nodal system in the isolated canine heart-lung preparation (5). Subsequently Mendez, Aceves, and Mendez found that pentobarbital and cardiac glycosides prolong this period and that at certain dose levels this action of the glycosides could be reduced by prior sympathetic denervation (7, 8).

Rosenbluth, Mendez, and their respective collaborators indicate that the functional refractory period of excitable tissue is the shortest attainable interval between two induced responses when the second stimulus is delivered at 2 or more times the threshold intensity (1, 3, 9). They emphasized, however, that this value is attended by an error equal to any difference that might occur in the propagation time from the point of stimulation to the point of recording between the first and second responses. In the present investigation on man it was observed that this error was inconsistent and frequently quite large. For example, in the tracing reproduced in Figure 2A, the ventricular depolarization resulting from the interpolated stimulus was delayed by 120 msec. This potential error results primarily from variations in the conduction velocity in the AV conduction system, which can be altered by influences that also modify the duration of the AVRP. Accordingly, rather than determining the shortest period between two induced responses, we measured the longest interval between two stimuli that could be achieved when the second stimulus failed to evoke a propagated ventricular depolarization. This measurement conforms closely to the "effective refractory period" defined by Hoffman and Cranefield (2), although the strength of the interpolated stimulus in our study was not maximal but only twice the diastolic threshold level. Since the stimulus was applied to the atrium, rather than directly to the AV node, alterations of conduction velocity within the atrium would modify the duration of the AVRP. However, in the present investigation no detectable alterations in atrial conduction, as determined from the duration of the atrial depolarization wave, occurred.

In the interpretation of changes in the AVRP observed in the present investigation the complexities of AV conduction, which have recently been reviewed (10, 11), must be considered. Thus, many findings are consistent with the hypothesis that decremental conduction occurs in the AV conduction system (10), and recordings of transmembrane action potentials have shown that conduction is slowest across a narrow zone at the atrial margin of the AV node (12). In the isolated rabbit heart, Cranefield, Hoffman, and Paes de Carvalho have shown that the failure of AV transmission caused by acetylcholine takes place in fibers at the atrial margin of the node (13), whereas lower portions of the node appear to be involved in the failure of transmission caused by tachycardia (10).

Thus, although determination of the AVRP
Fig. 4. Effects of A) methoxamine and phenylephrine and C) norepinephrine on the relationship between the AVRP and the R-R interval. B) Modification by atropine of the effect of phenylephrine on the AVRP and R-R interval. C = control; P = during phenylephrine; A = after atropine; A + P = during phenylephrine, after atropine.
does not simply provide a measurement of the period during which the AV node is totally inexcitable or unresponsive, the measurement does provide an assessment of the time interval following a stimulus during which the AV nodal conduction system is incapable of transmitting an impulse that can initiate a propagated ventricular depolarization. From the present investigation in man it is apparent that this period is not a static one, but that many common interventions can profoundly modify its duration. Tachycardia, produced by electrical stimulation of the atrium, tended to lengthen the AVRP slightly, a finding that correlates with the longer duration of the AVRP when it is measured by the increasing rate than when it is determined by the extra-stimulus method. It has previously been observed in normal human subjects that when the rate of stimulation of the right atrium is increased the interval between the stimulus and the QRS increases (14). Although the results of the present study are opposite to those described by Mendez, Gruhzit, and Moe in the anesthetized, open-chest, acutely denervated dog heart (3), they are consistent with the observation of a reduced amplitude and rate of rise of the transmembrane potential in the atrial part of the AV node during rapid electrical stimulation of the atrium (10). Therefore the shortening of the AVRP that occurred when tachycardia was induced by the administration of atropine and isoproterenol or by muscular exercise is even more striking when it is considered that tachycardia per se tends to lengthen this interval.

Alterations in the activity of both the sympathetic and parasympathetic nervous systems affect both the AV conduction system and the sinoatrial node. However, the relationship between the R-R interval and the duration of the AVRP was not a linear one, and indeed, the relative changes in the duration of the AVRP induced by atropine, isoproterenol, phenylephrine, methoxamine, and norepinephrine were often greater than the simultaneous changes that occurred in the duration of the R-R interval. The abbreviation of the AVRP associated with the administration of atropine and of isoproterenol is interpreted, respectively, as the effect of the withdrawal of parasympathetic influences from, and of the activation of adrenergic receptors in, the AV conduction system. Muscular exercise shortened the AVRP in a manner similar to atropine or isoproterenol, and it is likely that both stimulation of sympathetic receptors and reduction of parasympathetic inhibition were involved in the response. On the other hand, phenylephrine and methoxamine, two sympathomimetic pressor drugs that have no direct cardiac effects (15), simultaneously prolonged this period and slowed the sinus rate.

The finding in two patients that phenylephrine did not alter the AVRP after atropine supports the view that phenylephrine has no direct effect on the AV conduction system, although since only two patients were studied these results cannot be considered conclusive. It is well known that the slowing of heart rate produced by these drugs is reflexly mediated, and it has been shown, both in the dog and in man, that the efferent limb of the reflex to the sinoatrial node that is stimulated by elevating arterial pressure, is mediated only by the parasympathetic system (16). The finding that atropine completely prevented the prolongation of the AVRP induced by phenylephrine (Figure 4B) indicates that as a consequence of baroreceptor stimulation, the AV conduction system, like the sinoatrial node, is inhibited only by parasympathetic stimulation, withdrawal of tonic sympathetic activity playing no detectable role. Norepinephrine, like isoproterenol, stimulates adrenergic receptors in the heart, and it might therefore be expected to abbreviate the AVRP. However, the prolongation of the AVRP that occurred with norepinephrine was probably reflex in nature, consequent to its pressor action, and this effect probably masked any direct action of the catecholamine. Since only three patients received norepinephrine, however, these results must be considered to be tentative.

The chief finding that emerges from this study is that the AV conduction system in conscious human subjects is affected by many influences, both direct and reflex. In general, the effects of any given stimulus on the rhythmicity of the sinoatrial node (as reflected in the R-R interval) are directionally similar to those on the AV system (as reflected in the AVRP), although in many instances the relative effect on the latter is greater than on the former. Under most clinical circumstances, effects on the sinoatrial node, since they alter ventricular rate, are more obvious than effects on the AV system. Nevertheless, the
AVRP is of considerable clinical importance, since it limits the ventricular rate in patients with a variety of supraventricular tachycardias, including atrial fibrillation. Furthermore, the delay in conduction imposed by the AV system regulates the time interval between the contractions of the atria and ventricles and thereby plays a fundamental role in regulating the atrial contribution to ventricular filling (17).

Summary

A technique was devised for measurement of the refractory period of the atrioventricular conduction system (AVRP) in man. A bipolar electrode catheter was placed against the right atrial wall and stimulated the right atrium at a constant rate just above the sinus rate. Extra impulses were interposed between the regular stimuli; the time intervals between these extra impulses and the preceding stimuli were then shortened progressively. The time interval at which AV conduction just failed was taken as the AVRP. In 20 unanesthetized patients, studied in the basal state, the AVRP averaged 350 msec. In nine patients, atrial tachycardia, induced by rapid electrical stimulation of the right atrium, prolonged the AVRP. In seven patients muscular exercise shortened the AVRP by an average of 10 msec. Isoproterenol (seven patients) and atropine (11 patients) resulted in reductions of the AVRP similar to those observed during exercise. Elevations of arterial pressure with methoxamine or phenylephrine in eight patients prolonged AVRP by an average of 252 msec. Since this prolongation could be prevented by atropine, it was considered to result from reflex vagal stimulation. Thus, in any given patient the AVRP is affected profoundly by many influences, including sympathetic and parasympathetic stimuli.

References