Mechanism of Spontaneous Rhythmic Contraction in Isolated Rat Large Artery

Naoki Hayashida,*** Katsujî Okui,* and Yasuichiro Fukuda**

*Department of Surgery I and **Department of Physiology II,
School of Medicine, Chiba University, Chiba, 280 Japan

Abstract The mechanism of spontaneous contraction of vascular smooth muscles in the elastic artery was studied in a ring-shaped preparation isolated from the rat aorta. The observation of small changes in vascular tension with a high gain AC amplification of tension signal provided a reliable detection of spontaneous contractions. The spontaneous rhythmic contraction (RC) occurred consistently in the preparation taken from the thoracic aorta without external stimuli. The RC (frequency, 5–20 cycle/min; amplitude, 10–100 mg) was accompanied with small oscillatory changes in the membrane potential (2–5 mV, peak to peak). A reduction in temperature (below 30°C) or superfusing the preparation with Ca-free solution inhibited the generation of RC. Ca-entry blockers (verapamil and nifedipine) also inhibited the RC. The cessation of RC by these procedures reduced the vascular tension by about 40% of control baseline tension. The application of adrenergic blockers had little effect on the pattern of RC and on the vascular tension. The results suggest that the RC is generated by a synchronization of electrical and mechanical activities in relatively small groups of smooth muscle cells, which depends upon the temperature and requires the Ca-entry into the cells. The process of initiation of spontaneous RC in the rat aorta was discussed.

Key words: vascular smooth muscle, rat aorta, spontaneous rhythmic contraction, membrane potential, Ca.

The contraction of vascular smooth muscles alters the vessel diameter and regulates the vascular "tone." The contractile behavior or the pattern of contraction, however, differs between various segments of vascular tree. Many previous studies have shown that some isolated vessels, such as the portal vein of the rat, subcutaneous artery of the dog, and bovine mesenteric vein, generate spontaneous rhythmic contractions (Funaki and Bohr, 1964; Johansson and Bohr, 1966; Roddie and Scott, 1969). These rhythmic contractions are due to synchronization.

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of electrical and mechanical activities of many smooth muscle cells. In the large elastic arteries, on the other hand, spontaneous contractions occur only occasionally or can be initiated by application of chemical stimuli (high potassium, catecholamines, or low magnesium) (BiAMINO and KRUCKENBERG, 1969; ALTURA and ALTURA, 1974; ALTURA et al., 1980). Although the media of elastic artery contains plenty of smooth muscle cells, it is not certain whether these smooth muscles are inherently incapable of generating spontaneous rhythmic contractions. Furthermore, the mechanism of spontaneous contractions, if any, in the elastic arteries, has not been studied systematically. The present experiments aimed to observe the spontaneous contractions in the large conduit artery and to assess their generating mechanism and their contribution to the maintenance of vascular tension. The results showed that the large elastic artery of the rat is capable of initiating spontaneous rhythmic contractions.

METHODS

The aorta and its branches were isolated from the male Wistar rat weighing 200–300 g under pentobarbital anesthesia (50 mg/kg, i.p.). Ring-shaped vascular preparations (1–2 mm diameter, 2–4 mm width) were obtained from various regions of the large artery by transverse section and were placed in an incubation chamber (6 ml volume) (Fig. 1). The preparation was continuously superfused (6 ml/min) with the physiological salt solution (modified Tyrode solution) having the following composition in mM: NaCl 120, KCl 5.0, CaCl₂ 1.5, MgCl₂ 0.5, NaH₂PO₄ 1.0, NaHCO₃ 24.0, and glucose 10.0. The solution was bubbled with a gas mixture containing 5% CO₂ and 95% O₂ in a reservoir bottle (pH 7.40). The temperature of the solution was maintained at 37°C except when the effect of low temperature on the vascular contraction was examined. In the incubation chamber the preparation was suspended between two stainless steel rods, one for fixing the preparation and the other for connecting it to a strain-gauge transducer via a synthetic surgical thread (polyester fiber, TI-CRON® No. 5-0 or 6-0, Davis Geck), to register the isometric circular tension of the arterial segment (Fig. 1). The preparation was usually stretched transversely to apply baseline tension of about 200–400 mg. In some experiments the level of baseline tension was varied between 0 and 400 mg. The application of stretch tension higher than 500 mg was avoided because such a high baseline tension was not maintained constant but it was decreased gradually during the experiment. The output from a tension amplifier (carrier amplifier) was fed not only directly into a pen recorder (DC mode) for monitoring the baseline tension but also into a band pass filter (AC) amplifier (band width 0.05–10 Hz) for monitoring small changes in vascular tension due to spontaneous contraction. The noise level of the system for measuring the mechanical activity was reduced to about 1 mg (peak to peak) by using double mechanical shock absorbers (rubber pads), by the use of an appropriate band pass filter amplifier (see above) and by preventing the preparation-transducer system from touching directly with the wall of incubation.

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chamber which was connected to a circulating pump (Fig. 1).

The membrane potential of a single muscle cell was recorded by a conventional 3 m-KCl filled floating glass microelectrode (resistance, 10–20 MΩ). The microelectrode was inserted into the muscular layer from the adventitial surface or from the intimal layer. The transmembrane potential was monitored on an oscilloscope and was registered on a pen recorder together with mechanical activities and a temperature of the superfusing solution in the incubation chamber.

In some other experiments, effects on mechanical activities of a low temperature, nominally Ca-free solution or Ca-entry blockers (verapamil HCl, Eisai; nifedipine, Bayer) were investigated. Several adrenergic blocking agents (phentolamine mesylate, Ciba; prazosin HCl, Pfizer; yohimbine HCl, Wako; propranolol HCl, ICI) were also used to clarify whether the endogenous catecholamine contributed to the generation of spontaneous contractions. A total amount of 0.1–0.3 ml drug solution was injected for a period 1–3 min into a side tube through which the superfusing solution flowed into the incubation chamber. An approximate mean concentration of a drug in the incubation chamber was calculated. In some experiments, the physiological salt solution containing a known concentration of a drug was flowed into the incubation chamber. When nifedipine was used, the bath and solutions were protected from light. Circulating water which surrounded the incubation chamber and reservoir bottle was cooled by ice when the effects of
low temperature on the pattern of spontaneous contraction were studied.

RESULTS

1. Spontaneous rhythmic contraction in the rat large artery

The preparations isolated from the aorta and other large arteries showed spontaneous rhythmic contractions (RC) (Fig. 2). The typical RC was more consistently seen in the preparation taken from the thoracic aorta (including the ascending aorta and aortic arch) than that from more distal aorta or from some aortic branches (common carotid and common iliac arteries). The present experiment was, therefore, performed on the preparation obtained from the thoracic aorta. The RC was a relatively regular contraction usually seen in both the DC (low gain) and AC (high gain) tension registrations. The frequency of RC ranged from 4.7 to 19.6 cycle/min (11.9 ± 4.9 cycle/min, mean ± S.D., n = 20). When the baseline tension was increased up to about 400 mg by increasing passive stretch, the frequency remained fairly constant in the static phase of stretch (Fig. 2). Changes in the frequency in a dynamic phase of stretch were not systematically studied because a transient change in the frequency (i.e., increase) during or shortly after increasing the passive stretch was slight and it occurred only occasionally. The amplitude of RC, on the other hand, was increased with increasing the stretch load (Fig. 2). Generally the amplitude of RC (10–100 mg at a baseline steady state tension of about 300 mg) correlated inversely with the frequency, i.e., the higher the frequency,

![Figure 2](image-url)

Fig. 2. A typical example of spontaneous rhythmic contraction (RC) in the rat large artery (thoracic aorta). The RC at various levels of baseline tension (steady state condition) by increasing passive stretch is shown. DC, baseline vascular tension measured with low gain amplification in the DC mode; AC, tension registration with high gain amplification in the AC mode. These abbreviations are the same in following similar figures. Registration of changes in tension (AC mode) was partly saturated on a pen recorder.
the lower the amplitude. As shown in Fig. 2, the RC was clearly seen in the AC mode registration when it was not observed in low gain DC tension registration at a minimum passive stretch less than 20 mg.

2. Changes in membrane potential

The recording microelectrode was inserted into the smooth muscle layer from the adventitia or in a few cases from the intimal layer. The entry into the cell was identified by a sudden appearance of negative potentials of about 20–50 mV. Withdrawal or further penetration of the electrode abruptly abolished this negative potential. An important criterion for the successful recording was that the electrical potential was not shifted by rapid manual alteration of passive stretch tension (within 2–3 s) (Fig. 3, asterisk). This indicated that the changes in potential were not artifacts due to mechanical movements of tissue. The average membrane potential of the cells in the thoracic aorta was $-33.2 \pm 9.3$ mV (mean ± S.D., $n=44$). A continuous stable recording fulfilling the above criterion could be made only in 7
out of 90 cases of RC. In these 7 cases the RC was accompanied by small oscillatory changes in membrane potential (2-5 mV, peak to peak). The synchronization between each contraction and cellular depolarization was almost complete in 2 cases. However, in 5 other cases, only a few individual contractions occurred without accompanying the potential change in the recorded cell although the majority of contractions coincided with small depolarization (Fig. 3). Observations on an oscilloscope revealed that the potential was not associated with rapid action potentials or spikes. Although similar small changes in potential were recorded in other cases, the potentials from these cells were altered also by manual alteration of passive stretch so that they were deleted from analysis.

3. Effects of low temperature

The frequency of RC and the vascular tension were decreased with a reduction in temperature of superfusing solution from 37 to below 30°C (average rate of reduction in temperature, 2°C/min) (Fig. 4). The RC ceased at about 30°C, which was accompanied with a reduction in tension by 45 ± 14% (mean ± S.D., n = 16) of control baseline tension (37°C). In such low temperatures, the RC could not reappear by an artificial restoration of tension due to an increased passive stretch (Fig. 4, dot). This indicated that the reduction in the baseline tension itself did not inhibit the generation of RC but the inhibition of RC by low temperature might
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have reduced the baseline tension. As demonstrated in Fig. 4, small irregular changes in tension (2–8 mg) were always seen in the AC mode registration after the RC had ceased. The amplitude of this remaining small mechanical activity was larger than that of the noise level of the tension measuring system (1 mg). Furthermore, the preparations which were killed by alcohol fixation or severely deteriorated preparations showed neither the RC nor such small irregular mechanical activities. Although we did not analyze it systematically, this small mechanical activity seemed to originate in the vascular tissue. The small irregular activity was relatively refractory to low temperature and was abolished by extremely low temperature (below 10°C) only.

4. Effects of Ca-free solution and Ca-entry blockers

Superfusing the preparation with nominally Ca-free solution resulted in a decrease in the frequency of RC and consequent decline in tension (Fig. 5). The magnitude of reduction in tension by cessation of RC was 34 ± 22% (mean ± S.D., n = 6) of control baseline tension (normal solution). The RC recovered completely after switching the superfusing solution from Ca-free to a normal one. The application of verapamil (average concentration in the incubation chamber, above 5.2 x 10^{-6} M) or nifedipine (average concentration, above 5.8 x 10^{-7} M) also inhibited the generation of RC together with the reduction in tension by about 40% of control tension (Fig. 6). A restoration of RC after washing out the drug, however, took a long time, more than 30 min. A manual increase in stretch tension failed to
evoke the RC in Ca-free condition or during applicating the Ca-entry blocker (Figs. 5 and 6, dot). The inhibition of RC, therefore, seems to have reduced the vascular tension but not vice versa. The small irregular changes in tension, which remained during cessation of RC, were not abolished by prolonged exposure to Ca-free solution or to Ca-entry blockers in high concentrations.

5. Effects of adrenergic blockers

The effects of various adrenergic blockers (phentolamine, prazosin, yohimbine, and propranolol) on the RC were examined. An α-blocker phentolamine, when applied on the preparation in the average concentration of $1.0 \times 10^{-7} - 1.0 \times 10^{-5} \text{M}$, affected neither the frequency and amplitude of RC nor the vascular tension; while phentolamine in concentration, above $1.6 \times 10^{-4} \text{M}$, clearly inhibited the RC and reduced the vascular tension. This inhibitory effect, however, seemed to be due to nonspecific membrane action of the drug in an extremely high concentration. Applications of other adrenergic blockers (prazosin, $1.0 \times 10^{-6} - 1.0 \times 10^{-4} \text{M}$ (Fig. 7); yohimbine, $1.0 \times 10^{-6} - 1.0 \times 10^{-5} \text{M}$; propranolol, $1.0 \times 10^{-6} - 1.0 \times 10^{-5} \text{M}$) also had no obvious effects on the RC and tension. These results suggest that the adrenergic mechanism being activated by endogenous catecholamine is not involved in the initiation of RC in the isolated rat aorta.
In the present experiment spontaneous contractions were observed consistently in the isolated rat aorta even in the absence of external stimuli. This contrasts to the previous observations that rhythmic mechanical activities of large elastic arteries, especially those of the aorta, are seen only occasionally and are brought about by applications of norepinephrine and high potassium or by low Mg solution (BIAMINO and KRUCKENBERG, 1969; ALTURA and ALTURA, 1974; ALTURA et al., 1980). In the present experiment, we recorded not only the baseline tension of vascular preparation with a conventional low gain DC mode but also changes in tension with a high gain amplification of tension signals in the AC mode. As described previously (MASUDA et al., 1982), the latter method of tension registration may be effective in detecting the spontaneous mechanical activity if the amplitude of changes in vascular tension is small. Another possibility to explain the frequent occurrence of RC in the present study may be that the smooth muscle cells of the rat aorta are inherently characteristic of generating spontaneous contraction more than that of other mammalian species. This possibility remains to be verified. In the rat aorta, the smooth muscle cells are arranged diagonally in parallel array, but successive layers are oriented in different directions, i.e., high pitch or almost longitudinal and low pitch or almost transverse arrangements (PEASE, 1962; RHODIN, 1980). Since our preparations were made by transverse section, the RC was presumably generated in the muscle layer that was oriented in more or less transverse direction.

DISCUSSION

Fig. 7. Effect of α-adrenergic blocker (prazosin) on spontaneous rhythmic contraction in the thoracic aorta. The arrow indicates the starting time when the solution containing a given concentration of prazosin (1.0 × 10^{-6} M) was superfused continuously. Prazosin affected neither the frequency nor the amplitude of rhythmic contraction.
The RC was relatively small in amplitude and was accompanied by small oscillatory changes in membrane potential. Since slow potential changes recorded in the present study are small in amplitude and lack spike components, it is uncertain whether they are involved in triggering contraction. If this is the case, the cell to cell conduction of such a small depolarization may not spread widely and hence concomitant recruitment of contracting smooth muscle cells may be limited, initiating a relatively small rhythmic contraction. A complete synchronization between each contraction and slow potential change in the recorded cell was not always seen. This fact suggests that the RC may be initiated by membrane depolarization not only from a single pacemaker but also from a few but multiple pacemaker cells. The frequency of RC was not increased clearly in the static phase of passive stretch although in some cases there was a slight and transient increase in frequency in the dynamic phase of stretch. JOHANSSON and MELLANDER (1975) showed that the portal mesenteric vein exhibited a significant increase in the frequency of spontaneous action potentials only during the dynamic phase of stretch. A rapid stretch of the preparation did not influence the membrane potential in the present study. The lack of effect of stretch may be due to the stretch being applied only for a short period (within 2–3 s) or to the extent of stretch being small, or to both. Since effects of variation in the passive stretch on the activity of vascular smooth muscle are of physiological significance, influences of stretch on the membrane potential of large elastic artery should be more precisely examined in further experiments. The average membrane potential of cell in rat aorta was smaller than that reported in the rabbit large arteries (MEKATA, 1974). We could not, however, find any comparable data for the rat large arteries.

The frequency of RC was decreased by low temperature. Generally, low temperature inhibits the initiation of spontaneous activity in vascular smooth muscles and subsequently reduces the vascular tension in the deep vessels (KEATINGE, 1964; VANHOUTTE and LORENZ, 1970; VANHOUTTE, 1980). The cessation of RC by Ca-free solution or by Ca blocking agents means that the presence of extracellular Ca and/or its entry into the cell is necessary for the generation of spontaneous electrical activities or for the excitation (depolarization)-contraction coupling of smooth muscles. Rhythmic contractions of the rat portal vein and aorta have been shown to be blocked by Ca-free solution or by Ca-entry blockers (GOLENHOFFEN and WESTON, 1976; ALTURA et al., 1980; VANHOUTTE, 1981). Various adrenergic blockers were ineffective in suppressing the spontaneous RC. Furthermore, the aortic preparation isolated from the reserpine-pretreated rat showed also a typical RC (our unpublished observation). Thus the activation of vascular smooth muscles by catecholamines which may be released endogenously from sympathetic nerve terminals, if any, is not essential for the generation of spontaneous RC in the rat aorta. PATIL et al. (1972) described the absence of the adrenergic nerve terminals in the rat aorta.

After the RC had been inhibited by a low temperature (below 30°C) or by Ca-free solution, small irregular changes in tension remained in every preparation. This
small mechanical activity was suppressed by an extremely low temperature (below 10°C) but not by Ca blockers. The nature of this remaining activity is not clear. It may represent a compound form of spontaneous random contractions which occurs in a very localized group of smooth muscle cells because its amplitude was extremely small and its frequency was irregular. There is also a possibility that the small irregular activity represents the mechanical movements of structures other than the smooth muscle, such as the endothelial cells. Recent evidence suggests the ability of vascular endothelial cells to contract (Bevan et al., 1980).

An inhibition of RC by low temperature, Ca-free solution or by Ca blockers reduced the baseline vascular tension, which suggests that the generation of RC may increase the vascular tension in the large artery.

The present results do not provide any evidence that the RC occurs spontaneously in \textit{in situ} preparation and that it influences the vascular hemodynamics in the large artery. The presence of spontaneous contraction \textit{in situ} aorta has been recently shown by Mangel et al. (1981). Possible effects of spontaneous RC on the distensibility of elastic artery (Windkessel vessels) should be examined in further experiments using a more appropriate preparation such as a cylindrical segment of conduit artery.

REFERENCES


