研究成果の刊行に関する一覧表レイアウト

雑誌

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Yanagiya Y, Sato T. Kawada T, Inagaki M, Tatewaki T, Zheng C, Kamiya A, Takaki H, Sugimachi M, Sunagawa K.	Bionic epidural stimulation restores arterial pressure regulation during orthostasis.	J Appl Physiol	97	984-990	2004
Kakinuma Y, Zhang Y, Ando M, Sugiura T, Sato T	Effect of electrical modification of cardiomyocytes on transcriptional activity through 5'AMP-activated protein kinase		44	S435-S438	2004
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佐藤恭子, 山崎文靖, 古野貴志, 佐藤隆幸, 杉浦哲朗, 土居義典	高血圧治療における塩酸ベニジピンと AII 受容体拮抗薬併用療法の検討.	新薬と臨床	53	13-23	2004
佐藤隆幸	神経インターフェイス技法を用いた心不全 治療と術中血圧制御	循環制御	25	348-351	2004
佐藤隆幸	バイオニックラットからの提言:迷 走神経の電気刺激療法	循環制御	25	23-27	2004
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Bionic epidural stimulation restores arterial pressure regulation during orthostasis

Yusuke Yanagiya,^{1,2} Takayuki Sato,^{1,3} Toru Kawada,¹ Masashi Inagaki,¹ Teiji Tatewaki,^{1,4} Can Zheng,^{1,2} Atsunori Kamiya,¹ Hiroshi Takaki,¹ Masaru Sugimachi,¹ and Kenji Sunagawa¹

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Yanagiya, Yusuke, Takayuki Sato, Toru Kawada, Masashi Inagaki, Teiji Tatewaki, Can Zheng, Atsunori Kamiya, Hiroshi Takaki, Masaru Sugimachi, and Kenji Sunagawa. Bionic epidural stimulation restores arterial pressure regulation during orthostasis. J Appl Physiol 97: 984-990, 2004. First published May 7, 2004; 10.1152/japplphysiol.00162.2004.—A bionic baroreflex system (BBS) is a computer-assisted intelligent feedback system to control arterial pressure (AP) for the treatment of baroreflex failure. To apply this system clinically, an appropriate efferent neural (sympathetic vasomotor) interface has to be explored. We examined whether the spinal cord is a candidate site for such interface. In six anesthetized and baroreflex-deafferentiated cats, a multielectrode catheter was inserted into the epidural space to deliver epidural spinal cord stimulation (ESCS). Stepwise changes in ESCS rate revealed a linear correlation between ESCS rate and AP for ESCS rates of 2 pulses/s and above $(r^2, 0.876-0.979; slope, 14.3 \pm 5.8 \text{ mmHg} \cdot \text{pulses}^{-1} \cdot \text{s};$ pressure axis intercept, 35.7 ± 25.9 mmHg). Random changes in ESCS rate with a white noise sequence revealed dynamic transfer function of peripheral effectors. The transfer function resembled a second-order, low-pass filter with a lag time (gain, 16.7 ± 8.3 mmHg·pulses⁻¹·s; natural frequency, 0.022 ± 0.007 Hz; damping coefficient, 2.40 \pm 1.07; lag time, 1.06 \pm 0.41 s). On the basis of the transfer function, we designed an artificial vasomotor center to attenuate hypotension. We evaluated the performance of the BBS against hypotension induced by 60° head-up tilt. In the cats with baroreflex failure, head-up tilt dropped AP by 37 ± 5 mmHg in $5 \cdot 8$ and 59 ± 11 mmHg in 30 s. BBS with optimized feedback parameters attenuated hypotension to 21 \pm 2 mmHg in 5 s (P < 0.05) and 8 \pm 4 mmHg in 30 s (P < 0.05). These results indicate that ESCS-mediated BBS prevents orthostatic hypotension. Because epidural stimulation is a clinically feasible procedure, this BBS can be applied clinically to combat hypotension associated with various pathophysiologies.

baroreceptors; blood pressure; autonomic nervous system; Shy-Drager syndrome; orthostatic hypotension

THE ARTERIAL BAROREFLEX SYSTEM configures a negative feed-back system and reduces arterial pressure (AP) disturbances from external influences (9, 15, 22, 23). Sudden onset of hypotension by orthostatic change occurs as a result of baroreflex failure, despite normal functioning of the cardiovascular system and efferent sympathetic nervous system. This condition is seen in multiple-system atrophy (Shy-Drager syndrome) (21, 22, 30) as well as spinal cord injuries (7, 17). Current treatments, such as salt loading (19, 33), cardiac pacing (1, 14),

and pharmacological interventions (2, 3, 12, 20), fail to prevent the orthostatic hypotension. These therapies often result in an unwanted increase in AP in the supine position and neither restore nor reproduce the function of the feedback system that forms the basis of AP control (See DISCUSSION).

Previously, our laboratory developed a bionic baroreflex system (BBS) that substitutes the defective vasomotor center with an artificial controller (i.e., an artificial vasomotor center) to restore the native baroreflex function (24, 26). In these animal studies, the celiac ganglion was exposed by laparotomy and stimulated directly as the efferent neural interface in the BBS. However, for clinical application of the BBS, a less invasive and more stable electrical stimulation method is required.

In the present study, we examined the hypothesis that the spinal cord is a candidate site for the efferent neural interface in our bionic strategy. Epidural spinal cord stimulation (ESCS) has been used for the management of patients with malignant neoplasm, angina pectoris, and peripheral ischemia (6, 29). Stimulating the dorsal part of the spinal cord changes sympathetic nerve activity, AP in animals (11, 32) and heart rate in humans (18). If we can delineate how ESCS affects AP quantitatively, then this may lead to clinical application of the BBS. We studied the feasibility of ESCS-mediated BBS using an animal model of central baroreflex failure.

MATERIALS AND METHODS

Study design. BBS is a negative feedback system and consists of two components: peripheral effectors and the artificial vasomotor center (Fig. 1). Peripheral effectors change AP in response to ESCS. The artificial vasomotor center (controller) determines the ESCS rate in response to changes in AP. Using BBS, we computer programmed the artificial vasomotor center and substituted the defective vasomotor center with an artificial vasomotor center.

For this purpose, we first characterized the static as well as dynamic responses of the peripheral effectors. With this knowledge, we then designed an artificial vasomotor center using simulation to delineate the parameters for obtaining optimal AP response. Finally, we evaluated the performance of the ESCS-mediated BBS in cats during orthostatic AP changes.

Animals and surgical procedures. Animals were cared for in strict accordance with the "Guiding Principles for the Care and Use of Animals in the Field of Physiological Sciences," approved by the Physiological Society of Japan. Six adult cats of either sex, weighing

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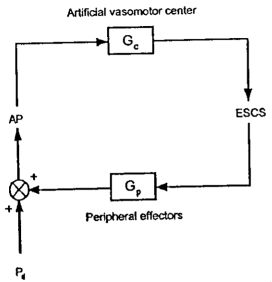


Fig. 1. Simplified diagram of bionic baroreflex system (BBS) using epidural spinal cord stimulation (ESCS). Peripheral effectors change arterial pressure (AP) in response to ESCS. The artificial vasomotor center determines ESCS rate in response to changes in AP. Transfer function of the peripheral effectors (G_p) cannot be controlled, but the vasomotor center transfer function (G_c) can be computer programmed as needed. P_d, pressure disturbance to AP.

1.6-3.7 kg, were premedicated with ketamine (5 mg/kg im) and then anesthetized by intraperitoneal injection (1.0 ml/kg) of a mixture of urethane (250 mg/ml) and α -chloralose (40 mg/ml).

For AP measurement, a high-fidelity pressure transducer (SPC-320, Millar Instruments, Houston, TX) was placed in the aortic arch via the right femoral artery. Pancronium bromide (0.3 mg/kg) was administered to prevent muscular activity. The cats were mechanically ventilated with oxygen-enriched room air. Body temperature was maintained at around 38°C with a heating pad. To produce the baroreflex failure model, the carotid sinus, aortic depressor, and vagal nerves were sectioned bilaterally. The rationale for using baroreceptor-deafferentiated animals as the baroreflex failure model is that, in patients with multiple-system atrophy, the reason for sudden hypotension induced by orthostatic change is a lack of reflex control of AP sensed at the baroreceptors.

A partial laminectomy was performed in the L₃ vertebra to expose the dura mater. A multielectrode catheter with interelectrode distance of 10 mm was introduced rostrally ~7 cm into the epidural space. The stimulating electrodes were positioned on the dorsal surface of the spinal cord within T₁₂, T₁₃, and L₁. These spinal levels were selected because our preliminary studies showed that ESCS to these levels produced greater AP response compared with other spinal levels. The catheter was connected to an isolated electric stimulator (SS-102J and SEN7203, Nihon Kohden, Tokyo, Japan) via a custom-made, constant-voltage amplifier. The stimulator was controlled with a laboratory computer (PC9801FA, NEC, Tokyo, Japan).

Data recording for characterizing AP response to ESCS. To characterize the static as well as dynamic AP responses to ESCS, we measured AP responses while changing ESCS rate. To estimate static response, we changed the ESCS rate sequentially in stepwise increments and decrements. Each stimulation step was maintained for 90 s. To estimate dynamic response, we randomly changed the ESCS rate, according to a binary white noise sequence with a minimum sequence length of 1 s. In both protocols, the stimulation voltage was fixed at 5 V, and the pulse width of the stimulus was 1 ms. While the stimulation was given, ESCS rate and AP were digitized at a rate of 200 Hz with a 12-bit resolution analog-to-digital converter [AD12-8 (PM), Contec, Osaka, Japan] and stored in another laboratory computer

(PC98-NX VA70J, NEC). In this study, "ESCS rate" is defined as the number of stimulation pulses per second and is distinguished from the term "frequency," which refers to how frequently ESCS rate changes.

Estimation of static AP response parameters. We parameterized the static AP response to ESCS. Each steady-state AP value was obtained by averaging the AP during the last 10 s of each ESCS step. Stimulation at low-ESCS rate decreased AP, but stimulation at higher rates increased AP. We fit both responses together to a linear regression model of AP vs. ESCS rate. We determined the rate at which the pressor and depressor effects balanced and designated it the offset stimulation rate (s_o).

Estimation of peripheral effector transfer function. The transfer function of the peripheral effector was estimated by using a white noise method described in detail elsewhere (10, 13, 16, 24-26, 31). Briefly, on the basis of the resampled data at 10 Hz, the linear transfer function from ESCS rate to AP was calculated as a quotient of the ensemble average of cross-power between the two and that of ESCS rate power. The transfer function was calculated up to 0.5 Hz with a resolution of 0.0098 Hz. We parameterized the transfer function by using an iterative, nonlinear, least squares fitting technique (10).

Design of central characteristics and implementation by the artificial vasomotor center. Based on the parameterized effector transfer function, we designed the vasomotor center transfer function using computer simulation. The characteristics of the vasomotor center have been identified as derivative characteristics in rabbits and rats (10, 13, 25). We did not have the corresponding data for cats but assumed that they resembled those in rabbits and rats. We adjusted the parameters by simulation, aiming to attenuate hypotension to ~20 mmHg within 5 s and to \$10 mmHg at 30 s. We implemented designed transfer function by the artificial vasomotor center with convolution algorithm (see APPENDIX).

Head-up tilt tests. The efficacy of the BBS against orthostatic stress was evaluated in each animal by the head-up tilt (HUT) test. We placed baroreflex failure animals in a prone position on a custom-made tilt table and measured AP responses to 60° HUT, with or without BBS activation. In the absence of BBS control, we fixed the ESCS rate at so, irrespective of AP changes. The BBS was activated by sending ESCS command to the stimulator, as calculated by the artificial vasomotor center in response to AP change. Tilt angle, ESCS rate, and AP were stored in a laboratory computer.

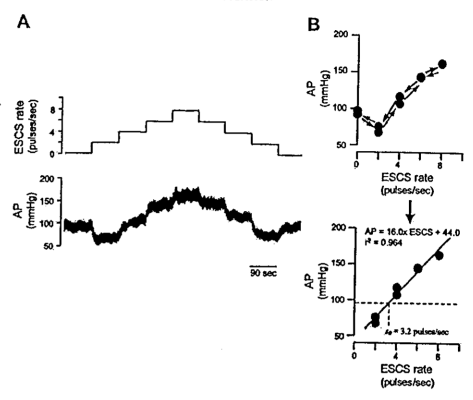
Statistical analysis. All data are presented as means \pm SD. We analyzed AP responses at 5 and 30 s after HUT. AP changes from control value were compared among protocols by using repeated-measures analysis of variance followed by Dunnett's multiple-comparison procedure (8). Differences were considered significant when P < 0.05.

RESULTS

Figure 2A is a representative example of static AP response to stepwise ESCS changes. Stepwise increases of ESCS rate produced a depressor response initially at low-ESCS rate and a pressor response at higher rates, and subsequent decreases of ESCS rate produced almost perfect reversal of AP changes. The relationship between AP and ESCS rate appeared nonlinear as a whole (Fig. 2B, top). However, for ESCS rates of 2 pulses/s and above, there is a linear relationship ($r^2 = 0.964$, AP = $16.0 \times ESCS + 44.0$, $s_0 = 3.2$ pulses/s; Fig. 2B). A linear relationship during ESCS was found in all animals [r^2 , 0.876-0.979 (median, 0.959) slope, 14.3 ± 5.8 mmHg·pulses⁻¹·s; pressure axis intercept, 35.7 ± 25.6 mmHg; s_0 , 4.9 ± 2.5 pulses/s]. The following protocols were performed by using this linear ESCS range.

Figure 3A is a representative example of dynamic AP response to ESCS. We selected low- and high-stimulation rates that produced depressor and pressor responses, respectively,

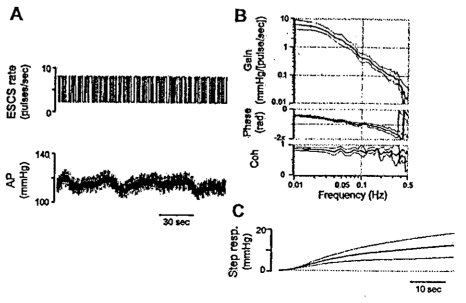
Fig. 2. A: representative time series of static AP response to ESCS. Stepwise increases of ESCS rate produced a depressor response initially at low-ESCS rate and a pressor response at higher rates, and subsequent decreases of ESCS rate produced almost perfect reversal of AP changes. B: procedures to estimate steady-state relationship between AP and ESCS rate. Each steady-state AP value was plotted against ESCS rate (top). Although the relationship between AP and ESCS rate appears nonlinear as a whole, linear regression analysis was conducted for the data obtained during ESCS (2, 4, 6, and 8 Hz), omitting the values before and after ESCS (0 Hz). A linear relationship is obtained. The value so represents the stimulation rate where pressor and depressor responses balance.



and stimulated the spinal cord, according to a binary white noise sequence. AP did not respond to fast changes in ESCS rate, but appeared to respond to slower changes, increasing with high-rate stimulation and decreasing with low-rate stimulation. The estimated transfer function indicated low-pass filter characteristics. Figure 3B shows the averaged transfer function from ESCS rate to AP in six animals. The gain decreased as the frequency increased and was attenuated to one-tenth of the lowest frequency at 0.1 Hz. The phase approached zero radian at the lowest frequency, reflecting in-

phase changes of ESCS rate and AP. The parameters obtained by the least squares fitting to the second-order, low-pass filter model are as follows: dynamic gain = 16.7 ± 8.3 mmHg·pulses⁻¹·s, natural frequency = 0.022 ± 0.007 Hz, damping coefficient = 2.40 ± 1.07 , and lag time = 1.06 ± 0.41 s. The dynamic gain was much higher than the gain at the lowest frequency in the transfer function (Fig. 3.4) but comparable to the slope obtained in the static protocol. The coherence function was close to unity between 0.01 and 0.4 Hz, indicating that the input-output relation was governed by almost linear

Fig. 3. A: representative example of dynamic AP response to ESCS. B: averaged transfer functions from ESCS rate to AP, i.e., G_P (gain and phase) and coherence function (Coh). C: estimated step response (resp) computed from the transfer function. Data are expressed as means \pm SD for 6 cats.



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dynamics in this range. To facilitate better understanding of the dynamic AP response to ESCS, the step functions were calculated by time integral of the inverse Fourier transform of the transfer functions. The estimated step functions are averaged and shown in Fig. 3C. An initial time lag and overdamped slow AP response to unit step ESCS are evident. The time courses of the estimated step functions were almost identical to but smoother than those of the actually observed AP responses to stepwise ESCS changes, indicating the ability to cancel out noises by the white noise method.

Figure 4 is a representative example of how we designed the vasomotor center transfer function. The steady-state gain is determined simply to match a total baroreflex loop gain of 5. This setting ensures that the steady-state AP fall will be attenuated to 10 mmHg in the case of a depressor stimulus of 60 mmHg. By changing the derivative corner frequency (f_c), we simulated various transient AP responses to orthostatic depressor stimulation. Lower f_c causes unstable oscillation in AP (Fig. 4B, left), and higher f_c slows AP restoration (Fig. 4B, right). On the basis of these simulations, $f_c = 0.02$ Hz was selected in this example (Fig. 4B, middle, 0.018 \pm 0.008 Hz in 6 cats).

Figure 5 shows a representative example of real-time application of BBS to a cat. In the control cat with baroreflex failure, abrupt HUT produced a rapid and then progressive fall in AP by 44 mmHg in 30 s (Fig. 5, left). Activation of the simulation-based BBS attenuated the AP fall (Fig. 5, middle; AP fall, 25 mmHg in 5 s and 19 mmHg in 30 s) but did not attain the predetermined target (gaps indicated by vertical bars in Fig. 5, middle and right). If the total loop gain of 5 were preserved, the AP fall should theoretically be attenuated to 44/(1 + 5), or \sim 7 mmHg, according to the linear control theory. The observed attenuation of 19/44 (=1/2.3) indicated that the actual gain was 1.3 (1 + 1.3 = 2.3). To achieve a total loop gain of 5, we increased the gain of the vasomotor center transfer function by

3.8-fold and reassessed the efficacy of BBS. As a result, AP returned to the predetermined target (Fig. 5, right; AP fall, 19 mmHg in 5 s, 7 mmHg in 30 s).

Figure 6 summarizes the results obtained from six cats, demonstrating the effectiveness of the BBS performance. In the cat model of baroreflex failure, HUT decreased AP by 37 \pm 5 mmHg in 5 s and by 59 \pm 11 mmHg in 30 s. In animals with simulation-based vasomotor center, the initial attenuation (AP fall: 32 \pm 7 mmHg in 5 s) was not significant, and the steady-state attenuation (17 \pm 8 mmHg in 30 s) did not satisfy the predetermined target. On the other hand, in animals with gain-adjusted vasomotor center (2.4 \pm 1.1-fold increase), the BBS achieved both initial and the steady-state targets (21 \pm 2 mmHg in 5 s, P < 0.05; 8 \pm 4 mmHg in 30 s, P < 0.05).

DISCUSSION

The present results indicate that AP can be controlled by ESCS and that ESCS-mediated BBS prevents orthostatic hypotension in anesthetized cats. Although the BBS based on simulation alone did not work as predicted during HUT, gain adjustments of the vasomotor center achieved quick and stable restoration of AP.

Necessity of BBS for treatment of central baroreflex failure. Conventional treatments for central baroreflex failure aim at increasing AP. Although they alleviate the hypotension to the extent of preventing syncope, they have adverse effects of causing supine hypertension and enhancing the risk of hypertensive organ disease. Recently, Shannon et al. (28) suggested well-timed water consumption as a treatment for orthostatic hypotension in patients with autonomic failure. Their strategy is superior to conventional treatments because it prevents AP fall if predicted in advance. However, at least a few minutes are necessary for their method to increase AP. This time lag makes it impossible to control AP against sudden or unpredictable AP

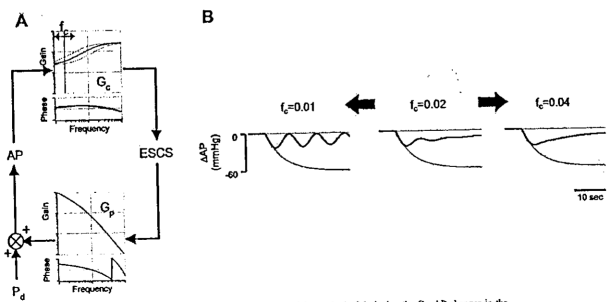


Fig. 4. Representative example of designing the G_c . A: schematic diagram of the method of designing the G_c . AP changes in the presence of P_d were simulated by changing the corner frequency (f_c) for derivative characteristic in the G_c . B: simulation results of this example. A vasornotor center with f_c of 0.02 Hz restores AP with sufficient speed and stability (center).

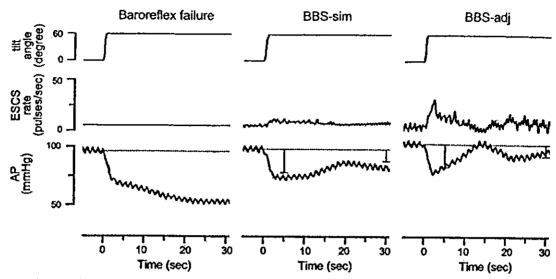


Fig. 5. Representative example of real-time application of the BBS during head-up tilt (HUT). Left: in the cat with baroreflex failure, ESCS rate was fixed at 5.0 pulse/s. HUT produced a rapid and then progressive fall in AP. Middle: simulation-based BBS (BBS-sim) attenuated the AP fall but did not attain the predetermined target. Right: gain adjustment of the vasomotor center (BBS-adj) resulted in quick and sufficient attenuation of AP fall (see text for detail). Vertical bars indicate the ranges of predetermined targets.

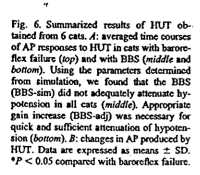
fall. None of the treatments attempted so far can prevent sudden orthostatic hypotension because the dynamics of the baroreflex remain impaired.

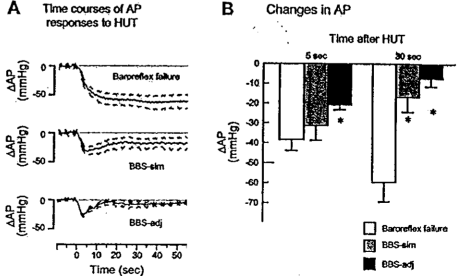
In contrast, the BBS continuously monitors and controls AP to achieve quick restoration of AP. Because AP is increased via sympathetic pathways, the AP response to the BBS vasomotor center command is as fast as that to the native vasomotor center control. Therefore, the quick, adequate, and stable nature of the native baroreflex system can be restored by the BBS with appropriate settings of the artificial vasomotor center.

Designing the vasomotor center transfer function and parameter adjustments. In a negative feedback system, closed-loop responses to external perturbation are dependent on the dynamic characteristics of the total open-loop transfer func-

tion. We assumed that the derivative characteristics in cats are identical to those in rabbits or rats, which have been delineated previously (10, 13, 25). We optimized gain and derivative parameters for each cat so that both speed and stability were achieved.

In animal experiments, however, the simulation results were not fully reproduced. Gain adjustments of the vasomotor center were necessary to attain quick and sufficient attenuation of the AP fall. This discrepancy cannot be explained without considering a possible slope decrease or nonlinearity in peripheral effector characteristics (AP-ESCS relationship) caused by the HUT, or both. Pooling of blood volume in the splanchnic and hindlimb circulation would be a cause for such attenuated AP response. If AP responses during posture change can be de-





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fined quantitatively, then a more elaborate artificial vasomotor center can be developed that automatically adjusts the parameters. Automated adjustments may also be accomplished with adaptive control systems that execute real-time system identification and self-tuning of controller. The present study suggests the necessity of such manipulation of the vasomotor center settings.

Pressor and depressor responses by ESCS. We demonstrated both depressor and pressor responses in AP to ESCS with a linear range (Fig. 2). Earlier studies have shown that dorsal column stimulation produces pressor and depressor responses. Depressor response is produced by a group of dorsal column fibers that project to the dorsal nuclei at the level of C8 to L1 and transmit to the fibers that ascend through the dorsal spinocerebeller tracts (4, 27, 32). Pressor response is produced by another group of fibers that ascends or descends through the terminal zone and enter the gray matter. Some of these fibers project to the intermedial lateral columns to activate sympathetic presynaptic fibers. AP responses to ESCS observed in the present study are the results of compound responses in these multiple pathways. Nevertheless, controllability of AP by the BBS is ascertained by a clear linearity between ESCS and AP response in both static and dynamic relationships.

Although we have not confirmed it in the closed-loop condition, depressor response shown in the open-loop condition indicates that ESCS-mediated BBS can attenuate hypertension as well as hypotension. Because an offset ESCS rate (s_o) is applied in the absence of pressure disturbance, lowering ESCS rate would attenuate hypertension to some extent. AP in conscious animals fluctuates between hypertension and hypotension, even in a quiet position (5). The speed of AP restoration is considered sufficient to control these fluctuations.

Future step for clinical application. Clearly, the next step toward clinical application is to demonstrate the safety and the effectiveness of the ESCS-mediated BBS during orthostasis in conscious patients and animals. To study BBS in conscious animals, we have been developing an implantable hardware that enables BBS. Elaboration of such devices is mandatory for its future clinical application by searching the optimal stimulating site and condition that do not cause uncomfortable sensation and muscle twitch. A control algorithm must be developed that overcomes the problem revealed in the present study. The developing implant would be as small and low power as a pacemaker, with the aid of recent LSI technologies, and would be telemetrically programmable. In parallel, we began collaboration with a clinical group to develop ESCSmediated BBS to suppress sudden hypotension in anesthetized humans during surgery. This study would prove the feasibility

Finally, new methods for long-term manometry are definitely required. Intravascular manometry can be achieved only with long-lasting antithrombotic material. Other indirect methods should be used until we are confident in antithrombotic ability.

Conclusions. As a step toward clinical application of BBS, we demonstrated that AP could be controlled with ESCS. We designed the artificial vasomotor center based on the dynamic characteristics of AP response to ESCS. Although there was a dissociation between the predicted and actual attenuation of AP fall, ESCS-mediated BBS with appropriate gain adjustment

was capable of preventing HUT-induced hypotension rapidly, sufficiently, and stably.

APPENDIX

Simulation of AP Restoration and Implementation by the Artificial Vasomotor Center

We modeled the vasomotor center transfer function (Gc) as

$$G_{c}(f) = K_{c} \frac{1 + \frac{f}{f_{c}i}j}{\left(1 + \frac{f}{f_{c}i}j\right)\left(1 + \frac{f}{f_{c}i}j\right)} \exp(-2\pi f j L_{c})$$
 (A1)

where f is frequency, K_c is the steady-state gain of the artificial vasomotor center, f_{c1} is f_c for derivative characteristics, f_{c2} and f_{c3} are corner frequencies for high-cut characteristics, f is an imaginary unit, and L_c is a pure delay. The value f_{c2} was set to $10 \times f_{c1}$, and f_{c3} was set to 1 Hz. These settings attenuate AP pulsation and preserve total baroreflex gain (13). L_c was introduced to simulate the possible time delay of 0.1 s in transforming AP to ESCS rate but was excluded in real-time application to improve AP stability.

The simulation was performed as follows. The block diagram in Fig. 4A is represented as

$$AP = G_p \cdot ESCS + P_d$$

$$ESCS = G_c \cdot AP$$

where G_p is transfer function of the peripheral effectors, and P_d is pressure disturbance to AP. Rearranging these formula yields

$$AP = G_c \cdot G_p \cdot AP + P_d \tag{A2}$$

The time domain representation of Eq. A2 is

$$\Delta AP(t) = \int g(\tau) \cdot \Delta AP(t-\tau)d\tau + P_{d}(t) \qquad (A3)$$

where $\Delta AP(t)$ is AP change from control value, and $g(\tau)$ is the impulse response function of the total open-loop transfer function $(G_c \cdot G_p)$. To simulate orthostatic hypotension, $P_o(t)$ is set as an exponential AP fall to -60 mmHg with a time constant of 5 s rather than a stepwise fall (see Fig. 6A, top). We simulated the transient AP response to depressor stimulus while changing f_{c1} in the presence of the negative feedback system.

To implement the designed vasomotor center transfer function, we programmed the artificial vasomotor center to calculate ESCS rate in response to AP changes, according to the following equation:

$$ESCS(t) = \int h(\tau) \cdot \Delta AP(t-\tau)d\tau + s_0$$

where $h(\tau)$ is the impulse response function of the designed vasomotor center transfer function, $\Delta AP(t)$ is AP change from the control value, and s_o is the offset ESCS rate obtained from the static parameterization.

GRANTS

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Effect of Electrical Modification of Cardiomyocytes on Transcriptional Activity through 5'-AMP-activated Protein Kinase

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Abstract: Endothelin-1 (ET-1) is known as an aggravating factor of the failing cardiomyocytes and, therefore, a therapeutic method is indispensable to decrease cardiac ET-1 expression. To study the mechanisms of how cardiac ET-1 gene expression can be modified, we investigated the effect of electrical stimulation against cardiomyocytes. Considering the physiology of cardiomyocytes, in vitro cultured cardiomyocytes demonstrate distinctive features from in vivo cardiomyocytes (i.e. the absence of a stretch along with electrical stimulation). In this study, we especially focused on the effect of electrical stimulation. The electrical stimulation reduced the gene expression of ET-1 mRNA in rat primary cultured cardiomyocytes. Furthermore, this effect on the transcriptional modification of ET-1 was also identified in H9c2 cells. Luciferase activity using H9c2 cells was decreased by electrical stimulation in the early phase, suggesting that the attenuation of the ET-1 gene transcription by electrical stimulation should be due to a transcriptional repression. To further investigate a trigger signal involved in the transcriptional repression, phosphorylation of 5'-AMP-activated protein kinase (AMPK) was evaluated. It was revealed that AMPK was phosphorylated in the early phase of electrical stimulation of H9c2 cells as well as in rat primary cultured cardiomyocytes, and that AMPK phosphorylation was followed by ET-1 transcriptional repression, suggesting that electrical stimulation directly regulates AMPK. This study suggests that AMPK activation in cardiomyocytes plays a crucial role in the transcriptional repression of ET-1.

Key Words: endothelin-1, cardiomyocytes, 5'-AMP-activated protein kinase

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The endothelin (ET) system is known to be indispensable for the development of the heart. In the developmental stage, ET-1 exerts the formation of the heart through the receptors endothelin-A and endothelin-B, and the system

exerts the biological function through either autocrine or paracrine fashion. It has been reported that deformity of the heart is produced in the absence of the ET system. Especially, cardiomyocytes have been reported to remarkably express and produce ET-1 in the pathophysiological condition (i.e. the failing heart) compared with the normal heart.1 The mechanisms have been extensively studied of how the failing heart expresses ET-1 in the progression of heart failure; however, our previous study clearly demonstrated one aspect of the mechanisms - impaired cardiac energy metabolism.2 The level of cardiac ET-1 gene expression is dependent on the condition of the cardiomyocytes in vivo; the failing cardiomyocytes produce more ET-1. However, even normal primary cultured cardiomyocytes in vitro could extraordinarily express ET-1 compared with in vivo. The abnormal pattern of cardiac ET-1 gene expression in vitro is also accompanied with a surprising switch of the myosin heavy chain isoform from α to β (a fetal pattern), suggesting that our cultured cardiomyocytes have already biologically changed their character.3 However, this finding might provide us with some therapeutic clue as to how cardiac ET-1 expression can be depressed using cultured cardiomyocytes. If we could obtain some tool to repress cardiac ET-1 gene expression in vitro, it might lead to clarification of one of the therapeutic strategies against heart failure. Consequently we have so far concentrated on searching for ways to decrease cardiac ET-1 gene expression. Among them, we have found several methods to modify the expression using not chemicals or drugs, but physical stimulation.3 In this study, using electrical stimulation (ES) we have successfully inhibited the cardiac ET-1 gene expression, and investigated the mechanisms by which such stimulation causes a depression of the gene expression.

METHODS

Cell Culture of Rat Cardiomyocytes and H9c2 Cells

According to our previous studies, cardiomyocytes were isolated from 2-day-old Wistar-Kyoto rats. 5-Amino-imidazole-4-carboxamide-1-β-D-ribofuranosyl 5-monophosphate (AICAR) was purchased from Sigma (St Louis, MO, U.S.A.), and H9c2 cells were transiently treated by AICAR.

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Electrical Stimulation

We have developed a specific ES device, which is modified to simultaneously provide multi-channels of ES and also to efficiently regulate bidirectional current. Our protocol for ES was performed as follows: 10 V, 10 milliseconds of duration and 4 Hz of frequency.

RNA Isolation and Reverse Transcription-Polymerase Chain Reaction

As previously described, total RNA was isolated, and 1 μg total RNA was reverse-transcribed and used for a polymerase chain reaction (PCR) template. PCR primers were prepared for preproET-1, hypoxia-inducible factor (HIF)-1 α , β -actin, 2,3 and glucose transporter 3.

Luciferase Assay

As previously reported, the 5'-regulatory region of preproET-1 gene was subcloned into a luciferase vector.² The reporter vector was transfected into H9c2 cells by a cationic reagent, Effecten (QIAGEN, Valencia, CA, U.S.A.), according to the manufacture's protocol. Forty-eight hours after transfection, cells were lysed for evaluation of luciferase activity.

Western Blot Analysis

Cells were harvested from dishes by scraping, were washed with phosphate-buffered saline, and cell lysates were mixed with sample buffer. The samples were fractionated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis and transferred onto membranes (Millipore Corp., Bedford, MA, U.S.A.). After transfer into the membrane, they were soaked in blocking buffer. The membranes were incubated with a monoclonal phosphor-5'-AMP-activated protein kinase- α (Thr172) antibody (1:1000; Cell Signaling Technology, Beverly, MA, U.S.A.). After the membranes were washed, horseradish peroxidase-conjugated secondary antibodies (Promega, Madison, WI, U.S.A.) were applied and the signal was detected using an enhanced chemiluminescence system (Amersham, Piscataway, NJ, U.S.A.).

RESULTS

Electrical Stimulation Affects Gene Expression of Rat Cardiomyocytes

To investigate transcriptional regulation of cardiomyocytes, primary cultured cardiomyocytes were subjected to ES. Even with a bi-directional current, cardiomyocytes could not be cultured for ES more than 16 hours. As demonstrated in Fig. 1, ES remarkably decreased gene expression of preproET-1 and HIF-1 α in the cardiomyocytes, compared with non-stimulated cardiomyocytes. However, the mRNA level of β -actin was not decreased by ES. Adversely, gene expression of glucose transporter 3 mRNA was increased by ES. These results suggested that

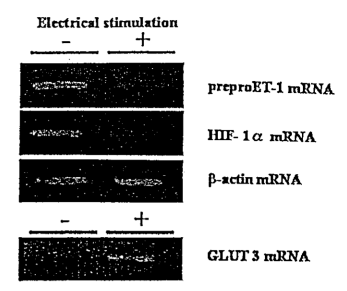


FIGURE 1. Electrical stimulation (ES) modifies gene expression of preproendothelin-1 (preproET-1), hypoxia-inducible factor- 1α (HIF- 1α), and glucose transporter (GLUT) 3 mRNAs. PreproET-1 and HIF- 1α gene expressions were decreased 5 hours after ES. In contrast, GLUT 3 gene expression was increased by ES. The level of β -actin mRNA was not affected by ES.

cardiomyocytes respond to ES with an increased gene expression of glucose transporter 3; however, in contrast, such an ES modified gene expression of cardiac preproET-1 and HIF- 1α .

Electrical Stimulation Transcriptionally Regulates PreproET-1 Gene Expression

Further to investigate especially the gene expression of preproET-1 mRNA, a reporter assay was performed using a reporter vector possessing the 5'-promoter regulatory region of the preproET-1 gene. H9c2 cells, a cell line of rat ventricular cardiomyocytes, were transfected by the reporter vector. As shown in Fig. 2, ES of H9c2 cells greatly decreased the luciferase activity of preproET-1. The phenomenon of the depressed luciferase activity was compatible with the decreased preproET-1 mRNA by reverse transcription-PCR.

Electrical Stimulation Elevates Phosphorylation of 5'-AMP-activated Protein Kinase

To investigate mechanisms to decrease a transcriptional level of preproET-1, we studied whether ES activates the phosphorylation of 5'-AMP-activated protein kinase (AMPK) using H9c2 cells. The time course study demonstrated that AMPK phosphorylation was detected soon after ES (Fig. 3). Furthermore, such an increase in AMPK phosphorylation was also observed in primary cultured cardiomyocytes. It was suggested that ES causes activation of

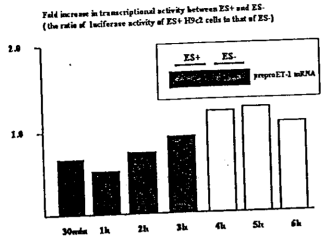


FIGURE 2. Electrical stimulation (ES) transcriptionally represses preproendothelin-1 (preproET-1) gene expression in H9c2 cells. Gene expression of preproET-1 in H9c2 cells was also decreased by ES. Using a reporter vector of the preproET-1 gene, luciferase activity was compared between electrical stimulated (ES+) H9c2 cells and non-electrical stimulated (ES-) cells. The ratio of luciferase activity of ES+ to that of ES- was measured, where a ratio > 1 suggests transcriptional activation, contrasting with a ratio < 1 suggesting repression.

AMPK with a comparable time course of suppression of the preproET-1 gene, and consequently that AMPK phosphorylation is profoundly related to the repression of transcriptional activity of preproET-1 gene expression.

An Activator of AMPK Causes a Decrease in PreproET-1 mRNA

H9c2 cells were treated by AICAR, which is known as an activator of AMPK. As demonstrated in Fig. 3, AICAR increased the phosphorylation of AMPK. The activation of AMPK occurred very rapidly with a comparable time course of ES. With treatment of AICAR, the mRNA level of preproET-1 was decreased (Fig. 4). It was suggested that AMPK activation was involved in the attenuated preproET-1mRNA gene expression.

DISCUSSION

ET-1 is one of the aggravating factors in heart failure, because ET-1 further activates the glycolytic system in the failing cardiomyocytes, resulting in aggravation of malfunction in the heart. Therefore, one of the therapeutic goals that inhibit the progression of heart failure might be to decrease cardiac ET-1 gene expression. There are many manipulations to decrease ET-1 gene expression, including blocking the renin-angiotensin system and ET receptor antagonists. However, we have further investigated whether other manipulations can modify the cardiac ET-1 expression,

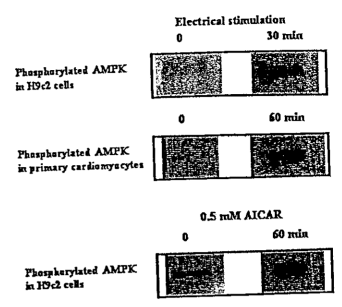


FIGURE 3. 5'-AMP-activated protein kinase (AMPK) is activated through phosphorylation by electrical stimulation (ES). ES caused activation of AMPK through phosphorylation in H9c2 cells. This phenomenon was also detected in rat primary cultured cardiomyocytes with a comparable time course. Also, 0.5 mM 5-aminoimidazole-4-carboxamide-1-β-D-ribofuranosyl 5-monophosphate (AICAR), an activator of AMPK, phosphorylated AMPK.

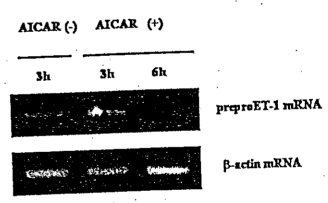


FIGURE 4.. 5'-AMP-activated protein kinase activator decreases preproendothelin-1 (preproET-1) gene expression in H9c2 cells. 5-Aminoimidazole-4-carboxamide-1-β-D-ribofuranosyl 5-monophosphate (AICAR) 1.5 mM treatment remarkably decreased the gene expression of preproET-1 mRNA in H9c2 cells.

and then finally we have identified that a mechanical stimulation (i.e. ES) decreases cardiac ET-1 expression through activation of AMPK.

AMPK is known as a fuel sensor kinase, which is activated through phosphorylation when a cellular adenosine triphosphate (ATP) level is decreased.⁵ Consequently, cells respond to a shortage of the energy and activate the

phosphorylation of AMPK. This kinase is therefore activated in the early phase to stimuli that might cause mitochondrial dysfunction, leading to ATP deprivation. This response through AMPK phosphorylation is considered an adaptation of cells to avoid cellular death with a shortage of ATP. However, this response is not only involved in pathological states, but also in physiological states. For example, exerciseinduced muscle hypertrophy leads to remarkable adaptation of the muscle to the more efficient utilization of energy (i.e. mitochondrial β-oxidation of fatty acid), and consequently to activation of the mitochondrial function.⁶ In the case of exercise, AMPK in skeletal muscle is known to be activated.7 Therefore, it is suggested that the activation of AMPK through phosphorylation is followed by enhancement of mitochondrial function in a physiological condition to obtain more adequate ATP.

It is known that the cardiomyocytes obtain ATP predominantly through mitochondrial B-oxidation of fatty acid; however, when cardiomyocytes were treated by hypoxia, the cardiac energy metabolic system was changed from \(\beta \)-oxidation to glycolysis, because fatty acid oxidation is impaired. In such a case, as our previous study demonstrated, HIF-1\alpha is induced for upregulation of glycolytic enzymes, and furthermore HIF-1\alpha transcriptionally activates preproET-1 gene expression in the failing heart.² It is suggested that cardiac ET-1 expression is accompanied with a cellular glycolysis-dominant energy system.8 With these findings in mind, further speculation is as follows: if the glycolysis-dominant energy system is inhibited, and alternatively mitochondrial \(\beta \)-oxidation of fatty acid is activated, ET-1 gene expression could be decreased. As our present study demonstrated, ES activates the phosphorylation of AMPK in a rapid fashion, followed by a decrease in the transcriptional activity of ET-1.

This is a first demonstration of transcriptional repression of cardiac preproET-1 gene expression using methods other than drugs. Moreover, this reaction is very rapid to decrease preproET-1 mRNA. Therefore, it is suggested that the manipulation of cardiomyocytes by ES is one candidate method to inhibit an increase in cardiac ET-1 gene expression.

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A self-calibrating telemetry system for measurement of ventricular pressure-volume relations in conscious,

freely moving rats Kazunori Uemura, Toru Kawada, Masaru Sugimachi, Can Zheng, Koji Kashihara, Takayuki Sato and Kenji Sunagawa AJP - Heart 287:2906-2913, 2004. doi:10.1152/ajpheart.00035.2004

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A self-calibrating telemetry system for measurement of ventricular pressure-volume relations in conscious, freely moving rats

Kazunori Uemura, ¹ Toru Kawada, ¹ Masaru Sugimachi, ¹ Can Zheng, ^{1,2,3} Koji Kashihara, ^{1,3} Takayuki Sato, ⁴ and Kenji Sunagawa ¹

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Uemura, Kazunori, Toru Kawada, Masaru Sugimachi, Can Zheng, Koji Kashihara, Takayuki Sato, and Kenji Sunagawa. A self-calibrating telemetry system for measurement of ventricular pressure-volume relations in conscious, freely moving rats. Am J Physiol Heart Circ Physiol 287: H2906-H2913, 2004; doi:10.1152/ajpheart.00035.2004.—Using Bluetooth wireless technology, we developed an implantable telemetry system for measurement of the left ventricular pressure-volume relation in conscious, freely moving rats. The telemetry system consisted of a pressure-conductance catheter (1.8-Fr) connected to a small (14-g) fully implantable signal transmitter. To make the system fully telemetric, calibrations such as blood resistivity and parallel conductance were also conducted telemetrically. To estimate blood resistivity, we used four electrodes arranged 0.2 mm apart on the pressure-conductance catheter. To estimate parallel conductance, we used a dual-frequency method. We examined the accuracy of calibrations, stroke volume (SV) measurements, and the reproducibility of the telemetry. The blood resistivity estimated telemetrically agreed with that measured using an ex vivo cuvette method (y = 1.09x - 11.9, $r^2 = 0.88$, n = 10). Parallel conductance estimated by the dual-frequency (2 and 20 kHz) method correlated well with that measured by a conventional saline injection method (y = 1.59x - 1.77, $r^2 = 0.87$, n = 13). The telemetric SV closely correlated with the flowmetric SV during inferior vena cava occlusions (y = 0.96x + 7.5, $r^2 = 0.96$, n = 4). In six conscious rats, differences between the repeated telemetries on different days (3 days apart on average) were reasonably small: 13% for end-diastolic volume, 20% for end-systolic volume, 28% for end-diastolic pressure, and 6% for end-systolic pressure. We conclude that the developed telemetry system enables us to estimate the pressure-volume relation with reasonable accuracy and reproducibility in conscious, untethered

conductance catheter; serial reproducibility; volumetric accuracy; dual-frequency method; Bluetooth

small experimental animals, such as rats and mice, are widely used in cardiovascular research. These animals can offer a variety of disease models, including heart failure and hypertension, and enable us to analyze the molecular mechanisms of the pathophysiology underlying such diseases (5, 7, 12, 21, 27). To interpret the molecular findings in terms of cardiac phenotype, an accurate assessment of cardiac function, including the contractile properties of the left ventricle (LV), is required. As a load-insensitive index of LV contractility, the end-systolic pressure-volume relation (ESPVR) has been estimated in small animal species with the use of a conductance

catheter technique or an ultrasonic crystal method in acute experimental settings (6, 9, 14, 15, 23). However, the anesthesia and thoracotomy required by these techniques inevitably exert adverse effects on the heart (13, 22, 30). In addition, the time course of disease progression or long-term drug effects cannot be assessed in acute experimental settings (7, 16). To overcome these problems, long-term experimental settings should be developed where the LV pressure-volume relation can be measured telemetrically in small experimental animals.

In the present study, we have developed a new telemetry system to measure LV volume, pressure, and electrocardiogram (ECG) in conscious, freely moving rats. In this system. the LV pressure-volume relation was obtained from a pressureconductance catheter chronically implanted in the rat LV. To calibrate the conductance signal and obtain absolute LV volume, measurements of blood resistivity (p) and parallel conductance (G_p) are required (3, 4). These calibration procedures require blood sampling and hypertonic saline infusion, but such ex vivo procedures are not applicable to conscious, freely moving small animals. To circumvent such ex vivo procedures in our new telemetry system (29), we adopted a self-calibrating method for the LV volume measurement, as reported in our previous study (28). The aim of the present study was therefore to develop a telemetry system and evaluate its performance. Our results indicate that we succeeded in measuring the LV pressure-volume relation in conscious, untethered rats with reasonable accuracy and reproducibility.

METHODS

Implantable Pressure-Volume Telemetry System

Figure 1A illustrates a newly developed pressure-volume telemetry system for rats; it consists of a pressure-conductance catheter, an analog processor-transmitter (weight = 14 g, volume = 11 ml), and a battery unit (lithium battery; weight = 12 g, volume = 10 ml).

Pressure-conductance catheter. Details of the pressure-conductance catheter are presented in Fig. 1B. To measure LV conductance, four platinum electrodes (0.25 mm wide) were used. Constant excitation current was applied to the two outermost electrodes while the voltage signal associated with LV conductance was measured from the two inner sensing electrodes. To measure LV pressure, a high-fidelity pressure transducer (Millar Instruments, Houston, TX) was mounted between the two sensing electrodes for the LV conductance measurement. To measure p, four smaller platinum electrodes (0.1 mm wide, 0.2 mm between centers of adjacent electrodes, 0.6 mm

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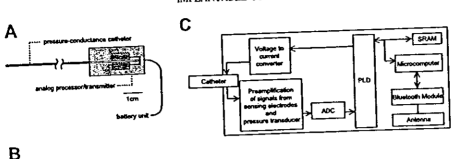


Fig. 1. A: schematic illustration of our pressure-volume telemetry system. A 10-cm-long pressure-conductance catheter obtains signals of left ventricular (LV) conductance and pressure, intraventricular blood resistivity, and an ECG. Signals are processed and transmitted by an analog processor transmitter, which is powered by a battery unit (lithium battery). B: schematic illustration of our pressure-conductance catheter. Catheter has 4 electrodes for measurement of LV conductance and 4 electrodes for measurement of intraventricular blood resistivity (inser). A high-fidelity pressure transducer is mounted between electrodes 2 and 3. C: block diagram of an analog processor transmitter. ADC, analog-to-digital converter, PLD, programmable logic device; SRAM, static random access memory.

between centers of excitation electrodes) were placed near the pressure transducer (Fig. 1B, inset). Constant excitation current was applied to the two outer electrodes while the voltage signal associated with p was measured from the two inner electrodes.

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Analog processor transmitter. A block diagram of the analog processor transmitter is presented in Fig. 1C. It was equipped with several functions. First, it delivered a dual-frequency (2 and 20 kHz) constant excitation current [20 μA root mean square (RMS)] for measurements of LV conductance and p. We validated the current output by injecting it into known resistors and examining the developed voltage. The resulting RMS current output was 20.4 μA (SD 0.2) and 19.3 μA (SD 0.2) at 2 and 20 kHz, respectively. These values were constant over different resistors (50–990 Ω). Second, it mea-

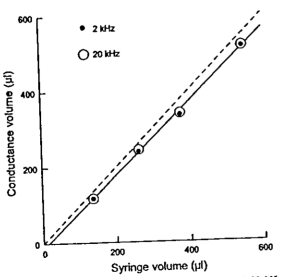


Fig. 2. Comparison of conductance-derived volumes at 2 and 20 kHz vs. known fluid volumes of syringes. Both conductance-derived volumes were essentially identical for each of the syringe volumes. Relation between conductance-derived volume and syringe volume was quite linear. Solid line, regression between conductance-derived volume at 20 kHz and syringe volume; dashed line, identity.

sured and processed the voltage signal from the sensing electrodes as follows: Analog signals were digitized (12 bits, 40-kHz sampling rate; model ADS7870, Texas Instruments, Dallas, TX) and then fed into a programmable logic device (model XC 2C256, Xilinx, San Jose, CA), which processed them to yield RMS digital signals corresponding to frequency components of 2 and 20 kHz and a low-frequency signal (<2 kHz; see APPENDIX). The circuit was connected to the larger or smaller electrodes in response to a command signal, so that LV conductance or p could be measured. Third, the analog processor-transmitter had a bridge amplifier for the LV pressure measurement. The LV pressure signal was also digitized (12 bits, 40-kHz sampling rate). All these functions were controlled by a microcomputer (model H8S, Hitachi, Tokyo, Japan).

Bluetooth technology was used to transmit the data (18). For real-time monitoring, all processed signals were resampled at 200 Hz by the microcomputer and transmitted to an external receiver (CA-SIRA, CSR, Cambridge, UK) by a Bluetooth module (model LMBTB027, Murata, Tokyo, Japan). For high-precision non-real-time analysis, signals recorded at 2,000 Hz over a 6-s interval were stored in a static random access memory (model HM62V16256, Hitachi) and then transmitted to the receiver by the Bluetooth module. The external receiver detected the radio-frequency signal from the transmitter and converted it to a serial bit stream.

Self-Calibration of Ventricular Volumetry

The principles of conductance volumetry have been described previously (3, 4). Briefly, the ventricular conductance signal (G) can be converted to absolute ventricular volume (V) as follows

$$V = (1/\alpha) (L^2 \cdot \rho) (G - G_\rho)$$
 (1)

where α is a volume calibration factor, L is the distance between the sensing electrodes, ρ is blood resistivity, and G_{ρ} is parallel conductance. L was 9 mm in the present catheter design.

In a preliminary experiment, when the catheter was placed in a series of graduated syringes filled with diluted saline, conductance-derived volumes at 2 and 20 kHz were close to the true syringe volume in the volume range of interest (Fig. 2). Conductance-derived volumes at the two frequencies were essentially identical for each of the syringe volumes. Hence, α was assumed to be unity in the present study (14, 23).

The four smaller electrodes were used to estimate ρ (Fig. 1B, inset). The distance between the excitation electrodes was set at 0.6 mm. In an in vitro experiment, we confirmed that the current distribution volume was confined to an ~4-mm diameter around the catheter with this electrode design (see APPENDIX). The end-diastolic LV diameter is 7-9 mm in normal rats and 9-12 mm in rats with left heart failure (17). Because the interelectrode distance between the excitation electrodes was small enough to confine the current distribution volume to within the end-diastolic ventricular blood pool in the rat LV, we estimated ρ at end diastole (10, 28).

 G_p was estimated by the dual-frequency excitation method (8, 9, 28) as follows

$$G_{p} = \kappa \times \Delta G_{20-2} \tag{2}$$

where ΔG_{20-2} is the difference in ventricular conductance values between the 20- and 2-kHz excitation frequencies and κ is an experimentally derived constant. Once κ is determined, $G_{\rm p}$ can be estimated from ΔG_{20-2} , obviating the need for saline infusion.

Instrumentation and Experimental Protocols

Thirty-three male Sprague-Dawley rats (350-400 g body wt) were used. Care of the animals was in strict accordance with the Guiding Principles for the Care and Use of Animals in the Field of Physiological Society as approved by the Physiological Society of Japan. The animals were anesthetized with pentobarbital sodium (50 mg/kg ip) and ventilated artificially. A vertical midline cervical incision was made to expose the right common carotid artery while the animal was in the supine position. The pressure-conductance catheter of the telemetry system was inserted into the LV retrogradely from the right common carotid artery. The position of the catheter was verified by monitoring the pressure-volume signal and by two-dimensional echocardiography. At the conclusion of the experiment, the animal was killed with an overdose of pentobarbital sodium, and the heart was examined to reconfirm the proper positioning of the catheter.

Group 1 (n = 23). We evaluated the accuracy of telemetric calibration of ρ and G_{ρ} under an esthetized, closed-chest conditions. Catheters (3-Fr) were inserted into the right and left jugular veins for blood sampling and saline injection, respectively. In 10 of the 23 rats, we compared ρ estimated telemetrically (ρ_{est}) with ρ measured from sampled blood by a conventional ex vivo cuvette method (pconv). In the remaining 13 rats, we estimated G_p by the dual-frequency excitation method $(G_{p,con})$ and by the hypertonic saline method $(G_{p,con})$. To obtain Gp.com, we injected 20 µl of saturated saline into the right jugular vein while continuously measuring LV conductance (14, 23). To obtain G_{p,est}, we measured LV conductance at 2- and 20-kHz excitation frequencies and derived ΔG_{20-2} by averaging the instantaneous conductance difference over ~10 cardiac cycles. We randomly selected 7 of the 13 rats and determined the proportionality constant (κ in Eq. 2) from the averaged ratio of $G_{p,conv}$ to ΔG_{20-2} . $G_{p,conv}$ were measured while the artificial ventilation was temporarily suspended at end expiration.

Group 2 (n = 4). Under anesthetized, open-chest conditions, we evaluated the accuracy of volumetry by comparing stroke volume (SV) measured by the telemetry system with SV measured by an ultrasonic flowmeter (model 2.5S273, Transonic Systems, Ithaca, NY). After median sternotomy, the aortic arch was dissected free from surrounding tissues. A flow probe was placed around the ascending aorta to measure the aortic blood flow. A string occluder was placed loosely around the inferior vena cava to decrease the LV preload and vary the SV over a wide range. We simultaneously measured the telemetric LV volume and the ultrasonic aortic blood flow while varying the preload. The measurements were done while the artificial ventilation was temporarily suspended at end expiration.

Group 3 (n = 6). Under conscious, closed-chest conditions, we evaluated the reproducibility of the telemetry on different days. Aseptic conditions were maintained throughout the surgical proce-

dure. The telemetry system was implanted in a subcutaneous pocket made at the right upper dorsum. The skin was closed, and the animal was allowed to recover from anesthesia. On the day after implantation surgery, the LV volume, pressure, and an ECG were measured telemetrically in the fully recovered, conscious animal (study 1). Each rat underwent a second set of telemetric measurements at 1-6 days after the initial study (study 2). Ambient barometric pressure was measured simultaneously and subtracted from the telemetric LV pressure to compensate for changes in atmospheric pressure.

Data Collection

We used the real-time mode (200-Hz sampling) of the telemetry system and recorded LV conductance, LV pressure, intraventricular ECG, and p on a hard disk of a dedicated laboratory computer system (model HFPA031003, Epson, Tokyo, Japan). In group 2, ultrasonic aortic blood flow was digitized at 1,000 Hz through a 12-bit analog-to-digital converter and stored on a hard disk for subsequent analyses.

Statistical Analysis

For the calculation of LV volume using Eq. 1, G and ρ were obtained from the 20-kHz frequency component. In group 1, we used linear regression analysis to compare the telemetric and conventional measurements of ρ (ρ_{est} vs. ρ_{conv}) and G_{ρ} ($G_{p,est}$ vs. $G_{p,conv}$). In group 2, we calculated the telemetric SV from the difference between the end-diastolic volume (EDV) and end-systolic volume (ESV) in each beat. The flowmetric SV was computed from the time integral of aortic blood flow. The telemetric SV was compared with the flowmetric SV by linear regression analysis. In group 3, we compared heart rate (HR), EDV, ESV, end-diastolic pressure (EDP), and endsystolic pressure (ESP) between study 1 and study 2 for each rat. Using the pressure-volume data, we calculated ejection fraction (EF), maximal pressure increase (+dP/dt_{max}) or decrease (-dP/dt_{max}) over time, and the time constant of isovolumic relaxation (7) and compared them between study I and study 2 for each rat. A nonparametric multiple comparison (Wilcoxon's signed-rank test) was used to examine the difference in each parameter between study 1 and study 2. Group data are expressed as means (SD). Differences were considered significant at P < 0.05.

RESULTS

Telemetric Calibration of p

Figure 3A is a representative time series showing LV pressure and ρ at 2 and 20 kHz derived from the telemetry. The bottom of the ρ waveform, which corresponded to end diastole, represents the time when there was sufficient blood volume around the catheter (10). The lowest ρ values at 2 kHz ($\rho_{2 \text{ kHz}}$) and 20 kHz ($\rho_{20 \text{ kHz}}$) were very close (197 and 207 Ω ·cm, respectively). This was the case for all the rats, indicating that ρ was frequency independent ($\rho_{2 \text{ kHz}} = 1.08 \rho_{20 \text{ kHz}} - 13.8$, $r^2 = 0.96$, SE of the estimate $= 6.7 \Omega$ ·cm) (6, 9). The lowest ρ at 20 kHz was treated as ρ_{est} .

Figure 3B summarizes the relation between $\rho_{\rm est}$ and $\rho_{\rm conv}$ obtained from 10 rats in group 1. $\rho_{\rm est}$ agreed with $\rho_{\rm conv}$ reasonably well ($\rho_{\rm est}=1.09\rho_{\rm conv}-11.9$, $r^2=0.88$, SE of the estimate = 10.7 $\Omega\cdot$ cm). The ratio of SE of the estimate to the mean of $\rho_{\rm est}$ was 0.046, indicating small variability around the regression line.

Telemetric Calibration of G.

Figure 4A illustrates a representative time series of telemetrically measured ECG, LV conductance signals at 2 and 20 kHz, and LV pressure. In this animal, ΔG_{20-2} was 0.56 mS

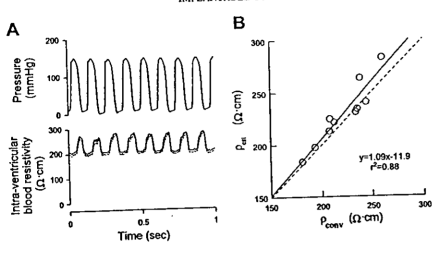


Fig. 3. A: waveforms of ventricular pressure and intraventricular blood resistivity at 2 kHz (dashed line) and 20 kHz (solid line) as a function of time obtained telemetrically. B: relation between blood resistivity as measured in a cuvette (ρ_{conv}) and as estimated via catheter electrodes (ρ_{conv}) in 10 rats. Solid line, regression; dashed line, identity.

and $G_{p,conv}$ was 3.27 mS. Therefore, κ was calculated to be 5.79 from Eq. 2 in this animal. The averaged κ from seven randomly selected rats was 5.14, which we used as the experimentally derived constant to obtain $G_{p,est}$ for all rats.

Figure 4B summarizes the relation between $G_{p,est}$ and $G_{p,conv}$ obtained from 13 rats in group 1. $G_{p,est}$ correlated well with $G_{p,conv}$ ($G_{p,est} = 1.59G_{p,conv} - 1.77$, $r^2 = 0.87$, SE of the estimate = 0.33 mS). The ratio of SE of the estimate to the mean of $G_{p,est}$ was 0.11, indicating that the estimation was reasonable around the mean of $G_{p,est}$.

Accuracy of the Televolumetry

Figure 5A depicts LV pressure and volume measured by telemetry and aortic blood flow measured by the ultrasonic flowmeter. Vena caval occlusion decreased LV pressure, volume, and aortic blood flow.

Figure 5B summarizes the relation between telemetric SV (SV_{Bow}) and flowmetric SV (SV_{Bow}) obtained from four rats in group 2. SV_{Ele} matched SV_{Bow} reasonably well in each of the four rats: $r^2 = 0.90 - 0.99$, slope = 0.86 (SD 0.16), intercept = 12.4 μ l (SD 10.4), and SE of the estimate = 4.3 μ l (SD 0.4). A linear regression analysis on the pooled data from all four rats also showed a highly linear relation between SV_{Ele} and

SV_{flow}: SV_{tele} = 0.96SV_{flow} + 7.5, r^2 = 0.96, SE of the estimate = $6.6 \mu l$. The ratio of SE of the estimate to the mean of SV_{tele} was 0.10.

Reproducibility of the Telemetry

Individual data obtained by the telemetry system for all six rats in group 3 are provided in Tables 1 and 2. The overall variability between repeated measurements in the same rat was reasonably small. There were no significant differences in repeated measurements of HR, EDV, ESV, EDP, and ESP between study 1 and study 2 (Table 1). There were no significant differences in repeated measurements of EF, +dP/dt_{max}, -dP/dt_{max}, and τ between study 1 and study 2 (Table 2).

Figure 6 illustrates the representative LV pressure-volume loops obtained from a rat in group 3. The pressure-volume loops in studies 1 and 2 were almost identical.

DISCUSSION

We have developed a novel telemetry system for measurements of LV volume, pressure, and ECG in conscious, freely moving rats. The system, for the first time to the best of our knowledge, has enabled measurement of the LV pressure-

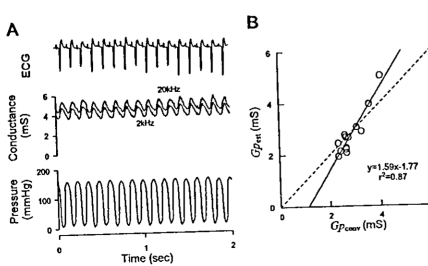
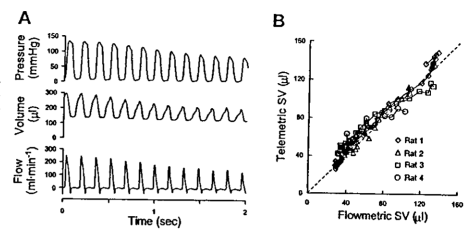


Fig. 4. A: waveforms of an ECG, conductance signals at 2 and 20 kHz, and ventricular pressure as a function of time, obtained telemetrically. B: relation between parallel conductance estimated by the saline infusion method $(G_{p,com})$ and by dual-frequency excitation method $(G_{p,col})$ in 13 rats. Solid line, regression; dashed line, identity.

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Fig. 5. A: representative traces of ventricular pressure, ventricular volume obtained telemetrically, and aortic flow measured by an ultrasonic flowmeter during vena cava occlusion in 1 rat. B: relation between telemetric stroke volume (SV) and flowmetric SV in 4 rats. Dashed line, identity.



volume relation in small experimental animals, such as rats, under completely conscious, unrestricted conditions with reasonably good accuracy and reproducibility.

Self-Calibrating Volumetry

In our conductance volumetric system, ρ and G_p were estimated using the telemetric signals alone (Figs. 3 and 4). We will be able to use the empirical constant κ (=5.14), determined in this study, in the future application of our telemetry system to rats. The self-calibrating feature made it possible to measure the LV pressure-volume relation in rats without tethering them for ex vivo calibration procedures, such as blood sampling and hypertonic saline infusion. Besides their impracticality in conscious, small animals, these procedures can alter hemodynamic conditions (6, 9). Frequent blood sampling can induce anemia. Concentrated saline injection depresses myocardial contractility and has volume-loading effects (6, 9). Our telemetry system is free of these problems.

The current used for resistivity measurements was distributed in a 2-mm radius around the catheter (see APPENDIX). The ratio of the radius (i.e., penetration depth) to the distance between the excitation electrodes was $\sim 3 \ (\rightleftharpoons 2/0.6)$. This ratio is at odds with previously reported values, which were around or less than unity (6, 10, 26). Penetration depth is affected by the relation between the resistivity of the target tissue and that of the surrounding structure (26). This relation in our study was

different from those in previous studies, which would be one reason for the discrepancy. Difference in shape and arrangement of the electrodes between our system and those previous studies would be another reason. Because the electrodes were placed very closely, stray capacitance between connecting wires could be a problem (31). The fact that resistivity values at 2 and 20 kHz were very close indicated that our titration method effectively removed the problem of stray capacitance (see APPENDIX). However, it might be better to incorporate techniques such as capacitance neutralization to completely prevent the problem, in case the capacitance were to significantly affect our titration accuracy in future long-term use, e.g., with increases in electrode impedance (31).

We used the dual-frequency excitation method previously described by Gawne et al. (8). Feldman et al. (6) combined measured resistivity of the myocardium with an analytic approach and estimated G_p from the conductance signals at 10 and 100 kHz. Although their method was completely independent of saline injection, it required measurement of myocardial resistivity with an additional four-electrode sensor.

Volumetric Accuracy and Reproducibility

We have verified the volumetric accuracy of our telemetry system by comparing SV_{tele} with SV_{flow} during inferior vena cava occlusions (Fig. 5A). The volumetric accuracy of the conductance catheter technique in the rat heart has been ex-

Table 1. Reproducibility of hemodynamic variables

Rat	HR, beats/min		EDV, μΙ		ESV, µI		EDP, mmHg		ESP, modify	
	SI	\$2	\$1	S 2	SI	S2	SI	\$2	SI	\$2
1	530	475	303	307	168	182	14	19	131	123
2	363	476	310	280	212	151	11	17	121	133
3	426	500	330	355	212	220	12	9	127	129
4	405	511	244	194	143	89	14	13	120	125
5	402	382	364	434	269	326	15	14	119	106
6	400	380	240	283	158	167	24	36	142	152
Mean (SD)	421 (57.3)	454 (58.2)	299 (49)	309 (81)	194 (47)	189 (80)	15 (5)	18 (10)		
Difference	,		()	007 (01)	471 (477	107 (80)	13 (3)	10 (10)	127 (9)	128 (15)
Mean (SD)	65 (40)		37 (23)		34 (26)		5 (4)		9.76	
Percent difference			2. (23)		34 (20)		3 (4)		8 (4)	
Mean (SD)	15	(9)	13 ((8)	20.6	(17)	28	(16)		(4)

S1, study 1; S2, study 2; HR, heart rate; EDV, lest ventricular end-diastolic volume; ESV, lest ventricular end-systolic volume; EDP, lest ventricular end-diastolic pressure; ESP, lest ventricular end-systolic pressure.