

No.	発表者氏名	論文タイトル名	発表誌名	巻号	ページ	出版年
168	Izawa KP, Watanabe S, Oka K, Osada N, <u>Omiya K</u>	Relation of the three-component model of short form-36 scores to disease severity in chronic heart failure outpatients.	International Journal of Cardiology	157	130-131	2012
169	Izawa KP, Watanabe S, Tochimoto S, Osada N, <u>Omiya K</u> , et al.	Relation between maximum phonation time and exercise capacity in chronic heart failure patients.	European Journal of Physical and Rehabilitation Medicine	48	1-7	2012
170	Izawa KP, Watanabe S, Oka K, Osada N, <u>Omiya K</u> , et al.	Upper and lower extremity muscle strength levels associated with an exercise capacity of 5 metabolic equivalents in male patients with heart failure.	Journal of Cardiopulmonary Rehabilitation and Prevention	32	85-91	2012
171	Uematsu M, Akashi YJ, Ashikaga K, Yoneyama K, Kida K, Suzuki K, <u>Omiya K</u> , Miyake F, et al.	Association between heart rate at rest and myocardial perfusion in patients with acute myocardial infarction undergoing cardiac rehabilitation – a pilot study.	Archives of Medical Science	4	622-630	2012
172	Izawa KP, Watanabe S, Oka K, Hiraki K, Morio Y, Kasahara Y, Takeichi N, Tsukamoto T, Osada N, <u>Omiya K</u>	Relation between physical activity and exercise capacity of >5 metabolic equivalents in middle- and older-aged patients with chronic heart failure.	Disability & Rehabilitation	34 (22-23)	2018-2024	2012
173	Izawa KP, Watanabe S, Hiraki K, Morio Y, Kasahara Y, Takeichi N, Oka K, Osada N, <u>Omiya K</u>	Determination of effectiveness of accelerometer use in the promotion of physical activity in cardiac patients: A randomized controlled trial.	Archives of Physical Medicine and Rehabilitation	93	1896-1902	2012
174	Itoh H, Ajisaka R, Koike A, Makita S, <u>Omiya K</u> , Kato Y, et al.	Heart rate and blood pressure response to ramp exercise and exercise capacity in relation to age, gender, and mode of exercise in a healthy population.	Journal of Cardiology	61	71-78	2013
175	西崎真里, 三河内弘, 他	肺高血圧症患者に対する心臓リハビリテーション	心臓	44	274-278	2012
176	安藤可織, 西崎真里, 他	経皮的肺動脈形成術を施行した慢性血栓塞栓性肺高血圧症患者に対する心臓リハビリテーション	心臓リハビリテーション(JJCR)	17(2)	261-265	2012
177	渡利 太, 横山茂樹, 西崎真里 等	心臓血管外科手術後患者の肺活量の回復に関する要因	Japanese Journal of Health Promotion and Physical Therapy	2(3)	113-118	2012
178	安藤可織, 西崎真里, 他	当院での肺高血圧症患者における退院時の運動処方	Therapeutic Research	33(10)	16-18	2012
179	安藤可織, 西崎真里, 他	肺高血圧症患者に対する呼吸筋トレーニングの有用性	心臓リハビリテーション(JJCR)	18(1)	124-129	2013

No.	発表者氏名	論文タイトル名	発表誌名	巻号	ページ	出版年
180	木村 穰	運動プログラムの効果と実際 動脈硬化における運動療法の臨床的検討	臨床スポーツ医学	28	1365-1370	2011
181	Tamura T, Mizukura I, Sekine M, <u>Kimura Y</u>	Monitoring and evaluation of blood pressure changes with a home healthcare system.	IEEE Trans Inf Technol Biomed.	15	602-607	2011
182	Satoshi Kurose, <u>Yutaka Kimura</u> , et al.	Improvement in endothelial function by lifestyle modification focused on exercise training is associated with insulin resistance in severely obese patients.	Obesity Research		Epub ahead of print	2012
183	木村 穰	肥満症治療チームに必要な行動変容理論と各構成要員の役割	肥満研究	18(2)	78-84	2012
184	木村 穰	生活改善を継続するための効果的なサポート	糖尿病ケア	9(8)	56-60	2012
185	木村 穰, 岩坂潤二	CKD 患者の心臓リハビリテーション効果とコメディカルの役割	心臓リハビリテーション (JJCR)	17(1)	33-36	2012
186	Ikeda N, <u>Yasu T</u> , Tsuboi K, Sugawara Y, Kubo N, Umemoto T, Arao K, Kawakami M, Momomura S.	Effects of submaximal exercise on blood rheology and sympathetic nerve activity.	Circ J	74	730-734	2010
187	Arao K, <u>Yasu T</u> , Ohmura N, Tsukamoto Y, Murata M, Kubo N, Umemoto T, Ikeda N, Ako J, Ishikawa S, Kawakami M, Momomura S.	Circulating CD34+/133+ progenitor cells in patients with stable angina pectoris undergoing percutaneous coronary intervention.	Circ J	74	1929-1935	2010
188	<u>安 隆則</u>	運動療法のメカニズム	Vascular Lab	7	89-93	2010
189	<u>安 隆則</u>	ガイドラインを読み解く TASC II	心臓リハビリテーション(JJCR)	15	86-88	2010
190	<u>安 隆則</u>	末梢動脈疾患に対する運動の効果	医学のあゆみ	232	842-846	2010
191	<u>安 隆則</u>	PAD 油断できない下肢の痛み	Heart View	15	132-135	2011
192	Sakima H, Isa K, <u>Yasu T</u> , Ohya Y	Recurrent embolic stroke due to nonbacterial thrombotic endocarditis followed by transesophageal echocardiography.	Arch Neurol	68	1604-1605	2011
193	Hoshina M, Wada H, Sakakura K, Kubo N, Ikeda N, Sugawara Y, <u>Yasu T</u> , Ako J, Momomura S	Determinants of progression of aortic valve stenosis and outcome of adverse events in hemodialysis patients.	J Cardiol	59	78-83	2012

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194	安 隆則	末梢動脈疾患患者のリハビリテーション	日本下肢救済・足病学会誌	4	113-116	2012
195	安 隆則	PADの心血管リハビリテーション	Heart	9	880-885	2012
196	安 隆則	C反応性蛋白	Heart View	16	153-156	2012
197	福永浩幸, 坂本浩樹, 榎木大介, 椎原香美, 堀千奈美, 鶴川俊洋	当院心臓血管外科術後におけるリハビリテーションの現状～ICU担当理学療法士の立場から～	鹿児島リハビリテーション 医学研究会会誌	22	45-48	2011
198	榎木大介, 山野朋博, 橋本睦美, 椎原香美, 鶴川俊洋	脳卒中急性期における再発予防を目標とした運動療法指導	鹿児島リハビリテーション 医学研究会会誌	23(1)	37-41	2012
199	鶴川俊洋	身体活動が少なかったために典型的な胸部症状を認めなかった大動脈弁閉鎖不全症の術後心臓リハビリテーション (症例報告)	鹿児島市医報	51(9)	23-29	2012

## V. 研究成果の刊行物・別刷

「IV. 研究成果一覧表」のうち、○を付けた刊行物の別刷りを掲載



## Efficacy of Out-Patient Cardiac Rehabilitation in Low Prognostic Risk Patients After Acute Myocardial Infarction in Primary Intervention Era

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**Background:** The efficacy of out-patient cardiac rehabilitation (OPCR) in patients with a low prognostic risk after acute myocardial infarction (AMI) is unclear in the recent primary intervention era.

**Methods and Results:** A total of 637 AMI patients who participated in in-hospital cardiac rehabilitation were divided into 2 groups; low prognostic risk group (n=219; age <65 years, successful reperfusion, Killip class I, peak serum creatine kinase <6,000 U/L, and left ventricular ejection fraction  $\geq$ 40%) and non-low prognostic risk group (n=418). The prevalence of coronary risk factors (CRF) was compared between the 2 groups. Then, in the low-risk group, the efficacy of OPCR was compared between active OPCR participants (n=52;  $\geq$ 20 sessions/3 months) and non-active participants (n=60; <6 sessions/3 months). Compared with the non-low prognostic risk group, the low prognostic risk group had a significantly higher prevalence of current smokers (72% vs. 49%,  $P<0.05$ ) and patients with multiple CRF (3 or more; 49% vs. 39%,  $P<0.05$ ). Among the low-risk group, active OPCR participants showed a significantly greater improvement in exercise capacity (peak  $\dot{V}O_2$ ,  $P<0.05$ ) and maintained a better CRF profile (total cholesterol, triglyceride and blood pressure, all  $P<0.05$ ) than inactive participants at 3 months.

**Conclusions:** Low prognostic risk AMI patients have a higher prevalence of multiple CRF than non-low risk patients. Even in this low risk group, active participation in OPCR is associated with improved exercise capacity and better CRF profile. (*Circ J* 2011; **75**: 315–321)

**Key Words:** Acute myocardial infarction; Cardiac rehabilitation; Coronary risk factors; Exercise capacity; Low prognostic risk

Cardiac rehabilitation (CR) is a comprehensive intervention including medically supervised exercise training, risk factor control, patient education, and psychosocial counseling. CR has been reported to be effective in improving numerous intermediate endpoints, including exertional ischemic symptoms, overall feelings of wellness, exercise tolerance, and coronary risk factors (CRF) in patients with coronary artery disease (CAD).<sup>1–6</sup> In addition, recent meta-analyses of randomized studies on the effects of exercise-based CR in patients with CAD have demonstrated a statistically significant reduction in total and cardiac mortality ranging from 20% to 32%<sup>7–9</sup> in patients undergoing CR compared with those receiving standard medical care. The guidelines from the American College of Cardiology/American Heart Association and Japanese Circulation Society recommend the use of CR after acute myocardial infar-

tion (AMI) as Class I.<sup>10–14</sup>

Recently, the widespread use of primary percutaneous coronary interventions (PCI) has enabled early ambulation of patients with AMI by reducing acute phase complications, resulting in minimal physical deconditioning. As a result, many AMI patients leave a hospital early without participating in a recovery phase (phase II) out-patient CR (OPCR) program.<sup>15</sup> However, the necessity and efficacy of OPCR remain unclear in AMI patients who are anticipated to be at low risk in terms of long-term prognosis (ie, non-elderly, successful reperfusion, absence of heart failure, and preserved left ventricular (LV) systolic function).

Accordingly, the purpose of the present study was to clarify the prevalence of CRF and to determine the efficacy of a 3-month OPCR program in such presumably low prognostic risk patients after AMI.

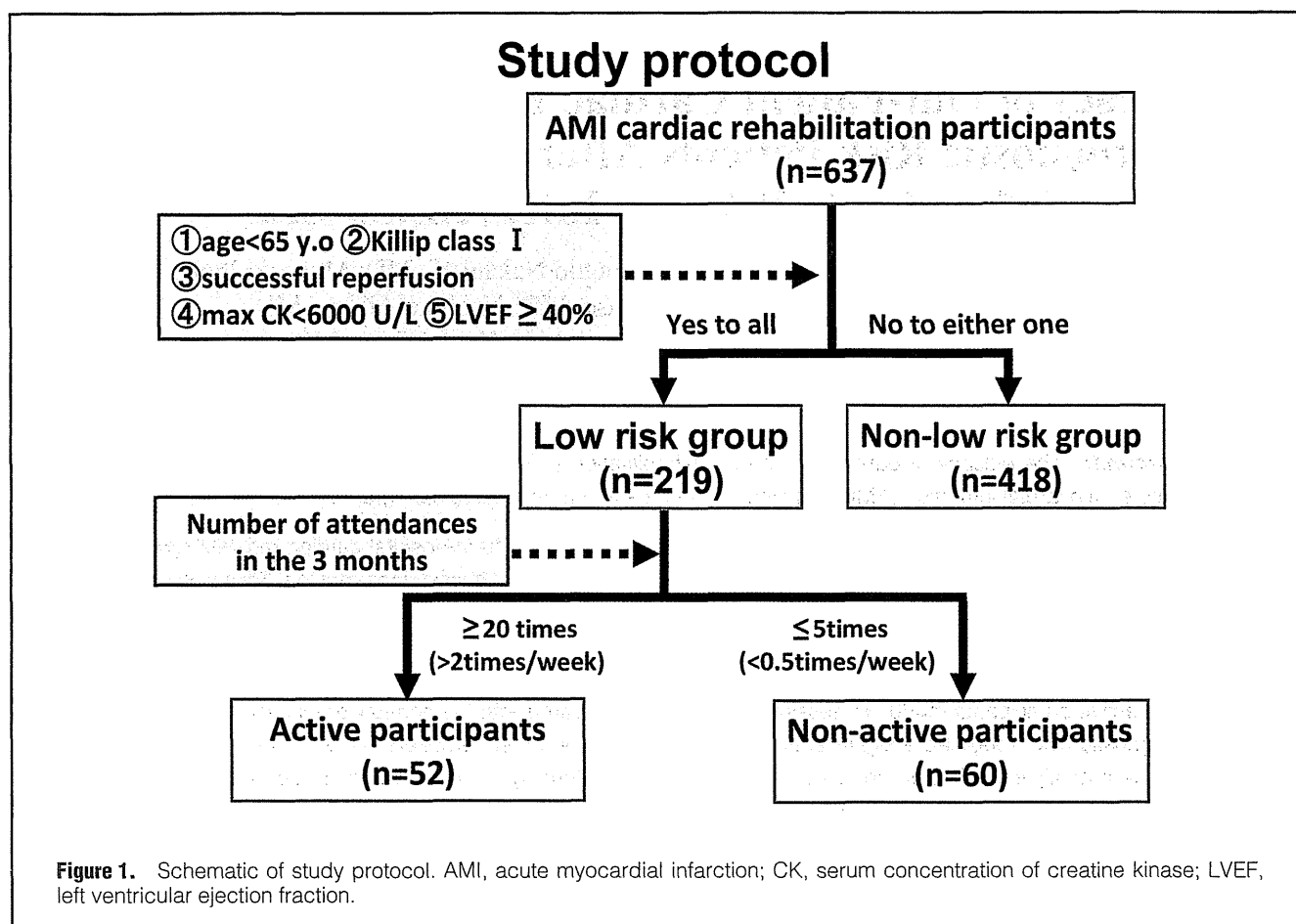
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## Methods

### Patients

We studied a total of 637 consecutive patients with AMI who participated in a recovery phase CR program and underwent cardiopulmonary exercise testing (CPX) at the beginning and end of a 3-month program in our hospital. The patients were divided into 2 groups: a low prognostic risk group and a non-low prognostic risk group. The low prognostic risk group comprised of 219 patients who fulfilled all of the following criteria indicative of favorable prognosis; age under 65 years, successful reperfusion, Killip class I (an indicator of absence of acute phase heart failure), peak serum creatine kinase (CK) <6,000 U/L, LV ejection fraction (LVEF)  $\geq 40\%$ . The remaining 417 patients who did not fulfill 1 or more of the above 5 criteria were referred to as the non-low prognostic risk group.

As the first step of data analysis, the prevalence each of the CRF (hypertension, hyperlipidemia, diabetes mellitus, obesity and smoking habit) was compared between the low prognostic group and the non-low prognostic group.

As the second step, the efficacy of OPCR in AMI patients at low prognostic risk was examined by comparing the data for exercise capacity and CRF between active participants and non-active participants in the low prognostic risk group. Active participants were defined as patients who attended the OPCR sessions at least 20 times in 3 months (ie, approximately >2times/week), and non-active participants were those who attended OPCR less than 6 times in 3 months (ie, approximately <0.5times/week). There were 52 active participants and 60 non-active participants in the low prognos-

tic group. We did not include the remaining 107 patients with intermediate attendance (patients with 6–19 attendances in 3 months) in the analysis, because the effect of OPCR in this patient group was considered to be modest, if any, and inclusion of this group in the analysis would dilute the measurable efficacy of OPCR. A schematic of the study protocol is provided in **Figure 1**.

### CR Program

The CR program began approximately 1 week after AMI and continued after hospital discharge for 3 months. Patients who had angina or evidence of ischemic changes in their electrocardiogram (ECG) at a low level of exercise (walking test), uncontrolled heart failure, and serious arrhythmia were excluded. Program components included supervised exercise sessions (walking, bicycle ergometer and calisthenics) and education, as previously described.<sup>16,17</sup> The exercise intensity was determined individually at 50–60% of heart rate reserve (Karvonen's equation,  $k=0.5-0.6$ )<sup>18,19</sup> or a heart rate of anaerobic threshold (AT) level obtained in a maximal symptom-limited CPX testing or at level 12–13 ('a little hard') of the 6–20 scale perceived rating of exercise (original Borg's scale).<sup>20</sup> The exercise program was started with supervised sessions for 2 weeks, followed by home exercise combined with once or twice-a-week supervised sessions for the remaining 10 weeks. Home exercise consisted mainly of brisk walking at a prescribed heart rate for 30 to 60 min, 3–5 times a week.

Patients were encouraged to attend the education classes that were held 4 times a week with lectures on CAD, secondary prevention, diet, smoking cessation, medication, and

	Low-risk group (n=219)	Non-low-risk group (n=418)	P value
Age (years)	55±7	65±9	<0.01
Male (%)	88	83	NS
Killip class ≥II (%)	0	13	<0.01
Peak CK (U/L)	2,458±1,444	3,339±2,639	<0.01
CK ≥6,000 U/L (%)	0	17	<0.001
Unsuccessful reperfusion (%)	0	24	<0.001
LVEF (%)	49.1±6.8	44.4±10.4	<0.01
LVEF <40% (%)	0	34	<0.001
BNP (pg/ml)	75.7±70.9	209.8±202.0	<0.001
HT (%)	57	56	NS
DM/IGT (%)	47	42	NS
HLP (%)	59	49	<0.05
Obesity (%)	28	27	NS
Smoking habit (%)	72	49	<0.001
Coronary risk factors ≥3 (%)	49	39	<0.05

CK, serum concentration of creatine kinase; LVEF, left ventricular ejection fraction; BNP, brain natriuretic peptide; HT, hypertension; DM, diabetes mellitus; IGT, impaired glucose tolerance; HLP, hyperlipidemia. Values are mean±SD.

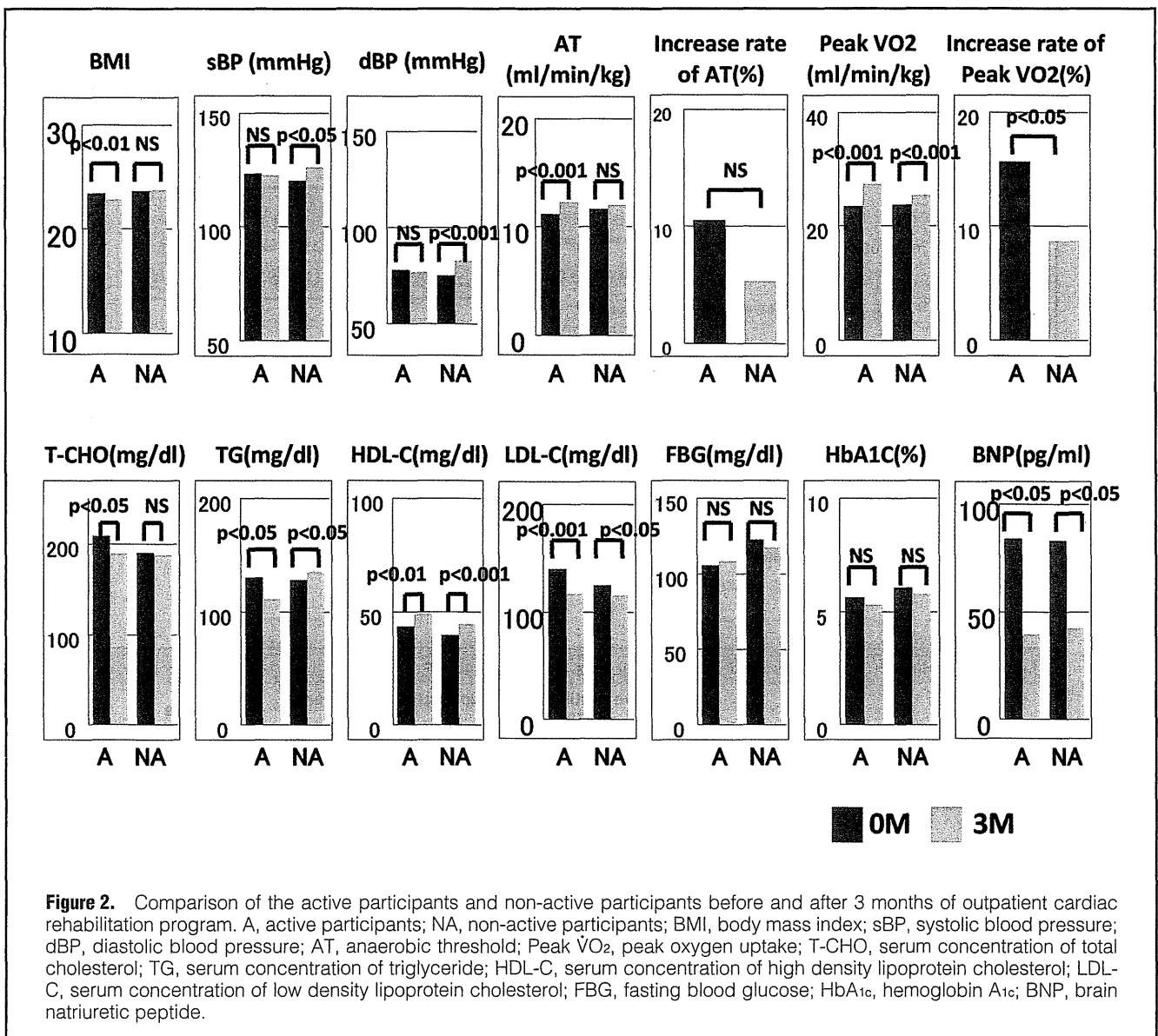
	Active participants (n=52)	Non-active participants (n=60)	P value
Age (years)	57.0±7.3	52.8±7.0	<0.01
Male (%)	83	95	<0.001
Peak CK (U/L)	2,361.1±1,264.2	2,419.5±1,357.1	NS
LVEF (%)	51.4±7.5	47.4±5.7	<0.01
BNP (pg/ml)	83.7±106.0	82.8±74.8	NS
OPCR attendance (times/3 months)	25.5±5.1	1.3±1.7	<0.001
HT (%)	58	52	NS
DM/IGT (%)	44	52	NS
HLP (%)	58	58	NS
Obesity (%)	29	30	NS
Smoking habit (%)	56	75	<0.05
ACE-I/ARB (%)	42	52	NS
β-blocker (%)	19	43	<0.01
Ca channel blocker (%)	40	40	NS
DM medications (%)	8	15	NS
Statin (%)	44	43	NS
Rest HR (/min)	72.6±10.8	71.5±14.9	NS
Rest sBP (mmHg)	123.1±20.2	119.8±21.0	NS
Rest dBP (mmHg)	77.7±11.0	74.7±12.0	NS
Peak WR (W)	132.3±25.2	136.0±31.3	NS
AT (ml·min <sup>-1</sup> ·kg <sup>-1</sup> )	11.1±2.5	11.6±2.6	NS
Peak $\dot{V}O_2$ (ml·min <sup>-1</sup> ·kg <sup>-1</sup> )	23.4±4.2	23.6±5.0	NS
Peak $\dot{V}O_2$ (%predict)	78.5±14.5	73.7±14.3	NS

Values are mean±SD.

OPCR, outpatient cardiac rehabilitation; ACE-I, angiotensin converting enzyme inhibitor; ARB, angiotensin II receptor blocker; HR, heart rate; sBP, systolic blood pressure; dBP, diastolic blood pressure; WR, work rate; AT, anaerobic threshold; Peak  $\dot{V}O_2$ , peak oxygen uptake. Other abbreviations see in Table 1.

physical activities given by physicians, nurses, dieticians, pharmacists and exercise instructors. In addition, all patients received individual counseling on exercise prescription, secondary prevention, and daily life activities by a physician and a nurse at the time of hospital discharge and the end of

the 3-month CR program. Patients were scheduled to undergo blood tests at the beginning and the end of the 3-month CR program.



**Figure 2.** Comparison of the active participants and non-active participants before and after 3 months of outpatient cardiac rehabilitation program. A, active participants; NA, non-active participants; BMI, body mass index; sBP, systolic blood pressure; dBP, diastolic blood pressure; AT, anaerobic threshold; Peak  $\dot{V}O_2$ , peak oxygen uptake; T-CHO, serum concentration of total cholesterol; TG, serum concentration of triglyceride; HDL-C, serum concentration of high density lipoprotein cholesterol; LDL-C, serum concentration of low density lipoprotein cholesterol; FBG, fasting blood glucose; HbA<sub>1c</sub>, hemoglobin A<sub>1c</sub>; BNP, brain natriuretic peptide.

## CPX

Patients were scheduled to undergo a symptom-limited CPX at the beginning and the end of the 3-month CR program.<sup>21</sup> After a 2-min rest on the bicycle ergometer in the upright position, the patients started pedaling at an intensity of 0 W for 1 min (warm-up), and then performed an incremental exercise test with a ramp protocol (10 or 15 W/min) until exhaustion. Twelve-lead ECG was continuously monitored and blood pressure (BP) was measured once-a-min with a sphygmomanometer. Expired gas was collected and analyzed continuously with an AE-300S gas analyzer (Minato Co, Osaka, Japan). Peak oxygen uptake (peak  $\dot{V}O_2$ ) was defined as the highest  $\dot{V}O_2$  value achieved at peak exercise. Ventilation ( $\dot{V}E$ ) and carbon dioxide output ( $\dot{V}CO_2$ ) were measured and the  $\dot{V}O_2$  value at AT or ventilatory threshold was determined as the point at which  $\dot{V}CO_2$  increased in a non-linear fashion relative to the rate of  $\dot{V}O_2$  (according to the  $\dot{V}E/\dot{V}O_2$  time trend, the respiratory exchange ratio flexion point, or the V-slope method).<sup>19,22</sup>

## Statistical Analysis

Baseline characteristics between the 2 groups were compared

using unpaired t-test and chi-square test. Data at baseline and after the 3-month OPCR were compared by paired t-test. A P-value less than 0.05 was considered statistically significant. Data are presented as the mean  $\pm$  standard deviation.

## Results

### Prevalences of CRF in Low Prognostic Risk Group vs. Non-Low Prognostic Risk Group

Clinical characteristics in the low prognostic risk group and the non-low prognostic risk group are summarized in Table 1. Compared with the non-low prognostic risk group, the low prognostic risk group was on average significantly younger, and did not have heart failure on admission or unsuccessful reperfusion, but had lower peak CK and B-type natriuretic peptide (BNP) concentrations and preserved LVEF. Although these findings were anticipated by the definition of the group, they reconfirm that the patients in the low prognostic group were undoubtedly at low prognostic risk. However, when the prevalence of CRF was compared between the 2 groups, the percentage of patients with dyslipidemia, smoking habit and multiple CRF (equal to or more



than 3) was significantly higher in the low prognostic risk group than in the non-low prognostic risk group.

### Efficacy of OPCR in Low Prognostic Risk Group: Comparison Between Active and Non-Active Participants

Baseline characteristics in active participants and non-active participants in the low prognostic risk group are summarized in **Table 2**. Although active participants were significantly older than the non-active participants, they were both non-elderly (less than 65 years old). Peak CK was low and LVEF was relatively preserved in both groups. These findings reconfirm that both active and non-active participants are apparently at low prognostic risk. Although there were minor differences in the prevalence of male patients, smokers and  $\beta$ -blocker use, there were no significant differences in exercise capacities at baseline between the 2 groups.

During the 3-month OPCR period, only a few patients experienced changes in medication; statins were introduced in 3 patients (5.8%) in the active participants and 2 patients (3.3%) in the non-active participants, and diabetic medications were started in 2 patients (3.3%) in the non-active participants. Thus, the baseline clinical characteristics of active and non-active participants were almost equivalent, except for the frequency of OPCR attendance.

**Figure 2** depicts comparisons of parameters before and after the 3-month OPCR between active and non-active participants in the low prognostic risk group. After the 3-month OPCR, only active participants, and not the non-active participants, showed significant improvements in body mass index (BMI;  $23.3 \pm 2.5$  to  $22.9 \pm 2.5$ ,  $P < 0.01$ ), AT ( $11.1 \pm 2.5$  to  $12.7 \pm 2.5$   $\text{ml} \cdot \text{min}^{-1} \cdot \text{kg}^{-1}$ ,  $P < 0.001$ ), total cholesterol ( $208.4 \pm 33.7$  to  $188.8 \pm 26.4$   $\text{mg/dl}$ ,  $P < 0.05$ ), and triglyceride ( $130.0 \pm 77.4$  to  $111.0 \pm 63.7$   $\text{mg/dl}$ ,  $P < 0.05$ ). In addition, while peak  $\dot{V}O_2$  increased in both groups (active participants  $23.4 \pm 4.2$  to  $27.3 \pm 5.0$   $\text{ml} \cdot \text{min}^{-1} \cdot \text{kg}^{-1}$ ,  $P < 0.001$ ; non-active participants  $23.7 \pm 5.0$  to  $25.3 \pm 5.3$   $\text{ml} \cdot \text{min}^{-1} \cdot \text{kg}^{-1}$ ,  $P < 0.001$ ), the magnitude of the increase was significantly greater in the active participants (15.6% vs. 8.6%,  $P < 0.05$ ). In contrast, only non-active participants showed significant worsening in systolic and diastolic BP (systolic BP: from  $119.8 \pm 21.0$  to  $126.1 \pm 20.4$   $\text{mmHg}$ ,  $P < 0.05$ , diastolic BP: from  $74.7 \pm 12.0$  to  $82.4 \pm 11.8$   $\text{mmHg}$ ,  $P < 0.001$ ) and triglyceride ( $128.0 \pm 57.1$  to  $135.3 \pm 63.9$   $\text{mg/dl}$ ,  $P < 0.05$ ). The following parameters showed significant improvements both in the active and non-active participants; high density lipoprotein cholesterol (HDL-C:  $43.6 \pm 14.1$  to  $49.0 \pm 12.2$   $\text{mg/dl}$ ,  $P < 0.01$ ;  $39.7 \pm 11.0$  to  $44.8 \pm 11.6$   $\text{mg/dl}$ ,  $P < 0.001$ ), low density lipoprotein cholesterol (LDL-C:  $140.1 \pm 31.9$  to  $117.6 \pm 25.9$   $\text{mg/dl}$ ,  $P < 0.001$ ;  $124.8 \pm 31.2$  to  $115.3 \pm 19.7$   $\text{mg/dl}$ ,  $P < 0.01$ ), and BNP ( $83.7 \pm 106.0$  to  $39.7 \pm 44.8$   $\text{pg/ml}$ ,  $P < 0.05$ ;  $82.9 \pm 74.8$  to  $42.4 \pm 51.7$   $\text{pg/ml}$ ,  $P < 0.05$ ).

## Discussion

The major findings of the present study are that the low prognostic risk AMI patients had a higher prevalence of smoking habit, dyslipidemia and multiple CRF than the non-low prognostic risk patients, and that in the low prognostic risk group, active participation in OPCR was associated with better CRF profile (ie, BP, dyslipidemia, and obesity) and exercise capacity. These findings suggest that, by actively participating in OPCR after AMI, even the low prognostic risk patients might gain clinical benefits such as better CRF modification and physical functioning.

## Previous Studies

Various guidelines for management of post-AMI (or established CAD) patients recommend aggressive modifications of CRF for secondary prevention,<sup>10,12,13</sup> and adherence to these recommendations and/or reduction of CRF have been shown to improve long-term prognosis.<sup>23–26</sup> In contrast, Thrombolysis In Myocardial Infarction (TIMI) risk score<sup>27</sup> and Controlled Abciximab and Device Investigation to Lower Late Angioplasty Complications (CADILLAC) risk score<sup>28</sup> have demonstrated that 1-year mortality is very low in AMI patients with age  $< 65$  years, successful reperfusion, absence of acute phase heart failure, and preserved LV function, which are compatible with the patient characteristics of the low prognostic risk group in the present study. However, little is known about the prevalence of CRF or clinical significance of accumulation of multiple CRF in such low prognostic risk patients. In relation to this, it is of note that, Lloyd-Jones and colleagues demonstrated that young subjects with accumulated CRF, despite low short-term risk, have a higher 'lifetime risks for CAD' and greater progression of subclinical coronary atherosclerosis compared with those at low lifetime risk.<sup>29,30</sup> These data suggest that apparently low prognostic risk patients stratified by TIMI or CADILLAC risk score are likely to have superb short-term (1 year) prognosis, but not necessarily favorable long-term or lifetime prognosis.

## Present Study

The present study has explicitly demonstrated that the low prognostic risk patients actually have higher prevalence of multiple CRF than the non-low prognostic risk patients. Although the finding that younger AMI patients have higher prevalences of smoking and hyperlipidemia than elderly patients is in accordance with previous studies,<sup>31</sup> there has been no report demonstrating higher prevalence of multiple CRF in low prognostic risk AMI patients with successful reperfusion and preserved LVEF. According to TIMI risk score<sup>27</sup> or CADILLAC risk score,<sup>28</sup> this finding might appear confusing or counterintuitive. However, from the viewpoint of lifetime CAD risk,<sup>29,30</sup> this finding might have a significant impact on the long-term prognosis of apparently low prognostic risk AMI patients.

The second major finding in the current study is that active participation in OPCR improved CRF (BP, dyslipidemia, and obesity) and exercise capacity even in the low prognostic risk group. There have been no studies that reported the effect of OPCR in the low prognostic risk AMI patients. Taylor et al<sup>9</sup> reported in a meta-analysis of randomized controlled trials that the effect of OPCR on total mortality did not differ between studies before and after year 1995 (odds ratio 0.84 before 1995 vs. 0.62 after 1995, NS), but they did not assess the effect of OPCR on the low prognostic risk patients after successful reperfusion. Witt et al recently reported that participation in OPCR after AMI was associated with improved survival and reduced recurrent myocardial infarction (MI) at 3 years, but the rate of reperfusion was only 33% in their patients.<sup>32</sup> Squires et al reported that a 3-year coronary disease management program in OPCR for CAD patients was effective in achieving the secondary prevention goals, but their assessment did not target the low prognostic risk patients.<sup>33</sup> Thus, the present study has demonstrated for the first time the favorable effects of OPCR on CRF and exercise capacity in the low prognostic AMI patients.

### Clinical Implications

It remains unknown whether the improvements in CRF profiles and exercise capacity achieved by active participation in OPCR can lead to an improved long term prognosis in the low prognostic risk AMI patients. However, Tani et al reported that successful life style modification with exercise, body weight reduction and smoking cessation for 6 months was associated with coronary plaque volume regression in low prognostic risk CAD patients.<sup>34</sup> Belardinelli et al reported in the ETICA (Exercise Training Intervention after Coronary Angioplasty) trial that a 6-month OPCR for the relatively low risk CAD patients after successful PCI (49% having AMI) reduced cardiac events and hospital re-admission during the follow-up period (33±7 months).<sup>35</sup> In addition, because the magnitude of the improvement in endothelial function afforded by OPCR does not correlate with the improvements in CRF,<sup>36</sup> the general consensus at present is that the favorable effect of OPCR on the long-term prognosis is mediated by a direct anti-atherosclerosis effect of exercise training rather than by improvements in CRF.<sup>4</sup> Therefore, further study is necessary to determine the long term effect of OPCR in AMI patients with low prognostic risk.

In the present study, significant differences were found between active and inactive OPCR participants in BMI, total cholesterol, triglyceride and BP, but not in LDL-C or glucose tolerance. One might argue that the prognostic impacts of BMI, total cholesterol, triglyceride and BP might be less powerful compared with those of LDL-C and diabetes. However, Nakatani et al reported that the metabolic syndrome, diagnosed from the combination of BMI, HDL-C, triglyceride, BP, and fasting blood glucose, was an independent predictor of subsequent combined cardiac events of cardiac death and non-fatal MI in Japanese patients after AMI.<sup>37</sup> Therefore, it is plausible that the improvements in BMI, triglyceride and BP observed in the present study might contribute to the improvement in the long-term prognosis in Japanese AMI patients.

### Future Direction

In the present study, the rate of active OPCR participation was only 24% (52/219 patients) in the low prognostic risk group. To reduce lifetime CAD risk in these low prognostic risk AMI patients, a substantial increase in participation rate in OPCR is necessary. However, according to a recent nation-wide survey in 526 Japanese Circulation Society authorized cardiology training hospitals,<sup>15</sup> the implementation rate was 92% for emergency PCI, but only 9% for OPCR. In addition, Ades et al reported that, by multivariate analysis, the strength of the physician's recommendation for participation was the most powerful predictor of OPCR participation.<sup>38</sup> Thus, to increase the participation rate in OPCR, it is critically important to greatly increase the number of CR facilities and to enhance physicians' understanding of the benefits of OPCR after AMI.

### Study Limitations

First, this study was a retrospective analysis and the number of patients was relatively small. The more active patients would be expected to participate in OPCR and this might have introduced a selection bias.

Second, the low prognostic risk group is anticipated to be at low risk in terms of short-term prognosis<sup>27,28</sup> and hence, whether improvements in CRF profile in such low prognostic risk patients are associated with actual improvements in outcome is uncertain. A longer follow-up in a larger number

of patients is necessary to increase the statistical power to demonstrate the beneficial effect of OPCR on the long-term prognosis.

### Conclusions

The low prognostic risk AMI patients have a higher prevalence of multiple CRF than the non-low risk patients. Active participation in OPCR program is associated with improved exercise capacity and CRF profile in such low prognostic risk patients. OPCR program can be effective in achieving secondary prevention goals even in the low prognostic risk AMI patients.

### Disclosure

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### References

- Ades P. Cardiac rehabilitation and secondary prevention of coronary heart disease. *N Engl J Med* 2001; **345**: 892–901.
- Wenger NK, Froelicher ES, Smith LK, Ades PA, Berra K, Blumenthal JA, et al. Cardiac rehabilitation as secondary prevention: Agency for Health Care Policy and Research and National Heart, Lung, and Blood Institute. *Clin Pract Guidel Quick Ref Guide Clin* 1995; **17**: 1–23.
- Iwanaga Y, Nishi I, Ono K, Takagi S, Tsutsumi Y, Ozaki M, et al. Angiotensin-converting enzyme genotype is not associated with exercise capacity or the training effect of cardiac rehabilitation in patients after acute myocardial infarction. *Circ J* 2005; **69**: 1315–1319.
- Leon AS, Franklin BA, Costa F, Balady GJ, Berra KA, Stewart KJ, et al; AHA Scientific Statement. Cardiac rehabilitation and secondary prevention of coronary heart disease. *Circulation* 2005; **111**: 369–376.
- Myers J, Prakash M, Froelicher V, Do D, Partington S, Atwood JE. Exercise capacity and mortality among men referred for exercise testing. *N Engl J Med* 2002; **346**: 793–801.
- Jegier A, Jegier A, Szmigielska M, Bilinska M, Brodowski L, Galaszek M, et al. Health-related quality of life in patients with coronary heart disease after residential vs ambulatory cardiac rehabilitation. *Circ J* 2009; **73**: 476–483.
- Graham I, Atar D, Borch-Johnsen K, Boysen G, Cifkova R, Dallongeville J, et al. European guidelines on cardiovascular disease prevention in clinical practice: Full text. Fourth joint task force of the European society of cardiology and other societies on cardiovascular disease prevention in clinical practice (constituted by representatives of nine societies and by invited experts). *Eur J Cardiovasc Prev Rehabil* 2007; **14**(Suppl 2): S1–S113.
- Stone JA, Arthur HM; Canadian Association of Cardiac Rehabilitation Guidelines Writing Group. Canadian guidelines for cardiac rehabilitation and cardiovascular disease prevention, second edition, 2004: Executive summary. *Can J Cardiol* 2005; **21**(Suppl D): 3D–19D.
- Taylor RS, Brown A, Ebrahim S, Jolliffe J, Noorani H, Rees K, et al. Exercise-based rehabilitation for patients with coronary heart disease: Systematic review and meta-analysis of randomized controlled trials. *Am J Med* 2004; **116**: 682–692.
- Antman EM, Hand M, Armstrong PW, Bates ER, Green LA, Halasyamani LK, et al: 2007 focused update of the ACC/AHA 2004 Guidelines for the Management of Patients With ST-Elevation Myocardial Infarction: A report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines (Writing Group to Review New Evidence and Update the ACC/AHA 2004 Guidelines for the Management of Patients With ST-Elevation Myocardial Infarction). *Circulation* 2008; **117**: 296–329.
- Mosca L, Banka CL, Benjamin EJ, Berra K, Bushnell C, Dolor RJ, et al. Evidence-Based Guidelines for Cardiovascular Disease Prevention in Women: 2007 Update. *Circulation* 2007; **115**: 1481–1501. Available at: <http://circ.ahajournals.org/cgi/reprint/CIRCULATIONAHA.107.181546> (accessed August 11, 2010).
- Smith SC, Allen J, Blair SN, Bonow RO, Brass LM, Fonarow GC, et al. AHA/ACC guidelines for secondary prevention for patients with coronary and other atherosclerotic vascular disease: 2006

- update. *Circulation* 2006; **113**: 2363–2372.
13. Takano T, Ogawa S, Kasanuki H, Kimura K, Goto Y, Sumiyoshi T, et al. Guidelines for the management of patients with ST-elevation myocardial infarction. *Circ J* 2008; **72**(Suppl IV): 1347–1442 (in Japanese).
  14. Nohara R, Adachi H, Itoh H, Ueshima K, Katagiri T, Kawakubo K, et al. Guidelines for rehabilitation in patients with cardiovascular disease (JCS2007) (in Japanese). Available at: [http://www.j-circ.or.jp/guideline/pdf/JCS2007\\_nohara\\_h.pdf](http://www.j-circ.or.jp/guideline/pdf/JCS2007_nohara_h.pdf) (accessed August 11, 2010).
  15. Goto Y, Saito M, Iwasaka T, Daida H, Kohzaki M, Ueshima K, et al. Poor implementation of cardiac rehabilitation despite broad dissemination of coronary interventions for acute myocardial infarction in Japan: A nationwide survey. *Circ J* 2007; **71**: 173–179.
  16. Takagi S, Sakuragi S, Baba T, Takaki H, Aihara N, Yasumura Y, et al. Predictors of left ventricular remodeling in patients with acute myocardial infarction participating in cardiac rehabilitation: Brain natriuretic peptide and anterior infarction. *Circ J* 2004; **68**: 214–219.
  17. Suzuki S, Takaki H, Yasumura Y, Sakuragi S, Takagi S, Tsutsumi Y, et al. Assessment of quality of life with 5 different scales in patients participating in comprehensive cardiac rehabilitation after acute myocardial infarction. *Circ J* 2005; **69**: 1527–1534.
  18. Karvonen M, Kentala K, Mustala O. The effects of training on heart rate: A longitudinal study. *Ann Med Exp Biol Fenn* 1957; **35**: 307–315.
  19. Fletcher GF, Balady GJ, Amsterdam EA, Chaitman B, Eckel R, Fleg J, et al. Exercise standards for testing and training: A statement for Healthcare Professionals From the American Heart Association. *Circulation* 2001; **104**: 1694–1740.
  20. Borg G. Perceived exertion as an indicator of somatic stress. *Scand J Rehabil Med* 1970; **2**: 92–98.
  21. Nishi I, Noguchi T, Furuichi S, Iwanaga Y, Kim J, Ohya H, et al. Are cardiac events during exercise therapy for heart failure predictable from the baseline variables? *Circ J* 2007; **71**: 1035–1039.
  22. Beaver WL, Wasserman K, Whipp BJ. A new method for detecting anaerobic threshold by gas exchange. *J Appl Physiol* 1986; **60**: 2020–2027.
  23. Newby KL, LaPointe NMA, Chen AY, Kramer JM, Hammill BG, DeLong ER, et al. Long-term adherence to evidence-based secondary prevention therapies in coronary artery disease. *Circulation* 2006; **113**: 203–212.
  24. Rasmussen JN, Chong A, Alter DA. Relationship between adherence to evidence-based pharmacotherapy and long-term mortality after acute myocardial infarction. *JAMA* 2007; **297**: 177–186.
  25. Ford ES, Ajani UA, Croft JB, Critchley JA, Labarthe DR, Kottke TE, et al. Explaining the decrease in U.S. deaths from coronary disease, 1980–2000. *N Engl J Med* 2007; **356**: 2388–2398.
  26. Chow CK, Jolly S, Rao-Melacini P, Fox KA, Anand SS, Yusuf S. Association of diet, exercise, and smoking modification with risk of early cardiovascular events after acute coronary syndromes. *Circulation* 2010; **121**: 750–758.
  27. Morrow DA, Antman EM, Charlesworth A, Cairns R, Murphy SA, de Lemos JA, et al. TIMI risk score for ST-elevation myocardial infarction: A convenient, bedside, clinical score for risk assessment at presentation: An intravenous nPA for treatment of infarcting myocardium early II trial substudy. *Circulation* 2000; **102**: 2031–2037.
  28. Halkin A, Singh M, Nikolsky E, Grines CL, Tchong JE, Garcia E, et al. Prediction of mortality after primary percutaneous coronary intervention for acute myocardial infarction the CADILLAC risk score. *J Am Coll Cardiol* 2005; **45**: 1397–1405.
  29. Lloyd-Jones DM, Leip EP, Larson MG, D'Agostino RB, Beiser A, Wilson PW, et al. Prediction of lifetime risk for cardiovascular disease by risk factor burden at 50 years of age. *Circulation* 2006; **113**: 791–798.
  30. Berry JD, Liu K, Folsom AR, Lewis CE, Carr JJ, Polak JF, et al. Prevalence and progression of subclinical atherosclerosis in younger adults with low short-term but high lifetime estimated risk for cardiovascular disease: The coronary artery risk development in young adults study and multi-ethnic study of atherosclerosis. *Circulation* 2009; **119**: 382–389.
  31. Imamura H, Izawa A, Kai R, Yokoseki O, Uchikawa S, Yazaki Y, et al. Trends over the last 20 years in the clinical background of young Japanese patients with coronary artery disease. *Circ J* 2004; **68**: 186–191.
  32. Witt BJ, Jacobsen SJ, Weston SA, Killian JM, Meverden RA, Allison TG, et al. Cardiac rehabilitation after myocardial infarction in the community. *J Am Coll Cardiol* 2004; **44**: 988–996.
  33. Squires RW, Montero-Gomez A, Allison TG, Thomas RJ. Long-term disease management of patients with coronary disease by cardiac rehabilitation program staff. *J Cardiopulm Rehabil Prev* 2008; **28**: 180–186.
  34. Tani S, Nagao K, Anazawa T, Kawamata H, Furuya S, Takahashi H, et al. Coronary plaque regression and lifestyle modification in patients treated with pravastatin. *Circ J* 2010; **74**: 954–961.
  35. Belardinelli R, Paolini I, Cianci G, Piva R, Georgiou D, Purcaro A. Exercise training intervention after coronary angioplasty: The ETICA trial. *J Am Coll Cardiol* 2001; **37**: 1891–1900.
  36. Green DJ, Walsh JH, Maiorana A, Best MJ, Taylor RR, O'Driscoll JG. Exercise-induced improvement in endothelial dysfunction is not mediated by changes in CV risk factors: Pooled analysis of diverse patient populations. *Am J Physiol* 2003; **285**: H2679–H2687.
  37. Nakatani D, Sakata Y, Sato H, Mizuno H, Shimizu M, Suna S, et al; Osaka Acute Coronary Insufficiency Study (OACIS) Group. Clinical impact of metabolic syndrome and its additive effect with smoking on subsequent cardiac events after acute myocardial infarction. *Am J Cardiol* 2007; **99**: 885–889.
  38. Ades PA, Waldmann ML, McCann WJ, Weaver SO. Predictors of cardiac rehabilitation participation in older coronary patients. *Arch Intern Med* 1992; **152**: 1033–1035.



## Effects of Exercise Training in Patients With Chronic Heart Failure and Advanced Left Ventricular Systolic Dysfunction Receiving $\beta$ -Blockers

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**Background:** It remains unclear whether patients with chronic heart failure (CHF) and advanced left ventricular (LV) dysfunction on  $\beta$ -blocker therapy benefit from exercise training (ET).

**Methods and Results:** We studied 45 CHF patients with advanced LV dysfunction [ejection fraction (LVEF) <25%] and impaired exercise tolerance [normalized peak oxygen uptake ( $\dot{V}O_2$ ) <70%] receiving a  $\beta$ -blocker: 33 patients participated in a cardiac rehabilitation program with ET (ET group) and 12 did not (inactive control group). Exercise capacity, LV dimension and plasma B-type natriuretic peptide (BNP) were assessed before and after a 3-month study period. At baseline, both groups had markedly reduced LVEF (ET group  $18\pm 4\%$  vs. Control group  $18\pm 5\%$ , NS) and impaired exercise capacity (normalized  $\dot{V}O_2$   $51\pm 10\%$  vs.  $55\pm 9\%$ , NS). Although one patient in the ET group withdrew from the program due to worsening CHF, no serious cardiac events occurred during the ET sessions. After 3 months, the ET group ( $n=24$ ) had significantly improved  $\dot{V}O_2$  by  $16\pm 15\%$  ( $1,005\pm 295$  to  $1,167\pm 397$  ml/min,  $P<0.001$ ), while the  $\dot{V}O_2$  of the control group was unchanged. LV end-diastolic dimension decreased in both groups to a similar extent, but plasma BNP was significantly decreased only in the ET group ( $432$  to  $214$  pg/ml,  $P<0.05$ ).

**Conclusions:** The data indicate that in CHF patients with advanced LV dysfunction on  $\beta$ -blocker therapy, ET successfully improves exercise capacity and BNP without adversely affecting LV remodeling or causing serious cardiac complications. (*Circ J* 2011; **75**: 1649–1655)

**Key Words:** Advanced left ventricular dysfunction; Beta-blocker; Chronic heart failure; Exercise capacity; Exercise training

In patients with chronic heart failure (CHF), exercise training (ET) improves exercise capacity, quality of life, and prognosis.<sup>1–4</sup> The ACC/AHA guidelines for the management of CHF recommend ET as an adjunct therapy to improve the clinical status of ambulatory patients with current or prior symptoms of heart failure (HF) and reduced left ventricular (LV) ejection fraction (LVEF).<sup>5</sup> However, previous studies of ET in patients with CHF have primarily examined patients with an LVEF in the range of 25–40%, and few studies enrolled patients with advanced LV dysfunction (LVEF <25%).<sup>6</sup> Thus, the therapeutic benefits and safety of ET in patients with advanced LV dysfunction remain unknown.

$\beta$ -blocker therapy improves the long-term prognosis of patients with advanced HF,<sup>7–9</sup> and  $\beta$ -blockers are recommended as a class I standard medication by the ACC/AHA and ESC guidelines.<sup>5,10</sup> However,  $\beta$ -blockers were prescribed in only

0–70% of eligible patients in previous studies of ET in patients with advanced LV dysfunction.<sup>6,11–14</sup> A large randomized controlled trial, HF-ACTION,<sup>15</sup> was designed to examine the efficacy and safety of ET in patients with CHF, and the results of this study were recently reported.  $\beta$ -blockers were prescribed in 95% of the patients in HF-ACTION, but the median LVEF of study participants was 25%, indicating that half of the patients had an LVEF higher than 25%. Thus, in the current era of widespread  $\beta$ -blocker use, it remains unclear whether patients with CHF and advanced LV dysfunction on  $\beta$ -blocker therapy benefit from ET. Accordingly, the purpose of the present study was to determine whether ET safely improves the exercise capacity of CHF patients on  $\beta$ -blocker therapy with advanced LV dysfunction.

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**Table 1. Clinical Characteristics and Baseline Data**

	ET group	Inactive control group	P value
n	33	12	
Age (years)	51±14	52±16	NS
Male (%)	88	83	NS
NYHA class (II/III) (%)	39/61	33/67	NS
NIsCM/IsCM (%)	64/36	83/17	NS
HT (%)	41	17	NS
HL (%)	36	67	NS
IGT/DM (%)	36	25	NS
BMI	21.4±3.1	23.1±2.6	NS
<b>Medications</b>			
Digitalis (%)	70	58	NS
Diuretics (%)	97	92	NS
ACEI/ARB (%)	91	92	NS
BB (Car/Meto/Biso [%])	88/9/3	92/0/8	
Ca antagonist (%)	9	8	NS
Nitrate (%)	24	8	NS
Cardioionic (%)	36	8	NS
Antiarrhythmic agent (%)	48	25	NS
<b>Baseline data</b>			
Plasma BNP (pg/ml)	365±390	215±129	NS
LVEDD (mm)	70±7	70±9	NS
LVEF (%)	18±4	18±5	NS
Peak $\dot{V}O_2$ (ml/min)	979±285	1,122±283	NS
Normalized peak $\dot{V}O_2$ (%)	51±10	55±9	NS
$\dot{V}E$ - $\dot{V}CO_2$ slope	35.3±10.1	31.7±5.0	NS

Data are mean ± SD.

ET, exercise training; NYHA, New York Heart Association; NIsCM, non-ischemic cardiomyopathy; IsCM, ischemic cardiomyopathy; HT, hypertension; HL, hyperlipidemia; IGT, impaired glucose tolerance; DM, diabetes mellitus; BMI, body mass index; ACEI, angiotensin-converting enzyme inhibitors; ARB, angiotensin II receptor blocker; BB,  $\beta$ -blocker; Car, carvedilol; Meto, metoprolol; Biso, bisoprolol; BNP, B-type natriuretic peptide; LVEDD, left ventricular end-diastolic diameter; LVEF, left ventricular ejection fraction;  $\dot{V}O_2$ , oxygen uptake;  $\dot{V}E$ , ventilation;  $\dot{V}CO_2$ , carbon dioxide output.

## Methods

### Patients

Forty-five CHF patients with advanced LV dysfunction who fulfilled the following inclusion criteria were included in the present study: 15–80 years old, advanced LV systolic dysfunction (LVEF <25%), reduced exercise tolerance with normalized peak oxygen uptake <70%, a well-controlled body fluid level (euvoletic), and no signs of worsening HF over the preceding 2-week period. LVEF was determined during left ventriculography using contrast medium or radioisotope. Of the 45 patients, we retrospectively assigned 33 patients who participated in a 3-month cardiac rehabilitation program with ET to the ET group and the remaining 12 patients who did not undergo ET to the inactive control group. All patients were taking a  $\beta$ -blocker and were in a stable state.

### The Cardiac Rehabilitation Program

The cardiac rehabilitation program with ET for CHF at our institute has been previously described.<sup>16</sup> Before entering the ET program, patients were confirmed not to have evidence of ischemia or severe arrhythmia during a level walking test. All patients gave written informed consent before entering

the program.

The exercise program consisted of walking, bicycling on an ergometer, and calisthenics of 40–60 min/session 3–5 sessions/week for 3 months. Exercise intensity was determined individually at 30–50% of heart rate (HR) reserve (Karvonen's equation:  $k=0.3-0.5$ ),<sup>17</sup> an anaerobic threshold (AT) level obtained in a maximal symptom-limited cardiopulmonary exercise test, or at levels 11–13 ("fairly light" to "somewhat hard") of the 6–20 scale rating of perceived exertion (the original Borg's score<sup>18,19</sup>). Care was taken to prescribe a slightly lower level of exercise intensity (30–40% of HR reserve or an AT level) and lower session frequency (3 sessions/week) to patients with very low LVEF (<20%). The exercise program usually began with supervised sessions for 2–4 weeks, followed by home exercise combined with once or twice weekly supervised sessions for the remaining 8–10 weeks. Home exercise consisted mainly of brisk walking at a prescribed HR for 30–50 min, 3–5 times a week.

Patients were encouraged to attend education classes, which were held 3 times each week with lectures given by physicians, nurses, dietitians, and pharmacists on coronary artery disease, secondary prevention, HF management, diet, smoking cessation, and medication. In addition, all ET group patients received individual counseling on exercise prescription, secondary prevention, and daily life activities by a physician and a nurse at the time of hospital discharge and at the end of the 3-month cardiac rehabilitation program.

The inactive control patients did not participate in any exercise program or perform regular home exercise.

### Cardiopulmonary Exercise Testing

Patients were scheduled to undergo a symptom-limited cardiopulmonary exercise test at the beginning and end of the 3-month study period.<sup>20</sup> After a 2-min rest period on the bicycle ergometer in an upright position, patients pedaled at an intensity of 0W for 1 min (warm-up), and were then subjected to an incremental exercise test with a ramp protocol (10 or 15W/min) until exhaustion. The 12-lead ECG was continuously monitored and blood pressure was measured every minute with a sphygmomanometer. Expired gases were collected and analyzed continuously with an AE-280S or AE-300S gas analyzer (Minato Co, Osaka, Japan). Peak oxygen uptake (peak  $\dot{V}O_2$ ) was defined as the highest  $\dot{V}O_2$  value achieved at peak exercise. Ventilation ( $\dot{V}E$ ) and carbon dioxide output ( $\dot{V}CO_2$ ) were measured and the gradient of the  $\dot{V}E$ - $\dot{V}CO_2$  relationship ( $\dot{V}E$  vs.  $\dot{V}CO_2$  slope) was determined.

### Clinical Data

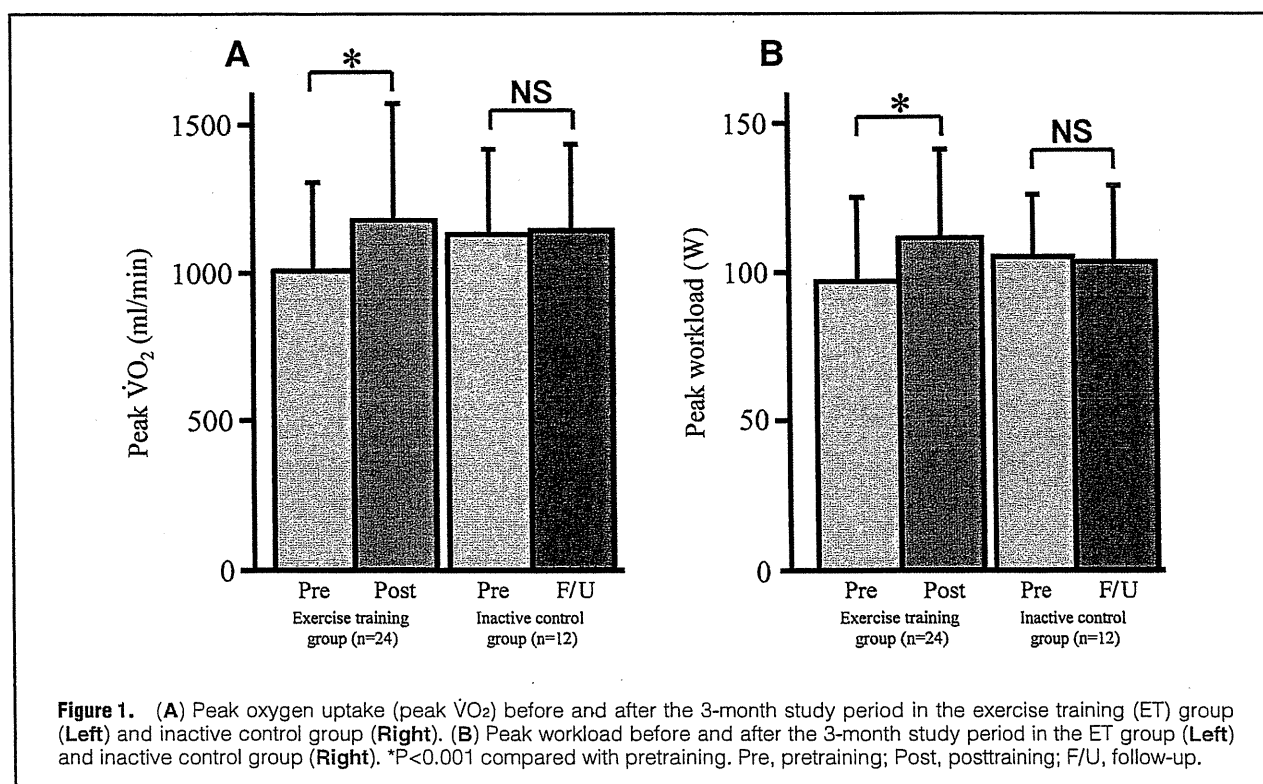
Patients were scheduled to undergo echocardiography and plasma B-type natriuretic peptide (BNP) measurements at the beginning and end of the 3-month study period. LV internal diameters were acquired from the parasternal short-axis view, at the approximate mitral chordae level, using direct 2-dimensional measurements or targeted M-mode echocardiography if the M-mode cursor could be positioned perpendicular to the septum and LV posterior wall. Plasma BNP levels were measured with a specific immunoradiometric assay for human BNP using a commercial kit (Shionoria). The upper limit of normal plasma BNP level was 18.4 pg/ml. The minimal and maximal detectable levels of BNP were 4 and 2,000 pg/ml, respectively.

### Statistical Analysis

Data are presented as the means ± standard deviations. Significant differences were determined with paired or unpaired

	ET group (n=24)			Inactive control group (n=12)		
	Pretraining	Posttraining	P value	Pretraining	Follow-up	P value
Plasma BNP (pg/ml)	432±451	214±232	<0.05	238±130	281±305	NS
LVEDD (mm)	73±6	66±11	<0.005	72±8	65±11	<0.01
LVFS (%)	12±4	16±6	0.065	10±3	19±9	<0.01
HR at rest (beats/min)	82±16	76±14	0.057	74±17	72±19	NS
Peak HR (beats/min)	129±28	134±22	NS	141±27	130±26	NS
Peak $\dot{V}O_2$ (ml/min)	1,005±295	1,167±397	<0.001	1,122±283	1,136±288	NS
Normalized peak $\dot{V}O_2$ (%)	50±11	58±14	<0.001	55±9	57±12	NS
Peak work rate (W)	96±28	111±30	<0.001	105±20	103±26	NS
AT (ml/min)	585±162	622±160	0.066	639±133	593±117	NS
$\dot{V}E-\dot{V}CO_2$ slope	34.8±11.0	32.9±10.9	NS	31.7±5.0	31.9±6.2	NS

Plasma BNP data were available for 21 patients in the ET group and 10 patients in the inactive control group. LVEDD data were available for 18 patients in the ET group and 11 patients in the inactive control group. LVFS data were available for 16 patients in the ET group and 11 patients in the inactive control group. AT data were available for 19 patients in the ET group and 11 patients in the inactive control group. LVFS, left ventricular fractional shortening; HR, heart rate; AT, anaerobic threshold. Other abbreviations as in Table 1.

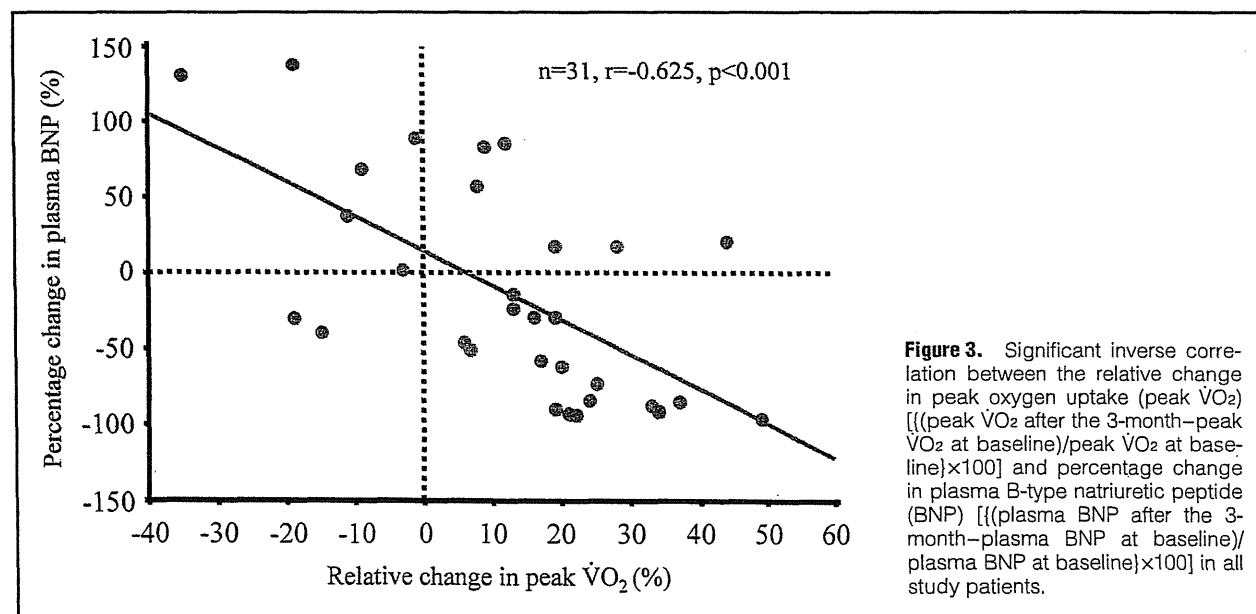
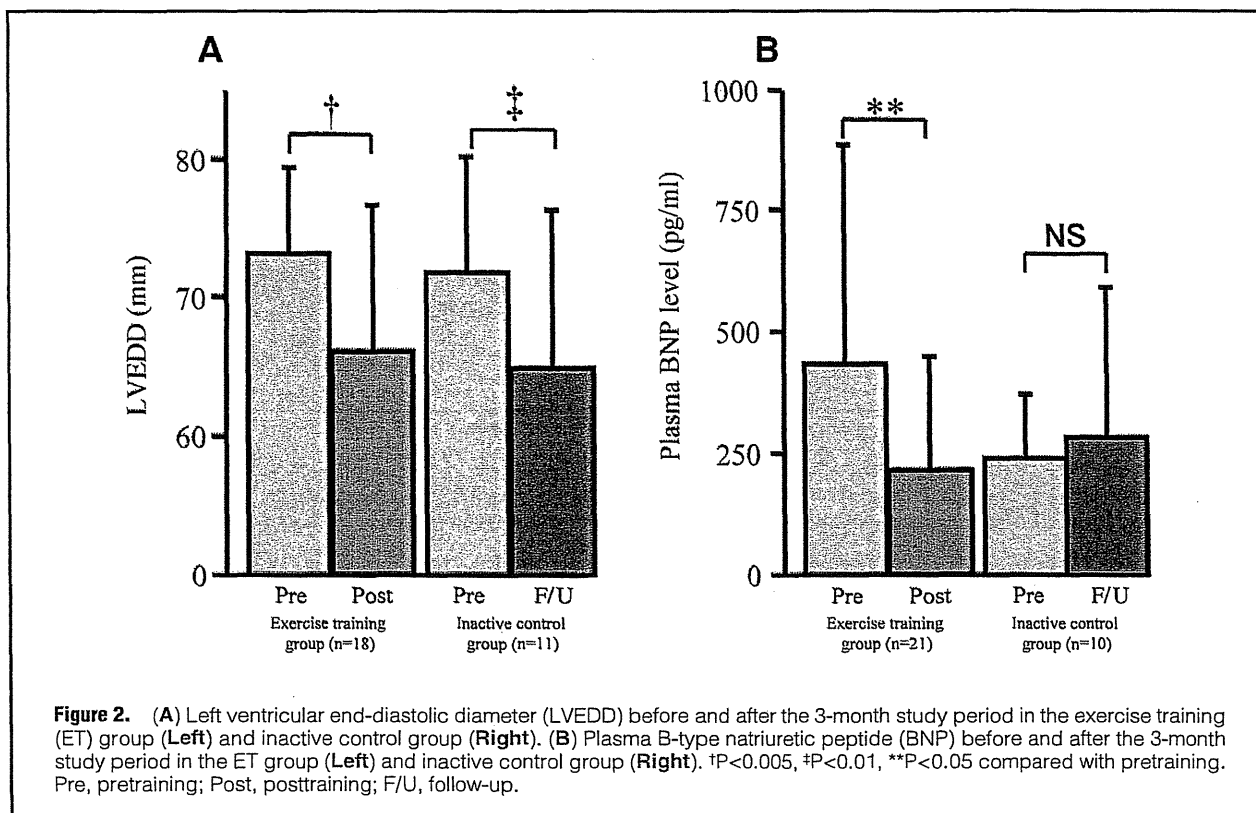


**Figure 1.** (A) Peak oxygen uptake (peak  $\dot{V}O_2$ ) before and after the 3-month study period in the exercise training (ET) group (Left) and inactive control group (Right). (B) Peak workload before and after the 3-month study period in the ET group (Left) and inactive control group (Right). \*P<0.001 compared with pretraining. Pre, pretraining; Post, posttraining; F/U, follow-up.

t-tests as appropriate. Considering the small number of inactive control patients, we also analyzed the differences between groups using a non-parametric Mann-Whitney U test, and the differences before and after ET in each group using Wilcoxon test. Differences in frequencies were analyzed with the Fisher exact probability test or the chi-square test. Pearson's correlation analysis was used to evaluate the correlation between the change in peak  $\dot{V}O_2$  and the change in BNP. Statistical calculations were performed using Statview software. A P value less than 0.05 was considered statistically significant.

### Results

Table 1 summarizes the clinical characteristics, baseline LV function and exercise tolerance in the ET and inactive control groups. There were no significant differences between the ET and control groups at baseline. The non-parametric Mann-Whitney U test yielded the same results (data not shown). All patients were already on  $\beta$ -blocker therapy at the time of study entry, but some patients were still being up-titrated. The proportion of patients taking  $\beta$ -blockers for more than 3 months was 24% (8/33 patients) in the ET group and 33% (4/12 patients) in the control group (NS). The doses of  $\beta$ -blockers were similar between the 2 groups at the time of



study entry (for carvedilol, 11.3 mg/day in the ET group and 13.2 mg/day in the control group, NS). The schedule of up-titration and the final dose of  $\beta$ -blockers were left to the attending physicians.

Of the 33 patients in the ET group, 24 (73%) completed the 3-month ET program, and among the 9 patients who did not complete the study, one had an exacerbation of HF (3%), one withdrew for a non-cardiac medical reason (claudication) (3%) and the remaining 7 withdrew for social reasons (the

distance to the institute was too far, return to work, etc) (21%). These 9 patients were excluded from the statistical analysis of ET effects. There were no significant differences in the baseline plasma BNP levels, LV end-diastolic dimensions (LVEDD), LVEF, or exercise capacity between the 24 patients who completed the 3-month ET program and the 9 patients who withdrew. No serious cardiac events, including death or cardiopulmonary arrest, occurred during the ET sessions. No patients developed new atrial fibrillation during

First author	Year	Number of patients	Age (years)	NYHA (II/III)	LVEF (%)	ACEI/ARB (%)	BB (%)	Duration	Increments of peak $\dot{V}O_2$ ( $\text{ml}\cdot\text{min}^{-1}\cdot\text{kg}^{-1}$ )
Coats AJS	1990	11	63	7/4	19	82	—	8 weeks	14.3→16.7 (17%†)
Coats AJS	1992	17	61.8	10/7	19.6	88	0	8 weeks	13.2→15.6 (18%†)
Adamopoulos S	1995	12	62.4	6/6	18.9	92	0	8 weeks	12.4→—
Sturm B	1999	13	55	6/7	17	92	23	12 weeks	15.9→18.5 (16%†)
Quittan M	1999	12	57	7/5	17	100	25	3 months	15.9→18.5 (16%†)
Van Berendoncks AM	2010	46	57.5	34/12	17	100	70	4 months	19→21 (11%†)
Nishi I	2011	24	51	11/13	17	88	100	3 months	16.3→18.7 (15%†)

Abbreviations as in Table 1.

the rehabilitation program.

### Exercise Variables

After the 3-month study period, resting HR was unchanged in the control group, but tended to decrease in the ET group ( $P=0.057$ , Table 2), and peak HR did not change in both groups (Table 2). The ET group showed significant improvements in peak  $\dot{V}O_2$  ( $1,005\pm 295$  to  $1,167\pm 397$  ml/min,  $16\pm 15\%$ ;  $16.3\pm 4.3$  to  $18.7\pm 5.1$   $\text{ml}\cdot\text{min}^{-1}\cdot\text{kg}^{-1}$ ,  $15\pm 14\%$ , both  $P<0.001$ ) and its normalized value ( $P<0.001$ ) after the study period, whereas the inactive control group did not (Table 2, Figure 1A). In addition, only the ET group achieved a significant increase in peak work rate after the 3-month ET ( $96\pm 28$  to  $111\pm 30$  W,  $P<0.001$ ) (Table 2, Figure 1B).  $\dot{V}E$  vs.  $\dot{V}CO_2$  slope did not significantly change in either group after the 3-month study period (Table 2).

The non-parametric Wilcoxon test yielded the same results, except that  $\dot{V}E$  vs.  $\dot{V}CO_2$  slope significantly decreased only in the ET group after the 3-month study period (median 31.1 to 29.4,  $P<0.05$ ).

### LV Function and BNP

LVEDD significantly decreased in both the ET group ( $P<0.005$ ) and the inactive control group ( $P<0.01$ ) (Table 2, Figure 2A), and the relative decrease in LVEDD was similar between the 2 groups (ET group  $-10\pm 11\%$  vs. inactive control group  $-10\pm 10\%$ , NS). LV fractional shortening (FS) increased in both groups, but the change reached a statistical significance only in the inactive control group ( $P<0.01$ ) and not in the ET group ( $P=0.065$ ) (Table 2).

Compared to their baseline levels, plasma BNP concentrations significantly decreased ( $P<0.05$ ) after the ET program, but did not change in the inactive control group ( $n=10$ ) (Table 2, Figure 2B).

When the percentage change in BNP [ $(\text{BNP after 3 months} - \text{BNP at baseline})/\text{BNP at baseline} \times 100$ ] was plotted against the relative change in peak  $\dot{V}O_2$  [ $(\text{peak } \dot{V}O_2 \text{ after 3 months} - \text{peak } \dot{V}O_2 \text{ at baseline})/\text{peak } \dot{V}O_2 \text{ at baseline} \times 100$ ] in the entire patient population, there was a significant inverse correlation ( $r=-0.625$ ,  $P<0.001$ ,  $n=31$ ), indicating that a greater increase in peak  $\dot{V}O_2$  was associated with a greater decrease in plasma BNP (Figure 3).

## Discussion

ET has been shown to favorably affect the outcomes and quality of life of many patients with CHF, but it has remained unclear whether ET is safe and beneficial for those CHF patients with advanced LV dysfunction and on  $\beta$ -blocker

therapy. We set out to examine that question, and our data indicate that, in this subset of CHF patients, an appropriate ET program safely increased exercise capacity and decreased plasma BNP levels without promoting deleterious LV remodeling. Additionally, there was a significant inverse correlation between plasma BNP and peak  $\dot{V}O_2$  during the study period, suggesting that improvements in exercise capacity are associated with amelioration of elevated LV load. Finally, the prescribed ET program was very well tolerated; the withdrawal rate from the exercise program due to cardiac reasons was only 3%.

### Previous Studies

Several other studies over the past 20 years have studied the role of ET in patients with advanced LV dysfunction (mean LVEF <20%) (Table 3), and demonstrated a significant improvement in peak  $\dot{V}O_2$  by 16–18% following ET.<sup>6,11,13,21</sup> However,  $\beta$ -blockers were prescribed to only 0–25% of enrolled patients in most of those studies,<sup>6,11–13</sup> and to 70% of the patients in one study.<sup>14</sup> In contrast, in the present study, all patients were prescribed  $\beta$ -blockers. Thus, our results are more relevant in the current era when  $\beta$ -blocker therapy is the standard of care.

$\beta$ -blocker therapy does not improve peak  $\dot{V}O_2$  in patients with CHF, despite the symptom amelioration, improvements in NYHA functional class, and increased LVEF associated with  $\beta$ -blocker therapy.<sup>22,23</sup> In contrast, ET can increase peak  $\dot{V}O_2$  in patients with moderate to severe HF who are taking a  $\beta$ -blocker.<sup>24–27</sup> However, most of the previous studies examining ET in patients using  $\beta$ -blockers recruited only patients with LVEF in the range of 23–35%. The study population of the recent HF-ACTION trial, in which  $\beta$ -blockers were prescribed to 95% of patients, did not contain more than 50% of patients with an LVEF <25%.<sup>15</sup> Thus, to our knowledge, the present study is the first to examine the effects of ET in CHF patients with advanced LV dysfunction (LVEF <25%) and concurrent  $\beta$ -blocker therapy.

### Exercise Capacity

There was concern that  $\beta$ -blockers could abrogate the benefits of ET in patients with CHF, but one study reported that  $\beta$ -blocker therapy did not affect ET-associated increases in exercise capacity.<sup>24</sup> Additionally, there were no differences in the effects of ET between patients receiving  $\beta$ -1 selective and non-selective  $\alpha$ - $\beta$  blockers.<sup>28</sup> The increase in peak  $\dot{V}O_2$  in the ET group in the present study (15%) was less than that seen in the study of Demopoulos et al (24–27%),<sup>24</sup> but was comparable to the increase reported by Forissier et al (14–17%).<sup>28</sup>

In the HF-ACTION trial,<sup>15</sup> the ET group had a significant



improvement in peak  $\dot{V}O_2$  at 3 months compared to the usual care group (0.6 vs. 0.2 ml·min<sup>-1</sup>·kg<sup>-1</sup>;  $P<0.001$ ), but this increase was relatively small (median 4%). The authors attributed the relatively small improvement to low adherence to the exercise protocol. Thus, baseline patient characteristics, the specific exercise program prescribed, and adherence to that program likely all contribute to the magnitude of change in exercise capacity observed in the different studies.

### LV Remodeling

In the present study, both study groups showed a decrease in LVEDD (ie, there was reverse LV remodeling). This result differs from those of Giannuzzi et al who found that LV remodeling was reversed by ET, but worsened in a control non-ET group with moderate CHF (average LVEF 25%).<sup>29</sup> These conflicting results may be due to the low rate of prescription of  $\beta$ -blockers in the study by Giannuzzi et al (20%).

In addition, experimental studies using a rat model showed that excessive exercise after a large myocardial infarction aggravated LV remodeling,<sup>30,31</sup> but moderate exercise had either no effect<sup>32</sup> or attenuated adverse LV remodeling.<sup>33</sup> Although the myocardial infarction models in the rat differ from human CHF, these findings suggest that exercise intensity in an ET program can affect LV remodeling. Based on this, exercise intensity was adjusted according to the severity of LV dysfunction in each patient in the present study. In addition to  $\beta$ -blocker therapy, this appropriate adjustment of exercise prescription may also have contributed to reverse LV remodeling.

### Plasma BNP

Passino et al reported that ET decreases plasma BNP levels in patients with moderate CHF.<sup>34</sup> The present results are consistent with that study, but we extended these findings to a patient population with more severe LV dysfunction (mean LVEF 18%) and higher baseline BNP (365 pg/ml) than Passino's study (mean LVEF of 35% and baseline BNP of 187 pg/ml).

The significant inverse correlation that we found between changes in BNP and exercise capacity is also consistent with the previous report,<sup>34</sup> which is somewhat counterintuitive because one could hypothesize that a greater increase in exercise capacity requiring a greater amount of ET would result in sustained LV wall stress and an associated increase in plasma BNP. However, the available data suggests that ET of an appropriate intensity and duration ameliorates the increased LV wall stress that stimulates BNP production. This hypothesis is consistent with the observation in a canine model of HF that regular ET lowers LV end-diastolic pressure at rest through enhanced nitric oxide production.<sup>35</sup> One remaining question is whether the observed decrease in BNP is attributable to ET alone or to the combination of ET and  $\beta$ -blocker therapy.

### Etiology of CHF

It is intriguing whether the response to ET differs between CHF due to ischemic and non-ischemic cardiomyopathies. Because the total number of patients was not sufficient, such subgroup analyses were beyond the scope of the present study. However, a tentative analysis of 26 patients with a non-ischemic etiology showed a consistent result with the main results: After the 3-month study period, the non-ischemic ET group (n=16) showed significant increases in peak  $\dot{V}O_2$  (1,019 to 1,210 ml/min,  $P<0.001$ ) and peak work rate (100 to 116 W,  $P<0.001$ ), whereas the non-ischemic inactive control group (n=10) did not (1,177 to 1,154 ml/min and 107 to 103 W, both NS), while having significant reductions in

LVEDD in both groups (74 to 66 mm and 71 to 65 mm, both  $P<0.05$ ). A larger study is necessary to explore the potential difference, if any, in response to ET between ischemic and non-ischemic CHF.

### Safety of ET in HF

Among the study patients undergoing ET, 1 (3%) had worsening of HF in the 3-month study period. The ELVD-CHF trial followed patients with a mean LVEF of 25% for 6 months, and clinical events (death, HF hospitalization, temporary worsening of symptoms not requiring hospitalization) occurred in 9% of patients in the exercise group, but in 18% of patients in the usual care group.<sup>29</sup> Additionally, HF-ACTION followed patients with a median LVEF of 25% for 30 months, and worsening HF occurred in 26% of the ET group and 29% of the patients in the usual care group.<sup>15</sup> Thus, the incidence of cardiac events observed in the present study for patients with severe LV dysfunction is comparable to that seen in patients undergoing conventional care in previous studies, suggesting that ET for HF patients with severe LV dysfunction on  $\beta$ -blockers may be safe.

### Clinical Implications

Our results extend previous findings<sup>11,12,21,24,28</sup> of the favorable effects of ET in CHF patients with moderate LV dysfunction<sup>24,28</sup> or not receiving  $\beta$ -blockers<sup>11,12,21</sup> to those with advanced LV dysfunction receiving  $\beta$ -blockers. Physicians are often reluctant to recommend ET to patients with HF, especially when the LVEF is markedly reduced. Our data suggest that, even in such patients with advanced LV dysfunction, appropriate ET/cardiac rehabilitation based on careful medical evaluation can lead to a significant improvement in exercise capacity and favorably affect LV remodeling and biomarkers of CHF progression.

### Study Limitations

Our data provide a good rationale for future studies of ET for patients with advanced LV dysfunction in CHF, but our study is limited because it is retrospective and from a single center with a relatively small number of patients. Therefore, a large controlled randomized study should be performed to confirm these results.

An additional confounding factor in our study is the exclusion from the analysis of the 9 patients who dropped out of the study. However, an additional comparison between the completion group (n=24) and dropout group (n=9) yielded similar backgrounds, and we expect little bias from the exclusion of these patients.

All the patients in the present study were clinically stable on  $\beta$ -blockers and the proportion of patients taking  $\beta$ -blockers for more than 3 months was similar between the 2 groups, but the time from the initiation of  $\beta$ -blocker therapy to the start of ET varied. Additionally,  $\beta$ -blocker therapy was not standardized across the study participants. Therefore, we cannot draw any conclusions regarding the optimal time to initiate ET after the start of  $\beta$ -blocker therapy or which  $\beta$ -blocker is best<sup>36</sup> combined with ET. Further studies are necessary to address these issues.

### Conclusion

In patients with CHF, advanced LV dysfunction and on  $\beta$ -blockers, ET can safely increase exercise capacity with favorable effects on LV remodeling and plasma BNP with a low incidence of cardiac complications (3%).

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### References

- European Heart Failure Training Group. Experience from controlled trials of physical training in chronic heart failure: Protocol and patient factors in effectiveness in the improvement in exercise tolerance. *Eur Heart J* 1998; **19**: 466–475.
- Hambrecht R, Fiehn E, Weigl C, Gielen S, Hamann C, Kaiser R, et al. Regular physical exercise corrects endothelial dysfunction and improves exercise capacity in patients with chronic heart failure. *Circulation* 1998; **98**: 2709–2715.
- Belardinelli R, Georgiou D, Cianci G, Purcaro A. Randomized, controlled trial of long-term moderate exercise training in chronic heart failure: Effects on functional capacity, quality of life, and clinical outcome. *Circulation* 1999; **99**: 1173–1182.
- Working Group on Cardiac Rehabilitation & Exercise Physiology and Working Group on Heart Failure of the European Society of Cardiology. Recommendations for exercise training in chronic heart failure patients. *Eur Heart J* 2001; **22**: 125–135.
- Hunt SA, Abraham WT, Chin MH, Feldman AM, Francis GS, Ganiats TG, et al. 2009 focused update incorporated into the ACC/AHA 2005 Guidelines for the Diagnosis and Management of Heart Failure in Adults: A report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines: Developed in collaboration with the International Society for Heart and Lung Transplantation. *Circulation* 2009; **119**: e391–e479.
- Sturm B, Quittan M, Wiesinger GF, Stanek B, Frey B, Pacher R. Moderate-intensity exercise training with elements of step aerobics in patients with severe chronic heart failure. *Arch Phys Med Rehabil* 1999; **80**: 746–750.
- Krum H, Sackner-Bernstein JD, Goldsmith RL, Kukin ML, Schwartz B, Penn J, et al. Double-blind, placebo-controlled study of the long-term efficacy of carvedilol in patients with severe chronic heart failure. *Circulation* 1995; **92**: 1499–1506.
- Goldstein S, Fagerberg B, Hjalmarson A, Kjekshus J, Waagstein F, Wedel H, et al; MERIT-HF Study Group. Metoprolol controlled release/extended release in patients with severe heart failure: Analysis of the experience in the MERIT-HF study. *J Am Coll Cardiol* 2001; **38**: 932–938.
- Packer M, Fowler MB, Roecker EB, Coats AJ, Katus HA, Krum H, et al; Carvedilol Prospective Randomized Cumulative Survival (COPERNICUS) Study Group. Effect of carvedilol on the morbidity of patients with severe chronic heart failure: Results of the carvedilol prospective randomized cumulative survival (COPERNICUS) study. *Circulation* 2002; **106**: 2194–2199.
- Task Force for Diagnosis and Treatment of Acute and Chronic Heart Failure 2008 of European Society of Cardiology, ESC Committee for Practice Guidelines. ESC Guidelines for the diagnosis and treatment of acute and chronic heart failure 2008: The Task Force for the Diagnosis and Treatment of Acute and Chronic Heart Failure 2008 of the European Society of Cardiology: Developed in collaboration with the Heart Failure Association of the ESC (HFA) and endorsed by the European Society of Intensive Care Medicine (ESICM). *Eur Heart J* 2008; **29**: 2388–2442.
- Coats AJS, Adamopoulos S, Radaelli A, McCance A, Meyer TE, Bernardi L, et al. Controlled trial of physical training in chronic heart failure: Exercise performance, hemodynamics, ventilation, and autonomic function. *Circulation* 1992; **85**: 2119–2131.
- Adamopoulos S, Ponikowski P, Cerquetani E, Piepoli M, Rosano G, Sleight P, et al. Circadian pattern of heart rate variability in chronic heart failure patients: Effects of physical training. *Eur Heart J* 1995; **16**: 1380–1386.
- Quittan M, Sturm B, Wiesinger GF, Pacher R, Fialka-Moser V. Quality of life in patients with chronic heart failure: A randomized controlled trial of changes induced by a regular exercise program. *Scand J Rehab Med* 1999; **31**: 223–228.
- Van Berendoncks AM, Beckers P, Hoymans VV, Possemiers N, Wuytss FL, Vrints CJ, et al. Exercise training reduces circulating adiponectin levels in patients with chronic heart failure. *Clin Sci (Lond)* 2010; **118**: 281–289.
- O'Connor CM, Whellan DJ, Lee KL, Keteyian SJ, Cooper LS, Ellis SJ, et al; HF-ACTION Investigators. Efficacy and safety of exercise training in patients with chronic heart failure: HF-ACTION randomized controlled trial. *JAMA* 2009; **301**: 1439–1450.
- Nishi I, Noguchi T, Furuichi S, Iwanaga Y, Kim J, Ohya H, et al. Are cardiac events during exercise therapy for heart failure predictable from the baseline variables? *Circ J* 2007; **71**: 1035–1039.
- Karvonen M, Kentala K, Mustala O. The effects of training on heart rate: A longitudinal study. *Ann Med Exp Biol Fenn* 1957; **35**: 307–315.
- Borg G. Perceived exertion as an indicator of somatic stress. *Scand J Rehabil Med* 1970; **2/3**: 92–98.
- Carvalho VO, Bocchi EA, Guimarães GV. The Borg scale as an important tool of self-monitoring and self-regulation of exercise prescription in heart failure patients during hydrotherapy: A randomized blinded controlled trial. *Circ J* 2009; **73**: 1871–1876.
- Kamakura T, Kawakami R, Nakanishi M, Ibuki M, Ohara T, Yanase M, et al. Efficacy of out-patient cardiac rehabilitation in low prognostic risk patients after acute myocardial infarction in primary intervention era. *Circ J* 2011; **75**: 315–321.
- Coats AJS, Adamopoulos S, Meyer TE, Conway J, Sleight P. Effects of physical training in chronic heart failure. *Lancet* 1990; **335**: 63–66.
- Olsen SL, Gilbert EM, Rehlund DG, Taylor DO, Yanowitz FD, Bristow MR. Carvedilol improves left ventricular function and symptoms in chronic heart failure: A double-blind randomized study. *J Am Coll Cardiol* 1995; **25**: 1225–1231.
- Gullestad L, Manhenke C, Aarsland T, Skardal R, Fagertun H, Wikstrand J, et al. Effect of metoprolol CR/XL on exercise tolerance in chronic heart failure: A substudy to the MERIT-HF trial. *Eur J Heart Fail* 2001; **3**: 463–468.
- Demopoulos L, Yeh M, Gentilucci M, Testa M, Bijou R, Katz SD, et al. Nonspecific beta-adrenergic blockade with carvedilol does not hinder the benefits of exercise training in patients with congestive heart failure. *Circulation* 1997; **95**: 1764–1767.
- Levinger I, Bronks R, Cody DV, Linton I, Davie A. Resistance training for chronic heart failure patients on beta blocker medications. *Int J Cardiol* 2005; **102**: 493–499.
- Fraga R, Franco FG, Roveda F, de Matos LN, Braga AM, Rondon MU, et al. Exercise training reduces sympathetic nerve activity in heart failure patients treated with carvedilol. *Eur J Heart Fail* 2007; **9**: 630–636.
- Wisløff U, Støylen A, Loennechen JP, Bruvold M, Rognum Ø, Haram PM, et al. Superior cardiovascular effect of aerobic interval training versus moderate continuous training in heart failure patients: A randomized study. *Circulation* 2007; **115**: 3086–3094.
- Forissier JF, Vernochet P, Bertrand P, Charbonnier B. Influence of carvedilol on the benefits of physical training in patients with moderate chronic heart failure. *Eur J Heart Failure* 2001; **3**: 335–342.
- Giannuzzi P, Temporelli PL, Corrà U, Tavazzi L; ELVD-CHF Study Group. Antiremodeling effect of long-term exercise training in patients with stable chronic heart failure: Results of the Exercise in Left Ventricular Dysfunction and Chronic Heart Failure (ELVD-CHF) Trial. *Circulation* 2003; **108**: 554–559.
- Oh BH, Ono S, Gilpin E, Ross J Jr. Altered left ventricular remodeling with beta-adrenergic blockade and exercise after coronary reperfusion in rats. *Circulation* 1993; **87**: 608–616.
- Gaudron P, Hu K, Schamberger R, Budin M, Walter B, Ertl G. Effect of endurance training early or late after coronary artery occlusion on left ventricular remodeling, hemodynamics, and survival in rats with chronic transmural myocardial infarction. *Circulation* 1994; **89**: 402–412.
- Alhaddad IA, Hakim I, Siddiqi F, Lagenback E, Mallavarapu C, Nethala V, et al. Early exercise after experimental myocardial infarction: Effect on left ventricular remodeling. *Coron Artery Dis* 1998; **9**: 319–327.
- Orenstein TL, Parker TG, Butany JW, Goodman JM, Dawood F, Wen WH, et al. Favorable left ventricular remodeling following large myocardial infarction by exercise training: Effect on ventricular morphology and gene expression. *J Clin Invest* 1995; **96**: 858–866.
- Passino C, Severino S, Poletti R, Piepoli MF, Mammì C, Clerico A, et al. Aerobic training decreases B-type natriuretic peptide expression and adrenergic activation in patients with heart failure. *J Am Coll Cardiol* 2006; **47**: 1835–1839.
- Wang J, Yi GH, Knecht M, Cai BI, Popskis S, Packer M, et al. Physical training alters the pathogenesis of pacing-induced heart failure through endothelium-mediated mechanisms in awake dogs. *Circulation* 1997; **96**: 2683–2692.
- Konishi M, Haraguchi G, Kimura S, Inagaki H, Kawabata M, Hachiya H, et al. Comparative effects of carvedilol vs bisoprolol for severe congestive heart failure. *Circ J* 2010; **74**: 1127–1134.

# 我が国における急性心筋梗塞後心臓リハビリテーション 実施率の動向：全国実態調査

Change in implementation of cardiac rehabilitation  
for acute myocardial infarction in Japan : a nationwide survey

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## 抄 録

【目 的】 全国における急性心筋梗塞 (AMI) 後心臓リハビリテーション (心リハ) 実施率の近年の推移は明らかでない。

【方 法】 2004 年の心リハ全国実態調査 (委託 15 指-2 研究班) および 2009 年の AMI 地域連携パス全国実態調査 (委託 20 公-7 研究班) の集計結果を用いて, AMI 後心リハ実施率を比較した。

【結 果】 2004 年調査 (2003 年実績) では 526 施設, 2009 年調査 (2008 年実績) では 597 施設の循環器専門医研修施設から回答が得られた (回答率: 2004 年 61%, 2009 年 64%)。2009 年調査で, 全病床数 (平均 456 床), 循環器科病床数 (平均 42 床), 冠動脈造影 (CAG) 年間件数 (平均 640 件) は 2004 年調査と同等で, 経皮的冠動脈形成術 (PCI) は 96% の施設で施行されていた。5 年間で PCI 年間件数は 200 件から 231 件へと増加し ( $p < 0.05$ ), 逆に冠動脈バイパス術は年間 55 件から 48 件と減少傾向を認めた ( $p = 0.06$ )。AMI 年間入院数に変化はなかったが, 平均在院日数は 19 日から 15 日へと短縮し ( $p < 0.0001$ ), 入院中の心リハ実施施設は 53% から 64%, 外来通院型心リハの実施施設は 9% から 21% と増加していた ( $p < 0.05$ )。

【結 論】 循環器専門医研修施設における AMI 後心リハは, 入院中および外来通院型とも 2003 年から 2008 年までに実施率が増加したものの, 在院日数の短縮を考慮すると, 外来通院型心リハの実施率は依然として極めて不十分であることが明らかになった。

(心臓リハビリテーション (JJCR) 16 (2): 188-192, 2011)

Key words : 急性心筋梗塞, 心リハ実施率, 外来通院型心リハ, 全国実態調査

## 1. はじめに

急性心筋梗塞 (AMI) 患者における心臓リハビリテーション (心リハ) は, 運動耐容能, 生活の質 (QOL), 再発予防, 長期予後に対する有効性が確立し<sup>1-3)</sup>, 日本循環器学会や米国の診療ガイドラインでもクラス I として推奨されている<sup>4-7)</sup>。欧米に比べ心リハの普及が遅れていた我が国でも, 近年の施設基準緩和や適応疾患拡大により心リハ実施施設は次第に増加している。

循環器病研究委託費 (15 指-2) 研究班の 2004 年全国実態調査によると, 日本循環器学会認定循環器専門医研

修施設 526 施設のなかで, AMI 入院を受け入れている施設は 97% であったのに対し, 急性期に何らかの心リハを実施しているのは 53%, 外来通院型心リハの実施施設は 9% と報告され, 外来通院型心リハの普及が著しく遅れていることが明らかにされたが<sup>8)</sup>, その後の心リハ実施率の推移は明らかでない。

平成 20 年度から, 循環器急性疾患の急性期から維持期に至るリハビリプログラムを組み込んだ地域連携パスの確立をめざして, 厚生労働省循環器病委託研究 (20 公-7) が 2 年間にわたって続けられ, その一環として 2009 年に〈地域連携パスに関する全国実態調査〉が施行され, 心リハ実施状況が調査された。ここでは同調査

表 1 2004 年および 2009 年調査での施設規模, カテーテル・手術件数, AMI 実績

	2004 年調査 (526 施設)	2009 年調査 (597 施設)	p
全病床数 (床)	469 ± 258	456 ± 241	NS
循環器科病床数 (床)	40 ± 19	42 ± 25	0.09
循環器科常勤医師数 (人)	6.4 ± 6.7	7.4 ± 7.5	< 0.05
CCU 有り	69%	77%	< 0.005
CAG 年間件数 (件)	655 ± 717	640 ± 611	NS
PCI 年間件数 (件)	200 ± 214	231 ± 209	< 0.05
CABG 年間件数 (件)	55 ± 48	48 ± 43	0.06
AMI 年間入院数 (人)	60 ± 50	65 ± 52	0.08
AMI 在院日数 (日)	19 ± 9 日	15 ± 6 日	< 0.0001

CAG: 冠動脈造影, PCI: 経皮的冠動脈形成術, CABG: 冠動脈バイパス術, AMI: 急性心筋梗塞, NS: 有意差なし

と 2004 年全国実態調査の集計結果を比較し, AMI 後の心リハ実施率の推移を調べた。

## 2. 方法

2009 年の〈急性心筋梗塞と脳卒中の地域連携パスに関する全国実態調査〉は, 厚生労働省循環器病研究委託事業 (20 公-7)「循環器疾患の地域連携パスの効果的運用システムの確立に関する研究」(後藤班)が, わが国における急性心筋梗塞および脳卒中の地域連携パスの普及実態・運用状況・メリットと問題点などを明らかにするため, 日本循環器学会認定循環器専門医研修施設・研修関連施設と脳卒中急性期・回復期診療施設を対象として実施された。

調査票は, 病院基本情報, 診療実績, 心臓リハビリテーション, 脳卒中リハビリテーション, 急性心筋梗塞地域連携パス, 脳卒中地域連携パスの項目から成り, 2008 年の実績を調べた。循環器関連施設では 2008 年 11 月に 1240 施設へ調査票を郵送し, 2009 年 6 月までに 780 施設から有効回答が得られた (回答率 63%)。そのうち日本循環器学会認定循環器専門医研修施設が 597 施設, 循環器専門医研修関連施設が 183 施設であった。

2004 年全国実態調査 (15 指-2) では, 2003 年の実績について, 循環器専門医研修施設 526 施設, 循環器専門医研修関連施設 194 施設から有効回答が得られており (回答率 59%), 今回の検討は循環器専門医研修施設から得られた回答 (2004 年調査 526 施設, 2009 年調査 597 施設) をもとに行った。両調査とも回答が得られた循環器専門医研修施設は 291 施設で, 循環器専門医研修施設に限定した回答率は, 2004 年調査 61%, 2009 年調査 64%であった。

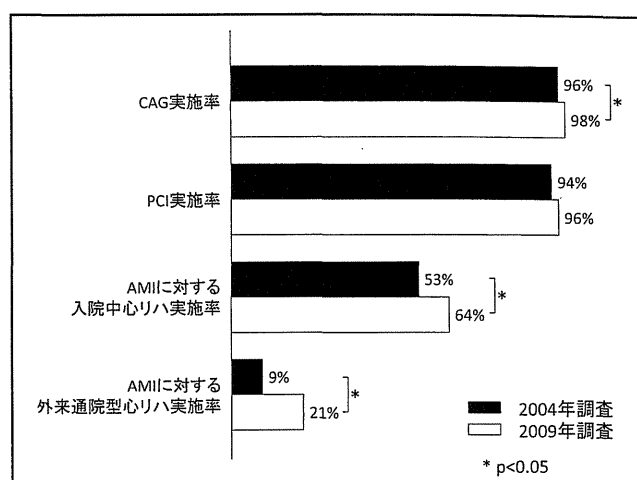


図 1 2004 年および 2009 年調査での心臓カテーテルと AMI 心リハの実施率

各指標は平均 ± 標準偏差で表示し, 二群間の比較には, 対応のない t 検定および  $\chi^2$  検定を用いた。統計学的有意水準は 5% 未満とした。

## 3. 結果

全病床数は平均 450~470 床, そのうち循環器科病床数は全体の 1 割程度で, 2004 年調査と 2009 年調査で有意差はなかった。循環器科常勤医師数は平均 6.4 人から 7.4 人へ, CCU を有する施設の割合は 69% から 77% へと, ともに 2004 年調査より 2009 年調査で有意な増加を認めた (表 1)。

冠動脈造影 (CAG) と経皮的冠動脈形成術 (PCI) の実施率は, 2004 年調査でそれぞれ 96%, 94% であったが, 2009 年調査ではそれぞれ 98%, 96% とさらに増加傾向を認めた (図 1)。CAG 件数は年間 650 件程度で, 両調査間で有意差はなかったが, PCI の年間件数は平均 200 件から 231 件へと有意に増加した。一方, 冠動脈バ