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Alteration of Growth Responses in Established Cardiac Pressure Overload Hypertrophy in Rats with Aortic Banding

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Abstract

We examined the acute effects of elevated wall stress, norepinephrine, and angiotensin II on cardiac protein synthesis as well as protooncogene expression in hearts with established pressure overload left ventricular hypertrophy. Isolated rat hearts with chronic hypertrophy (LVH) were studied 12 wk after ascending aortic banding when systolic function was fully maintained. New protein synthesis (incorporation of [3H]phenylalanine [Phe]) was analyzed in isolated perfused rat hearts after a 3-h protocol; c-fos, c-jun, c-myc, and early growth response gene-1 (EGR-1) mRNA levels (Northern blot) were studied over a time course from 15 to 240 min of perfusion. Under baseline conditions (i.e., before mechanical or neurohormonal stimulation), [3H]-Phe-incorporation (280 nmoles/gram protein/h) and protooncogene mRNA levels were similar in age-matched control and LVH hearts. However, hearts with chronic LVH were characterized by a markedly blunted or absent [3H]-Phe-incorporation after acute imposition of isovolumic systolic load (90 mmHg/gram left ventricle), as well as norepinephrine (10–6 M), or angiotensin II infusion (10–6 M plus prazosin 10–7 M) compared with nonhypertrophied control hearts. Similarly, stimulation of LVH hearts with acute systolic load or norepinephrine was associated with a significantly blunted increase of protooncogene mRNA levels relative to control hearts. The blunted induction of c-fos mRNA in LVH hearts was not due to feedback inhibition, since cycloheximide perfusion of hearts exposed to elevated wall stress further increased the differences between age-matched control and LVH hearts.

The data suggest that acute molecular growth responses to mechanical or neurohormonal stimulation are altered in rat hearts with established LVH relative to nonhypertrophied control hearts. This alteration of molecular adaptations in hearts with compensatory hypertrophy may prevent inappropriate excess cardiac growth in response to mechanical and neurohormonal stimuli. (J. Clin. Invest. 96:2768–2774.) Key words: hypertrophy rat protooncogene angiotensin norepinephrine

Introduction

In hypertension or valvular heart disease the myocardium is exposed to a long lasting pressure overload. Since cardiac myocytes are terminally differentiated cells without the ability to divide, cellular hypertrophy is the only mechanism to normalize the persisting elevation of systolic wall stress under these conditions. In normal myocardium, pathways involved in the induction of cardiac hypertrophy may be activated within a few minutes after the onset of pressure overload, suggesting that myocardial cells are continuously sensing changes in loading conditions (1–4). Early signals of the cellular growth response include induction of protooncogenes and stimulation of overall cardiac protein synthesis (2, 4). Pressure overload persisting for days or weeks induces multiple changes in the pattern of myocardial gene expression that finally result in a substantial remodeling of the heart’s geometry and a fetal pattern of cellular biology (5, 6).

In later stages of chronic pressure overload or in the aging heart the capacity to respond to hypertrophic stimuli may diminish (7–10). The down-regulation of acute molecular responses to growth stimuli may prevent excess myocyte growth in hearts with chronic concentric hypertrophy. On the other hand, exhaustion of compensatory hypertrophy may contribute to cell loss, progressive fibrosis, impairment of diastolic and systolic function, as well as other sequelae of hypertensive heart disease (7, 11–13). However, little is known about signal transduction or molecular adaptations to mechanical or neurohormonal stress in established pressure overload hypertrophy. Thus, in the present investigation, we studied the immediate dynamic molecular responses to well-described mechanical and neurohormonal growth inducing stimuli on cardiac protooncogene induction and protein synthesis in isolated hearts from ascending aortic-banded adult rats with chronic left ventricular (LVH) and age-matched normal controls.

Methods

Preparation of animals. Male Wistar rats (100 gram) were obtained from Charles River Breeding Laboratories (Wilmington, DE). Aortic stenosis was created in anesthetized weaning rats by placing a stainless-steel clip of 0.6 mm internal diameter on the ascending aorta (14, 15). Sham-operation was performed in age-matched controls that underwent a left thoracotomy. Animals were fed a standard rat chow (Purina) and water ad libitum for 8, 12, and 20 wk until used for experimentation.

1. Abbreviations used in this paper: LVH, left ventricular hypertrophy; Phe, phenylalanine.
Perfusion of isolated hearts

Rats were injected intraperitoneally with 25 mg pentobarbital and the thorax was rapidly opened. Next, the hearts were quickly removed and placed in a water-jacketed constant temperature chamber. Retrograde perfusion of the coronary arteries was restored through a short cannula inserted in the aortic root, using a modified Krebs-Henseleit buffer as previously described in detail (14, 15). For measurement of protein synthesis, the buffer contained 0.5% albumin and a mixture of amino acids in the following concentrations: aspartic acid 38 μM, asparagine 64 μM, glutamic acid 207 μM, glutamine 656 μM, glycine 328 μM, alanine 559 μM, valine 226 μM, leucine 184 μM, isoleucine 99 μM, serine 285 μM, threonine 371 μM, methionine 57 μM, proline 246 μM, phenylalanine 400 μM, tyrosine 119 μM, tryptophane 84 μM, histidine 77 μM, lysine 532 μM, and arginine 157 μM (16). The perfusate was equilibrated with a 5% CO2-95% O2 gas mixture such that the perfusate pH was between 7.36 and 7.44 and the PO2 was approximately 550 mmHg. Temperature of the hearts was kept constant at 35°C and monitored with a thermistor probe. Cardiac performance was allowed to stabilize for 10 min of buffer perfusion. Flow was then adjusted to achieve a coronary perfusion pressure of 80 mmHg in control and 100 mmHg in LVH hearts and then kept constant throughout the experimental protocol such that a similar flow per gram left ventricular weight was achieved in control and LVH hearts (10, 14, 15).

Experimental protocols

Time course of functional and morphological adaptations after aortic banding. Groups of 5 to 10 age-matched control hearts, as well as groups of five to ten hearts with LVH were studied after 8, 12, and 20 wk after banding of the ascending aorta. The time points were selected because we have previously established that this encompasses the period from compensated hypertrophy through the transition to failure (17–20). Left ventricular pressure was measured using a fluid-filled latex balloon, slightly larger than the left ventricular chamber, that was inserted into the left ventricle (14, 15) and connected to a Statham P23D8 pressure transducer (Statham Instruments Inc., Puerto Rico). Maximal LV systolic pressure was measured at a balloon volume that resulted in a left ventricular end-diastolic pressure of 10 mmHg.

Effects of wall stress on amino acid incorporation. Groups of six age-matched control hearts, as well as groups of six hearts with LVH, 12 wk after banding, were subjected to an acute elevation of isovolumic systolic wall stress by distention of a fluid-filled balloon in the left ventricular chamber (10). The hearts were initially perfused for 10–15 min with flaccid left ventricles and no left ventricular pressure generation. LV balloon volume was adjusted to achieve a left ventricular developed pressure of 150 mmHg in LVH and 100 mmHg in control hearts resulting in a left ventricular developed pressure per gram of ~90 mmHg/g in both groups. Using previous comparisons of LVH with control hearts, these adjustments allowed the abrupt generation of a peak left ventricular systolic wall stress that was similar in control and LVH groups, using an estimation of peak meridional left ventricular systolic wall stress which was calculated using the method of Isoyama et al. (8).

After 60 min, the generation of systolic pressure overload was stopped by deflation of the intracardiac balloon and hearts were perfused for another 120 min with the modified Krebs-Henseleit buffer in which 0.5 mCi/liter [3H]phenylalanine was added. Thus, the hearts were allowed to incorporate tritiated phenylalanine into newly synthesized proteins for 2 h. Morgan et al. have previously shown that phenylalanine incorporation in isolated hearts is linear for 2 h (21). Unlabeled phenylalanine, present in a defined concentration in the buffer, allowed the calculation of the incorporation of phenylalanine into cardiac proteins on a molar basis (4, 16, 21).

Effects of angiotensin II and norepinephrine on amino acid incorporation. Separate groups of six isolated age-matched control hearts, as well as groups of six hearts with LVH were perfused with (1) norepinephrine (1 × 10^{-8} M); (2) a mixture of angiotensin II (1 × 10^{-8} M) and prazosin (1 × 10^{-7} M) (Sigma Chemical Co., St. Louis, MO); or (3) vehicle. Angiotensin II was combined with the α1-blocker prazosin to prevent any indirect stimulus of protein synthesis via activation of the postynaptic sympathetic nervous system (22). Concentrations used in this experiment were derived from previous dose finding studies (14, 23–27). These experiments were carried out without a balloon in the left ventricle such that the left ventricle was flaccid to avoid stimulation by mechanical load (left ventricular systolic pressure <20 mmHg).

Effects of wall stress and norepinephrine on c-fos, c-jun, c-myc, and early growth response gene-1 expression. The time course of cardiac protooncogene induction in response to an abrupt elevation of systolic wall stress at a level of 90 mmHg/gram was first examined in groups of three isovolumic hearts that were stimulated for 15, 30, 60, 120, and 240 min. The abrupt development of this level of systolic pressure was achieved by inflation of a balloon in the left ventricle to increase left ventricular preload with secondary increase in the level of systolic pressure generation (10). As previously reported, the magnitude of passive diastolic wall stretch employed does not cause protooncogene induction in the absence of systolic pressure generation (10). In additional experiments, the comparison of wall stress or norepinephrine-mediated protooncogene induction in LVH and control hearts was then carried out at the time point of maximal increase in mRNA levels. Groups of five control hearts, as well as groups of five hearts with LVH were stimulated with a left ventricular systolic developed pressure per gram LV of 90 mmHg/gram. Separate groups of five control hearts as well as five LVH hearts without a balloon in the left ventricle were perfused with norepinephrine (1 × 10^{-6} M) for 60 min (c-fos, c-jun), or 120 min (c-myc, EGR-1). Thus, to avoid hemodynamic perturbations, left ventricular pressure was <20 mmHg, and coronary perfusion pressure was adjusted to comparable levels between hearts perfused with norepinephrine or vehicle.

To study whether negative feedback regulation might explain reduced induction of protooncogenes in LVH hearts, four normal control hearts as well as four hearts with LVH were subjected to elevated wall stress in the presence of cycloheximide to allow superinduction of protooncogene mRNA levels (28).

Biochemical analysis

Protein synthesis. After the perfusion protocols, atria and great vessels were quickly removed. Left and right ventricles were blotted dry, balanced, and snap-frozen in liquid nitrogen. For measurement of protein synthesis, the methods of Morgan et al. (21) with modifications by Kent et al. (4) were used. An aliquot (~100 mg) was minced and homogenized in 1 ml ice cold 5% perchloric acid to denature proteins and to remove unincorporated [3H]phenylalanine. After centrifugation, the pellet was washed with 5% perchlorate, resuspended, and heated to 80°C to remove transfer RNA-bound [3H]phenylalanine. After centrifugation, the pellet was washed with 5% perchlorate and then resuspended in 0.2 N NaOH. A small aliquot (50 μl) of this solution was used for protein assay and a second aliquot (500 μl) was used for liquid scintillation counting. Data were corrected for quench by extrapolation. The net protein synthesis by left or right ventricles during the 120 min of perfusion with [3H]phenylalanine was calculated as phenylalanine incorporation (mol/g protein/h) = phenylalanine (dpm/g protein/h)/perfuase phenylalanine specific activity (dpm/mol), where dpm stands for deceleraations per minute.

RNA measurements. Homogenization of left ventricles and RNA extraction was carried out using standard procedures (10, 14, 29). Comparison of relative mRNA levels were made in reference to the same
Table 1. Weight and Function of Isolated Hypertrophied and Normal Hearts at 8, 12, and 20 wk after Ascending Aortic Banding

<table>
<thead>
<tr>
<th></th>
<th>8 wk</th>
<th>12 wk</th>
<th>20 wk</th>
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<tbody>
<tr>
<td></td>
<td>Control</td>
<td>LVH</td>
<td>Control</td>
</tr>
<tr>
<td>n</td>
<td>5</td>
<td>5</td>
<td>10</td>
</tr>
<tr>
<td>BW (grams)</td>
<td>362±7</td>
<td>347±13</td>
<td>386±6</td>
</tr>
<tr>
<td>LV weight (grams)</td>
<td>0.85±0.03</td>
<td>1.10±0.10*</td>
<td>1.05±0.03</td>
</tr>
<tr>
<td>LV/B weight (grams/kg)</td>
<td>2.35±.18</td>
<td>3.11±.19</td>
<td>2.75±.09</td>
</tr>
<tr>
<td>LVP (mmHg)</td>
<td>102±8</td>
<td>147±12*</td>
<td>100±6</td>
</tr>
<tr>
<td>LV dev P (mmHg/gram)</td>
<td>120±8</td>
<td>143±17*</td>
<td>95±7</td>
</tr>
<tr>
<td>LVEDP (mmHg)</td>
<td>10.3±.3</td>
<td>10.5±.2</td>
<td>9.5±.2</td>
</tr>
<tr>
<td>Coronary flow (ml/min per gram)</td>
<td>13.6±.7</td>
<td>13.2±.8</td>
<td>13.1±.6</td>
</tr>
<tr>
<td>LV balloon vol (ml)</td>
<td>.16±.01</td>
<td>.16±.01</td>
<td>.18±.01</td>
</tr>
</tbody>
</table>

Measurements were made in isolated perfused hearts from age-matched control and hypertrophied hearts from rats 8, 12, and 20 wk after ascending aortic banding. Values are expressed in mean±SEM; BW (grams), body weight; LV weight (grams), left ventricular weight; LV/B weight (grams/kg), left ventricular weight in grams per kilogram body weight; LVP (mmHg), left ventricular systolic pressure; LV dev P (mmHg/gram), left ventricular developed pressure in mmHg per gram of left ventricle; LVEDP (mmHg), left ventricular end diastolic pressure; coronary flow in ml per min per gram of left ventricle; LV balloon vol. (ml), left ventricular constant balloon volume. * P < 0.05 vs age-matched controls; † P < 0.01 vs age-matched controls.

Results

Functional and morphological characteristics after aortic banding. The model of ascending aortic banding in weanling rats allows a gradual adaptation of left ventricular function and development of concentric geometry since severity of aortic stenosis increases during growth of the animals (17–20). Table I demonstrates the time course of functional and morphological adaptations in this model. Corroborating our previous observations in this model (17–20), the present experiments show that left ventricular systolic function is preserved in LVH hearts studied 8 and 12 wk after aortic banding; in contrast, by 20 wk post banding contractile function assessed as left ventricular developed pressure per gram is depressed in comparison with age-matched controls at comparable preload. The present study on molecular adaptations to acute mechanical or neurohumoral stimulation was carried out after 12 wk of aortic banding. At this time point, significant left ventricular hypertrophy was achieved but left ventricular systolic function was fully maintained (Table I).

Effects of wall stress on amino acid incorporation. Under previous conditions, i.e. without stimulation by wall stress or neurohormones, the rate of phenylalanine incorporation into newly synthesized proteins was similar in isolated age-matched control rat hearts (287±48 nmoles phenylalanine/g protein/hr) and hearts with LVH after 12 wk of aortic banding (277±49 nmoles phenylalanine/g protein/hr). In response to the abrupt elevation of left ventricular systolic wall stress for 60 minutes (LV developed pressure of 90 mmHg/gram LV; LV systolic meridional wall stress of 287±30×10^3 dyn/cm² in control and 301±30×10^3 dyn/cm² in LVH, respectively, NS) followed by 120 min of [3H]phenylalanine perfusion, the control hearts exhibited a 65% increase of phenylalanine incorporation (P < 0.05 vs baseline), whereas a similar high level of systolic wall stress resulted in a markedly blunted response in LVH hearts (35% increase, P = NS vs baseline) (Fig. 1).

Effects of angiotensin II and norepinephrine on amino acid incorporation. 60 min of angiotensin II/prazosin infusion followed by 120 min of vehicle perfusion resulted in a 390% increase of phenylalanine incorporation in left ventricles of control hearts (P < 0.005 vs baseline). In contrast, the increase in phenylalanine incorporation in left ventricles of LVH hearts was limited to a 53% increase (P = NS vs baseline), that was significantly less than in age-matched controls (P < 0.005 vs controls) (Fig. 2, right panel). Similarly, norepinephrine infusion resulted in a significant increase of phenylalanine incorporation in control hearts, albeit smaller than that seen with angiotensin II (Fig. 2, left panel). As compared with control hearts, the induction of protein synthesis in response to norepinephrine infusion was significantly blunted in the LVH hearts (Fig. 2, left panel).

Effects of wall stress and norepinephrine on protooncogene expression. Since protooncogenes may be important participants in signalling pathways leading to induction of protein synthesis, we studied the effects of the abrupt elevation of wall stress and
Figure 1. Effects of elevated wall stress on amino acid incorporation in normal and hypertrophied rat hearts. LVH hearts had been subjected to 12 wk of aortic banding whereas control hearts had been subjected to sham operation. Each bar represents the mean±SE for six hearts per group; statistics were carried out by ANOVA for multiple comparisons. The symbols − and + represent the absence or presence of the imposition of identical levels of systolic wall stress. Under baseline perfusion conditions, the incorporation of [3H]phenylalanine into cardiac proteins was similar in age-matched control and LVH hearts. However, the increase of new protein synthesis after stimulation with the acute imposition of elevated systolic wall stress was markedly blunted in hypertrophied hearts.

norepinephrine on the induction of the c-fos, c-jun, c-myc and EGR-1 mRNA levels. When hearts were studied at the time point of maximal induction, i.e., 60 min for c-fos and c-jun, and 120 min for EGR-1 and c-myc (data not shown), hypertrophied hearts were characterized by a significantly blunted induction of these protooncogenes relative to the normal control hearts (Fig. 3, left, and Table II). We confirm our previous observation that norepinephrine infusion results in a marked induction of these protooncogenes in normal adult isolated hearts (31). However, the present study shows that this response was significantly blunted or completely absent in hearts with established LVH (Right panel Fig. 3, right, and Table II). We confirm our previous observation that upon wall stress imposition, LVH hearts display a blunted response of c-fos and c-jun mRNA induction (10). In addition, LVH hearts failed to augment EGR-1 and c-myc levels after imposition of left ventricular wall stress (Table II).

To avoid the potential for negative feedback of c-fos protein on c-fos mRNA levels (23), additional hearts were perfused with the inhibitor of protein synthesis cycloheximide in parallel to the abrupt imposition of wall stress or infusion of norepinephrine. The superinduction of protooncogene mRNA in response to relief of feedback inhibition by cycloheximide resulted in significantly higher c-fos levels in normal control as compared to LVH hearts (Fig. 4).

Discussion

LVH is an adaptive cardiac response to the imposition of long-term pressure overload of the heart (6, 32). However, clinical observations suggest that the heart’s capacity to hypertrophy is limited (33–35). The initial benefits of cardiac hypertrophy such as normalization of wall stress and preservation of systolic force generation may be offset during the late stages of chronic hemodynamic overload (6). Progressive cell loss, myocardial fibrosis, and deterioration of cardiac function may ensue (7, 12, 13, 36), suggesting exhaustion or failure to initiate and sustain adaptive cell growth mechanisms. Despite careful investigations that have identified a variety of factors that initiate hypertrophy, little is known about the molecular response to hypertrophic stimuli in hearts with established hypertrophy undergoing the
Elevated systolic wall aortic to chronic pressure of transition controls. perfusion, c-fos (c-fos/GAPDH ratio) c-jun (c-jun/GAPDH ratio) c-myc (c-myc/GAPDH ratio) EGR-1 (EGR-1/GAPDH ratio) have of the disease. progression that the adaptive response was elevated after 8 wk of aortic banding. At 12 wk of aortic banding, the maximal left ventricular to body weight ratio was reached. Therefore, the increase in left ventricular weight was less pronounced and slower than that of body weight, suggesting that the rate of adaptive cardiac growth may slow down during progression of the disease. Further, recent studies from our laboratory have rigorously characterized left ventricular function and alterations of steady state gene expression in this model (17, 18, 20). In particular, we have shown that left ventricular systolic function is well preserved at 12 wk after banding. In contrast, transition to failure with cardiac dilatation and premature death is apparent 20–22 wk after banding (18–20). Thus, cardiac function in this model of left ventricular pressure overload hypertrophy takes a similar course over a compressed time span as that described in human disease (6).

In the present study, the effect of established left ventricular hypertrophy on factors critical for stimulation of cardiac growth was thus studied before cardiac decompensation and interstitial fibrosis were evident. The acute elevation of left ventricular systolic wall stress to similar levels in LVH and control hearts, as well as no norepinephrine and angiotensin II infusions were used to stimulate the early growth responses such as induction of protein synthesis and stimulation of early immediate genes which have been well characterized in normal hearts and isolated myocytes without chronic hypertrophy. In our study, a defined level of elevated systolic wall stress failed to induce protein synthesis in rat hearts with chronic hypertrophy that had been exposed to 12 wk of pressure overload. In contrast, sham operated control hearts responded to similar levels of meridional systolic wall stress with a significant stimulation of amino acid

<table>
<thead>
<tr>
<th>Sham Wall Stress</th>
<th>LVH</th>
<th>Sham Norepinephrine</th>
<th>LVH</th>
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Figure 3. Effects of elevated wall stress and norepinephrine on protooncogene expression in normal and hypertrophied rat hearts. The figure shows representative Northern blot analyses of early immediate gene mRNA levels of age-matched control and LVH hearts stimulated with the abrupt elevation of systolic wall stress (left) or norepinephrine infusion (right). Both systolic wall stress and norepinephrine resulted in significant stimulation of c-fos, c-jun, and c-myc and EGR-1 mRNA levels in age-matched control hearts, whereas a significantly blunted response was observed in LVH hearts.

Table II. Left Ventricular Protooncogene mRNA Levels of Age-matched Control and LVH Hearts at Baseline and after Stimulation with Elevated Systolic Wall Stress or Norepinephrine

<table>
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<tr>
<th>Systolic wall stress</th>
<th>Control</th>
<th>LVH</th>
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<td>- + - +</td>
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<table>
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<tr>
<th>No norepinephrine</th>
<th>Control</th>
<th>LVH</th>
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<td>- + - +</td>
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All values are expressed as normalized relation to GAPDH mRNA levels and expressed in mean ± SEM. Stimulation with elevated systolic wall stress or norepinephrine infusion was carried out for 60 min, respectively; - and + represent without and with stimulation. After 60 min of perfusion, c-fos and c-jun mRNAs were quantified. Hearts for measurements of c-myc and EGR-1 were perfused for another 60 min, without stimulation before quantification of these mRNAs was carried out. * P < 0.05 vs baseline; † P < 0.01 vs baseline; †† P < 0.01 vs age-matched controls.

Figure 4. Effects of cycloheximide on c-fos mRNA after stimulation with elevated wall stress in normal and hypertrophied rat hearts. The figure shows representative Northern blot analyses of c-fos mRNA levels of age-matched control and LVH hearts stimulated with elevated isovolumic wall stress after administration of cycloheximide. Cycloheximide was infused in parallel with the elevation of systolic load to inhibit new protein synthesis, including c-fos protein, and thereby relieve feedback inhibition of c-fos protein on c-fos mRNA transcription. Expression of c-fos mRNA levels was markedly higher in age-matched control hearts as compared to LVH, suggesting that negative feedback mechanisms do not account for differences in c-fos expression between control and LVH hearts.
incorporation into cardiac proteins, corroborating our prior ob-
servations in normal adult perfused rat hearts (31). The data
in the present study extend recent findings of in vivo measure-
ments of cardiac protein synthesis that demonstrated maximal
induction of protein synthesis within 24 h of cardiac pressure
overload but impaired rates of protein synthesis and degradation
during long-term compensatory hypertrophy (36–39). In con-
junction with the data presented here, it may be concluded that
chronic left ventricular hypertrophy is characterized by a
depressed capacity for new cardiac protein synthesis that occurs
eVEN when the imposed stimulus, e.g., meridional wall stress,
is similar in control and LVH hearts.

Neurohormonal activation may substantially contribute to
adaptive growth of the heart (5). In particular, both norepineph-
rine and angiotensin II have been suggested to participate in
the initiation of cardiac hypertrophy (1, 40–43). In agreement
with this hypothesis, the present study corroborates our previous
finding that normal isolated control hearts respond to norepi-
nephrine and angiotensin II with marked stimulation of cardiac
protein synthesis even when the ventricle is completely un-
loaded (31). In contrast, hearts with established LVH displayed
no effect of norepinephrine and angiotensin II on new protein
synthesis, suggesting a markedly blunted growth response to
these neurohormonal stimuli.

We also studied the effects of growth stimuli on the induc-
tion of early immediate genes. We have previously shown that
acute imposition of systolic load induces expression of c-fos
and c-jun protooncogenes in normal isolated perfused hearts,
whereas the response in hypertrophied hearts appeared blunted
(10). The present study extends these findings. In particular, in
contrast to age-matched controls, hearts with established LVH
inhibit markedly blunted increases in c-myc and EGR-1 mRNA
levels when stimulated with elevated wall stress. Likewise, we
now demonstrate that stimulation with norepinephrine results
in a muted increase of c-fos, c-jun, EGR-1 and c-myc levels in
hearts with established LVH. The data reinforce the concept that
the molecular response to major growth stimuli is pro-
doundly down-regulated in hearts with chronic left ventricular
hypertrophy.

A limitation of the present study is that the mechanism of
impaired effects of acute imposition of systolic wall stress and
norepinephrine on early immediate gene induction in LVH can-
not be elucidated. In particular, the question of whether an
intrinsic alteration of myocardial responsivity or receptor down-
regulation may explain our findings in hearts with LVH remains
open. Differences in hemodynamic responses to neurohormonal
stimuli cannot explain findings in the present study since neu-
rohormonal stimulation was carried out in isolated hearts in which
the flaccid left ventricle did not develop pressure, and coronary
vascular pressurization was comparable between groups. Nega-
tive feedback inhibition (28) of mRNA expression in LVH does
not appear to contribute to the differences between the groups,
since differences in c-fos mRNA levels between LVH and con-
trol hearts were still evident when cycloheximide was adminis-
tered to block synthesis of c-fos protein before stimulation.
Similarly, differences in the intracellular response to changes
in systolic signalling may not explain the different levels of
protooncogene expression, since we have previously shown
(10) that calcium ionophore administration results in similar
levels of c-fos and c-jun protooncogenes in normal and hypertro-
phied rat hearts.

With respect to data on down-regulation of the alpha-adren-
ergic and AT1-angiotensin II receptors in LVH (44), it isattrac-
tive to speculate that the impaired response to growth stimuli
may be in part related to alterations at the level of membrane
receptors or receptor coupling. In addition, downstream re-
 sponses to neurohormone receptor activation appear to differ in
hypertrophied versus normal myocytes. For example, we have
recently shown a significant difference in angiotensin and endo-
the lin-induced changes in intracellular pH and calcium in iso-
lated control and hypertrophied myocytes from this model (45).
Thus, the failure of the intact hypertrophied heart to mount a
robust early growth response to these stimuli may reside in
changes in intracellular receptor signaling pathways as well as
membrane receptor availability. The mechanism receptor that
allows sensing of elevated systolic cardiac load has not been
identified yet. Thus further studies are needed to identify the
molecular mechanisms of the blunted acute response to growth
stimuli responses in hearts with established LVH.

In summary, these observations indicate that major growth
stimuli such as the acute elevation of systolic wall stress, norepi-
nephrine, and angiotensin II are far less effective in triggering
the immediate cardiac response in hypertrophied as compared
with normal adult rat hearts. Impaired induction of protein syn-
thesis in rat hearts with established LVH was preceded by a
failure to induce protooncogenes such as c-fos, c-jun, c-myc,
and EGR-1. The data suggest a marked down-regulation of early
signaling pathways involved in cellular adaptation in response
to mechanical and neurohormonal stimulation in hearts with
chronic hypertrophy. In this model, the molecular down-regula-
tion of the response to these growth stimuli occurs before the
development of mechanical decompensation in the hypertro-
phied hearts, suggesting that it may contribute to the transition
from compensatory hypertrophy to failure.

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