Multiparametric Molecular Imaging Provides Mechanistic Insights into Sympathetic Innervation Impairment in the Viable Infarct Border Zone

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Short Title: Impaired Innervation After MI

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ABSTRACT

Impaired catecholamine handling in the viable infarct border zone may play an important role in ventricular remodeling and lethal arrhythmia. We sought to get further biologic insights into cardiac sympathetic neuronal pathology after myocardial infarction, using multiple tomographic imaging techniques.

Methods – In a porcine model of myocardial infarction (n=13), positron emission tomography (PET) and magnetic resonance imaging (MRI) were performed after 4-6 weeks and integrated with electrophysiologic testing and post-mortem histology.

Results - PET with the physiologic neurotransmitter carbon-11 epinephrine, which is sensitive to metabolic degradation unless it is stored and protected in neuronal vesicles, identified a defect exceeding the perfusion defect (defined by nitrogen-13 ammonia; defect size in all animals: 42±12 vs 35±12% of left ventricle, p<0.001). In a subgroup of 7 animals, defect of the metabolically resistant catecholamine C-11 hydroxyepedrine was smaller when compared to epinephrine(41±8 vs 47±6% of left ventricle, p=0.004), while defect of a third catecholamine, C-11 phenylephrine, which is sensitive to metabolic degradation, was similar to epinephrine(48±6 vs 47±6%, p=0.011). Histology confirmed presence of nerve fibers in the infarct border zone. Tagged MRI identified impaired peak circumferential wall strain and wall thickening in myocardial segments with epinephrine/perfusion mismatch (n=6). Confirmatory of prior work, inducible ventricular tachycardia was associated with a larger epinephrine/perfusion mismatch (n=11).
**Conclusion** – In the viable infarct borderzone, neuronal vesicular catecholamine storage and protection from metabolic degradation are more severely altered than catecholamine uptake. This may reflect an intermediate state between normal innervation and complete denervation in advanced disease.

**KEY WORDS**

Positron emission tomography, myocardial infarction, sympathetic innervation, magnetic resonance imaging
INTRODUCTION

Previous studies have suggested that myocardial sympathetic innervation is compromised after myocardial infarction(1-3). High sensitivity of nerve terminals to ischemic damage leads to impaired innervation in the infarct region but also in the viable border zone(4). A detailed understanding of the pathophysiology of poorly innervated but viable myocardium is important, because it may precede arrhythmia or contractile failure, and hence may be useful for guidance of preventive therapy(5-7).

A common limitation of prior cardiac innervation studies is the use of a single neuronal imaging agent. This provides only a snapshot of neuronal integrity. Reduced tracer uptake may be due to absence of nerve terminals, or due to impaired function(8). These two conditions may not be equal with regards to disease progression or regeneration.

Positron emission tomography (PET) with multiple catecholamine tracers may provide mechanistic insights into the state of sympathetic nerve terminals(9). If uptake of one catecholamine analogue is reduced, but another one is avidly retained, functional impairment of a molecular mechanism which defines the kinetics of the first, but not the second agent, is likely, and complete denervation is unlikely. C-11 epinephrine, e.g., is a physiologic neurotransmitter which is sensitive to degradation by cytosolic monoaminooxidase (MAO)(10). Retention may be diminished if it is not taken up by nerve terminals, or if it is taken up but not stored efficiently in neuronal vesicles (where it is protected from MAO). Combination with C-11 hydroxyephedrine, a MAO-resistant catecholamine analogue(11) may distinguish between impaired catecholamine uptake in general, versus functional impairment of vesicular storage. Additional combination with
C-11 phenylephrine, a MAO-sensitive catecholamine analogue which is inefficiently stored in vesicles and shows degradation even under normal physiologic conditions([12]), may be used to confirm presence or absence of the ability to protect neurotransmitter from metabolic degradation.

Hence, it was our goal to obtain more detailed insights into the molecular alterations underlying impaired innervation in the normally perfused infarct border zone. We studied a porcine model of regional myocardial infarction, in which we previously confirmed the feasibility of PET for detecting perfusion/innervation mismatch and its association with ventricular arrhythmia([13]). We speculated that PET with the above described 3 radiolabeled catecholamines, integrated with magnetic resonance imaging (MRI) and ex vivo tissue analysis, would provide mechanistic insights beyond those that are currently available from single neuronal tracer studies.

MATERIALS AND METHODS

Animal Model and Study Protocol

The experimental protocol was approved by the Johns Hopkins Institutional Animal Care and Use Committee. Animals were maintained in accordance with the guiding principles of the American Physiological Society.

Myocardial infarction was induced in 13 young farm pigs (25-35kg), as previously described([13,14]). Under general anesthesia, balloon occlusion of the mid left anterior descending coronary artery (LAD), immediately distal to the second diagonal branch, was performed for 150min. Post-operative treatment included narcotics and non-steroidal anti-inflammatory drugs.
After 4-6 weeks, animals underwent two imaging sessions within 2-3 days: A basic perfusion/innervation PET session was performed in all animals. This was followed by a second PET session with 2 additional catecholamine analogues (n=7), or by MRI (n=6). After the second session, invasive electrophysiology was performed. Immediately thereafter, animals were sacrificed and hearts were excised for post-mortem analysis. Eleven animals completed the protocol. The remaining 2 animals completed non-invasive imaging, but died either before or during electrophysiology.

**Positron Emission Tomography (PET)**

PET was performed under general anesthesia using a GE Advance or a Discovery Rx camera. After supine positioning, a transmission scan was acquired using external sources or X-ray CT, for correction of photon attenuation.

**First Session.** To measure myocardial perfusion at rest, 400-600MBq of N-13 ammonia were injected, and dynamic imaging (21 frames; 12x10, 6x30, 3x300sec) was acquired for 20 min. After a break of 40 min to allow for radioactivity decay, 600-800MBq of C-11 epinephrine, synthesized with specific activity $>1\text{Ci/mmol}$ as previously described(15), were injected. Dynamic imaging (14 frames; 6x30, 2x60, 2x150, 2x300, 2x600sec) was acquired over 40 min.

**Second Session.** After 2-3 days to allow for recovery, 7 animals were re-positioned in the scanner using external skin markers and the laser system of the gantry. C-11 phenylephrine and C-11 hydroxyephedrine were synthesized at specific activity of $>1\text{Ci/mmol}$ as previously described(16,17). 600-800MBq of C-11 phenylephrine were injected first, followed by dynamic imaging over 40min (14 frames; 6x30, 2x60, 2x150, 2x300, 2x600sec). After a break of 60min, 600-800MBq of C-11 hydroxyephedrine were...
injected, followed by dynamic imaging of 40min (14 frames; 6x30, 2x60, 2x150, 2x300, 2x600sec).

For the second PET scan in each session, residual activity from the first injection was ruled out by checking the first frame of the dynamic series, which was acquired before the second dose was injected. The sequence of phenylephrine and hydroxyephedrine scans in session 2 was not randomized due to overall lower myocardial retention and physiological myocardial washout of phenylephrine, reducing the likelihood of residual activity.

To determine contribution of C-11 labelled metabolites to blood activity, venous blood samples were drawn at 1,5,10,20 and 40min after injection of each catecholamine and plasma metabolites were assayed using Sep-Pak cartridges(18).

**Quantitative PET Data Processing.** Attenuation-corrected transaxial PET images were reconstructed by filtered backprojection. PET images were analysed by an observer blinded to MRI and electrophysiology. Using volumetric sampling of the last frame of the perfusion study, myocardial radioactivity was defined in 460 left-ventricular sectors, and depicted in a polar map(19). For reproducible and comparable quantitative analysis, the so defined cardiac long axis and myocardial segments were automatically transferred to dynamic series of all other tracers, and time-activity curves were obtained, as previously established (20,21). Additionally, for each tracer, arterial input function was defined by a small region of interest in left ventricular cavity. Absolute myocardial blood flow at rest was quantified using a validated 3-compartment model for N-13 ammonia(17). For the C-11 labelled tracers, as previously described(13,18), a retention index was calculated by normalizing myocardial activity at 40min to the integral under the metabolite-corrected
arterial input function. Finally, to determine clearance, monoexponential fitting of myocardial curves between 5 and 40 min yielded washout rate constants (22).

**Polar Map Analysis.** In order to define defect size in percent of left ventricular (LV) myocardium, a threshold of 60% of the maximum was used (23). The reproducibility of this approach has been previously reported (19). Also, areas of the polar map which showed a matched perfusion/epinephrine defect (infarct), an epinephrine defect but no perfusion defect (mismatch, border zone), and no defect at all (remote myocardium) were compared.

**Magnetic Resonance Imaging (MRI)**

**Image Acquisition.** MRI was performed in 6 animals, using a 3T human scanner (Achieva, Philips Medical Systems, Best, NL) equipped with vector ECG technology and a 6-element phased array cardiac coil. Spiral CSPAMM grid-tagged images (24) were acquired for motion vector analysis throughout the cardiac cycle. To avoid off-resonance effects due to fat, a spectral spatial excitation was used. Additionally, cine imaging and delayed enhancement imaging of gadolinium-DTPA was performed.

**Data Analysis.** MRI was analysed by a reader blinded to PET results. In cine and delayed enhancement images, endocardium and epicardium were manually contoured at each short-axis level in endsystole and enddiastole in order to calculate LV volumes, ejection fraction and mass, as well as regional wall thickening and thickness. Harmonic phase analysis was conducted for tagged studies to determine peak circumferential myocardial strain throughout the cardiac cycle and time to peak strain in eight segments of left ventricular myocardium (25). For accurate matching with PET, septum on all MRI studies was defined as the area between anterior and posterior insertion of the RV, and
the other myocardial walls were defined accordingly. The apex, which is not appropriately imaged by short-axis images, was excluded.

**Electrophysiologic Testing**

An abbreviated invasive electrophysiologic workup consisted of programmed stimulation to identify inducibility of ventricular tachycardia (VT) as previously described in more detail(13).

**Post-Mortem Analysis**

After completion of the study protocol, anesthetized pigs were euthanized (4mmol/L potassium chloride IV). Hearts were immediately removed and gross macroscopic left ventricular short-axis slices were created. Under guidance by PET images and gross morphology, tissue samples were collected from the central infarct region, the infarct border zone at the location of PET perfusion/innervation mismatch, and remote non-infarct region, for further analysis. Samples were frozen and sliced for general histology by hematoxylin/eosin (HE) staining. Additionally, immunohistochemical staining was performed using anti-tyrosine hydroxylase (TH) antibody in order to detect sympathetic nerve fibers, and anti-growth-associated protein 43 (GAP43) antibody for detecting nerve sprouts, as previously described (monoclonal mouse anti-GAP43 and anti-TH, respectively, 1:50 dilution; Chemicon International, Inc)(26). Image J software (NIH, Bethesda, MD) was used to quantify the percentage of intact tissue in HE stains, and to determine the number of positively stained nerve fibers per field of view in TH and GAP43 stains(26).
Statistical Analysis

Statistical analysis was carried out using Medcalc (Mariakerke, Belgium). All data are shown as mean±standard deviation. P<0.05 was considered statistically significant. Mann-Whitney U-test was used to compare groups with and without VT inducibility. Wilcoxon test was used to compare global perfusion and innervation defect sizes and mismatch and remote myocardium intraindividually. Kruskal-Wallis test was used for comparison of PET and MRI variables between three types of segments and Friedman test was employed to compare defects from 3 different catecholamine tracers. Pearson’s correlation was used to describe univariate relationship between continuous variables. Multivariate stepwise regression analysis was performed to identify independent determinants of MRI-derived strain parameters. To support the assumption of independence of myocardial segments, a correlation analysis was performed using the most important PET variable, the mismatch between perfusion and innervation abnormality. Values in the distal anterior wall (the predominant site of mismatch), did not correlate with corresponding values from any of the eight other segments.

RESULTS

Innervation/Perfusion Mismatch In The Infarct Border Zone

In all animals, global perfusion defect size was 35±12% LV and epinephrine retention defect size was significantly larger at 42±12% (p<0.001). This resulted in an innervation/perfusion mismatch comprising 7±4% LV in the infarct border zone (figure 1). Programmed stimulation showed VT inducibility in 5/11 animals. Consistent with
prior work(13), animals with inducible VT had significantly larger innervation/perfusion mismatch (11±4 vs 5±3% LV; p=0.03).

**Innervation/Perfusion Mismatch and Regional Strain**

In the MRI subgroup (n=6), LVEF was 35±9%, and there was mostly anteroseptal transmural delayed enhancement. 6/48 myocardial segments showed a matched reduction of perfusion and relative epinephrine retention, and 10 showed innervation/perfusion mismatch. In mismatch segments, perfusion, absolute flow, and enddiastolic wall thickness were similar to normal segments, while epinephrine retention, wall thickening and peak strain were significantly impaired when compared to normal segments (table 1, figure 2). Peak strain correlated significantly with regional perfusion (r=-0.55; P<0.001) and epinephrine retention (r=-0.64; P<0.001), but only epinephrine retention was an independent correlate at stepwise multiple regression analysis.

**Epinephrine, Hydroxyephedrine and Phenylephrine Handling**

In the dual PET session subgroup (n=7), defect size for the physiologic neurotransmitter epinephrine was larger than the perfusion defect (figure 3). Hydroxyephedrine defect, however, was significantly smaller than epinephrine and not different from perfusion defect size. Phenylephrine defect was similar to epinephrine, but larger than hydroxyephedrine and perfusion defect.

Table 2 summarizes retention and washout kinetics of the 3 tracers in myocardial regions. Reduced regional retention in the viable infarct border zone was due to significantly elevated washout of epinephrine and phenylephrine, while there was no washout of hydroxyephedrine.
Ex Vivo Analysis

Histology confirmed extensive tissue damage in the infarct, and mild damage but mostly preserved tissue integrity in the border zone (figure 4). Immunostaining in the infarct area was not conclusive due to lack of viable tissue. In the border zone, immunohistochemistry revealed presence of TH-positive nerve fibers and GAP43-positive nerve sprouts. Quantitatively, the number of TH- (0.58±0.14 vs 0.79±0.24 per field of view; p=0.28) and GAP43-positive fibers (0.83±0.29 vs 1.52±0.56 per field of view; p=0.08) were not different from remote myocardium.

DISCUSSION

Our study confirms prior experimental and clinical work which showed impaired sympathetic innervation in the viable infarct border zone that has electrophysiologic implications and contributes to ventricular arrhythmia(3,7,13). Integrated analysis of PET and MRI also shows that an innervation/perfusion mismatch after myocardial infarction is associated with altered regional contractility and wall strain despite preserved perfusion and viability. Most importantly, however, interrogation of the myocardial kinetics of multiple different catecholamine analogues suggests that the innervation/perfusion mismatch in the infarct border zone is associated with functionally impaired innervation in our model. Neuronal vesicular catecholamine storage and protection from metabolic degradation are more severely altered than catecholamine uptake.

In healthy myocardium, the physiologic catecholamine epinephrine is taken up by sympathetic nerve terminals through the norepinephrine transporter, then stored in neuronal vesicles, where it is protected from monoaminoxidase (MAO) degradation
In our study, epinephrine retention is reduced in the normally perfused infarct border zone. In contrast, hydroxyepinephrine, a catecholamine analogue which is an avid substrate for neuronal transport but is resistant to MAO degradation and therefore depends less on vesicular storage for neuronal retention, showed a smaller defect when compared to epinephrine. This suggests that neuronal uptake mechanisms may at least be partially preserved in the infarct border zone, while vesicular storage is more severely impaired. The presence of increased washout of epinephrine in the border zone provides further support, as do the findings using a third catecholamine, phenylephrine. This MAO-sensitive neurotransmitter analogue is taken up by nerve terminal and stored in vesicles, but even under physiologic conditions there is vesicular leakage and subsequent washout. Concordant with epinephrine, regional phenylephrine retention was reduced in the infarct border zone and washout was increased, confirming the presence of impaired vesicular storage rather than complete absence of intact nerve terminals in this region in our model of well defined, regional myocardial damage.

Immunohistologic detection of intact nerve fibers and nerve sprouting in the infarct border zone of our model is consistent with prior work and provides more support of a functional impairment of sympathetic innervation. Of note, the histologic presence of nerve fibers alone does not allow for conclusions about their functional state. Molecular imaging provides further biologic insights and suggests that neuronal transport may still be intact in histologically detectable nerves, while vesicular storage is substantially impaired.

It should be noted that some of our findings may be specific to our animal model. Firstly, myocardial infarction was induced in otherwise healthy, young pigs without
concomitant disease. In the clinical situation, infarction is associated with more or less extensive coronary artery disease. There, repetitive ischemic insults may result in more severe neuronal damage in otherwise viable myocardium and hence, in more severe defects for the uptake marker hydroxyephedrine. Consistently, a prior study using hydroxyephedrine in humans showed retention defects exceeding the perfusion defect in a fraction, but not all patients early after myocardial infarction (29). Also, the recent PAREPET study, which identified hydroxyephedrine defects exceeding perfusion defects and predicting arrhythmia risk (30), was conducted in subjects with ischemic cardiomyopathy and heart failure and thus covered a more severe state of ischemic damage. Our animal model does not reflect chronic ischemia, nor multi-vessel disease or longstanding heart failure. The present results thus probably reflect a less profound state of neuronal impairment which may precede a more severe impairment in advanced ischemic heart disease.

The primary purpose of our work was to highlight the fact that impaired sympathetic innervation is not a black and white phenomenon of denervation or innervation, but that there are transitional stages of incomplete molecular impairment which can be identified using multiple catecholamine imaging agents. The proof of feasibility for this approach is provided in the present study. The practical implications of molecularly impaired but not completely denervated myocardium for the clinical course of ischemic heart disease will need to be clarified in subsequent work. Our initial results may serve as a basis for this.

Secondly, species differences in catecholamine kinetics have been reported. In pigs, extraneuronal clearance of catecholamines by the uptake-2 system is more
pronounced than in primates(31). Also, there may be differences in MAO activity(32) which may interfere with intraneuronal washout rates. But we combined 3 tracers to determine nerve terminal function, and relative comparison of their kinetics can be done independent of species properties. Thirdly, the dependency of hydroxyephedrine and epinephrine on vesicular storage requires some additional discussion. It has been reported that reserpine, a blocker of vesicular storage, reduces hydroxyephedrine retention in vivo by up to 50%(33), but this may be partially attributed to systemic effects (resulting from competition by increased norepinephrine levels and effects on ventricular loading). Other studies in isolated perfused hearts have shown a more severe, almost complete blockage of epinephrine retention by reserpine(10). Taken together with the effect of a desipramine chase on hydroxyephedrine only(9) and the MAO-sensitivity of epinephrine, necessitating vesicular storage for protection from metabolism, epinephrine retention is clearly more dependent on vesicular storage than that of hydroxyephedrine. Forth, all scans were performed during isoflurane anesthesia. Isoflurane increases sympathetic tone(34), but the net effect on kinetics of all 3 tracers is expected to be similar so that relative comparison of their kinetics is not affected. And finally, spatial correlation of histology with imaging is challenging. We used the infarct region as landmark, but misalignment cannot be completely excluded. It should be considered, however, that our histologic analysis confirms prior work, which provides some validation to the approach.

Our work may have practical implications: Speculatively, although not proven in this study, recovery of a state of impaired nerve function may be easier to achieve than in case of complete denervation. Also, while the current clinical focus for innervation imaging is on heart failure(30, 35), studies such as ours suggest that additional focus on
acute myocardial infarction as a precursor of arrhythmia and heart failure, and on the analysis of regional abnormalities rather than global impairments could be valuable. And finally, while a more widespread use of the carbon-11 tracers used in this project cannot be expected due to the dependency on a cyclotron and on-site radiopharmacy, our work suggests that the kinetics of more widely available neuronal imaging agents such as iodine-123 metaiodobenzylguanidine, or a new fluorine-18 labeled neuronal PET imaging agent(36), should be carefully evaluated in order to identify the major molecular mechanisms which determine their kinetics and the respective imaging signal.

CONCLUSION

Using a complex PET imaging protocol with multiple catecholamines and adjunct functional and electrophysiologic tests in an established pig model of myocardial infarction, the present study suggests that neuronal vesicular catecholamine storage and protection from metabolic degradation are more severely altered than catecholamine uptake in the viable infarct borderzone. This may reflect an intermediate state between normal innervation and complete denervation. These results provide valuable pathobiologic insights and they may serve as a foundation for the design of future studies investigating the clinical value of neuronal imaging in more depth.
FUNDING SOURCES

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DISCLOSURES

There are no conflicts of interest to disclose in relation to the subject matter of the article.
REFERENCES


**Figure 1.** Perfusion/innervation mismatch after myocardial infarction. (A.) Representative PET polar maps of myocardial perfusion (N-13 ammonia, left) and innervation (C-11 epinephrine, right). Innervation defect (white area) is larger than the perfusion defect (blue area), resulting in normally perfused myocardium with impaired innervation in the infarct border zone. (B.) Bar chart showing defect sizes in all animals (n=13). (C.) Bar chart comparing perfusion/innervation mismatch in subgroups with (n=5) and without (n=6) inducible ventricular tachycardia (VT).
Figure 2. Circumferential myocardial strain analysis. (A.) Matched short axis slices of the mid/distal left ventricle in a representative animal. PET perfusion (using N-13 ammonia) and MRI delayed enhancement (scar) show anteroseptal infarct. PET innervation (using C-11 epinephrine) shows defect which exceeds perfusion defect (innervation/perfusion mismatch). Color-coded tagged MRI of circumferential strain is shown on the right. (B.) Regional strain in representative segments. Curves throughout the cardiac cycle are shown for scar (red), mismatch (blue), and normally perfused remote myocardium (green). Infarct curve is irregular with attenuated and delayed nadir. Nadir in mismatch region is attenuated versus remote. (C.) Bar charts comparing peak strain and time to peak in 3 different groups of segments (*p<0.05 vs remote).
**Figure 3.** Characterization of myocardial innervation using multiple radiolabeled catecholamines. (A.) Polar maps of a representative infarct animal showing retention of 4 tracers (top), along with color-coded defect areas. Epinephrine and phenylephrine defects exceed perfusion defect, while hydroxyephedrine defect is comparable to perfusion. (B.) Bar chart showing group results of defect sizes.
**Figure 4.** Ex vivo tissue analysis. (A.) Gross morphologic short axis slice (top left) indicates location of sampling for microscopic analysis from scar (1), border zone (2) and remote area (3). Hematoxylin/eosin (HE) stains show extensive tissue damage in (1), mild damage/fibrosis in (2) (arrows), and preserved integrity in (3). Quantitative analysis (bar chart, right) confirms visual impression (*p<0.01 vs remote region). (B.) Immunohistochemical stains for tyrosine hydroxylase (TH) and growth-associated protein (GAP) 43 show brown-stained nerve fibers/sprouting nerves (arrows) in infarct border zone (2) and remote region (3).
### Table 1. PET and MRI Results in Myocardial Segments

<table>
<thead>
<tr>
<th>Segmental Pattern</th>
<th>Normal (n=32)</th>
<th>Mismatch (n=10)</th>
<th>Scar (n=6)</th>
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<tr>
<td>Relative Perfusion (% of LV max)</td>
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<td>78±7</td>
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<td>Absolute Flow (ml/min/g)</td>
<td>0.76±0.21</td>
<td>0.79±0.21</td>
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<td>Relative Epinephrine Uptake (% of LV max)</td>
<td>84±9</td>
<td>66±10*</td>
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<td>17.3±2.0*</td>
<td>9.6±1.3*†</td>
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<td>Enddiastolic Wall Thickness (mm)</td>
<td>7±1</td>
<td>7±1</td>
<td>5±1*†</td>
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<tr>
<td>Wall Thickening (%)</td>
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<td>24±25*</td>
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<td>Peak Strain (%)</td>
<td>-14±4</td>
<td>-8±5*</td>
<td>-4±5*</td>
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<tr>
<td>Time to Peak Strain (% of RR Interval)</td>
<td>50±7</td>
<td>52±8</td>
<td>70±7*†</td>
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</table>

*P<0.05 vs normal; † P<0.05 vs mismatch; LV, left ventricular; RR, cardiac cycle
Table 2. Kinetics of 3 Radiolabeled Catecholamines After Myocardial Infarction

<table>
<thead>
<tr>
<th></th>
<th>Myocardial Region</th>
<th>Normal</th>
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<tr>
<td>Retention Index</td>
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<td>18.1±3.1*</td>
<td>9.9±1.3*†</td>
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<td>1.5±0.4*</td>
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<td><strong>Hydroxyephedrine</strong></td>
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<tr>
<td>Retention Index</td>
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<td>13.5±4.7</td>
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<tr>
<td>Washout Rate</td>
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<td>- (§)</td>
<td>n/a (‡)</td>
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<tr>
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<tr>
<td>Retention Index</td>
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<td>11.5±1.5*</td>
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<td>Washout Rate</td>
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<td>1.2±0.3*</td>
<td>n/a (‡)</td>
<td></td>
</tr>
</tbody>
</table>

*P<0.05 vs normal; †P<0.05 vs mismatch; ‡due to lack of specific uptake, washout rates in scar do not reflect catecholamine turnover; §hydroxyephedrine uptake was stable over time, washout rates were 0 or <0