In Vivo PET Imaging of Cardiac Presynaptic Sympathoneuronal Mechanisms in the Rat

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The sympathetic nervous system of the heart plays a key role in the pathophysiology of various cardiac diseases. Small-animal models are valuable for obtaining further insight into mechanisms of cardiac disease and therapy. To determine the translational potential of cardiac neuronal imaging from rodents to humans, we characterized the rat sympathetic nervous system using 3 radiotracers that reflect different subcellular mechanisms: ¹¹C-meta-hydroxyephedrine (HED), a tracer of neuronal transport showing stable uptake and no washout in healthy humans; ¹¹C-phenylephrine (PHEN), a tracer of vesicular leakage and intraneuronal metabolic degradation with initial uptake and subsequent washout in humans; and ¹¹C-epinephrine (EPI), a tracer of vesicular storage with stable uptake and no washout in humans. Methods: We used a small-animal PET system to study healthy male Wistar rats at baseline, after desipramine (DMI) pretreatment (DMI block), and with DMI injection 15 min after tracer delivery (DMI chase). The rats were kept under general isoflurane anesthesia while dynamic emission scans of the heart were recorded for 60 min after radiotracer injection. A myocardial retention index was determined by normalizing uptake at 40 min to the integral under the arterial input curve. Washout rates were determined by monoexponential fitting of myocardial time-activity curves. Results: At baseline, HED showed high myocardial uptake and sustained retention, EPI showed moderate uptake and significant biphasic washout, and PHEN showed moderate uptake and monoexponential washout. The average (\pm SD) left ventricular retention index for HED, PHEN, and EPI was 7.38% \pm 0.82%/min, $3.43\% \pm 0.45\%$ /min, and $4.24\% \pm 0.59\%$ /min, respectively; the washout rate for HED, PHEN, and EPI was 0.13% \pm 0.23%/min, 1.13% \pm 0.35%/min, and 0.50% \pm 0.24%/min, respectively. The DMI chase resulted in increased washout only for HED. DMI block decreased myocardial uptake of all tracers by less than 90%. Conclusion: Kinetic profiles of HED in the rat myocardium were similar to those of HED in humans, suggesting comparable neuronal transport density. Unlike in humans, however, significant washout of EPI and faster washout of PHEN were encountered, consistent with high intraneuronal metabolic activity, high catecholamine turnover, and reduced vesicular storage. This evidence of increased neuronal activity in rodents has implications for translational studies of cardiac neuronal biology in humans.

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The sympathetic and parasympathetic cardiac neuronal networks play an important role in the regulation of cardiovascular performance. Although parasympathetic innervation exerts inhibitory effects via activation of muscarinic receptors, sympathetic innervation exerts a cardiac stimulatory effect (increased force of contraction and acceleration of heart rate and conduction) via activation of adrenergic receptors by norepinephrine (1,2). Impairments of the functional integrity of sympathetic nerve terminals in the heart have been shown to be a key component of diseases such as congestive heart failure, dysrhythmia, ischemia, dysautonomia, Parkinson's, and diabetes (3–8). PET allows for the noninvasive assessment of cardiac sympathetic neuronal integrity using several radiolabeled catecholamine analogs (9).

¹¹C-*meta*-hydroxyephedrine (HED), a metabolically resistant catecholamine analog, is recognized as a tracer of neuronal transport (*10*). ¹¹C-phenylephrine (PHEN) is another catecholamine analog that, in contrast to HED, is a substrate to monoamine oxidase (MAO) and thus metabolized intraneuronally. It is considered a marker of leakage from intraneuronal storage vesicles and subsequent metabolism (*11*). ¹¹C-epinephrine (EPI) is a true catecholamine that, in contrast to HED and PHEN, is efficiently stored in intraneuronal vesicles, where it is protected from metabolic degradation under normal conditions. It is thus considered to be a marker of neuronal uptake and vesicular storage (*12*). Figure 1 illustrates the neuronal pathways tracked by these [**Fig. 1**] 3 tracers.

All 3 radiotracers have been used in humans to assess the sympathetic nervous system. HED and EPI showed stable uptake and no washout, whereas PHEN showed uptake and subsequent washout in healthy volunteers (13-15). The kinetic profiles of these radiotracers were also studied in

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FIGURE 1. Schematic display of neuronal pathways tracked by 3 different sympathoneuronal PET tracers.

vitro in the isolated perfused rat heart model. However, tracer kinetics in the living rat may not be similar to isolated heart perfusion studies because of systemic effects and central-peripheral nervous system interaction.

The rat is a well-established animal model for cardiovascular research and drug discovery. Although the use of a single tracer provides a snapshot of the integrity of innervation, we speculated that the combination of multiple tracers with different kinetic properties would provide further insights into sympathetic nerve terminal biology. Thus, to determine the translational potential of in vivo cardiac neuronal imaging from rats to humans and vice versa, we sought to characterize the rat sympathetic nervous system using HED, PHEN, and EPI together with small-animal PET methodology.

MATERIALS AND METHODS

Radiopharmaceuticals

HED, PHEN, and EPI were synthesized as previously described (11,16,17). The specific radioactivities at the end of synthesis were in the range of 370–740 GBq/ μ mol and the radiochemical purities were greater than 98%.

Animal Preparation

All animal procedures were performed in strict accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals (18) and approved by Johns Hopkins University Animal Care and Use Committee. Male Wistar rats (250-350 g) were housed in pairs, allowed food and water ad libitum, and studied. For PET, animals were anesthetized with 1.5% of isoflurane in oxygen and kept in the prone position on the scanner bed with the heart region centered in the gantry. The radiotracers were injected intravenously via the tail vein, and an aqueous solution of desipramine (DMI) hydrochloride was injected intraperitoneally before or after radiotracer injection, if applicable. Throughout the scanning procedure, body temperature was monitored by a rectal probe and maintained between 36.5°C and 37.5°C by a heating lamp. A single PET scan with 1 tracer under one of the subsequently described conditions was performed per animal, and after the PET scan the animals were euthanized with an overdose of isoflurane.

Animal Groups

Animals were assigned to 1 of 4 groups. In group 1 (control), each radiotracer—HED, PHEN, and EPI—was used in 4 animals without pharmacologic intervention. In group 2 (DMI chase), 4 animals per tracer were injected with radiotracer, and after 15 min into the PET scan, they received a 2-mg dose of DMI solution per kilogram intraperitoneally to inhibit reuptake of radiotracer by norepinephrine transporter (NET). In group 3 (DMI pretreatment), experiments were performed on a single rat per tracer. The animal received a 2-mg dose of DMI solution per kilogram intraperitoneally 15 min before injection of each of the 3 radiotracers to inhibit uptake of radiotracer by NET. In the fourth group, 4 animals were injected with low-specific-radioactivity EPI (9–18 MBq/µmol) without pharmacologic intervention.

PET

Imaging was performed using the Johns Hopkins University small-animal PET system (eXplore Vista; GE Healthcare). The characteristics of this scanner have been described previously (19). In brief, the system has a transaxial field of view (FOV) of 6.7 cm and an axial FOV of 4.8 cm, achieving an in-plane spatial and volume resolution of approximately 1.4 mm and 2.9 mm³, respectively.

Under anesthesia, animals were placed in the prone position in the scanner gantry, and radiotracer (18.5 MBq) was injected via the tail vein, 10–15 min after synthesis. Image acquisition and radiotracer injection started simultaneously, and a dynamic sequence (40 frames: 6×5 s, 10×15 s, 12×30 s, 6×60 s, 3×300 s, and 3×600 s) was acquired over 60 min.

Image Reconstruction and Time-Activity Curves

Transaxial images were reconstructed by a 2-dimensional iterative algorithm (ordered-subset expectation maximization, 1 iteration, and 16 subsets). Image data were not corrected for attenuation and scatter. For an organ-based analysis, heart and liver were identified on PET images, and regions of interest (ROIs) associated with the left ventricle (LV) chamber, LV lateral wall, LV septum, LV apex, and liver were drawn. ROIs initially indicate values in radioactivity per cubic centimeter. We used the injected amount of radioactivity to transform values to percentage injected dose per cubic centimeter. Then, assuming a relative density of 1 g/cm³, multiplication with the true animal weight in grams was performed to obtain percentage of standardized uptake value (SUV). This was

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done to normalize for the differences in rat weight and administered dose and to allow for comparison of intergroup with interorgan data. In animal groups 1, 2, and 4, ROIs were drawn on the tomographic images on frame 40 (last frame). In group 3, because of absence of specific uptake in later frames, an earlier frame (frame 10) was selected. Decay-corrected time–activity curves were presented as percentage SUV; all percentage SUVs in the study presented a mean of measurements in 4 rats with each of the 3 radiotracers in group 3, was obtained from a single rat.

Retention Index and Washout Rate

For detailed analysis of myocardial kinetics, volumetric sampling was used to create polar maps of activity distribution throughout the left ventricular myocardium (20,21). A non-metabolite-corrected input function was derived from a circular ROI in the LV cavity. Then, a non-metabolite-corrected radiotracer retention index was determined by normalizing myocardial tracer activity from 30 to 40 min to the integral of the arterial input function. The clearance of radiotracers was defined from data between 10 and 60 min after injection that were fitted to a monoexponential function and expressed as %/min. The average non-metabolite-corrected LV myocardial retention index and washout rate were computed from septal, lateral, anterior, inferior, and apical regions.

Statistical Analysis

Data are reported as mean \pm SD, which was calculated by averaging all ROIs for all animals in a given study to include both subject and group variability for SD. Comparisons of 2 mean values were performed with an unpaired Student *t* test. A *P* value of less than 0.05 was considered significant.

RESULTS

The LV and liver were visualized clearly after HED, [Fig. 2] PHEN, and EPI injection in untreated control rats (Fig. 2). The contrast and quality of HED tomographic images were better than those for PHEN and EPI (radioactivity ratio of LV lateral wall to liver at 55 min was 0.96, 0.44, and 0.48 for HED, PHEN, and EPI, respectively). Tracer kinetics in heart and liver differed significantly among tracers, as described in the following sections.

HED

HED showed rapid uptake and sustained retention in the rat myocardium without medical intervention (Fig. 3A). [Fig. 3] Using the DMI chase protocol, initial retention was followed by rapid washout after DMI administration (Fig. 3B). After DMI pretreatment, rapid washout without retention in the myocardium was observed (Fig. 3C). The average non-metabolite-corrected LV retention index at the end of the scan was $7.38\% \pm 0.82\%/\text{min}$ and $3.28\% \pm 0.60\%/\text{min}$ for the control group and DMI chase. In the single animal after DMI pretreatment, it was $0.21\% \pm 0.35\%/\text{min}$ (Table 1). The LV [Table 1] retention index was lower in DMI chase (P < 0.05) than in untreated rats. The average LV washout rate of HED was significantly faster after DMI chase ($1.86\% \pm 0.57\%/\text{min}$) than in controls ($0.13\% \pm 0.23\%/\text{min}$) (Table 1).

In the liver, HED displayed slow uptake in control rats, slow initial uptake followed by more rapid uptake after DMI chase, and rapid uptake immediately after radiotracer injection after DMI pretreatment. The PET-measured HED liver uptake values at 30 s and 55 min in controls, DMI chase, and DMI pretreated rats were 300% and 500% SUV, 300% and 600% SUV, and 600% and 800% SUV, respectively (Figs. 3A–C).

PHEN

PHEN showed rapid uptake followed by monoexponential washout in the myocardium of control rats (Fig. 4A). Myo- [Fig. 4] cardial kinetics of PHEN in controls and after DMI chase were mostly similar (Figs. 4A and 4B). DMI chase did not enhance washout of PHEN in the myocardium. In the single DMI pretreated rat, PHEN showed rapid washout without retention in the myocardium (Fig. 4C). The average non-metabolite-corrected LV retention index of PHEN radio-activity at the end of the scan was $3.43\% \pm 0.45\%/min$,



FIGURE 2. Representative static coronal PET images and reangulated, smoothed cardiac images of rats, acquired between 50 and 60 min after intravenous injection of HED, PHEN, or EPI. Shown on right are midventricular short-axis (SA), vertical long-axis (VLA), and horizontal long-axis slices (HLA). Liver is visualized inferior to left ventricle.

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FIGURE 3. Myocardial time–activity curves for LV chamber (\Box), LV lateral wall (\triangle), LV septum (∇), LV apex (\diamond), and liver (*) in rats after intravenous injection of HED, without pharmacologic intervention (A), with intraperitoneal administration of 2 mg of DMI solution per kilogram 15 min after tracer injection (DMI chase) (B), and after intraperitoneal administration of 2 mg of DMI solution per kilogram 15 min before tracer injection (DMI block) (C). A and B show data averaged from 4 rats, and C shows observation from 1 rat (LV septum and LV apex were included in FOV but not detectable in C; kinetics in anterior wall were also analyzed but did not differ from other myocardial regions and are therefore not shown).

 $2.95\% \pm 0.51\%$ /min, and $0.24\% \pm 0.25\%$ /min, respectively, for control group, DMI chase group, and the single DMI pretreated rat (Table 1). The average LV washout rate of PHEN was not significantly faster in DMI chase rats $(1.51\% \pm 0.16\%$ /min) than in the control group $(1.13\% \pm 0.35\%$ /min) (Table 1).

In the liver, PHEN showed rapid uptake and subsequent washout between 0 and 15 min followed by sustained retention until 60 min. The PET-measured PHEN liver uptake at 30 s and 55 min in the control group, DMI chase, and DMI pretreated rats were 600% and 450% SUV, 600% and 450% SUV, and 650% and 450% SUV, respectively (Figs. 4A–C).

EPI

EPI showed rapid uptake and subsequent biphasic washout from the myocardium with a first rapid phase followed by slower clearance and retention of radioactivity in the second did not affect myocardial tracer kinetics significantly (Figs. 5B and 5C). Myocardial retention was not observed in the single DMI pretreated rat (Fig. 5D). The average non-metabolite-corrected LV retention index at the end of the scan was $4.24\% \pm 0.59\%$ /min, $3.87\% \pm 0.81\%$ /min, $4.02\% \pm 0.99\%$ /min, and $0.22\% \pm 0.30\%$ /min, respectively, for the control group, low-specific-radioactivity group, DMI chase group, and the single DMI pretreated rat (Table 1). The average LV washout rate of EPI was not significantly faster in DMI chase rats ($0.50\% \pm 0.24\%$ /min) and the low-specific-radioactivity group ($0.61\% \pm 0.28\%$ /min) than in the control group ($0.69\% \pm 0.17\%$ /min) (Table 1).

In the liver, EPI showed rapid uptake, followed by sustained retention until 60 min. The PET-measured EPI liver uptake at 30 s and 55 min in control, low-specific-radioactivity, DMI chase, and DMI pretreated rats was 400% and 550% SUV, 500% and 700% SUV, 450% and 550% SUV, and 550% and 700% SUV, respectively (Figs. 5A–5D).

[Fig. 5] phase (Fig. 5A). Lower specific radioactivity and DMI chase

 TABLE 1

 Non–Metabolite-Corrected Myocardial Retention Index and Washout Rate of Radiotracers With and Without Treatment With DMI

	Retention index (%/min at 40 min)			Washout rate (%/min between 10 and 60 min)		
Radiotracer	Without intervention (n = 4)	DMI chase (n = 4)	DMI pretreatment (n = 1)	Without intervention (n = 4)	DMI chase (n = 4)	DMI pretreatment (n = 1)
HED	7.38 ± 0.82	$3.28 \pm 0.60^{*}$	$0.21\pm0.35^\dagger$	0.13 ± 0.23	1.86 ± 0.57*	‡
PHEN	3.43 ± 0.45	$2.95 \pm 0.51^{*}$	$0.24\pm0.25^\dagger$	1.13 ± 0.35	1.51 ± 0.16	ŧ
EPI	4.24 ± 0.59	4.02 ± 0.99	$0.22\pm0.30^{\dagger}$	0.50 ± 0.24	0.61 ± 0.28	‡
EPI (low specific radioactivity)	3.87 ± 0.81	_	_	0.69 ± 0.17	_	_

*P < 0.05 vs. animals without intervention.

[†]Single-case observation precludes statistical testing, but values are markedly reduced.

[‡]Complete washout of radiotracer from myocardium in first 10 min in DMI pretreated rats.

Values are average ± SD from septal, lateral, inferior, anterior, and apex regions of left ventricle.

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FIGURE 4. Myocardial time–activity curves of radioactivity in LV chamber (\Box), LV lateral wall (\triangle), LV septum (∇), LV apex (\diamond), and liver (*) in rats after intravenous injection of PHEN, without pharmacologic intervention (A), with intraperitoneal administration of 2 mg of DMI solution per kilogram 15 min after tracer injection (DMI chase) (B), and after intraperitoneal administration of 2 mg of DMI solution per kilogram 15 min before tracer injection (DMI block) (C). A and B show data averaged from 4 rats, and C shows observation from 1 rat (LV septum and LV apex were included in FOV but not detectable in C; kinetics in anterior wall were also analyzed but did not differ from other myocardial regions and are therefore not shown).

DISCUSSION

Our dynamic in vivo PET studies of the sympathetic nervous system of the rat heart confirm the previously described kinetic properties of the catecholamine analog ¹¹C-HED but showed differences in the handling of the physiologic catecholamine ¹¹C-EPI and, to a lesser degree, of ¹¹C-PHEN, characterized by high metabolic activity, high turnover, and clearance from the myocardium. This has important implications for the use of the rat heart as a model of cardiovascular disease, and it provides further insights into the biology of the rat sympathetic nervous system.

In general, the rat is an attractive species to study cardiovascular biology; compared with larger animals (pigs, dogs), they are easy to handle and reduce logistic complexity and costs. Rats are also preferred over smaller animals (mice), in which the small size of the heart still poses challenges for the robustness of noninvasive nuclear imaging. When compared with observations in humans (13-15) and isolated perfused rat hearts (10,12,22), our multitracer characterization suggests that the cardiac sympathetic nervous system of the living rat is different and that conclusions from one species to the other must be interpreted with caution. For interpretation of our results, it is also important to emphasize that in vivo studies of sympathetic neuronal tracer kinetics may differ from studies of the isolated perfused heart because the peripheral nervous system and central nervous system are not preserved, and effects of systemic tracer degradation do not play a role in isolated perfused hearts.



FIGURE 5. Myocardial time-activity curves of radioactivity in LV chamber (\Box), LV lateral wall (\triangle), LV septum (\heartsuit), LV apex (\diamond), and liver (*) in rats after intravenous injection of EPI, without pharmacologic intervention (A), with lower specific radioactivity without pharmacologic intervention (B), after intraperitoneal administration of 2 mg of DMI solution per kilogram 15 min after tracer injection (DMI chase) (C), and after intraperitoneal administration of 2 mg of DMI solution pre kilogram 15 min before tracer injection (DMI block) (D). A, B, and C show data averaged from 4 rats, and D shows observation from 1 rat (LV septum and LV apex were included in FOV but not detectable in D; kinetics in anterior wall were also analyzed but did not differ from other myocardial regions and are therefore not shown).

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Pretreatment with the specific NET blocker DMI was done in single animals in our study. In these observations, a significantly decreased radioactivity concentration of each of the 3 mentioned radiotracers in the myocardium was observed, a finding that is consistent with earlier, more extensive work (10,12,22) and confirms their specific neuronal uptake via uptake-1. This supports the validity of these PET sympathoneuronal imaging agents for assessing myocardial sympathetic innervation.

For the MAO-resistant catecholamine analog HED, PETmeasured in vivo kinetics in the rat myocardium were comparable with isolated rat heart studies and cardiac PET in humans (10,15,23,24). Avid uptake and subsequent stable retention was observed without DMI. The acceleration of myocardial clearance after DMI chase confirms earlier studies and suggests that HED is continuously taken up and released from nerve terminals. Vesicular storage does not seem to be a critical determinant of tracer retention, and myocardial retention seems to depend significantly on uptake-1-mediated reuptake into the nerve terminal. In the liver, accumulation of HED mainly reflects nonspecific uptake, which is accelerated and increased by DMI.

For the MAO-sensitive catecholamine analog PHEN, specific uptake was followed by myocardial washout as previously described. This washout has been attributed to leakage from intraneuronal vesicles and subsequent MAO metabolism, as confirmed by MAO blockade and comparison with the MAO-resistant analog D2-PHEN (13,22). In humans, a mean clearance half-time of 55 min has been observed (13), whereas in untreated rats our clearance rates translate into a mean half-time of 44 min, suggesting a somewhat more rapid metabolic degradation of PHEN due to either higher MAO activity or more pronounced vesicular leakage in rats than in humans. DMI chase did not significantly affect myocardial washout, suggesting that release and reuptake into the neuron are not significant contributors to PHEN kinetics. In the liver, PHEN-derived radioactivity shows an early peak with rapid clearance followed by stable retention that cannot be blocked by DMI. Early kinetics thus appear to reflect hepatic MAO metabolism, whereas sustained retention may be a combined effect of metabolite-derived activity, efflux from the liver, and influx from peripheral organs to the liver.

For the physiologic catecholamine EPI, kinetics differed from those previously described in humans and isolated perfused rat hearts, in which rapid initial uptake was followed by stable retention (12,14). Our in vivo PET studies showed initial myocardial uptake followed by significant washout, suggesting a high myocardial turnover of catecholamines.

Several factors may contribute to this rapid turnover. For interpretation of these results, it is important to emphasize that EPI as a natural catecholamine is a substrate to all enzymatic metabolic pathways for catecholamines. The observation of stable retention in isolated perfused rat hearts and human hearts is explained by efficient storage of the tracer in neuronal vesicles, in which the tracer is protected from degradation after initial neuronal uptake via uptake-1/ NET (12,14). The washout and lack of stable retention in the heart of living rats may theoretically be explained by intraneuronal vesicular leakage and subsequent metabolic degradation. The somewhat faster washout rates for PHEN (as a tracer of vesicular leakage) in rats than in humans may partially support this hypothesis. But the difference between species was subtle and not sufficient to explain the marked difference of EPI kinetics alone. Also, the lack of washout in isolated perfused rat hearts, in which axons are transsected and intraneuronal vesicular leakage may be even more pronounced, does not support this hypothesis. Second, nonneuronal degradation via the nonspecific uptake-2 and extraneuronal MAO may account for the washout of EPI. The very efficient blockade of myocardial EPI uptake by DMI pretreatment does not support this hypothesis, although very rapid extraneuronal degradation in the early phase, which may be difficult to separate from delivery and equilibrium effects, cannot be ruled out. Third, anesthesia may have influenced catecholamine metabolism and autonomic tone in rats and thereby explain the different results demonstrated in humans. Norepinephrine levels can rise during isoflurane anesthesia (25), but an increase in systemic norepinephrine should also affect tracer kinetics of HED, which was not observed in our study. Also, an isoflurane-induced increase in norepinephrine at the presynaptic site should reduce uptake of EPI, but EPI showed high initial uptake and then biphasic washout. Finally, the rapid washout of EPI radioactivity might be because of high neuronal activity and nerve firing, which is supported by the high heart rate found in rats. Rapid vesicular release from neurons may contribute to clearance from myocardium and subsequent extraneuronal metabolic degradation. We were not able to accelerate release of EPI by blockade of reuptake using a DMI chase, but the lack of influence of reuptake on the kinetics does not rule out vesicular release from neurons followed by rapid extraneuronal degradation.

We observed no difference in the EPI myocardial kinetic profile between higher (370–740 GBq/ μ mol) and lower (9–18 MBq/ μ mol) specific radioactivity, suggesting that mass effects do not play a significant role in tracer kinetics (26,27) and that neuronal uptake and storage are most likely high-capacity systems.

Although the myocardium was our primary target organ, we included the liver as a noncardiac organ in our analysis because it was in the FOV and showed high uptake that may interfere with cardiac image quality. Liver uptake was not blocked but rather increased by DMI, supporting the role of the liver as an organ of metabolic degradation in which radioactivity is mostly determined by nonspecific uptake of metabolites. Metabolism of catecholamines and its degradation products in hepatocytes, and the relative contribution of specific neuronal versus nonspecific nonneuronal uptake to liver kinetics, is not well understood but may be a subject of future dedicated studies.

Some limitations of our study need to be emphasized. First, we described kinetics of our tracers by simple measures of uptake and washout. Compartmental modeling of these tracers is not well established, and it is further complicated by inaccuracies of noninvasive definitions of the arterial input function in small-animal studies. Validation of kinetic modeling was thus considered beyond the scope of this project. Second, we were not able to measure plasma metabolites and obtain metabolite-corrected retention indices. Repeated blood sampling is difficult to perform in rodents because of the low total blood volume. Systemic tracer degradation may thus have contributed to inaccuracies of our retention measures. But the overall kinetic properties observed in our studies are unlikely to be a consequence of systemic effects only, and metabolite-independent measures such as the SUV are available for additional interpretation. Finally, we did not conduct our studies with multiple tracers in the same animals because we wanted to avoid residual effects of DMI or tracerderived radioactivity from prior imaging on a subsequent study. Intersubject differences thus may have contributed to higher variability between groups, but our results remain conclusive despite small sample sizes.

CONCLUSION

Our multitracer PET study suggests high turnover of physiologic catecholamines in the heart of living rats. Although the underlying molecular events are not fully elucidated, this may be because of rapid vesicular release and subsequent metabolic degradation. This observation has important implications for imaging studies of the cardiac sympathetic nervous system. First, the rat heart has limitations as a translational model of cardiac neuronal disease due to different pharmacokinetics of catecholamines. Second, conclusions from autonomic innervation studies in isolated perfused rat heart studies, in which this high turnover was not observed, to the in vivo environment may also be difficult because of lack of systemic and central nervous effects. And third, a single catecholaminergic tracer may not necessarily reflect the complete biology of sympathetic nerve terminals. The false neurotransmitter HED, for example, yielded highquality myocardial images in this study and comparable kinetics to humans but would not have allowed for identification of the turnover kinetics of true catecholamines such as epinephrine.

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