Norepinephrine transporter variant A457P knock-in mice display key features of human postural orthostatic tachycardia syndrome

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SUMMARY

Postural orthostatic tachycardia syndrome (POTS) is a common autonomic disorder of largely unknown etiology that presents with sustained tachycardia on standing, syncope and elevated norepinephrine spillover. Some individuals with POTS experience anxiety, depression and cognitive dysfunction. Previously, we identified a mutation, A457P, in the norepinephrine (NE; also known as noradrenaline) transporter (NET; encoded by SLC6A2) in POTS patients. NET is expressed at presynaptic sites in NE neurons and plays a crucial role in regulating NE signaling and homeostasis through NE reuptake into noradrenergic nerve terminals. Our in vitro studies demonstrate that A457P reduces both NET surface trafficking and NE transport and exerts a dominant-negative impact on wild-type NET proteins. Here we report the generation and characterization of NET A457P mice, demonstrating the ability of A457P to drive the POTS phenotype and behaviors that are consistent with reported comorbidities. Mice carrying one A457P allele (NET+/P) exhibited reduced brain and sympathetic NE transport levels compared with wild-type (NET+/+) mice, whereas transport activity in mice carrying two A457P alleles (NET+/P) was nearly abolished. NET+/P mice exhibited elevations in plasma and urine NE levels, reduced 3,4-dihydroxyphenylglycol (DHPG), and reduced DHPG:NE ratios, consistent with a decrease in sympathetic nerve terminal NE reuptake. Radiotelemetry in unanesthetized mice revealed tachycardia in POTS patients. NET is expressed at presynaptic sites in NE neurons and plays a crucial role in NE inactivation and homeostasis (Schömig et al., 1989; Xu et al., 2000). Altered sympathetic nerve terminal NET uptake sites and/or activity are found in hypertension, diabetes, cardiomyopathy, heart failure and ischemia (Böhm et al., 1998; Merlet et al., 1992; Rumantir et al., 2000; Schnell et al., 1996; Schömig et al., 1991).

We identified a link between a NET genetic mutation with a familial incidence of postural orthostatic tachycardia syndrome (POTS) (Shannon et al., 2000; Hahn et al., 2003). POTS is characterized by an increase in standing heart rate that is not accompanied by hypotension (Robertson, 1999). Idiopathic POTS is accompanied by features that suggest a failure of NET, including increased NE spillover, and decreased NE clearance, intraneuronal NE metabolism and response to tyramine (Furlan et al., 1998; Jacob et al., 1999). In normal human subjects, pharmacological NET blockade produces orthostatic tachycardia much like that observed in POTS (Mayer et al., 2006; Schroeder et al., 2002). Some POTS patients experience anxiety, depression and cognitive dysfunction (Raj et al., 2009). In a search for a molecular basis for the POTS diagnoses of twin probands, Shannon et al. identified a nonsynonymous mutation in the NET gene that produces the amino acid substitution A457P in transmembrane domain 9 of the NET protein (Shannon et al., 2000). In a within-family analysis, A457P carriers demonstrated standing-induced increased HR and plasma NE, and a decreased plasma 3,4-dihydroxyphenylglycol (DHPG):NE ratio, relative to family members not carrying the mutation (Shannon et al., 2000). Subsequent in vitro expression studies revealed that A457P disrupts normal plasma membrane targeting,

INTRODUCTION

Norepinephrine (NE; also known as noradrenaline) is the primary signaling neurotransmitter of the autonomic sympathetic nervous system (SNS), where it acts to control heart rate (HR). Compromised NE signaling has been associated with autonomic disease, particularly impacting cardiovascular function (Billman et al., 1997; Esler and Kaye, 2000; Exner et al., 1999; Parati and Esler, 2012). NE signaling is also implicated in disorders of anxiety, depression and cognition (Beane and Marrocco, 2004; Pliszka, 2005; Ressler and Nemeroff, 1999). Released NE is inactivated through NE reuptake into noradrenergic nerve terminals. Our in vitro studies demonstrate that A457P reduces both NET surface trafficking and NE transport and exerts a dominant-negative impact on wild-type NET proteins. Here we report the generation and characterization of NET A457P mice, demonstrating the ability of A457P to drive the POTS phenotype and behaviors that are consistent with reported comorbidities. Mice carrying one A457P allele (NET+/P) exhibited reduced brain and sympathetic NE transport levels compared with wild-type (NET+/+) mice, whereas transport activity in mice carrying two A457P alleles (NET+/P) was nearly abolished. NET+/P mice exhibited elevations in plasma and urine NE levels, reduced 3,4-dihydroxyphenylglycol (DHPG), and reduced DHPG:NE ratios, consistent with a decrease in sympathetic nerve terminal NE reuptake. Radiotelemetry in unanesthetized mice revealed tachycardia in NET+/P mice without a change in blood pressure or baroreceptor sensitivity, consistent with studies of human NET A457P carriers. NET+/P mice also demonstrated behavioral changes consistent with CNS NET dysfunction. Our findings support that NET dysfunction is sufficient to produce a POTS phenotype and introduces the first genetic model suitable for more detailed mechanistic studies of the disorder and its comorbidities.

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NET A457P mice display POTS

RESULTS

NET A457P knock-in mice were generated through a homologous recombination strategy that introduced A457P into exon 9 of the mouse gene at the position orthologous to the identified human A457P mutation (Fig. 1A). Embryonic stem (ES) cell clones were confirmed for correct homologous recombination of both the 5' and 3' arms through Southern blot (Fig. 1B) and PCR. The presence of A457P was also determined through dyeoxy sequencing. Heterozygous (\( \text{NET}^{+/\text{P}} \)) and homozygous (\( \text{NET}^{\text{P/P}} \)) A457P mice were identified by PCR genotyping strategy to detect a residual \( \text{loxP} \) site in intron 11 (Fig. 1C), and by sequencing of the A457P mutation. A ‘speed congenics’ breeding strategy was used to backcross mice to the C57BL/6 strain (Jackson Laboratories, Bar Harbor, ME) (Wong, 2002). Mice appeared healthy, bred with expected genotype distributions and exhibited no differences in body weight among genotypes for either male or female mice (data not shown).

\( \text{NET}^{+/\text{P}}, \text{NET}^{\text{P/P}} \) and \( \text{NET}^{\text{P/P}} \) mice were assessed for NET expression and activity to determine whether the deficits observed for A457P under heterologous expression would translate to noradrenergic neurons. Initial experiments determined similar changes in male and female A457P mice, and thus western blot and single point transport data from male and female A457P mice were combined for analysis. In assays assessing the \text{ex vivo} \ transport of radiolabeled NE in peripheral noradrenergic terminals, \( \text{NET}^{+/\text{P}} \) mice demonstrated significant transport decreases in the atria and ventricle of the heart, vas deferens and femoral vein (Fig. 2A: \( n=5 \)–6). This is supportive of the hypothesis that a deficit in NET activity in the SNS, particularly in the heart, drives the phenotype in heterozygous human A457P carriers.

We next sought to determine whether these transport deficits extended to CNS noradrenergic neurons. In synaptosomes prepared from cortex and hippocampus, transport levels in \( \text{NET}^{+/\text{P}} \) mice were \( \sim60\% \) of \( \text{NET}^{+/+} \) mouse levels. Saturation kinetic experiments in cortical and hippocampal synaptosomes from male \( \text{NET}^{+/\text{P}} \) mice demonstrated a decrease in maximal velocity of reaction (\( V_{\text{MAX}} \)) values, with no change in the substrate concentration required for 50% \( V_{\text{MAX}} \) (\( K_{S} \)) for NE (Fig. 2D; \( P<0.05, n=8 \)). In \( \text{NET}^{\text{P/P}} \) mice, transport was nearly abolished (Fig. 2C). Thus, the A457P protein exhibits very little if any activity in the CNS, as we previously observed \text{in vitro} (Hahn et al., 2003).

NET expression was determined in total and surface fractions of synaptosomal preparations using a membrane-impermeant biotinylation assay. \( \text{NET}^{+/\text{P}} \) and \( \text{NET}^{\text{P/P}} \) mice demonstrated decreased NET expression in both the cortex and hippocampus (Fig. 2E,F). \( \text{NET}^{+/\text{P}} \) mice demonstrated \( \sim60\% \) of \( \text{NET}^{+/+} \) total NET levels, and displayed similar decreases in surface NET pools (Fig. 2E). \( \text{NET}^{\text{P/P}} \) demonstrated loss of NET protein in cortical and hippocampal terminals similar to that of the \( \text{NET}^{+/\text{P}} \) (Fig. 2F). Thus, unlike a NET gene deletion, A457P mice retain a high level of protein expression of a nonfunctional transporter that might play a role in influencing normal NET function or trafficking in \( \text{NET}^{+/\text{P}} \) mice. Overall, these results are consistent with the pattern of loss of A457P expression and plasma membrane trafficking previously demonstrated using transfected cells.
We next assessed whether the NET deficit altered cardiovascular function in NET+/P mice similar to that of human A457P carriers. Because A457P carriers are heterozygotes, it was crucial to determine whether NET+/P mice could drive the cardiovascular phenotypes. Indeed, we demonstrated that the presence of one A457P allele in NET+/P mice is sufficient to produce deficits in NET expression and uptake. Here, we ascertained cardiovascular phenotypes in NET+/P mice, using radiotelemetry to monitor blood pressure (BP) and HR in conscious, freely behaving mice. Fig. 3A depicts HR over the course of 3 days of data collection. HR of both NET+/+ and NET+/P mice followed a diurnal rhythm with lower and higher HRs during rest (during the daytime/lights on in the colony room) and active (night time/lights off) periods, respectively. Two-way ANOVA revealed a significant effect of both genotype and time of day on HR [$F(1,27)=15.59$, $P<0.001$ for genotype and $F(2,27)=18.26$ for time of day]. NET+/P mice exhibited higher HR over both the entire 24-hour period and at night (Fig. 3B; $P<0.05$, $n=5$). A457P carriers, and other individuals with POTS that also exhibit excessive plasma NE, do not exhibit the hypotension that accompanies other forms of orthostatic intolerance (Shannon et al., 2000). In this group of patients, BP remains unchanged or becomes somewhat hypertensive during standing. Mean arterial BP, measured during the same recording period as HR, was unchanged in NET+/P compared with NET+/+ mice (Fig. 3C). Neither diastolic nor systolic BPs were different between genotypes (data not shown). We also determined whether there was a change in the baroreflex sensitivity in A457P knock-in mice. Here, we measured baroreceptor-mediated changes in the HR response following intravenous administration of phenylephrine (PHE) or nitroprusside (NTP) in anesthetized NET+/+ or NET+/P mice. Two-way ANOVA revealed effects of dose for both PHE and NTP [$F(4,50)=61.78$, $P<0.01$, $n=3-6$ for NTP and $F(5,60)=23.53$, $P<0.01$, $n=3-6$ for PHE] (Fig. 4A,B). However, there were no differences in the BP response between NET+/+ and NET+/P mice following PHE or NTP administration. Baroreceptor sensitivity (BRS), the ratio of the change in HR to the change in BP at an effective dose of both PHE and NTP, was not different between genotypes (Fig. 4C,D; $n=6$; PHE, 20 µg/kg body weight; NTP, 30 µg/kg body weight).
Disease Models & Mechanisms

**RESEARCH ARTICLE**

**Fig. 2. [3H]NE transport and expression in A457P knock-in mice.** Data are expressed as mean percent ± s.e.m. of NET^{+/P} mice (A–C,E,F) or as raw values ± s.e.m. (D). *P<0.05, **P<0.01 and ***P<0.001 compared with NET^{+/P} mice (two-tailed t-test). (A) [3H]NE transport in the SNS, n=5–6. Vent., ventricle; VD, vas deferens. (B,C) Single point [3H]NE transport in brain, n=6–7. (D) Saturation kinetics of [3H]NE transport in brain, n=8. Cortex NET^{+/P} (black squares), V_{MAX}=6.6±0.40 moles×10^{-13}/μg/minute, K_{M}=87.8±20.9 nM; cortex NET^{+/P} (white squares), V_{MAX}=4.6±0.50 moles×10^{-13}/μg/minute, K_{M}=96.4±38.1 nM; hippocampus NET^{+/P} (black triangles), V_{MAX}=8.2±0.77 moles×10^{-13}/μg/minute, K_{M}=118.3±40.6 nM; hippocampus NET^{+/P} (white triangles), V_{MAX}=5.5±0.54 moles×10^{-13}/μg/minute, K_{M}=136.8±46.3 nM. (E,F) NET expression in total and surface fractions of cortex (CTX) and hippocampus (HPC) synaptosomes in NET^{+/P} (n=7; E) and NET^{+/P} (n=7; F) mice compared with NET^{+/P} mice.

Fig. 3. Radiotelemetry in NET^{+/P} versus NET^{+/P} mice. (A) 3-day collection of HR measures; data were analyzed and graphed in 30-minute increments. Traces shown start at lights off in the colony room. (B) HR during 3 days of data collection, expressed as means ± s.e.m. averaged over the 24-hour period and during 12-hour day and night periods. Significance was determined by two-way ANOVA. Genotype: F(1,27)=15.59, P<0.001. Time of day: F(2,27)=18.26, P<0.0001. *P<0.05, post-hoc t-test, n=5. (C) Mean arterial BP measured during the same 3-day period as in A and B (n=5).

One of the defining characteristics that we attribute to NET deficiency in A457P carriers that is also a salient feature in other POTS patients, is the disturbance in the homeostasis of peripheral NE indicated by changes in the DHPG:NE ratio. We attribute this to a lack of NE reuptake that results in both elevated NE in plasma and urine and a decrease in the intraneuronal metabolism of NE to DHPG. Other individuals with POTS, although they do not harbor the A457P mutation, have also been hypothesized to have faulty NE uptake as part of the disease phenotype. Indeed, recent studies support loss of NET protein in other POTS patients (Lambert et al., 2008). We sought to determine whether the introduction of A457P into mice was sufficient to recapitulate this phenotype. Plasma and urinary NE and DHPG were measured in mice under resting conditions. Two-way ANOVA revealed significant effects in both male and female mice [male plasma genotype, F(2,135)=50.77, P<0.001, n=6–12; male urine genotype, F(2,215)=10.82, P<0.001, n=9–20; female plasma genotype, F(2,110)=20.21, P<0.001, n=8–9; female urine interaction, F(8,85)=2.88, P=0.01, n=6–7]. NE levels were elevated, DHPG levels were decreased and the DHPG:NE ratio was decreased, highly consistent with those abnormalities observed in human A457P carriers (Fig. 5A,C, P<0.05-0.01). These effects were observed in NET^{+/P} mice and NET^{+/P} mice, consistent with the ability of a single copy of the mutant allele, as seen in the POTS patients where it was identified, to confer these changes in NE homeostasis. In some cases, NE increases were not significant in NET^{+/P} mice and this might be attributable to the collection of both plasma and urine under anesthetized and non-stressful conditions, respectively. Indeed, family members with A457P exhibit elevated standing plasma NE levels, whereas supine NE levels are much less affected. Dopamine (DA) and epinephrine (EPI; also known as adrenaline) levels were also increased in male and female NET^{+/P} mice in plasma but not urine (Fig. 5B,D; P<0.05-0.01).

We also investigated whether impaired NE reuptake influences tissue levels of catecholamines and metabolites. There was a trend for female NET^{+/P} mice to have smaller hearts compared with wild-type females (data not shown; P=0.08). DHPG levels were decreased in heart of NET^{+/P} and NET^{+/P} mice, and NE levels and the DHPG:NE ratio were decreased in NET^{+/P} mice (Table 1). There was a trend for a decrease in the DHPG:NE ratio in male NET^{+/P} heart (P=0.08). EPI levels were also decreased in hearts of A457P knock-in mice, with a 30% decrease in NET^{+/P} (P=0.09) mice and a significant 60% decrease in NET^{+/P} mice. Tissue DA levels were unchanged.

We also investigated the impact of impaired NE reuptake to produce dysfunction in the CNS. There were no differences
between \textit{NET}\textsuperscript{+/+} mice and either \textit{NET}\textsuperscript{+/P} or \textit{NET}\textsuperscript{P/P} mice in brain weights (data not shown). Consistent with the transport activity deficits observed, NE and DHPG levels and the DHPG:NE ratio were decreased in the brain of male and female \textit{NET}\textsuperscript{P/P} mice, although there were no significant changes in \textit{NET}\textsuperscript{+/P} mice (Table 1).

The role of NET in anxiety and depression, and the presence of these symptoms in some POTS subjects, led us to explore whether A457P mice might exhibit phenotypes related to these disorders. Because A457P carriers are heterozygotes and \textit{NET}\textsuperscript{+/P} mice showed deficits in CNS NET expression and NE uptake, we focused behavioral tests on \textit{NET}\textsuperscript{+/P} mice. First, mice were exposed to the elevated plus maze (EPM), which measures an animal’s anxiety based on the conflict between the motivation to approach and explore novel situations versus a fear of open, brightly lit spaces (open arms). Additionally, freezing and head dips, ethological measures of fear- and anxiety-related behaviors sensitive to anxiolytic compounds, were assessed in the EPM (Blanchard et al., 2003; Gerits et al., 2007; Gross et al., 2000; Rosen and Schulkin, 1998; Wall et al., 2004; Weiss et al., 1998). We detected a highly significant interaction effect in the EPM between genotype and compartment \([F(2,96)=13.74, \quad P<0.0001]\). \textit{NET}\textsuperscript{P/P} mice spent less time in the open arms and more time in the closed arms of the maze compared with \textit{NET}\textsuperscript{+/+} mice (Fig. 6A; \(P<0.05\) and 0.01, respectively, \(n=16-18\)). Additionally, \textit{NET}\textsuperscript{P/P} mice exhibited a significant increase in freezing incidence as well as durations of freezing as compared with \textit{NET}\textsuperscript{+/+} mice and also exhibited fewer head dips, all of which are behaviors that are indicative of anxiety (Fig. 6B,C; \(P<0.01\), \(n=16-18\)). We detected no differences in the number of entries into any of the zones, indicating a lack of alteration in general motor activity level (data not shown) (Braun et al., 2011). Anxiety-like behavior was also detected in the open field, where \textit{NET}\textsuperscript{P/P} mice spent significantly less time in the center of the field and increased time along the periphery (thigmotaxis) (Fig. 6D; \(P<0.01\), \(n=16-17\)). As with indications of normal general activity in the EPM, \textit{NET}\textsuperscript{P/P} mice exhibited no difference in total distance travelled in the open field (Fig. 6E). Finally, compared with \textit{NET}\textsuperscript{+/+} mice, \textit{NET}\textsuperscript{P/P} mice exhibited decreased immobility in the tail suspension test (TST), a test that is sensitive to NE-enhancing antidepressant medication (Fig. 6F; \(P<0.05\), \(n=7-9\)).

**DISCUSSION**

\textit{NET}\textsuperscript{P/P} mice exhibited elevated HRs, supporting our prior contention based on within-family measure of phenotype associations that a single copy of the A457P allele is responsible for the tachycardia that is evident in human heterozygous A457P carriers. We suspect that the larger increase in HR during the active period reflects increased sympathetic activity during this time. Greater NE release during the active period probably renders synaptic NE levels, NE spillover and HR more dependent on NE reuptake.
affected by the A457P allele, similar to the phenotype observed in human A457P carriers and most idiopathic POTS subjects (Shannon et al., 2000). This finding could be explained by the fact that NET removes greater amounts of NE released in heart compared with other sympathetically innervated organs (Goldstein et al., 1988). We did not observe a change in baroreflex sensitivity. Some studies have reported altered baroreflex sensitivity in POTS patients whereas others have found no differences (Farquhar et al., 2000; Furlan et al., 1998; Jacob et al., 1999; Jordan et al., 2002; Muenter Swift et al., 2005). Regardless, our results indicate that the tachycardia induced by A457P does not seem to be directly mediated by changes in baroreflex sensitivity.

Previously, we demonstrated both trafficking and transport deficits of A457P in vitro, with a decrease in surface NET expression and a more severe, near total, loss of transport activity (Hahn et al., 2003). Consistent with these studies, A457P mice are deficient in both NET surface expression and NE transport activity in native SNS and CNS noradrenergic nerve terminals, with NET+/P mice exhibiting near total loss of transport. Previous in vitro studies also demonstrated that NET A457P physically associates with wild-type NET transporters and hinders them from reaching the cell surface, serving to decrease wild-type NET transport activity (Hahn et al., 2003). Evidence from other studies supports that neurotransmitter transporters form oligomeric complexes early in the secretory pathway that are maintained in the plasma membrane, suggesting the likelihood of both physical and functional interactions among wild-type and mutant transporter species (Bartholomäus et al., 2008; Hahn et al., 2003; Hastrup et al., 2001; Kilic and Rudnick, 2000; Scholze et al., 2002; Sitte et al., 2004; Torres et al., 2003). Data presented here suggest that a dominant negative effect of A457P might contribute, in part, to deficits in vivo in NET+/P mice. First, observations from our laboratory indicate that heterozygous NET knockout mice do not express significant changes in NE transport (data not shown), whereas NET+/P mice exhibited significant transport deficits in the present study. Additionally, changes in DHPG levels and DHPG:NE ratios were altered in plasma and urine in NET+/P mice in our study, whereas no changes were detected in DHPG levels in the heterozygous NET knockout mouse (Keller et al., 2004). The consequences of expression of an impaired NET protein in NE neurons might additionally impact the interaction of NET with its multiple protein partners (Bauman et al., 2000; Bartholomäus et al., 2004). The consequences of expression of an impaired NET protein in NE neurons might additionally impact the interaction of NET with its multiple protein partners (Bauman et al., 2000; Bartholomäus et al., 2004). The consequences of expression of an impaired NET protein in NE neurons might additionally impact the interaction of NET with its multiple protein partners (Bauman et al., 2000; Bartholomäus et al., 2004). The consequences of expression of an impaired NET protein in NE neurons might additionally impact the interaction of NET with its multiple protein partners (Bauman et al., 2000; Bartholomäus et al., 2004). The consequences of expression of an impaired NET protein in NE neurons might additionally impact the interaction of NET with its multiple protein partners (Bauman et al., 2000; Bartholomäus et al., 2004).

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<tr>
<th>Genotype</th>
<th>Heart</th>
<th>Brain</th>
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<tr>
<td></td>
<td>DHPG</td>
<td>NE</td>
</tr>
<tr>
<td>Male</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NET+/+</td>
<td>64.3±5.3</td>
<td>3454±353.3</td>
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<tr>
<td>NET+/−</td>
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<td>3711±152.5</td>
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<tr>
<td>NET+/P</td>
<td>30.6±2.7***</td>
<td>3063.7±182.3</td>
</tr>
<tr>
<td>Female</td>
<td></td>
<td></td>
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<tr>
<td>NET+/+</td>
<td>62.8±7.5</td>
<td>4570.8±353.8</td>
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<tr>
<td>NET+/−</td>
<td>58.9±8.3</td>
<td>4810.4±208.5</td>
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<tr>
<td>NET+/P</td>
<td>34.0±6.8*</td>
<td>3686.1±269.4</td>
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Values shown are mean ± s.e.m., pmol/g. *P<0.05; **P<0.01; ***P<0.001.

Table 1. Tissue content of DHPG and NE in NET A457P mice

Fig. 6. NET+/+ versus NET+/P mice in behavioral tests. Data are expressed as mean ± s.e.m. (A–C) Behavior in the EPM. For time spent in the different zones (A), significance was determined by two-way ANOVA. Interaction: F(2,96)=13.74, P<0.0001. n=16-18. Freezing behavior (B) and head dips (C) were also assessed. n=16-18. (D,E) Open field exploration. Time spent in the center versus surrounding zones (D) and distance travelled (E) were measured. n=16-18. (F) Time immobile in the last 4 minutes of the TST. n=7-9. *P<0.05, **P<0.01, post-hoc t-test.
released in heart compared with other sympathetic targets (Goldstein et al., 1988), collection of whole body blood might not detect increased cardiac NE that is responsible for the observed tachycardia. Plasma DA and EPI were also increased in NET<sup>P/P</sup> mice. A similar effect is observed in NET knockout mice (Keller et al., 2004). Plasma DA probably originates from neuronal release with NE and loss of NET would also prevent reuptake of DA (Goldstein and Holmes, 2008). Excessive EPI might derive similarly from the lack of reuptake, or might be a result of increased EPI release from the adrenal gland. Differential sources of DA and EPI in urine versus plasma could explain the observed selective increases in plasma (Goldstein et al., 2003).

Our results indicate that heart NE content was unchanged in NET<sup>P/P</sup> mice, although it was decreased in NET<sup>P/P</sup> mice in the brain. The fact that heart NE content was largely unaffected by the loss of reuptake and its contribution to NE vesicular stores suggests that neuronal NE synthesis rates are sufficient to compensate for this loss, or that synthesis is upregulated to maintain stores in A457P mice. In support of this possibility, the activity of tyrosine hydroxylase, the rate-limiting enzyme in NE synthesis, is enhanced in both DA transporter and NET knockout mice to maintain appropriate monoamine release levels (Jones et al., 1998; Xu et al., 2000). Interestingly, we show that NET A457P results in reduced heart EPI content, which is believed to rely mainly upon uptake rather than local synthesis (Goldstein et al., 2003). It will be important to determine whether changes in NE synthesis rates in A457P mice account for the maintenance of NE stores despite loss of reuptake.

Although A457P was identified in a cardiovascular syndrome, A457P mice also demonstrate behaviors that are associated with anxiety in both EPM and open field exploration. Our findings are in good agreement with evidence that noradrenergic activity is anxiogenic and that adrenergic receptor antagonists can reverse anxiety behavior (Goddard et al., 2010; Katayama et al., 2010; Kukolja et al., 2008; Morilak et al., 2005; Schank et al., 2008). Indeed, NET knockout mice exhibit increased conditioned freezing (Keller et al., 2006). We did not observe changes in locomotor activity in either the open field or EPM, supporting that the observed behavioral changes reflect increased anxiety (Braun et al., 2011). Our results demonstrate decreased immobility in the TST, consistent with results in NET knockout mice (Xu et al., 2000).

Comorbidity of cognitive and mood disorders with cardiovascular disease is a serious and prevalent problem that is not well understood. Patients with anxiety complain of heart palpitations and patients with tachycardia disorders have a higher incidence of anxiety (Fleet et al., 2003; McCrank et al., 1998). Activation of the SNS has been associated with depression and patients with major depression are at an increased risk for heart disease (Musselman et al., 1998). Interestingly, a cardiac NE uptake deficiency, characterized by decreased clearance of NE in the heart, and decreased DHPG:NE ratios, like that seen in POTS, has been described in individuals with depression (Barton et al., 2007). Individuals with POTS experience increased ratings of depression, anxiety and inattention/ADHD symptoms (Raj et al., 2009). These findings indicate that some comorbidity of cardiovascular and psychiatric disease might involve common underlying genetic determinants, such as NET.

Although a role for CNS NE in stress and anxiety is well established and we predict that central NET dysfunction contributes to anxiety phenotypes in A457P mice, there might also be a cardiovascular influence on brain neurotransmission. Autonomic nervous system input is integrated within brain regions to direct behavioral response to stress, autonomic homeostasis, and emotional and goal-directed behavior. Central noradrenergic neurotransmission is crucial to many of these responses. For example, peripheral cardiovascular and other autonomic stimuli are strong activators of locus coeruleus noradrenergic neuronal firing activity, which in turn drives changes in forebrain activity (Page and Valentino, 1994; Valentino et al., 1991).

Understanding how NET variants can contribute to both cardiovascular disease and CNS dysfunction could have implications for the development of novel treatment strategies. Therapeutics with NET-blocking activity are routinely used to treat mental health disorders (Klimek et al., 1997; Leonard, 1997; Morilak and Frazer, 2004; Ressler and Nemeroff, 1999). Such compounds can have adverse effects on the heart, including drug-induced long QT syndrome (Alvarez and Pahissa, 2010; Vitiello, 2008). Thus, although genetic variation in NET might predispose individuals to disorders indicating the use of NET blockers, such genetic predispositions might also contribute to increased sensitivity to the cardiovascular side effects of medication.

In summary, the present results in NET A457P mice support the contention that NET dysfunction can produce the POTS phenotype as well as behavioral comorbidities reported in POTS (Barton et al., 2007; Fleet et al., 2003; McCrank et al., 1998; Musselman et al., 1998; Raj et al., 2009). Coupled with the emerging evidence for a more widespread incidence of NET deficiency in POTS, our studies argue for our model as an important tool for the assessment of NET-derived plasticities in heart and brain that contribute to patient phenotypes and might provide a path to the development of improved therapeutics.

**MATERIALS AND METHODS**

**Generation of NET A457P knock-in mice**

Knock-in mice carrying the NET A457P allele (GenBank accession no. AY188506), the orthologous sequence to the human NET A457P allele (GenBank accession no. NM_001043.3), were generated as follows and as shown in Fig. 1. The targeting construct and Southern strategy were generated by Gene Dynamics, LLC. A 5.0 kb 5' arm and a 5.0 kb 3' arm were cloned by PCR from 129S6/SvEv genomic DNA. The 5' arm comprised NET exon and intron sequences from exon 6 through intron 11 and the 3' arm comprised sequences from exon 11 through the 3' untranslated region. The A457P mutation was generated in the 5' arm by site-directed mutagenesis of nucleotide 1387 in exon 9 from a guanosine to a cytosine. The arms were subcloned into a targeting vector, resulting in the placement of a CRE recombinase/neomycin selection cassette flanked by loxp sites in intron 11 of NET. The CRE was driven by a testis-specific angiotensin converting enzyme promoter to produce excision of the loxp-flanked region with germline targeting of the construct. The targeting construct was electroporated into 129S6/SvEv mouse ES cells in the Vanderbilt Transgenic Mouse/ES Cell Shared Resource. Positive ES cell clones were confirmed for homologous recombination by Southern blot and PCR analyses. For Southern blot analysis, ES cell DNA was digested with BglII, and hybridized...
with a probe 3’ to and outside of the 3’ arm, with predicted bands of 7.8 and 11.6 kb for the untargeted and recombined NET gene, respectively. For PCR of the 3’ end of the targeted vector, a forward primer located within the CRE region and a reverse primer located outside of the 3’ arm (5’-GAGGTCCTCTGTGAGGCTGTG-3’ and 5’-CCCCCTCCTCAAATCTAAG-3’, respectively) were used to generate a 5.1 kb fragment in recombined clones only. Additionally, PCR of the 5’ arm was performed using a forward primer located 5’ to the 5’ arm and a reverse primer located in the 3’ arm (5’-GGGCAAGACACACACTAGAC-3’ and 5’-CAGCCCCAGAGATGACAG-3’, respectively), producing a product that includes the A57P site. The PCR products were subjected to dideoxy sequencing for A57P. Chimeras generated from selected ES cells were intercrossed with C57BL/6J mice. Offspring were genotyped using PCR primers spanning the position of the loxP site remaining after excision of the floxed region (5’-TGTGTGTTGAAGTTCTGAGC-3’ and 5’-GGGCAAGACACACACTAGAC-3’, respectively) to generate PCR products of 346 and 312 bp for wild-type and targeted mice, respectively. This PCR strategy was used subsequently to genotype all offspring. Additionally, mice in the initial litters were also sequenced for the presence of A57P through PCR and dideoxy sequencing of a region spanning the A57P mutation (5’-CTATAGATGGGAGGCATGGAG-5’ and 5’-CAGCCCCAGAGATGACAG-5’). Speed congenics were used to backcross A57P mice to the C57BL/6J strain in six generations at Jackson Laboratories (Bar Harbor, ME). Speed congenics is a procedure used to decrease the number of generations of breeding required to produce congenic mice on the desired background strain. A panel of single nucleotide polymorphism (SNPs) is genotyped to verify congenicity. Mice were bred during the speed congenics process, mice were bred and maintained in the Vanderbilt University animal facilities. Mice were anesthetized with 2-3% isoflurane, and body temperature was determined using a mouse thermocouple (DSI) was performed on adult, male NET A57P knock-in mice. Mice were anesthetized with 2-3% isoflurane, and body temperature was monitored using radiotelemetry.

Net A457P mice display POTS

Radiolabeled transport assay

[3H]NE transport was assayed in KRH buffer. Synaptosomes were preincubated for 10 minutes at 37°C, with or without 1 μM desipramine, to assess nonspecific accumulation, followed by the addition of 150 nM (single-point) or varying concentrations of [3H]NE (10 nM to 1 μM for kinetic analysis) (~45 Ci/mmol; PerkinElmer Life Sciences). Transport assays were terminated 10 minutes following the addition of [3H]NE by filtration of synaptosomes over 0.3% polyethylenimine-coated glass fiber filters (Whatman GF/B; Whatman) using a cell harvester (Brandel Inc., Gaithersburg, MD). Accumulated [3H]NE was determined by liquid scintillation counting (Beckman LS 6000). Peripheral tissues were finely minced and [3H]NE (300 nM or 1 μM) was used for a 30-minute transport assay. Following incubation with [3H]NE, tissue minces were washed 3x with KRH buffer and accumulated [3H]NE determined by liquid scintillation counting.

Surface biotinylation

To investigate the level of NET surface expression, biotinylation was performed on intact synaptosomes. Synaptosomes were incubated with cell-impermeant sulfo-NHS-2-(biotinamido)ethyl-1,3-dithiopropropionate (sulfo-NHS-SS-biotin; 3.0 mg/ml; Pierce) for 30 minutes at 4°C, washed, quenched with 100 mM glycine, extracted in radioimmunoprecipitation assay (RIPA) buffer (10 mM Tris, 150 mM NaCl, 1 mM EDTA, 0.1% SDS, 1% Triton X-100, 1% sodium deoxycholate, pH 7.4, with 250 μM PMSE, 1 μg/ml aprotinin, 1 μg/ml leupeptin, and 1 μM pepstatin) for 30 minutes at 4°C, and then incubated with a 50% slurry of immobilized streptavidin agarose beads (Pierce) for 30 minutes at room temperature. Beads were washed three times in RIPA buffer, and proteins that were bound to streptavidin beads were eluted in 1x Laemmli sample buffer containing 2-mercaptoethanol. Total NET levels were measured in synaptosomes extracted in RIPA buffer. Samples were separated by 10% SDS-PAGE and immunoblotted as described for western blot analyses.

Western blots

Proteins were electrophoretically transferred overnight at 4°C to Immobilon-P polyvinylidene fluoride membrane (Millipore). Membranes were incubated in blocking solution (PBS, 2% Tween 20, 5% dry milk), followed by incubation with a monoclonal antibody directed against NET at a dilution of 1:500 (NET50; Mab Technologies, Stone Mountain, GA) and incubation with goat anti-mouse HRP-conjugated secondary antibody at a dilution of 1:5000 for analysis (Jackson ImmunoResearch). Antibody visualization was performed using Western Lightning chemiluminescence reagents (PerkinElmer Life Sciences). Quantitation of band density was performed using ImageJ (NIH). Gel analyses were performed in replicates, and data are presented with representative blots and graphs of mean optical density from replicate experiments. No differences were observed between male and female A457P mice for NET expression and male and female results were combined for analysis.

Radiotelemetry

Mouse blood pressure radiotelemetry [Data Sciences International (DSI)] was performed on adult, male NET A457P knock-in mice. Mice were anesthetized with 2-3% isoflurane, and body temperature...
was maintained with an isothermal pad (Braintree Scientific, Inc., Braintree, MA). After antiseptic preparation, the left carotid artery was isolated, and the lumen was cut to allow insertion of the transmitter catheter to the point of bifurcation. The transmitter body was placed in a lateral subcutaneous pocket. The skin was sutured and secured with adhesive (Nexaband, Veterinary Products Laboratories). The singly-housed mice were monitored in their home cage. BP and HR were recorded continuously in 1-second intervals using flatbed radiofrequency receivers (PhysioTel Receiver RPC-1; DSI) and a digital acquisition system (Dataquest A.R.T.; DSI). Data were analyzed by customized MatLab software (Mathworks) in 30-minute increments, averaged over 24-hour time periods or 12-hour time periods, to coincide with the light/dark cycle. Although data was collected continuously, data presented were collected from mice allowed to recover for at least 10 days following surgery.

**ECG recording from anesthetized mice**

Under isoflurane anesthesia, mice were instrumented with arterial and jugular vein catheters for electrocardiogram (ECG) recording of HR and BP, and intravenous drug delivery, respectively. Mice were maintained on an isothermal pad throughout the experiment to maintain body temperature at 36-37°C. The arterial catheter was Micro-Renathane tubing (0.025-inch internal diameter (ID); 0.040-inch outer diameter (OD); Braintree Scientific) stretched over hot air to give a tip 300-500 μm in diameter. This tubing was coupled with a smoothed segment of 23 G needle to a 25-cm piece of polyethylene tubing (PE−50; 0.023-inch ID, 0.038-inch OD; Clay-Adams). A small incision was made in the femoral artery and the catheter was advanced into the vessel so that the tip was 0.5 cm above the aortic bifurcation, within the abdominal aorta. Next, an incision was made in the right jugular vein, and a 4- to 5-cm length of Micro-Renathane tubing (0.12-inch ID; 0.25-inch OD) prefilled with heparinized saline (1:1) was inserted and surgically secured. The pharmacological protocol began at the completion of surgery with the mouse remaining under continuous anesthesia. To determine the baroreceptor-mediated cardioinhibitory response, mice underwent a challenge with phenylephrine (PE; 5-40 μg/kg body weight) or nitroprusside (10-40 μg/kg) given intravenously using a syringe pump (CMA 400 pump, CMA Microdialysis, Stockholm, Sweden) in a dose-response manner. Baseline ECG and BP were recorded for 1 minute before drug administration, and the post-drug response was recorded for 3 minutes after administration. The ratio of the maximal change in HR over the change of mean systolic BP was calculated and averaged at each dose in each animal. BRS was determined by the averaged ratio of HR change [change in the R-wave to R-wave interval (RRI) over systolic BP (SBP) change (ΔRRI/ΔSBP)].

**Catecholamine assay**

Catecholamines and metabolites were measured in plasma, urine and tissue. Blood was collected via cardiac puncture in isoflurane-anesthetized mice and preserved with EGTA and reduced glutathione, and organs were harvested immediately thereafter. Spot urine was collected at ~9:00 am from awake, unstressed mice placed in individual weighboats. 6 N HCl was added to urine samples to prevent catechol breakdown. Urine and plasma were alumina extracted and tissues were extracted in 0.1 N perchloric acid. Catecholamine and metabolite concentrations were determined by high-performance liquid chromatography combined with electrochemical detection (HPLC-EC). 3,4-dihydroxybenzylamine (DHBA) was used as an internal standard in all samples. Urine catecholamine and metabolite measures were normalized to urine creatinine levels.

**Behavior**

**Open field activity**

All behavior was performed in the Vanderbilt Murine Neurobehavioral Laboratory. For the open field test, mice were placed in open field activity chambers (29×29×20.5 cm) that were housed inside environmental chambers (Med Associates, Inc., St Albans, VT). The environmental chambers remained closed for the duration of the test. The testing room was kept in normal lighting (250-300 lux) during testing. Testing was begun at ~9:00 am. Mice were exposed to the open field for 1 hour. Movement and location were detected by the interruption of infrared photoelectric beams by the mouse and distance travelled in center and surrounding zones was calculated in 5-minute blocks (Med Associates).

**Elevated plus maze**

The elevated plus maze (EPM) consisted of two closed and two open arms connected by a center zone (dimensions are 67 cm length×6.5 cm width×15 cm height in the closed arms). The EPM was set on a table during testing. The testing room was kept in normal lighting (250-300 lux). Testing was begun at ~9 am. Mice were brought into the testing room, in home cages, 30 minutes prior to testing with free access to food and water. To initiate the test, mice were lowered by the tail onto the center of the maze, directly next to and facing a closed arm. Test duration was 5 minutes and was video-recorded by an overhead camera. Time spent in each zone, number of entries into each zone and freezing behavior were analyzed using ANY-maze software (Stoelting, Wood Dale, IL). Head dips were counted by an observer blind to mouse genotype.

**Tail suspension**

Mice were brought into an anteroom 30 minutes prior to testing. For the tail suspension test, mice were suspended for 7 minutes by the tail from a vertical aluminum bar (bar size: 11.5×2.3 cm) attached to the top of a box-like enclosure (33×33×32 cm) open on the front side (Med Associates). Mice were attached to the bar by tape placed ~1.5 cm from the tip of the tail. Mice were monitored visually throughout the test to ensure they did not tail climb. The testing room was kept in normal lighting (250-300 lux) during the test. Testing was begun at ~9:00 am. The level and duration of force placed on the bar by the mouse was measured, with a force level below the lower threshold being counted as immobilization (Med Associates). The following settings were used: lower threshold of 7, upper threshold of 20, gain of 8, and resolution of 220 milliseconds.

**Statistics**

Data were analyzed by one- or two-way ANOVA followed by post-hoc testing, or by two-tailed t-tests, as appropriate for experimental design and a priori predictions (SPSS20, IBM; Prism version 5, GraphPad Software Inc.). P-values <0.05 were considered.


