NPBWR1 and NPBWR2: implications in energy homeostasis, pain, and emotion

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INTRODUCTION

Receptors for neuropeptide B (NPB)/neuropeptide W (NPW) were originally identified as orphan receptors GPR7 and GPR8, which share 70% nucleotide identity and 64% amino acid identity with each other. They have relatively high similarities with opioid and somatostatin receptors. GPR8 was not found in the rodent genomes, while GPR7 was highly conserved in both human and rodents (O’Dowd et al., 1995). This suggests the GPR8 gene was relatively recently generated through gene duplication.

In 2002–2003, two endogenous peptide ligands for these receptors were identified and named NPB/NPW (Fujii et al., 2002; Shimomura et al., 2002; Brezillon et al., 2003; Tanaka et al., 2003). Following the deorphaning of these receptors, GPR7 and GPR8 were reclassified by IUPHAR as NPB/W receptor-1 (NPBWR1) and NPB/W receptor-2 (NPBWR2), respectively (Davenport and Singh, 2005).

In rodents, Npbwr1 mRNA is localized in some discrete brain regions, including the hypothalamus (dorsomedial hypothalamus and suprachiasmatic nucleus), hippocampus, ventral tegmental area (VTA), and extended amygdala (CeA and bed nucleus of the stria terminalis; BNST) (Lee et al., 1999; Tanaka et al., 2003), suggesting this receptor is involved in regulatory mechanisms of homeostasis, reward system, and emotion.

This review discusses recent findings concerning the pharmacology, histology, and phenotypic analysis of genetically engineered mice of the NPB/NPW system.

IDENTIFICATION OF NPB AND NPW

In 2002–2003, three groups independently identified endogenous peptide ligands for GPR7 and GPR8 by so-called “reverse pharmacology” in combination with bioinformatics approaches. To identify the cognate endogenous ligands for GPR7 (NPBWR1) and GPR8 (NPBWR2), Shimomura et al. (2002) expressed these receptors in Chinese Hamster Ovary (CHO) cells and measured the decrease in forskolin-induced cAMP production in these cells as the read-out for receptor activation. While screening the bovine hypothalamic extract fractions, they detected activities that specifically inhibit cAMP production in those cells, and purified the activities. During their purification process, they identified two forms of NPW with different peptide lengths of 23 and 30 amino acid residues, and named them NPW23 and NPW30, respectively.

In a similar manner, three groups (Fujii et al., 2002; Brezillon et al., 2003; Tanaka et al., 2003) independently purified and identified an additional endogenous ligand for NPBW1 and NPBW2. Fujii et al. first screened the Celera database to identify novel secretory peptides and then expressed the cDNAs of the putative secretory peptides to find novel peptide ligands. Subsequent pharmacological studies and purification of the peptide from bovine hypothalamic extracts led to identification of the second ligand for GPR7 and GPR8, which was named NPB due to the unique modification, namely, bromination of the first tryptophan residue. Brezillon et al. (2003) also identified human NPB mRNA using a bioinformatics approach by searching the EST database using the NPW sequence as a query.

Tanaka et al. used a unique melanin-pigment aggregation assay in Xenopus melanophore cells expressing GPR7 as the assay system to purify NPB from bovine hypothalamus. Through EST database searches they also identified NPW as a putative paralogous peptide (Tanaka et al., 2003).

STRUCTURES OF NPW AND NPB

Neuropeptide W and NPB do not display any significant sequence similarity to other known peptide families, while sharing a high degree of sequence similarity with each other, constituting a distinct family of peptides (Figure 1A) (Fujii et al., 2002; Shimomura et al., 2002; Brezillon et al., 2003; Tanaka et al., 2003).

As mentioned earlier, sequence analysis of purified peptides showed that NPW has two isoforms with lengths of 23 and 30 amino acid residues, i.e., neuropeptide W23 (NPW23) and...
neuropeptide W30 (NPW30), respectively. NPW23 is produced as a result of proteolytic processing at a pair of arginine residues in the 24th and 25th positions in NPW30 (Fujii et al., 2002; Brezillon et al., 2003; Tanaka et al., 2003). In vitro experiments using cells expressing these receptors showed that synthetic NPW23 activates and binds to both NPBW1 and NPBW2 at similar effective doses, while NPW30 shows slightly lower affinity to both receptors as compared with NPW23 (Figure 1B) (Brezillon et al., 2003; Tanaka et al., 2003).

As mentioned earlier, NPB has a unique modification at the N-terminus tryptophan residue, C-6-bromination (Fujii et al., 2002; Tanaka et al., 2003). While this represents the first evidence of bromination of the protein in mammals, the biological significance of this modification is unclear, as it has been demonstrated that des-bromo-NPB is equipotent to brominated NPB in in vitro CAMP inhibition assays (Tanaka et al., 2003). By analogy with NPW, Brezillon et al. (2003) predicted that two isoforms of NPB, NPB23 and NPB29, could be produced from the processing of a 125-amino acid human precursor through the alternative usage of a dibasic amino acid pair. However, the dibasic motif, Arg24–Arg25, which is seen in human NPB, does not exist in NPB of other mammalian species including bovine, rat, and mouse. Therefore, it is unlikely that the NPB23 isoform exists as a mature peptide in non-human species. Consistently, both Fujii et al. and Tanaka et al. were only able to isolate NPB29 from bovine hypothalamus extracts during their purification procedures.

In X. laevis melanophore pigment aggregation assay, NPB29 binds and activates human NPBW1 or NPBW2 with median effective concentrations (EC50) of 0.23 and 15.8 nM, respectively (Tanaka et al., 2003), suggesting that NPB is a relatively selective agonist for NPBW1 (Figure 1B).

The preferred conformations of des-bromo-NPB and -NPW have been determined by a combination of 1H NMR, CD, and molecular modeling (Lucyk and Miskolzie, 2005).

**STRUCTURE-ACTIVITY RELATIONSHIPS OF NPW AND NPB**

The rank order of potency of NPB and NPW isoforms has been determined in cell lines expressing NPBW1 and NPBW2. Both NPB and NPW bind and activate NPBW1 and NPBW2 receptors with varying degrees of affinity. NPBW1 has slightly higher affinity for NPB as compared with both forms of NPW, whereas NPBW2 shows a potency rank order of NPW23 > NPW30 > NPB (Figure 1B) (Fujii et al., 2002; Shimo-mura et al., 2002; Brezillon et al., 2003; Tanaka et al., 2003). NPW23 and NPB29 have similar affinity for both receptors. NPW23 and NPW30 bind to both receptors almost equally, while NPB acts as a relatively selective agonist for NPBW1.
Tanaka et al. (2003) showed that deletion of Trp-1 from NPB or NPW markedly decreased their activity, suggesting that the N-terminus is involved in receptor binding. This is consistent with the fact that NPB and NPW have similarity in their sequences in the N-terminal region.

**STRUCTURES AND FUNCTIONS OF NPBWR1 AND NPBWR2**

The human NPBWR1 and NPBWR2 genes are localized on chromosome 10q11.2–121.1 and 20q13.3, respectively. Human NPBWR1 and NPBWR2 have 328 and 333 amino acid residues, respectively, and share 64% sequence homology with each other (Figure 2). Among other family members of GPCRs, NPBWR1 and NPBWR2 are most closely related to opioid and somatostatin receptors (Figure 2) (O’Dowd et al., 1995). Amino acid analysis of NPBWR1 orthologs in other mammalian species has revealed a high degree of conservation throughout evolution (Lee et al., 1999). In contrast, while the gene encoding NPBWR2 has been discovered in several mammalian species such as monkey, lemur, bat, shrew, and rabbit, it has not been found in rodent genomes (Lee et al., 1999), suggesting that these two receptors were produced by relatively phylogenetically recent gene duplication (Lee et al., 1999).

As suggested by the assay systems used during their purification process and structural similarities to somatostatin/opioid receptors, both NPBWR1 and NPBWR2 couple to the Gi-class of G-proteins (Tanaka et al., 2003). This suggests that these neuropeptides have inhibitory properties on neurons via activation of GIRK (Kir3) channels. NPB and NPW were also shown to stimulate Erk p42/p44 activity in human adrenocortical carcinoma-derived NCI-H295 cells (Andreis et al., 2005). This activation is probably mediated by beta/gamma subunits released from G-proteins (Tim van et al., 1995). Recently, a synthetic low molecular weight antagonist for NPBWR1 has been reported (Anthony Romero et al., 2012).

**DISTRIBUTION OF NPB/NPW AND NPBWR1/NPBWR2 IN BRAIN AND OTHER TISSUE**

**NEUROPEPTIDE B**

Npb mRNA was found in several specific regions in the mouse brain such as the paraventricular hypothalamic nucleus (PVN), CA1–CA3 fields of the hippocampus, and several nuclei in the midbrain and brainstem, including the Edinger–Westphal (EW) nucleus, the sensory and motor nuclei of the trigeminal nerve, locus coeruleus (LC), inferior olive, and lateral parabrachial nucleus (Figure 3) (Tanaka et al., 2003; Jackson et al., 2006).

Schulz et al. reported that NPB-immunoreactive cell bodies were observed in many regions within the hypothalamus. High levels of NPBwr1 and Npb mRNA expression were also observed in the hypothalamus, including the ventromedial hypothalamic (VMH) and the lateral hypothalamic (LH) regions.

**GPR7**

This receptor is highly expressed in the hippocampus and amygdala and may be involved in the regulation of food intake and energy homeostasis.

**GPR8**

This receptor is expressed in the hypothalamus and may be involved in the regulation of body temperature and energy expenditure.

**SOMATOSTATIN RECEPTOR SSTR3**

This receptor is expressed in the hypothalamus and may be involved in the regulation of food intake and energy homeostasis.

**DELTAP OPIOID RECEPTOR**

This receptor is expressed in the hypothalamus and may be involved in the regulation of body temperature and energy expenditure.

**KAPPA OPIOID RECEPTOR**

This receptor is expressed in the hypothalamus and may be involved in the regulation of food intake and energy homeostasis.

**M1 OPIOID RECEPTOR**

This receptor is expressed in the hypothalamus and may be involved in the regulation of body temperature and energy expenditure.

**FIGURE 2** | Primary structures of NPBWR1/NPBWR2 compared to somatostatin and opioid receptors. Conserved amino acid residues are shown in red. Each transmembrane domain is boxed. Modified from Hondo et al. (2008).
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FIGURE 3 | Schematic representation of distribution of NPB/NPW and Npbwr1 mRNA in mouse brain coronal sections. Receptor distribution is shown in the right hemisphere, while ligand distribution is shown in the left. Npb mRNA (red regions) was observed in the hippocampus (CA1, CA2, CA3), lateral habenular nucleus (LHb), paraventricular hypothalamic nucleus, medial parvicellular part (PaMP), Edinger–Westphal (EW) nucleus, motor root of the trigeminal nerve (m5), sensory root of the trigeminal nerve (s5), lateral parabrachial nucleus alpha part (Sub CA), locus coeruleus (LC), noradrenergic cell group A5 (A5), and inferior olive subnucleus B (IOB) (Tanaka et al., 2003). Npw mRNA (blue regions) was observed in the periaqueductal gray (PAG) matter, EW nucleus (EW), ventral tegmental area (VTA), and dorsal raphe nucleus (DR). NPBWR1 mRNA (yellow regions) was observed in the claustrum (Cl), dorsal endopiriform nucleus (DEn), bed nucleus of the stria terminals, laterodorsal part (BSTLD), bed nucleus of the stria terminals, mediodorsal part (BSTMD), suprachiasmatic nucleus (Sch), magnocellular preoptic nucleus (MCPo), paraventricular hypothalamic nucleus, posterior part (PaPo), dorsomedial hypothalamic nucleus (DM), central amygdala (CeA), CA1 field, hippocampus (CA1), ventral tegmental area (VTA), sensory root trigeminal nerve (Su5), subiculum (S), anterior hypothalamic area, posterior part (AHp), and arcuate hypothalamic nucleus (Arc). Modified from Hondo et al. (2008).

nucleus, dorsomedial hypothalamic nucleus, arcuate nucleus, supraoptic retrochiasmatic nucleus, and in the area ventral to the zona incerta (Schulz et al., 2007). Although Npb mRNA was detected in several regions outside the hypothalamus such as the hippocampus and brain stem (Tanaka et al., 2003; Jackson et al., 2006; Schulz et al., 2007), the existence of NPB-immunoreactive neurons in these regions in the brain has not been clearly demonstrated, since no good antibody for NPB is available.

In peripheral tissues, expression of human Npb mRNA was detected by RT-PCR in the kidney, uterus, ovary, testis, and placenta, while murine Npb mRNA was detected by Northern blot at high levels in the stomach, spinal cord, and testis and at lower levels in the liver and kidney (Brezillon et al., 2003). The biological significance of this expression remains unknown.

NEUROPEPTIDE W

The existence of NPW in the mouse brain was more confined to several nuclei in the midbrain and brainstem including the EW, VTA, periaqueductal gray (PAG), and dorsal raphe nucleus (DR) (Figure 3) (Tanaka et al., 2003; Kitamura et al., 2006). Most of these neurons expressed tyrosine hydroxylase (TH), suggesting that these cells are also catecholaminergic (our unpublished observation). In humans, high levels of Npw mRNA were detected in the substantia nigra, and moderate expression levels were detected in the amygdala and hippocampus (Fujii et al., 2002). In peripheral tissues, expression of human Npw mRNA was confirmed by RT-PCR in the progenital system, comprising the kidney, testis, uterus, ovary, and placenta, and also in the stomach and respiratory system, while murine Npw mRNA was detected by Northern blot at
high levels in the lung and lower levels in the stomach (Brezillon et al., 2003).

Consistent with the mRNA distribution, NPW-immunoreactive (ir) cells were also exclusively detected in the EW, VTA, PAG, and DR in rats (Kitamura et al., 2006). NPW-ir fibers were observed in several brain regions in rats including the lateral septum, bed nucleus of the stria terminalis (BNST), dorsomedial and posterior hypothalamus, CeA, CA1 field of hippocampus, interpeduncular nucleus, inferior colliculus, lateral parabrachial nucleus, facial nucleus, and hypoglossal nucleus. Among these regions, NPW-ir fibers were most abundantly observed in the CeA and BNST, the output nuclei of the extended amygdala. These observations suggest that NPW-producing neurons are exclusively localized to the midbrain, and they project mainly to the limbic system, especially the CeA and BNST (Figure 3).

Some reports showed the existence of NPW-ir cell bodies in the PVN in rats and mice (Dun et al., 2003). However, the staining of NPW-like immunoreactivity-positive cells in the PVN is likely to be non-specific staining, for two main reasons (Kitamura et al., 2006). First, PVN immunoreactivity is also observed in Npw−/− mice using many of the commercially available antibodies. Second, Npw mRNA is not expressed in the PVN in either mice or rats (Kitamura et al., 2006).

**NPBWR1 (GPR7)**

*In situ* hybridization study showed that the CeA and BNST express the highest levels of NPBWR1 mRNA expression in the mouse brain (Figure 3) (Tanaka et al., 2003; Singh et al., 2004; Jackson et al., 2006). Other nuclei with high levels of Npbwr1 are the suprachiasmatic (SCN) and ventral tuberomammillary nuclei of the hypothalamus. Moderate levels of expression are seen in the CA1–CA3 regions of the hippocampus, dorsal endopiriform, dorsal tenia tecta, bed nucleus, and red nucleus. Low level expression of Npbwr1 is seen in the olfactory bulb, parastrial nucleus, hypothalamus, laterodorsal tegmentum, superior colliculus, LC, and nucleus of the solitary tract.

**NPBWR2 (GPR8)**

Because Npbwr2 does not exist in rodent genomes, very limited information about the tissue distribution of NPBWR2 has been available thus far. RT-PCR analysis showed that Npbwr2 mRNA is strongly expressed in the human amygdala and hippocampus. Lower levels of expression were also detected in the corpus callosum, cerebellum, substantia nigra, and caudate nucleus (Brezillon et al., 2003).

**BIOLOGICAL ACTIVITIES OF NPB AND NPW**

Several papers reported pharmacological actions of NPB and NPW *in vivo* (Table 1). In this section, I discuss biological activities of these peptides, referring the pharmacological actions and phenotypes of genetically-modified mice on NPB/W systems (Table 2).

**FEEDING AND ENERGY HOMEOSTASIS**

The NPB/W system was initially thought to be involved in regulation of feeding behavior (Shimomura et al., 2002; Tanaka et al., 2003), and therefore many of the currently available reports have focused on the roles of NPB/W in this function. The first physiological study on the action of NPW showed acute hyperphagia in male rats when NPW was administered intracerebroventriculatly (i.c.v.) (Shimomura et al., 2002; Tanaka et al., 2003). However, Tanaka et al. (2003) showed that the effect of NPB on feeding behavior in mice is very complex. When NPB was i.c.v. injected during the light period, no significant effect of NPB on feeding was observed (Tanaka et al., 2003). In contrast, in the dark period, i.c.v. administration of 3 nmol NPB increased feeding only within the first 2 h. A higher dose of NPB suppressed food intake in this period. After 2 h, both doses of NPB decreased food intake. Because rodents only have NPBWR1 as a receptor for NPB, the biphasic, i.e., earlier orexigenic and later anorexic, effect cannot stem from the different potency rank orders of the two peptides on NPBWR1 and NPBWR2. In fact, we also observed a similar biphasic action of NPW on feeding behavior when administered i.c.v. in mice or rats (our unpublished observation). Mondal et al. (2003) also reported anorexic effects of NPW. These findings suggest a complex role of NPB and NPW in the regulation of food intake.

Interestingly, the anorexic effect of NPB was markedly enhanced when corticotrophin-releasing factor (CRF), a known anorexigenic peptide, was co-administered (Tanaka et al., 2003). The i.c.v. administration of these two peptides almost completely suppressed food intake over 4 h. The biphasic effects of NPB/W on feeding behavior and the synergistic anorexic effects of NPB and CRF suggest complex roles of these peptides in the regulation of feeding behavior in relation to the stress response. The synergic effect of NPB with CRF in suppression of feeding suggests that this neuropeptide is likely to be implicated in inhibition of feeding under stressful conditions.

Continuous i.c.v. infusion of NPW was reported to suppress feeding and body weight gain over the infusion period (Mondal et al., 2003). Conversely, i.c.v. administration of anti-NPW IgG stimulated feeding, suggesting that endogenous NPW plays an inhibitory role in feeding behavior (Mondal et al., 2003). However, unlike the results of continuous i.c.v. infusion of NPW, bolus intra-PVN microinjection of NPW23 at doses ranging from 0.1 to 3 nmol increased feeding for up to 4 h, and a bolus dose ranging from 0.3 to 3 nmol was reported to increase feeding for up to 24 h (Levine et al., 2005). These observations suggest that the orexigenic versus anorectic effects of NPB/W could result from different sites of action. When these peptides are injected into the lateral ventricle, they might initially act on the PVN, followed by diffusion to other regions implicated in the suppression of feeding. Alternatively, delayed inhibition of feeding by NPB/W might result from the production of other anorectic factors that are stimulated by NPB or NPW.

The i.c.v. administration of NPW also increased body temperature and heat production (Mondal et al., 2003). These effects suggest that endogenous NPB/W might increase energy expenditure, which is consistent with the late-onset obesity seen in male NPBWR1−/− mice and NPB−/− mice (Ishii et al., 2003; Kelly et al., 2005).

Male NPBWR1−/− mice have been shown to develop adult-onset obesity that progressively worsens with age and is greatly exacerbated when animals are fed a high-fat diet (Ishii et al., 2003). The obesity is caused by hyperphagia along with decreased
Table 1 | In vivo pharmacological effects of NPB/NPW.

| Substance | Effects | Animal | Preference |
|-----------|---------|--------|------------|
| NPB (i.c.v.) | Food intake ↑ | Rat (male) | Shimomura et al. (2002) |
| NPB (i.c.v.) | Body weight ↑ | Rat | Tanaka et al. (2003) |
| NPB (i.c.v.) | Body temperature ↑ | Rat | Mondal et al. (2003) |
| NPB (i.c.v.) | ACTH ↑ | Rat | Hoehol et al. (2007) |
| NPB (i.c.v.) | Heat production ↑ | Rat | Yu et al. (2007) |
| NPW30 (i.c.v.) | Arterial blood pressure (ABP) ↑ | Rat | Tanaka et al. (2003) |
| NPW (i.c.v.) | Food intake (light period–dark period) ↑ | Mouse | Samson et al. (2004) |
| NPW (i.c.v.) | Prolactin ↑ | Rat (male) | Samson et al. (2004) |
| NPW (i.c.v.) | Growth hormone ↑ | Rat | Taylor et al. (2005) |
| NPW23/NPB (i.t) | Inflammatory pain ↓ | Rat | Yamamoto et al. (2005) |
| NPW/NPB (i.c.v.) | Corticosterone in circulation ↑ | Rat | Samson et al. (2004) |
| NPW/NPB (i.p) | Plasma level parathyroid hormone ↑ | Rat | Taylor et al. (2005) |
| NPW/NPB (i.c.v.) | Circadian rhythm → | Rat/mouse | Our unpublished observations |

Table 2 | Summary of behavioral phenotypes of NPBWR1 knockout mice (modified from Nagata-Kuroiwa et al., 2011).

| Behavioral test | Parameter | Results |
|-----------------|-----------|---------|
| Open field test | Anxiety | Normal time spent in center of arena |
| Elevated plus maze test | Anxiety | Normal time spent and number of entries in open arms |
| Light-dark exploration test | Anxiety | Decrease in escape latency and time spent in light box |
| Porsolt forced swim test | Depression, learning helplessness | Normal time spent swimming |
| Prepulse inhibition test | Sensory motor reactivity | Normal percentage of prepulse inhibition |
| Marble burying behavior test | Compulsive behavior | Normal number of marbles buried |
| Cued and contextual fear conditioning test | Fear and memory | Decrease in time of freezing behavior during contextual testing while normal during auditory-cued testing |
| Morris water maze test | Spatial memory | Normal escape latency |
| Resident-intruder test | Social interaction | Abnormal social interaction (see text) |
| Stress-induced hyperthermia | Stress response | Exaggerated hyperthermia |
| Daily locomotor activity | Circadian rhythm | Normal in both light/dark cycle and constant dark condition. Normal entrainment by food or light |
| Sleep-wake behavior (EEG/EMG) | Sleep/wake cycle | Normal in each episode duration, times spent in each state in hourly sleep/wake analysis |

energy expenditure. Npbwr1−/− male mice showed decreased hypothalamic neuropeptide Y mRNA level and increased proopiomelanocortin mRNA level, a set of effects opposite to those evident in ob/ob mice. Furthermore, ob/ob;NPBWRI−/− and Ay/a;NPBWRI−/− double mutant male mice had an increased body weight compared with normal ob/ob or Ay/a male mice, suggesting that the obesity of NPBWR1−/− mice is independent of leptin and melanocortin signaling. Female mice did not show any significant weight increase or associated metabolic defects. These data suggest a potential role for NPBWR1 in regulating energy.
homeostasis independent of leptin and melanocortin signaling, in a sexually dimorphic manner (Ishii et al., 2003). Consistent with the phenotype of Npbwr1−/− mice, Nprl−/− mice were also reported to exhibit mild adult-onset obesity (Kelly et al., 2005).

PAIN REGULATION
Intracerebroventricular administration of NPB produces analgesia to pain induced by subcutaneous formalin injection in rats (Tanaka et al., 2003). Intrathecal (i.t.) injection of either NPW23 or NPB also decreased agitation behaviors induced by formalin injection into the paw and attenuated the level of mechanical allodynia (Yamamoto et al., 2005). The effects were not antagonized by naloxone, suggesting that this effect is not mediated through the opioid receptor system. While i.t. injection of either NPW23 or NPB did not show any effect in the hot plate test or mechanical nociceptive test, i.t. injection of either NPW23 or NPB significantly suppressed the expression of Fos-like immunoreactivity of the L4–5 spinal dorsal horn induced by formalin injection into the paw. These data suggest that spinally applied NPW/B suppressed the input of nociceptive information to the spinal dorsal horn induced by formalin injection into the paw. These data suggest that spinally applied NPW/B suppressed the input of nociceptive information to the spinal dorsal horn, producing an analgesic effect on inflammatory pain, but not mechanical or thermal pain (Yamamoto et al., 2005).

Consistent with the pharmacological studies suggesting an analgesic action of NPB on inflammatory pain, NPB−/− mice exhibited hyperalgesia to inflammatory pain, while they showed normal responses to mechanical or thermal pain (Kelly et al., 2005). These observations suggest that NPB in the brain and/or spinal cord modulates pain in a modality-specific manner. These observations suggest that NPB might play a role in the pathophysiology of allodynia. Unfortunately, however there are no available data about expression of NWP, NPB, or NPBWR1 in the spinal cord or dorsal root ganglia.

A low level of NPBWR1 receptor expression was observed in Schwann cells in both normal human and rat nerves as well as in primary rat Schwann cell cultures. Peripheral nerve samples taken from patients exhibiting inflammatory/immune-mediated neuropathy showed a marked increase of NPBWR1 receptor expression restricted to myelin-forming Schwann cells. Complementary animal models of immune-inflammatory and ligation-induced nerve injury and neuropathic pain similarly exhibited increased myelin-associated expression of NPBWR1 (Zaratin et al., 2005). These observations suggest that NPBWR1 is involved in regulation of inflammatory pain at least in part by modulating Schwann cell function.

NEUROENDOCRINE REGULATION
Central administration of NPB or NPW elevated the plasma corticosterone level in rats (Samson et al., 2004; Taylor et al., 2005). NPB also increased prolactin and decreased growth hormone levels (Samson et al., 2004). Pretreatment with a polyclonal anti-CRF antiserum or CRF antagonist completely blocked the effect of NPB or NPW to stimulate ACTH release, and significantly inhibited the effect of NPW/W on plasma corticosterone level (Samson et al., 2004; Taylor et al., 2005). These findings suggest that NPW and NPB play a role in the neuroendocrine response to stress via activation of the hypothalamus-pituitary-adrenal (HPA) axis. Consistent with these observations, whole cell patch-clamp recording from hypothalamic slice preparations showed that bath application of NPW depolarized and directly increased the spike frequency of neuroendocrine PVN neurons (Taylor et al., 2005).

A whole cell patch-clamp study showed that NPW23 exhibited effects on oxytocin, vasopressin, and thyrotrophin-releasing hormone neurons in the PVN, although both depolarizing and hyperpolarizing effects were observed in each of these cell groups (Price et al., 2009). Corticotrophin-releasing hormone cells were unaffected. Further subdivision of chemically phenotyped cell groups into magnocellular, neuroendocrine, or pre-autonomic neurons, using their electrophysiological fingerprints, revealed that neurons projecting to medullary and spinal targets were predominantly inhibited by NPW, whereas those that projected to the median eminence or neural lobe showed almost equivalent numbers of depolarizing and hyperpolarizing cells (Price et al., 2009).

The intracellular mechanisms of these electrophysiological effects of NPW have remained unclear. Since NBPWR1 primarily couples to Gi subclass of G-proteins, activation of this receptor should lead to inhibition of neurons. Depolarizing effect of NPW described in above mentioned report might be indirect effect due to inhibition of inhibitory interneurons of recorded cells.

AUTONOMIC REGULATION
Intracerebroventricular administration of NPW to rats was reported to increase arterial blood pressure (ABP), heart rate (HR), and plasma catecholamine concentration (Yu et al., 2007). The same report showed that most of the PVN neurons were excited, while a subset of a smaller population of PVN neurons was inhibited by NPW30, although the chemical identities of these neurons were not determined. These observations suggest that NPW/W modulate PVN neuronal activities, which play an important role in regulation of the autonomic nervous system as well as the HPA axis. The expression of NPW mRNA in the PVN suggests that NPW is more likely to be involved in HPA axis regulation. However, NPBWR1−/− mice have normal blood pressure and HR in a basal state (Nagata-Kuroiwa et al., 2011). The physiological relevance of the NPW/W system in regulation of the autonomic nervous system remains unclear.

SLEEP AND WAKEFULNESS
Neuropeptide B is reported to induce slow wave sleep in mice when injected intracerebroventricularly (Hirashima et al., 2011). However, since NPBWR1−/− mice did not show any sleep abnormality, the biological relevance of this finding remains unclear.

EMOTION AND BEHAVIOR
Npbwr1 mRNA is abundantly expressed in discrete brain regions in rodents, including the hypothalamus (dorsomedial hypothalamus and suprachiasmatic nucleus), hippocampus, VTA, and extended amygdala (the CeA and bed nucleus of the stria terminalis; BNST) (Lee et al., 1999; Tanaka et al., 2003). These regions are implicated in mood and emotion. Especially, the particularly strong expression of Npbwr1 in the CeA and BNST, together with the robust projections of NPW-containing axons to these regions (Kitamura et al., 2006), suggest that this receptor might be an important modulator or regulator of the output signal from the extended amygdala, which is a well-defined subcortical nuclear group that is the center of emotion including fear (Phelps and LeDoux, 2005).
A sensory stimulus associated with an aversive outcome will change neural transmission in the amygdala to produce somatic, autonomic, and endocrine signs of fear, as well as increased attention to that stimulus. Fear learning involves the lateral and basolateral amygdala (BLA), where the association between incoming sensory stimuli leads to potentiation of synaptic transmission. The BLA receives sensory information from the thalamus, hippocampus, and cortex, and then activates or modulates synaptic transmission in target areas appropriate for the reinforcement signal with which the sensory information has been associated. The BLA projects to the CeA and BSNT, whose efferents to the hypothalamus and brainstem trigger the expression of fear. Histological and electrophysiological studies have shown that NPBWR1 acts as an inhibitory regulator on a subpopulation of GABAergic neurons in the lateral division of the CeA and terminates stress responses (Nagata-Kuroiwa et al., 2011). These anatomical and functional findings suggest that the NPB/W systems might have important roles in the modulation of output from the extended amygdala.

The role of NPBWR1 in social behavior was recently investigated using Npbwr1<sup>−/−</sup> mice (Nagata-Kuroiwa et al., 2011). When presented with an intruder mouse, Npbwr1<sup>−/−</sup> mice showed impulsive contact with the strange mice, produced more intense approaches toward them, and had longer contact and chasing time along with greater and sustained elevation of HR and blood pressure compared to wild type mice. Npbwr1<sup>−/−</sup> mice also showed increased autonomic and neuroendocrine responses to various physical stresses, suggesting that NPBWR1 might play a role in coping with stress.

Another interesting phenotype of NPBWR1<sup>−/−</sup> mice is the impairment of contextual fear conditioning and inversion of “safety conditioning.” In the safety conditioning paradigm, mice received unpaired presentations of a CS (tone) and a US (electrical shock). Because the shock never occurs during the CS, mice typically learn to treat the CS as a safety signal, so that fear-related behavior is inhibited during presentation of the CS. Npbwr1<sup>−/−</sup> mice show an inversion of this learning in that the safety-trained CS elicits fear rather than a reduction in fear. This observation suggests that Npbwr1<sup>−/−</sup> mice undergo trace conditioning rather than safety conditioning in this procedure, in that they associate the CS and US across a long trace interval, while the control mice treat CS and US as unpaired. These results suggest that Npbwr1<sup>−/−</sup> mice are more stimulus-bound, meaning that they preferentially attend to discrete stimuli, to the exclusion of more complex, conjunctive stimuli such as context. This hypothesis would also explain the deficit in contextual fear conditioning. To determine the neural mechanisms and the possible developmental and/or extra-amgdalar origins of the phenotype, further investigation using spatially restricted knockout mice and/or genetic rescue approaches toward them is needed.

The human NPBWR1 gene has a frequent single nucleotide polymorphism at nucleotide 404 (SNP rs33977775) in the coding region (404A→T). Importantly, this polymorphism causes an amino acid substitution (Y135F) within the highly conserved DRY motif of G-protein-coupled receptors at the junction of the third transmembrane domain and second intracellular loop, which is known to play an important role in G-protein coupling. Recently, Watanabe et al. (2012) demonstrated that this change partially impairs receptor function as measured by inhibition of cAMP production. Furthermore, this variation affects the emotional response to stimuli showing human faces with four categories of emotional expression (anger, fear, happiness, and neutral). The subject’s emotional levels at seeing these faces were rated on scales of hedonic valence, emotional arousal, and dominance (V–A–D).

A significant genotype difference was observed in valence evaluation; the 404AT group (position 404 in one allele is replaced by T) perceived the face as more pleasant than did the 404AA group (position 404 in both alleles of NPBWR1 is A), regardless of the category of facial expression. The 404AT group tended to feel less submissive to an angry face than did the 404AA group. These results suggest that a single nucleotide polymorphism of NPBWR1 seems to affect human behavior in a social context. Future studies using functional brain imaging of human subjects are needed to clarify the mechanisms of the effects of the polymorphisms.

**EFFECTS ON CIRCADIAN RHYTHM**

The abundant expression of NPBWR1 in the suprachiasmatic nucleus suggests the possibility that this neuropeptide/receptor system has a role in regulating circadian rhythm (Lee et al., 1999; Tanaka et al., 2003; Singh et al., 2004). However, we did not observe any effects of NPB/W on circadian activity in rats or mice when administered by i.c.v. injection. Npbwr1<sup>−/−</sup> mice displayed a normal circadian pattern of behavior in both light-dark and constant dark conditions (Uchio et al., 2009). Both light-entrainable and food-entrainable oscillation were also normal in these mice (our unpublished observations). The function of NPBWR1 in the SCN remains unclear thus far.

**PERIPHERAL ACTIONS**

Expression of Npb, Npw, and Npbwr1 mRNA in both the adrenal cortex and adrenal medulla has been reported (Andreis et al., 2005; Mazzocchi et al., 2005; Hochol et al., 2007). NPB and NPW were shown to pharmacologically stimulate adrenal glucocorticoid secretion by an ACTH-independent mechanism. It was also reported that NPW stimulates in vitro aldosterone secretion by enhancing the release of medullary catecholamines, which activate beta-adrenoceptors located on zona glomerulosa cells (Hochol et al., 2007).

Bolus intraperitoneal (i.p.) injection of NPB or NPW increased the plasma levels of parathyroid hormone, corticosterone, and testosterone. NPB was also reported to increase the blood concentration of thyroxine, and NPW was shown to increase ACTH and estradiol levels. These findings suggest that NPB and NPW play a role in regulation of the endocrine system (Hochol et al., 2006).

The existence of NPW in rat gastric antral cells was reported. The level of NPW in the stomach was decreased in fasted animals, while it was increased by re-feeding (Mondal et al., 2003), which is consistent with the notion that NPW may act as a suppressant of feeding. However, we did not observe any effects on feeding in mice when NPB or NPW was intravenously administered, suggesting that peripheral NPB/W has limited, if any, significant role in modulating feeding behavior (our unpublished observations).

Neuropeptide B and NPW are reported to inhibit proliferative activity of rat calvarial osteoblast-like (ROB) cells (Ziolkowska
CONCLUSION

Neuropeptide B and NPW are likely to be multi-tasking factors. Both Npb−/− and Npbwr1−/− mice show late-onset obesity and hyperphagia, suggesting that the endogenous NPB-NPBWR1 pathway negatively regulates feeding behavior and positively regulates energy expenditure (Ishii et al., 2003; Kelly et al., 2005). This notion is further supported by several pharmacological studies that showed that i.c.v. NPB/W increased heat production and sympathetic outflow (Yu et al., 2007). However, the biphasic (early orexigenic followed by anorexic) effects of NPB/NPW suggest complex actions of NPB and NPW in the regulation of food intake. orexigenic followed by anorexic) effects of NPB/NPW suggest complex actions of NPB and NPW in the regulation of food intake. Many studies have also shown that the NPB/W system is involved in the modulation of inflammatory pain. Consistent with these results, Npb−/− mice are hypersensitive to inflammatory pain but display no significant difference in chemical or thermal pain. These data together strongly support a physiological role of central NPB in pain regulation, and agonists for NPBWR1 or NPBR2 might be good candidates for analgesic drugs for chronic inflammatory pain.

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