Identification of oxytocin receptor activating chemical components from traditional Japanese medicines

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Identification of oxytocin receptor activating chemical components from traditional Japanese medicines

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Abstract

Oxytocin (Oxt) is known to regulate social communication, stress and body weight. The activation of Oxt receptors (OTR) has clinical potential to abate stress disorders and metabolic syndrome. Kamikihito (KKT) is a traditional Japanese medicine used to treat psychological stress-related disorders. We investigated the effects of KKT, its ingredients and chemical components on Oxt neurons and OTR. C-Fos expression was examined after oral and peripheral administration of KKT in rats. Electrophysiological change of Oxt neurons and Oxt release upon application of KKT were measured in rat brain slice. The direct effect of KKT, its ingredients and its chemical components were examined by cytosolic Ca2⁺ ([Ca2⁺]i) measurement in Oxt neurons and OTR-expressing HEK293 cells. Both intraperitoneal and oral administration of KKT in rats induced c-Fos expression in neurons of the paraventricular nucleus (PVN) including Oxt neurons. Application of KKT induced activation of Oxt neurons and Oxt release. KKT increased [Ca2⁺]i in OTR-expressing HEK293 cells, and failed to activate with OTR antagonist. KKT-induced PVN Oxt neuron activation was also attenuated by OTR antagonist. Seven chemical components (rutin, ursolic acid, (Z)-butylidenephtalide, p-cymene, senkunolide, [6]-shogaol, [8]-shogaol) of three ingredients (Zizyphi Fructus, Angelicae Acutilobae Radix, Zingiberis Rhizoma) from KKT had potential to activate OTR. KKT can directly activate PVN Oxt neurons by interacting with OTR. The interaction of seven chemical components from KKT may contribute to activate OTR. Effect of KKT on Oxt neurons and OTR may contribute to the treatment of Oxt related disorders.

Keywords: Kamikihito, Oxytocin neuron, Oxytocin receptor, Traditional Japanese medicine

1. Introduction

Oxytocin (Oxt) is a neuropeptide known to have important peripheral actions, such as the contraction of the uterus during labor and milk ejection [1]. Oxt is produced in the hypothalamus, mainly in the paraventricular nucleus (PVN) and supraoptic nucleus (SON), and is released within the central nervous system or circulation through the body via the posterior pituitary gland. Recent studies have revealed that in the brain, Oxt is considered to be involved in mood control and psychological stress adaptation [2–4]. Genetic deletion of Oxt in rodents was reported to increase anxiety-like behavior, which was reversed by Oxt treatment [5]. In humans, polymorphism of the Oxt receptor (OTR) has been reported to be associated with stress vulnerability [6], and epigenetic alteration of the OTR gene caused by stress in early life was found to be related to psychological malfunction [7]. We have also shown in the past that Oxt can treat metabolic syndrome and obesity in mice [8–10]. In addition to this, plasma Oxt levels are known to decrease with age and are considered to

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be related to the pathogenesis of sarcopenia and osteoporosis [11,12]. Importantly, application of Oxt is reported to contribute to the recovery of muscle regeneration and osteoporosis as well.

Recently, external application of Oxt to the central nervous system (CNS) through nasal spray was reported to be clinically effective in treating patients with neuropsychological disorders such as autism and obesity [9–11,13–15].

As the external application of Oxt is effective, a drug that induces secretion of endogenous Oxt from Oxt neurons in the hypothalamus would be effective in the treatment of neuropsychological disorders. Although such a drug is very much expected, it is as yet to be discovered.

Kamikihito (KKT), a traditional Japanese (Kampo/herbal) medicine composed of 14 crude drugs, is clinically used in Japan for the treatment of neuropsychological stress disorders including neurosis, amnesia, and insomnia [16]. As Oxt and KKT have common clinical effects, it is possible that KKT has an effect on Oxt neuron or OTR positive cells. Here, we analysed the effect of KKT on Oxt neurons in the hypothalamus and OTR expressing cells. We found that KKT can directly activate PVN Oxt neurons by interacting with OTR. We further elucidated seven chemical components of three ingredients from KKT that induce PVN Oxt neuron activation.

2. Materials and methods

2.1. Animals

8- to 10-week old Male Wistar rats, transgenic rats expressing Oxt-monomeric red fluorescent protein 1 fusion gene (Oxt-mRFP rats) [17] aged 6 weeks, and 10-week old male OTR-Venus mice [18] were used in this study. The animals were kept on a 12-h light/dark cycle and given conventional food (CE-2; Clea, Osaka, Japan) and water ad libitum in individual cages. The lights were turned on at 07:00 am and turned off at 19:00. All experimental procedure and care of animals were carried out according to relevant guidelines and regulations and approved by Fukushima Medical University Institute of Animal Care and Use Committee.

2.2. Reagents

KKT, Hochuekkito (HE), Shikunshito (SKS) and Ninjinto (NJ) were kindly provided by Tsumura & Co (Tokyo, Japan). These Kampo medicines were manufactured under strict scientific and quality control, and approved for ethical clinical use by the Ministry of Health, Labor, and Welfare of Japan. KKT comprises 14 herbal medicines in fixed proportions (see Table 1). KKT, HE, SKS and NJ were prepared as spray-dried powders from hot-water extracts. For in vitro studies, these Kampo medicines were dissolved in an experimental solution and, after centrifugation at 3000 rpm for 5 min, the supernatant was filtrated and used. For animal studies, KKT was orally administered or intraperitoneal (ip) injected to rats, as described below.

<table>
<thead>
<tr>
<th>Crude drugs in KKT, Hochuekkito, Shikunshito, Ninjinto.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>KKT</strong></td>
</tr>
<tr>
<td>Astragali Radix</td>
</tr>
<tr>
<td>Ginseng Radix</td>
</tr>
<tr>
<td>Zizyphi Fructus</td>
</tr>
<tr>
<td>Angelicae Acutiloba Radix</td>
</tr>
<tr>
<td>Zingbeis Rhizoma</td>
</tr>
<tr>
<td>Polygalae Radix</td>
</tr>
<tr>
<td>Atractylosis Lanceae Rhizoma</td>
</tr>
<tr>
<td>Saussweae Radix</td>
</tr>
<tr>
<td>Gardenia Fructus</td>
</tr>
<tr>
<td>Longa Atillus</td>
</tr>
<tr>
<td>Buplewi Radix</td>
</tr>
<tr>
<td>Zizyphi Semen</td>
</tr>
<tr>
<td>Glycyrrhizae Radix</td>
</tr>
<tr>
<td>Poria</td>
</tr>
<tr>
<td>Citii Unshiu Pericarpium</td>
</tr>
<tr>
<td>Cimicifugae Rhizoma</td>
</tr>
</tbody>
</table>

Abbreviations

KKT Kamikihito
Oxt Oxytocin
OTR Oxytocin receptor
PVN paraventricular nucleus
Six chemical components of *Zizyphi Fructus* (ZF), ten chemical components of *Angelicae Acutilobaet Radix* (AR) and seven chemical components of *Zingiberis Rhizoma* (ZR) were provided by Tsumura & Co (Tokyo, Japan) and listed in Table 2. These chemical components were dissolved in DMSO or ethanol and used after over 1:1000–1:10000 dilution.

2.3. Immunostaining of c-Fos after KKT administration

Twenty-two male Wistar rats were used in this experiment. The rats kept in individual cages were habituated for 10 days. On the experiment day, food was removed at 09:00, and oral or ip administration of KKT was performed at 11:00. The KKT for oral administration was dissolved in distilled water, and the dose and concentration was 500 mg/kg/5 mL. KKT for ip administration was dissolved in saline and centrifuged for 20 min at 3000 rpm, and the obtained supernatant was then filtered. The dose and concentration of KKT for ip administration was 500 mg/kg/10 mL. The dose of KKT was decided based on the prior studies on effects of KKT (200–2000 mg/kg oral administration in mice or rats) [19–22].

Two hours after injection, the rats were deeply anesthetized by a mixture of three types of anesthetic agents (5 mL/kg) (composed of z2 adrenaline receptor agonist (medetomidine), GABAA receptor agonist (midazolam), and opioid k receptor agonist (butorphanol)) and, perfused with 4% PFA containing 0.2% picric acid. Then, 40-μm-thick brain sections were cut using a freezing microtome. Sections at 160 μm intervals between −1.3 and −2.1 mm from the bregma were used for c-Fos immunostaining. C-Fos staining was performed similar with the previous article [23]. Briefly, sections were rinsed in phosphate buffered saline (PBS: 0.01 M, pH 7.4) and incubated for 20 min with 0.3% H2O2. Then, sections were incubated in PBS containing 0.3% Triton-X, 2% normal goat serum (Cat# 005-000-121; Jackson Laboratories Inc. PA) and 2% bovine serum albumin (Cat# B2064, Sigma Aldrich, CA) (BSA) and then incubated with rabbit anti-c-Fos polyclonal IgG (Cat# sc-52; Santa Cruz, CA; 1:5000) (BSA) and then incubated with rabbit anti-c-Fos polyclonal IgG (Cat# BA-1000; Vector Laboratories Inc., CA; 1:500), and then with avidin-biotin complex (Cat# PK-6100; ABC kit; Vector Laboratories Inc., CA). Immunoreactions were visualized by incubation in a 0.02% diaminobenzidine solution containing 0.3% nickel ammonium and 0.015% H2O2 for 5 min. The number of c-Fos-positive cells per section was counted from the bregma were used for c-Fos immunostaining. C-Fos staining was performed similar with the previous article [23]. Briefly, sections were rinsed in phosphate buffered saline (PBS: 0.01 M, pH 7.4) and incubated for 20 min with 0.3% H2O2. Then, sections were incubated in PBS containing 0.3% Triton-X, 2% normal goat serum (Cat# 005-000-121; Jackson Laboratories Inc. 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### Table 2. Chemical components of Zizyphi Fructus, Angelicas Acutilobaet Radix and Zingiberis Rhizoma.

<table>
<thead>
<tr>
<th>Ingredients</th>
<th>Chemical components</th>
<th>Concentrations</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zizyphi Fructus</td>
<td>Rutin</td>
<td>0.8-8 μM</td>
<td>[56]</td>
</tr>
<tr>
<td></td>
<td>Ursolic acid</td>
<td>0.05-2 μg/mL</td>
<td>[57,58]</td>
</tr>
<tr>
<td></td>
<td>(0.1-4.4 μM)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Scopoletin</td>
<td>N/A</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Jujubeside A</td>
<td>1-25 μM</td>
<td>[59]</td>
</tr>
<tr>
<td></td>
<td>Jujubeside B</td>
<td>0.1-100 μM</td>
<td>[60]</td>
</tr>
<tr>
<td></td>
<td>Betulonic acid</td>
<td>5 μM</td>
<td>[61]</td>
</tr>
<tr>
<td>Zizyphi Fructus</td>
<td>(r)-butylenepthalalide</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Bergapten</td>
<td>2-50 μM</td>
<td>[63]</td>
</tr>
<tr>
<td>Angelicas Acutilobaet Radix</td>
<td>Umbeliferone</td>
<td>N/A</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Xanthotoxin</td>
<td>25-100 μM</td>
<td>[64]</td>
</tr>
<tr>
<td></td>
<td>p-Cymene</td>
<td>50-250 μM</td>
<td>[65]</td>
</tr>
<tr>
<td></td>
<td>β-Cytoasterol</td>
<td>N/A</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Senkyunolide H</td>
<td>6.25-100 μM</td>
<td>[66]</td>
</tr>
<tr>
<td></td>
<td>Senkyunolide A</td>
<td>13-80 μg/mL</td>
<td>[67]</td>
</tr>
<tr>
<td></td>
<td>(68.3-420 nM)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Falcarinol</td>
<td>5-25 μM</td>
<td>[68]</td>
</tr>
<tr>
<td></td>
<td>Levistolide A</td>
<td>12.5-100 μM</td>
<td>[69]</td>
</tr>
<tr>
<td>Zizyphi Fructus</td>
<td>[6]-gingerol</td>
<td>1-10 μM</td>
<td>[70]</td>
</tr>
<tr>
<td></td>
<td>[8]-gingerol</td>
<td>10 μM</td>
<td>[71]</td>
</tr>
<tr>
<td></td>
<td>[10]-gingerol</td>
<td>5-200 μM</td>
<td>[72]</td>
</tr>
<tr>
<td></td>
<td>Zingerone</td>
<td>5-15 μM</td>
<td>[73]</td>
</tr>
<tr>
<td></td>
<td>[6]-shogaol</td>
<td>1-10 μM</td>
<td>[70]</td>
</tr>
<tr>
<td></td>
<td>[6]-paradol</td>
<td>1-20 mg/mL</td>
<td>[74]</td>
</tr>
<tr>
<td></td>
<td>(3.7-7.45 μM)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>[8]-shogaol</td>
<td>10-50 μM</td>
<td>[75]</td>
</tr>
</tbody>
</table>

Concentrations means the used concentration in vitro experiment in the each reference indicate right.

N/A: Scopoletin and Umbeliferone failed to measure [Ca2+]i due to toxicity by interaction of excitation light.

N/A: There were no previous reports (in vitro) about β-Cytoasterol.
avidin-biotin complex (Cat# PK-6100; ABC kit; Vector Laboratories Inc., CA). Immunoreactions were visualized by incubation in a 0.02% diaminobenzidine solution containing 0.015% H2O2 for 5 min. The number of c-Fos-positive cells, Oxt-positive cells and double positive cells per section were counted for PVN (5-6 sections/rats) and SON (4-5 sections/rats) under light microscope. The average of c-Fos in these sections are treated as individual data.

2.4. Preparation of single PVN neurons and measurement of [Ca2+]i, and subsequent immunocytochemistry in single neurons

Nine male Wistar rats were used in this experiment. Single neurons were prepared from the PVN by using papain, accordingly to the previous report [8]. Cytosolic Ca2+ concentration ([Ca2+]i) was measured by ratiometric fura-2 microfluorometry combined with digital imaging, as previously reported [8]. Briefly, prepared single neurons on a glass-bottom dish were incubated with 2 mM fura-2/AM (Dojin chemical, Kumamoto, Japan) for 30 min at room temperature, placed in a chamber, and superfused with KRB at 1 mL/min at 30 °C. Cells loaded with fluorescent dye were illuminated by alternating 340 and 380 nm, and the resultant fluorescent images were captured. The ratio (F340/F380) images were produced by Aquacosmos version 2.6 (Hamamatsu Photonics, Hamamatsu, Japan). The concentrations of KKT in the present study (5–50 μg/mL) for [Ca2+]i measurements in single neurons were based on the prior studies (0.01–10 μg/mL) [19,20]. Therefore, the concentrations of ZF, AR and ZR were also set at 5–50 μg/mL, accordingly to concentrations used for KKT.

After the [Ca2+]i measurements, single neurons were fixed in 4% PFA overnight. For immunocytochemistry, staining was carried out as previously reported [8]. Briefly, cells were incubated for 20 min with PBS containing 0.3% H2O2. Then, sections were incubated in PBS containing 2% normal goat serum (Cat# 005-000-121; Jackson Laboratories Inc., PA) and 2% BSA (Cat# B2064, Sigma Aldrich, CA) and then incubated with rabbit anti-oxytocin polyclonal IgG (Cat# 20068; Immunostar Inc., WI; 1:5000) for overnight at 4 °C. Subsequently, cells were washed in PBS and incubated with biotinylated goat anti-rabbit polyclonal IgG (Cat# BA-1000; Vector Laboratories Inc., CA; 1:500), and then with avidin-biotin complex (Cat# PK-6100; ABC kit; Vector Laboratories Inc., CA). Immunoreactions were visualized by incubation in a 0.02% diaminobenzidine solution containing 0.015% H2O2 for 5 min.

2.5. Criteria for [Ca2+]i responses

Amplitudes of [Ca2+]i responses to agents were calculated by subtracting the prestimulatory basal [Ca2+]i ratio from the peak [Ca2+]i ratio. When increases in [Ca2+]i took place within 5 min after the addition of agents, and their amplitudes were 0.4 or larger, they were considered responses. For calculation of 0 mg/mL of KKT, ZF, AR and ZR, basal [Ca2+]i ratio from peak [Ca2+]i ratio for 5 min within 10 min from onset of measurement was analysed.

To combine individual [Ca2+]i imaging with immunocytochemical data, at the end of [Ca2+]i imaging, we took photographs of the cell in which [Ca2+]i was recorded and correlated [Ca2+]i data with immunocytochemical results.

2.6. Electrophysiology

Coronal brain slices (300 μm) containing the PVN were prepared from transgenic rats expressing Oxt-monomeric red fluorescent protein (Oxt-mRFP rats). Seven male Oxt-mRFP rats were used in this experiment. All electrophysiological experiments were performed as previously described [25]. KKT, HE, SKS and NJ were prepared as solution of 500 μg/mL. The concentrations of KKT (500 μg/mL) for patch clamp in brain slice were decided based on the concentration of KKT used in single neurons (5–50 μg/mL). The concentration of KKT in brain slice patch clamp was set 10 times higher than concentration of KKT used for [Ca2+]i measurement in single neurons. Thus, the concentration of HE, SKS and NJ were also set at similar concentration (500 μg/mL). OTR inhibitor, H4928 (BACHEM, Bubendorf, Switzerland) was applied at 10−7 M. The concentration of H4928 (10−7 M) was decided based on the previous study [8].

2.7. Measurement of oxytocin secretion from PVN slices

Eighteen male Wistar rats were used in this experiment. The measurement of oxytocin from PVN was performed accordingly to previous reports [8,25]. Briefly, Wistar rats aged eight weeks were deeply anesthetized by a mixture of three types of anesthetic agents, decapitated, and their brains were removed. Three 400 μm thick slices were prepared using a vibrating microtome, and the PVN-containing area was punched out.
These processes were performed in a buffer composed of (in mM) 229 mannitol, 3 KCl, 26 NaHCO₃, 1H₃PO₄, 0.5 CaCl₂, 7 MgCl₂, 7 glucose, and 1 kynurenic acid to pH 7.4 with 95% O₂ and 5% CO₂ mixed gas.

Prepared PVN slices were incubated at 37 °C for 30 min in artificial cerebrospinal fluid (aCSF) composed of (in mM) 126 NaCl, 3 KCl, 26 NaHCO₃, 1H₃PO₄, 2 CaCl₂ and 1 MgSO₄ at pH 7.4, with 95% O₂ and 5% CO₂ mixed gas for recovery. Next, the PVN slices were incubated in aCSF with or without KKT (500 μg/mL) for 1 h. The concentrations of KKT (500 μg/mL) for Oxt secretion from brain slice were decided based on the concentration of KKT used in brain slice patch clamp experiments (500 μg/mL). These incubations were performed in pairs (control and KKT). Secreted Oxt in the supernatant was determined using an oxytocin ELISA kit (Enzo Life Sciences) [10].

2.8. Immunostaining of Oxt in OTR-Venus mice

Three male OTR-Venus mice were deeply anesthetized by a mixture of three types of anesthetic agents, and perfused and 40-μm-thick brain sections were cut using a freezing microtome. PVN containing sections at 160 μm intervals between −0.58 and −1.22 mm from the bregma were used for Oxt immunostaining. The sections were incubated with in PBS containing 0.1% Triton-X, 2% normal goat serum (Cat# 005-000-121; Jackson Laboratories Inc. PA) and 2% BSA (Cat# B2664, Sigma Aldrich, CA), and then incubated with mouse anti-oxytocin monoclonal IgG (Cat# MAB5296; Millipore, CA, 1:1000) [25] for overnight at 4 °C, and incubated with Alexa flour 594-labelled goat anti-mouse IgG (Cat# A11005; Life Technologies, CA, 1:400) for 30 min. Fluorescence images of double positive of oxytocin and OTR were acquired by using a confocal laser-scanning microscope (Fluoview FV10i; Olympus, Tokyo, Japan). Oxt, OTR and double-positive neurons were counted from these confocal images.

2.9. Plasmid construct

The mouse oxytocin receptor (mOTR) gene was amplified by using the primers 5'-ATA GGG AGA CCC AAG CTG GCT AGC GTT TAA ATG GAG GGC ACG CCC GCA GCC -3' and 5'-GCG ATG GCA GTC GCG TCG GTA TCT CGT CTA ATT AAG TTT GTG CCC CAG-3'. The amplified DNA fragment was incorporated into the multicloning site in the pcDNA™3.1(+) plasmid by homogeneous recombination with primers 5'-ATG TCT AAA GGT GAA GAA TTA TTC ACT GGC -3' and 5'-TTA AAC GCT AGC CTT GGG TCT CCC TAT -3'. The RFP-encoding gene, which was kindly provided by Prof. Kazuto Kobayashi of Fukushima Medical University, was amplified with primers 5'-CAG AGG AGC TGT TCTCAA CCA TCC TCG GCA ATG GTG TCT AAG GGC GAA GAG -3' and 5'-CGA GGG GCC GCC ACT GTG CTG GAT ATC TGC CTA ATT AAG TTT GTG CCC CAG-3'. The amplified fragment was incorporated into the C-terminus of the mOTR gene by homogeneous recombination with primers 5'-GCA GAT ATC CAG CAC AGT GGC GGC CGC TCG -3' and 5'-TGC CTA ATT AAG TTT GTG CCC CAG-3'. Homogeneous recombination was performed by using the In-Fusion HD Cloning Kit (Takara Bio, Shiga, Japan).

2.10. Transfection of OTR to HEK293 cells and measurement of [Ca²⁺]i

Cells were cultured in 35 mm a non-coated glass-bottom dish with antibiotics (0.5%/v/v penicillin/streptomycin) in an incubator at 37 °C, 5% CO₂. When the cell confluency reached approximately 50–70% after 48 h incubation, 2.5 μg of OTR plasmid DNA and 5.0 μL of Lipofectamine2000 reagent were mixed and incubated for 10 min according to the manufacturer’s instructions, and applied to the cells.

HEK293 cells, which were transfected with OTR-RFP, were incubated with 2 mM fura-2/AM (Dojin chemical, Kumamoto, Japan) for 30 min at room temperature, placed in a chamber and superfused with KRB containing 10 mM glucose at 1 mL/min at 30 °C. The ratio (F340/F380) images were produced by an Aquacosmos version 2.6. These cells were applied Oxt (Peptide institute Inc. Osaka, Japan) (10⁻⁹–10⁻⁵ M) or KKT (0.5–75 μg/mL). OTR inhibitor, H4928 (10⁻⁷ M), was applied under treatment of Oxt or KKT. Amplitudes of [Ca²⁺]i responses to each agent were calculated by subtracting the prestimulatory basal [Ca²⁺]i ratio from the peak [Ca²⁺]i ratio within 5 min after onset of agent application.

The concentration of Oxt used for the measurement of [Ca²⁺]i responses in cultured HEK cells was decided based on the prior study (10⁻¹¹–10⁻⁶ M) [26]. The concentration of KKT in measurement of [Ca²⁺]i responses in cultured HEK cells was decided based on the results shown in Fig. 2d-f (5–50 μg/mL). The concentration of H4928 (10⁻⁷ M) was decided based on the previous study [8].

The OTR antagonizing experiment using H4928 with Oxt or KKT administrations, the concentrations of Oxt (10⁻⁷ M, which are enough dose to increase [Ca²⁺]i in OTR expressing HEK293 cells) and KKT
studies (Table 2). As shown in Table 2, these provided by Tsumura & Co. and the prior in vitro studies (Table 2). As shown in Table 2, these chemical components were used in ranges of μM. Therefore, the concentrations of chemical components in ZF, AR and ZR are all used in concentration of 10 μM in this study.

After measurement of \([Ca^{2+}]_o\), the cells were fixed in 4% PFA for 2 h. The cells were mounted on a cover glass with a mount medium with DAPI (Vector laboratories, CA). The confocal images were captured by FV10i (Olympus, Tokyo, Japan). The percentage of OTR expression was analysed from 15 areas (200 μm \(\times\) 200 μm/area) from three plates.

2.11. Measurement of plasma concentrations of carvacrol and quercetin

Three male wistar rats (aged 9 weeks) were used. Food was deprived at 19:00 before the day of experiment. On the experiment day, KKT (500 mg/kg/5 mL) was orally administered at 9:00. Blood samples were collected from tail vein into potassium EDTA-treated tubes (BD Microtainer®, blood collection tubes, Becton Dickinson, Tokyo, Japan) before oral administration (0 min), and at 20, 60 and 120 min after oral administration. Blood samples were centrifuged at 10,000 g for 10 min, and plasma samples were stored at \(-80^\circ\)C until analysis.

Plasma concentrations of carvacrol and quercetin were determined using ultra-performance liquid chromatography (UPLC) system (Waters Corp., Milford, MA) with UV and fluorometric detection. Myricetin and thymol were used as internal standards (I.S.) for the quantification of quercetin and carvacrol, respectively. Sample preparation was performed using a Monolithic solid-phase extraction column (MonoSpin® C18, GL Sciences, Tokyo, Japan) according to the manufacturer’s instruction with minor modifications. After filtration thorough a 0.22 μm membrane filter (Millex LG, Millipore), 10 μL of the sample solution was injected to the UPLC system. The mobile phase consisted of 0.2% formic acid (A) and acetonitrile (B), and chromatographic separation was achieved on an ACQUITY BEH Shield RP18 column (particle size 1.7 μm; 2.1 mm \(\times\) 50 mm, Waters) at 40 °C using gradient elution with a flow rate of 0.6 mL/min. The gradient program for carvacrol determination was set as follows: 0–4.0 min, 30–40% B; 4.0–5.0 min, 40–90% B; and 5.0–8.0 min, 30% B. The gradient program for quercetin determination was set as follows: 0–4.0 min, 10–30% B; 4.0–5.0 min, 30–80% B; and 5.0–8.0 min, 10% B. Carvacrol was detected with the excitation wavelength set at 278 nm and the emission at 306 nm. Quercetin was detected at 370 nm. The limits of quantification for carvacrol and quercetin were 6 nM and 3 nM, respectively. The mean intra- and inter-assay coefficients of variance (CV) for quercetin were 9.9% and 2.3%, respectively. The mean intra- and inter-assay CVs for carvacrol were 4.9% and 5.2%, respectively. The accuracies for quercetin and carvacrol were 100 ± 0.6% and 100 ± 6.6%, respectively. Typical chromatograms of quercetin and carvacrol with the respective I.S. were shown in Fig. 9b.

2.12. Statistical analysis

All data are presented as mean ± SEM. Numbers for every experiment (group) are given in the figure legends and refer to the number of every individual mouse in the respective group.

Animals used for c-Fos expression after ip or oral KKT treatment (Fig. 1), were randomly divided into two group. Animal and cells were given to experimenter lacking the information regarding conditions of cells or treatment done to rats in order to make experimenter blind to treatment or condition. All data were not treated as technical replicates, treated as independent values.

For statistical analysis, GraphPad Prism 7.0 (GraphPad Software, San Diego, USA) were used. After ANOVA, post hoc test were only performed if...
F had achieved the significance level of p < 0.05. Differences are considered statistically significant when p-values are lower than 0.05. Assessment of significance was only done for numbers ≥5.

All data were analyzed without any transformation except the data from the effect of chemical components of ZF, AR, ZR on measurements of \([\text{Ca}^{2+}]_i\) in OTR-expressing HEK293 cells (Fig. 8). This is because in order to compare the effect of Oxt itself to each chemical components on OTR, we calculated ratio of average of \([\text{Ca}^{2+}]_i\) treated with Oxt as “1”. Thus, the data shown in Fig. 8 are presented as fold of control mean ratio. The differences in fold of control mean ratio between OTR(−) and OTR(+) were analyzed using unpaired t-test. Concerning with sample size in experiment using cultured cells, sample size was decided based on the previous study [27]. Fujitsuka et al. measured \([\text{Ca}^{2+}]_i\) in 100% GHS-R expressing COS cells at n = 8–12. In our experiment 46% HEK293 cells were transfected OTR. Thus, it was estimated three times sample size were considered to be reliable. Therefore we decided to measure \([\text{Ca}^{2+}]_i\) at least n > 30.

3. Results

3.1. Induction of c-Fos expression by peripheral administration of KKT

Oral administration of KKT (500 mg/kg) in rats induced c-Fos expression localized in the PVN (control: 74.6 ± 12.8/section, KKT: 177.9 ± 13.1/section, Fig. 1a–c). There were also significant differences in the number of c-Fos positive neurons in the SON.
(control: 55.2 ± 6.7/section, KKT: 95.1 ± 4.2/section). The percentage of Oxt neurons among c-Fos positive neurons in the PVN were significantly higher in the KKT group (15.4 ± 2.4%) compared to the water-administered control group (5.0 ± 0.6%) (Fig. 1d–h).

It has been known that oral Kampo administration can change its state of ingredients by the gastrointestinal factors such as gastric digestion or gut flora. In order to examine weather activation of PVN neurons by KKT is induced by the components of KKT or by the gastrointestinal factors mediated end products, we performed i.p injection of KKT to skip the gastrointestinal factors. Similar results were obtained when the same dose of KKT was i.p administered (Fig. 1i–k). The number of c-Fos positive neurons after i.p injection of KKT in the PVN and SON were significantly higher (PVN: 156.8 ± 36.5/section, SON: 159.9 ± 20.9/section) compared with the saline-injected control (PVN: 47.2 ± 20.6/section, SON: 30.2 ± 14.5). The percentage of Oxt neurons among c-Fos-positive neurons in the PVN were significantly higher than the KKT group (12.8 ± 1.8%) compared to the saline-administered control group (1.4 ± 0.4%) (Fig. 1l–p). These results indicate that KKT can affect the PVN with both oral and i.p administration, indicating that gastrointestinal factors such as gastric digestion or gut flora has only small effect if any.

3.2. KKT directly activates PVN Oxt neurons with secretion of Oxt

Application of 500 μg/mL KKT dramatically increased the electrical activity of identified Oxt neurons in the PVN from Oxt-mRFP rats (Fig. 2a). KKT significantly increased the membrane potential from a hyperpolarized (−48.2 ± 2.1 mV) to depolarized (−36.3 ± 2.3 mV) state (n = 5; p < 0.01) (Fig. 2b). In parallel with the increase in the membrane potential, the action potential frequency increased from 0.3 ± 0.2 Hz to 8.4 ± 1.2 Hz (n = 5; p < 0.001) (Fig. 2c). An increase in firing frequency was reversed after washing out KKT for 5 min (0.6 ± 0.5 Hz after washout; n = 5; p < 0.001) (Fig. 2c).

To examine whether KKT had a direct effect on PVN Oxt neurons, we monitored [Ca^{2+}]_i levels in isolated single neurons. After finishing [Ca^{2+}]_i measurements, these neurons were stained immunocytochemically with a specific antiserum for Oxt. Administration of 5 and 50 μg/mL KKT to single PVN neurons increased [Ca^{2+}]_i (Fig. 2d) which neuron was subsequently shown to contain Oxt. The amplitude of [Ca^{2+}]_i (Fig. 2e) and the percentage of KKT responsive neurons (Fig. 2f) increased dose dependently. Seven (38.9%), and eight (44.4%) out of 18 neurons responded to 5 μg/mL and 50 μg/mL of KKT, respectively. Three out of eight (37.5%) KKT (50 μg/mL)-responsive neurons were identified as Oxt neurons.

Data from the electrophysiological experiment and [Ca^{2+}]_i imaging indicate that KKT directly acts on PVN neurons including Oxt neurons, and induces its activation.

Next, in order to see whether Oxt neuron activation by KKT is reflected in the secretion of Oxt, we measured the amount of secreted Oxt from brain slices containing PVN with and without KKT by using a batch incubation technique [25]. As shown in Fig. 2g, 1-h application of KKT (500 μg/mL) significantly increased the amount of secreted Oxt, indicating that activation of Oxt neurons by KKT is indeed reflected in Oxt secretion.

3.3. OTR expression in the PVN Oxt neurons

Taking the results of c-Fos expression after KKT administration (oral and i.p) and measurement of [Ca^{2+}]_i under KKT application into consideration, we hypothesized that KKT may activate Oxt neurons through OTR expressed in the Oxt neurons. Therefore, we examined the colocalization of Oxt and OTR in the PVN. By staining Oxt immunologically in OTR-Venus mice, which express Venus fluorescent proteins in OTR-expressing neurons, abundant co-localization of Oxt and OTR in the neurons of the PVN were identified (Fig. 3). Further analysis showed that, 52.5 ± 4.2% (696/1240 cells) of Oxt neurons expressed OTR.
Fig. 5. The effect of OTR inhibitor on the effect of KKT in HEK293 cells and PVN Oxt neurons. a: Representative change of $[\text{Ca}^{2+}]_i$ image applying with $10^{-7}$M Oxt in HEK293 cells, which was not transfected OTR. b, c: Representative change of $[\text{Ca}^{2+}]_i$ image applied with $10^{-7}$M Oxt (b) or $10^{-7}$M Oxt pre-treatment of $10^{-7}$M H4928 (c) in OTR-transfected HEK293 cells. d: Representative change of $[\text{Ca}^{2+}]_i$ image applying with $50\mu$g/mL KKT in H4928.
3.4. KKT stimulates cells with OTR expression

Since we found that 50% of PVN Oxt neurons express OTR, we examined the effect of KKT on OTR-transfected cultured cells. As shown in Fig. 4a–h, OTR was expressed in HEK293 cells, and we monitored \([Ca^{2+}]_i\) levels upon application of KKT. As shown in Fig. 4i–l, HEK293 cells with OTR showed increase in \([Ca^{2+}]_i\) level upon application of \(10^{-9}\) M to \(10^{-5}\) M Oxt in a dose-dependent manner, indicating that the HEK293 cells with OTR expression that we used in this study were properly functional. We then applied KKT in these cells and measured the change in \([Ca^{2+}]_i\) level. The application of KKT from 0.5 \(\mu\)g/mL to 50 \(\mu\)g/mL induced a dose-dependent increase in \([Ca^{2+}]_i\) level (Fig. 4m–q). Because the application

HEK293 cells, which was not transfected OTR. e, f: Representative change of \([Ca^{2+}]_i\) image applying with 50 \(\mu\)g/mL KKT (e) or 50 \(\mu\)g/mL KKT pre-treatment of \(10^{-7}\)M H4928 (f) in OTR transfected HEK293 cells. g, h: Mean of amplitude of \([Ca^{2+}]_i\) in each cell and treatment. (g; from left bar, n = 91, 104, 49; from left bar, n = 97, 103, 138). **p < 0.01, one-way ANOVA followed by Tukey multiple range test. i, j: Representative recording of membrane potential with application of 500 \(\mu\)g/mL KKT with pre-treatment of \(10^{-7}\)M H4928 in the Oxt neurons in Oxt-mRFP rat. Two from eight neurons exhibited slight action potentials, as shown in i, and six from eight exhibited no action potential, as shown in j. k, l: Mean membrane potential (+SEM) and Mean firing frequency (+SEM) before (control), with H4928 and with both H4928 and KKT treatment (n = 8).
Fig. 7. The effect of Zizyphi Fructus (ZF), Angelicae Acutiloba Radix (AR), Zingiberis Rhizome (ZR) on [Ca$^{2+}$]$_i$ in PVN Oxt neurons. a, d, g: Representative [Ca$^{2+}$]$_i$ image under treatment of 5 µg/mL (5) and 50 µg/mL (50) ZF (a), AR (d) and ZR (g) in Oxt neurons. b, f, i: Mean of amplitude of [Ca$^{2+}$]$_i$ under treatment of 5 µg/mL (5) and 50 µg/mL (50) ZF (b, n = 83), AR (f, n = 44) and ZR (i, n = 15) measured in the PVN neurons. 0 indicates the basal control before treatment of each crude drug. *p < 0.05, one-way ANOVA followed by Tukey’s multiple range test. b, e, h: The incidence of each dose of ZF- (c), AR- (f) and ZR- (i) responsive neurons in the PVN neurons. The criteria of responsive neurons were described in detail in the Methods section. j: Representative [Ca$^{2+}$]$_i$ image under treatment of 50 µg/mL ZF-AR-ZR mixture (MIX) and 50 µg/mL KKT in PVN Oxt neuron. k: Mean of amplitude of [Ca$^{2+}$]$_i$ under treatment of MIX and KKT (n = 16). There were no significant differences. Unpaired t-test.
Fig. 8. The effect of chemical components of Zizyphi Fructus (ZF), Angelicae Acutilobae Radix (AR), Zingiberis Rhizome (ZR) on [Ca\(^{2+}\)]\_i in OTR-expressing HEK293 cells. a: Fold of control ((+Oxt)) mean ratio under treatment of each chemical component of ZF. (from left bar; n = 244, 274, 35, 35, 35, 245, 208, 98, 35, 105, 35, 65, 34, 35, 33). b: Fold of control ((+Oxt)) mean ratio under treatment of each chemical components of AR. (from left bar; n = 208, 103, 35, 68, 70, 65, 65, 67, 65, 68, 69, 102, 33, 34, 35, 31, 35, 32, 68, 60, 99, 102). c: Fold of control ((+Oxt)) mean ratio under treatment of each chemical components of ZR. (from left bar; n = 206, 210, 35, 65, 67, 35, 35, 35, 35, 34, 70, 104, 102, 102, 70, 67, 32, 35). Each ratios were normalized by the ratio for amplitude of [Ca\(^{2+}\)]\_i under treatment of 10\(^{-5}\) M Oxt in each experiment. (\(-\)) indicates HEK293 cells without OTR expression. (+) indicates OTR-expressing HEK293 cells. **p < 0.01, *p < 0.05, unpaired t-test.
Fig. 9. The change of plasma quercetin levels after oral KKT administration and the effects of quercetin on [Ca^{2+}]_i in OTR-expressing HEK293 cells. a: Plasma quercetin concentration after oral KKT (500 mg/kg) administration at 0, 20, 60 and 120 min (n = 3). b: Typical
The bars in figures c-h indicate the term of treatment. **: Mean of amplitude of $[Ca^{2+}]_i$ in the OTR-transfected cells (OTR transfection) compared with single application of Oxt on OTR cells (Fig. 5f). Pretreatment of H4928 significantly reduced $[Ca^{2+}]_i$ level by 50% upon application of Oxt and there were no significant differences in $[Ca^{2+}]_i$ amplitude when compared with a single application of KKT on OTR(-- cells (Fig. 5g). Similar results were obtained from experiments using KKT. In addition, KKT 50 μg/mL slightly increased $[Ca^{2+}]_i$ in the OTR(-- cells (Fig. 5d), and resulted in an apparent increase in $[Ca^{2+}]_i$ in the OTR(+) cells (Fig. 5e). Pre-treatment of H4928 reduced the increase in $[Ca^{2+}]_i$ level compared with the increase in $[Ca^{2+}]_i$ level upon application of a single Oxt in the OTR (+) cells (Fig. 5c). Pretreatment of H4928 significantly reduced $[Ca^{2+}]_i$ level by 50% upon application of Oxt and there were no significant differences in $[Ca^{2+}]_i$ amplitude when compared with single application of Oxt on OTR(-- cells (Fig. 5g). Similar results were obtained from experiments using KKT. In addition, KKT 50 μg/mL slightly increased $[Ca^{2+}]_i$ in the OTR(-- cells (Fig. 5d), and resulted in an apparent increase in $[Ca^{2+}]_i$ in the OTR(+) cells (Fig. 5e). Pre-treatment of H4928 reduced the increase in $[Ca^{2+}]_i$ level compared with a single KKT application (Fig. 5f). Pretreatment of H4928 significantly reduced $[Ca^{2+}]_i$ level by 50% upon application of KKT, and there were no significant differences in $[Ca^{2+}]_i$ amplitude when compared with a single application of KKT on OTR(-- cells (Fig. 5h).

To determine whether this observed effect in OTR(+) cells also occurs in rat PVN Oxt neurons, we used brain slices containing PVN from Oxt-mRFP rats, and measured the change in membrane potential and firing frequency of Oxt neurons upon applying KKT with H4928. As shown in Fig. 5i and j, the PVN Oxt neurons were less activated (Fig. 5i; 25% of examined PVN Oxt neurons), or failed to be activated (Fig. 5j; 75% of examined PVN Oxt neurons) by KKT (500 μg/mL) application in the presence of H4928. There were no changes in firing frequency (Fig. 5k) or membrane potential (Fig. 5i) upon applying KKT on H4928 pre-treated PVN Oxt neurons. These results indicate that KKT activates PVN Oxt neurons directly via OTR.

### 3.6. Clarification of crucial crude drugs that activate Oxt neurons in KKT

The clinically used Japanese Kampo/herbal medicines are based on a systematic combination of crude drugs. KKT is composed of 14 different crude drugs: Astragali Radix, Ginseng Radix, Zizyphi Fructus (ZF), Angelicae Acutiloba Radix (AR), Zingiberis Rhizoma (ZR), Polygalae Radix, Atractylodis Lanceae Rhizoma, Saussurea Radix, Longan Arillus, Bupleuri Radix, Zizyphi Semen, Glycyrrhiza Radix and Poria. There are three other Kampo/herbal medicines that include some of the above crude drug components of KKT (Hochuekkito, Shi-kunshito, Ninjinto). As shown in Table 1, in order of Hochuekkito, Shikunshito and Ninjinto, they gradually lose the crude drug components that are present in KKT. Therefore, we considered that comparing the activating effects of these three Kampo/herbal medicines on PVN Oxt neurons would allow us to pinpoint the crucial crude drugs that activate Oxt neurons. When measuring Oxt neuron activity using a brain slice patch clamp, Hochuekkito activated PVN Oxt neurons similar to that of KKT (Fig. 6a–c). On the other hand, Shikunshito did activate PVN Oxt neurons, although activation took a longer time compared to KKT (Fig. 6d–f). Ninjinto failed to activate the PVN Oxt neurons (Fig. 6g–i). Therefore, by comparing the components of these three Kampo/herbal medicines (Table 1), we narrowed down the candidates to three crude drugs; ZF, AR, and ZR. These three crude drugs are absent in Ninjinto, which failed to activate Oxt neurons.
We then applied these three crude drugs individually and measured the [Ca$$^{2+}$$] level in single PVN Oxt neurons (Fig. 7a–i). The administration of ZF and ZR to single PVN neurons increased [Ca$$^{2+}$$]. The amplitude of [Ca$$^{2+}$$], and the percentage of each responsive neuron were increased followed by ascending concentration of each crude drug. Upon application of 50 µg/mL ZF, 47% PVN neurons responded to ZF, and 33% of ZF-responsive neurons were identified as Oxt neurons. On the other hand, upon application of 50 µg/mL ZR, 60% of PVN neurons responded to ZR, and 89% of ZR-responsive neurons were identified as Oxt neurons. With regards to AR, 5 µg/mL but not 50 µg/mL AR increased [Ca$$^{2+}$$]. There was no dose-dependent response in [Ca$$^{2+}$$], amplitude or incidence of AR-responsive neurons. However, upon application of 50 µg/mL AR, 32% of PVN neurons responded to AR, and 29% of AR-responsive neurons were identified as Oxt neurons.

In order to examine whether the combination of these three crude drugs explains the activation of PVN Oxt neurons by KKT, the amplitude of [Ca$$^{2+}$$], upon application of mixture of three crude drugs (50 µg/mL each) (MIX) and KKT (50 µg/mL), was compared in the PVN neurons. The MIX increased the amplitude of [Ca$$^{2+}$$] to same level as that of 50 µg/mL KKT (Fig. 7j, k). Approximately 44% of the measured PVN neurons responded to MIX and KKT. Among these neurons, over 50% were Oxt-positive neurons. These results indicate that the three crude drugs in MIX are indeed the key factors to activate PVN Oxt neurons, and also that a combination of these three crude drugs is important to induce KKT’s activating effect.

3.7. Identification of chemicals that activate OTR

Since we found KKT activates Oxt neurons by directly interacting with OTR and that the crucial crude drugs for the activation being ZF, AR and ZR, we attempted to identify the chemicals that activate OTR within ZF, AR and ZR. We examined the effects of chemicals composing ZF, AR and ZR by monitoring [Ca$$^{2+}$$] levels in OTR expressed HEK 293 cells. Based on information provided by the manufacturer (Tsumura & Co.) and previous reports from the past, we have analyzed the chemical compounds included in ZF as rutin, ursolic acid, scopoletin, jujuboside A, B, betulinic acid [28–30] and AR as (Z)-butillidenephatalide, bergapten, umbelliferone, xanthotoxin, p-cymene, β-sitosterol, sensyynolide H, senkyunolide A, falcarnidol, levisitolide A [31–35] and ZR as of [6]-gingerol, [8]-gingerol, [10]-gingerol, zingerone, [6]-shogaol, [8]-shogaol, [6]-paradol [36,37]. As shown in Fig 8, 2 chemicals (rutin, ursolic acid) in ZF, 3 chemicals ((Z)-butillidenephatalide, p-cymene, senkyunolide A) in AR and 2 chemicals ([6]-shogaol, [8]-shogaol) in ZR showed increase in [Ca$$^{2+}$$], levels upon its application, indicating combination of these chemicals may contribute to the effects of KKT on Oxt neurons. However, we failed to measure [Ca$$^{2+}$$] under application of scopoletin and umbelliferone due to toxicity induced by interaction of excitation light.

3.8. The change of plasma quercetin and carvacrol after KKT administration

In order to examine the bioactivity of metabolites of rutin and p-cymene [38,39], plasma concentrations of quercetin and carvacrol were measured at 20, 60 and 120 min after oral administration of KKT (500 mg/kg). Carvacrol was not detected in plasma at any time points in all rats. However, quercetin was detected in all time points. As shown in Fig. 9a, plasma concentration of quercetin reached approximately 15 nM at 60 min after oral KKT administration. In addition, the of activation of OTR by quercetin was confirmed using OTR-expressing HEK 293 cells. As expected, both 10$$^{-5}$$ M and 10$$^{-8}$$ M quercetin increased [Ca$$^{2+}$$]i in OTR-expressing cells, with no effect on non-OTR-expressing cells (Fig. 9c–h). 10$$^{-8}$$ M quercetin, which corresponded to plasma quercetin levels at 20 min after oral KKT administration, showed ability to increase [Ca$$^{2+}$$]i by 30% OTR expressing cells compared to increase [Ca$$^{2+}$$]i observed by the application of 10$$^{-7}$$ M Oxt (Fig. 9i).

4. Discussion

The current study demonstrated for the first time that traditional Japanese medicine, KKT, has a potential to activate Oxt neuron through OTR. Compared with the activation effect of Oxt itself, the activation of OTR by KKT was moderate (Fig. 4). However, we have found that KKT have certain potential to activate OTR.

Because approximately half of Oxt neurons in the PVN express OTR, Oxt neurons with OTR may be specifically activated by KKT and secrete Oxt. In support of this idea, all Oxt neurons were not activated by KKT in vivo or in vitro. Oral or ip administration of KKT induced c-Fos in approximately 15% of Oxt neurons, and 37.5% of KKT-responsive neurons were Oxt neurons.

KKT is clinically used in Japan, and is known to be effective in treating psychological stress disorders. Recently, KKT’s effectiveness in the treatment of
memory impairment and Alzheimer’s disease was also reported [19,20,39,40]. These reports indicate that KKT has a therapeutic effect by modulating CNS functions.

Several studies have investigated the pharmacological mechanisms induced by KKT on the CNS. Watari et al. reported that KKT may improve tau phosphorylation and axonal atrophy accompanied by increased protein phosphatase 2A activity in a mouse model of Alzheimer’s disease [20]. Several other reports have indicated that KKT may alter the binding properties of neurotransmitters to receptors [41–43]. However, all these suggested mechanisms are based on the chronic effect of KKT.

Oxt neurons are distributed in the PVN and SON. However, in the present study, c-Fos expressions were found after both oral and ip administration of KKT in the PVN. Oxt neurons in the PVN are considered to be involved in the regulation of psychological stress and mood. This is because PVN Oxt neurons project to areas such as the amygdala, hippocampus, and lateral septum, which are considered to be involved in mood control and stress adaptation [44]. In addition, abundant expressions of OTR were found in these brain areas [45,46]. The activation of OTR induced by KKT shown in the present study raises the possibility that, aside from effects reported by others, the novel effect mediated by OTR shown in this study may also underlie the clinical effect of KKT.

Recently, evidence is being gathered for the clinical use of Oxt. Among the suggested clinical uses, Oxt for the treatment of autism has been gaining most attention. Autism is a neurodevelopmental disorder characterized by restricted interest and impaired social communication. A number of human studies were performed to evaluate the effect of Oxt for the treatment of autism, and most of these studies revealed a positive outcome [13,15]. In addition, Oxt may also be effective in treating obesity due to overeating [14]. These studies suggest that manipulation of the Oxt system in the CNS may be clinically beneficial. However, such treatment is limited to the application of Oxt itself (mainly nasal application), and there have been no reports of pharmaceutical drugs that can reach and activate OTR, thereby inducing the release of endogenous Oxt in the brain. Development of an Oxt analogue or drug that can stimulate Oxt release is still controversial, but such a drug may have strong clinical potential for Oxt-related disorders like autism or obesity. Our study suggests that KKT can directly activate Oxt neurons by acting on the OTR and stimulate Oxt release in the brain. Therefore, KKT may have potential as a new pharmacological agent in targeting Oxt neurons and OTR in the CNS. Another advantage of KKT as an Oxt neuron activator is that it can be orally administered. Because Oxt is neuropeptide, it can only be administered peripherally by injection or nasal administration. The present study showed that oral administration of KKT induced c-Fos expression on PVN Oxt neurons in a similar way to that of ip injection. Therefore, orally administered KKT (at least its critical ingredient that activates Oxt neurons) can overcome the blood–brain barrier (BBB) and reach the CNS to show its effect. Supporting this hypothesis, it was reported that the metabolites of Glycyrrhizae and Atractylodes, which are both components of KKT, can overcome the BBB [47,48].

In the present study, we have also clarified that a combination of the three crude drugs (ZF, AR and ZR) composing KKT is the key factors in activating PVN Oxt neurons. Of these crude drugs, ZR may be the one that specifically contributes to the activation of the Oxt neurons, since 89% of ZR-responsive neurons were Oxt neurons. The chemical components of these crude drugs that were capable of activating OTR were rutin, ursolic acid for ZF and (Z)-butylenephthalide, 3-cymene, senkyunolide A for AR and [6]-shogaol [8], [6]-shogaol for ZR. Because there are 54 compounds identified in only ZF [49], we concentrated in testing part of the components in these crude drugs. Thus, it is possible that other compounds in KKT may also activate OTR.

In order to evaluate from tested compounds as potentials of being agonists of OTR, we calculated response of each components to that of control (Supplemental Fig. 1). Although, the response of KKT to control was approximately 5 times in each experiment of testing each crude drug (red arrows), the response of AR was only with high response with approximately 18 times to that of control. The crude drugs, which responses was over 2.5 times (half response to KKT) were only 3 chemical components, rutin (2.8 times), (Z)-butylenephthalide (3.2 times) and senkyunolide A (3.9 times). These data suggested that AR is the strongest OTR agonist among 3 crude drugs, and a potential OTR agonists were rutin, (Z)-butylidenephthalide and senkyunolide A. Supporting this, only 5 μg/mL AR was able to significantly increase the response of [Ca2+]i as shown in Fig. 7.

On the other hand, although the response to ZR were 3.5 times to control, there were no chemical component, which responded with more than 2.5
times to control in ZR. The data suggested ZR as a crude drug with the strongest synergistic effects. The contribution of synergistic effect is reported in the mechanism of Kampo medicine. For example, in the Kampo medicine Daikenchuto, its chemical component [6]-shogaol and [6]-gingerol are unable to activate voltage-activated cation channels, but become able to activate these voltage-activated cation channels after other chemical component, hydroxy-α-sanshool, block two-pore-domain potassium leak channels (KCNK3 and KCNK9) [50]. Similar to synergistic mechanism reported in components of Daikenchuto, it is possible to consider that such mechanisms are present in chemical components of ZR.

Although further study is required to clarify the underlying mechanism of these three crude drugs, and seven chemical components to induce Oxt neuron activation, our present data indicate the clinical possibility of these crude drugs or chemical components for treating Oxt-related disorders. It is possible that these seven chemical components may fulfill the criteria of bioactive chemical markers, defined as a group of chemo-markers with pharmacological activities comparable to whole KKT. Detail study to determine the bioactive chemical markers for KKT is required for the quality control, correct safety use of KKT in the future [51,52].

Common crude drugs, Astragali Radix and Bupleuri Radix, which are contained in KKT and Hochuekkito are not examined in the present study. It remains the possibility that these two crude drugs are also related to OTR activation. Further study about Astragali Radix and Bupleuri Radix are needed in the future.

Although our present data is mainly based on ex vivo condition and further functional studies are required, this is the first study to show clearly in cellular level that KKT has potential to activate Oxt neurons through OTR. Considering the effects of Oxt to treat so many disorders, the effects of KKT shown in this study has strong potential for future clinical use.

Although the present study focused on the effect of KKT on the CNS, OTR is expressed in various peripheral tissues, such as muscle [12,53], osteoblasts [54] and the pancreas [55]. Oxt contributes to the maintenance and regeneration of muscle cells [12], increases trabecular bone volume [11], and promotes insulin secretion from pancreatic β-cells [23]. Taking our present findings, that KKT can activate the OTR, into consideration, KKT may have a positive effect in peripheral tissues, as well as the CNS. These are important issues that should be investigated in the future.

5. Conclusion

In conclusion, we have identified a novel effect of KKT on the activation of PVN Oxt neurons via the OTR. We suggest that this acute effect of KKT on Oxt neurons and OTR may contribute to the treatment of psychological and neurological disorders. As manipulation of the Oxt system in the CNS may be a therapeutic target for various diseases such as autism, KKT could be introduced as a new potential pharmacological agent.

Author Contributions

Y.M and K.S conceived and designed the project. Y.M performed in vivo experiment and the histological analysis. K.S. performed electrophysiological experiments. S. Y. performed cytosolic Ca2+ measurements. S.H designed the OTR vector for transfection. S. H and T.O prepared the OTR-transfected cultured cells, and provide the critical observation. S.M. performed experiments of pharmacokinetics. K. N. and H. Y. K provided valuable comments and suggestions. Y. U provided the Oxt-mRFP rats as well as valuable comments on the study. Y.M, P. P and K.S wrote the manuscript. All authors discussed the results and contributed to the final manuscript.

Conflicts of interests

The authors declare no competing financial or non-financial interests.

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Appendix.

Fig. S1.

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