Model of Chronic Itch in Aged Mice: Beneficial Effects of Drugs Affecting Descending Modulatory Systems

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Pruritus in the elderly, particularly those cases without skin dryness or other identifiable causes, makes treatment challenging due to the lack of evidence regarding the therapeutic effects of antipruritics. This study proposes an age-related alloknesis mouse model for an evaluation system for such cases, and aimed to investigate the effectiveness and mechanisms of actions of several drugs commonly used as antipruritics in Japan, utilizing this model. Mice 69–80 weeks old were used as aged mice, and the level of mechanical alloknesis was counted as the number of scratching behaviours in response to innocuous stimuli. Bepotastine, norrotropin, pregabalin, baricitinib, and abrocitinib were used as antipruritics, and yohimbine and methysergide as inhibitors of the descending inhibitory pathway. The findings suggest that mechanical alloknesis in aged mice is a suitable animal model for assessing pruritus in the elderly without xerosis, and pregabalin, norrotropin, baricitinib, and abrocitinib may be effective antipruritics in the elderly through activating both the noradrenergic and serotonergic descending inhibitory pathways. These findings may be useful for the selection of antipruritics for pruritus in the elderly without skin lesions or dryness.

Key words: pruritus in the elderly; mechanical alloknesis; descending inhibitory system.

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Pruritus is a common symptom that can affect individuals of all races and ages. However, it is more frequently observed in the elderly (1, 2). Pruritus in the elderly can be defined as idiopathic chronic itching in a person aged over 65 years old (1). Its reported prevalence ranges from approximately 11% to 64% among elderly individuals (3–5). This condition can manifest with or without concomitant visible skin changes (2, 6), and often has a significant impact on the quality of life and sleep (1, 6). Although the pathophysiology remains largely unclear, it is believed to be associated with age-related factors such as (i) epidermal barrier disorders, (ii) changes in the immune system characterized by a shift towards an allergic phenotype (Th2 dominance), or (iii) central or peripheral neurodegenerative diseases (2, 6).

The most common cause of pruritus in the elderly is senile xerosis (6, 7). Studies have reported that dry skin is prevalent in 38–85% of elderly individuals (8–10). However, in addition to cases caused by factors other than xerosis, instances of senile pruritus without primary skin rash and without the presence of xerosis or other identifiable causes have been documented (1, 7). Given the wide array of potential causes and the absence of a definitive cure, treatment of this condition remains a challenge (1, 2, 6).

Various antipruritic treatments have been used for pruritus in the elderly. However, data regarding their therapeutic efficacy are not always complete, as some are limited, and some are effective for pruritus other than that in the elderly (1, 6). Recently, it was reported that aged mice without skin lesions displayed mechanical alloknesis, which is itch hypersensitivity caused by normally innocuous mechanical stimuli, with no increase in spontaneous scratching behaviour (11, 12).

In this study, we utilized this mechanical alloknesis phenomenon as a model for senile pruritus with no rash or xerosis to investigate the effectiveness and mechanisms of the actions of several drugs currently used for pruritus treatment in Japan (7). Here we describe the effects of neuraxial blockers on mechanical alloknesis in aged mice.

SIGNIFICANCE

Many elderly people suffer from itch, and the cause and appropriate treatment are unknown, especially for those without redness or dryness. We aimed to find better medications for such clinical cases and to clarify how these work. We employed the phenomenon called “mechanical alloknesis”, the itch hypersensitivity by innocuous mechanical stimuli, using a mouse model, tested several drugs and found that some of them are effective via activating neural pathways from the brain to the spinal cord, which is called “the descending inhibitory system”. Our findings could improve the quality of life of elderly patients suffering from itch.

MATERIALS AND METHODS

Animals

C57BL/6 mice (Oriental Yeast Co., Ltd., Tokyo, Japan) were bred in-house and used for behavioural analysis. Male mice aged 69
to 80 weeks were designated as aged mice, and those aged 8 to 12 weeks as young. Mice were housed under controlled temperature (23–25°C) and light (12 h of light, 08:00 a.m.–08:00 p.m.) conditions. Food and water were provided ad libitum. The animal experiments were approved by Junendo University Animal Ethics Committee (numbers 2021102, 2022068, and 2023106).

Drugs
Bepotastine besilate, a selective histamine H1-receptor antagonist (FUJIFILM Wako Pure Chemical Corporation, Osaka, Japan); neurotropin, an extract from inflamed rabbit skin following vaccination with vaccinia virus (Nippon Zoki Co. Ltd, Oosaka, Japan); baricitinib, the JAK1/2 inhibitor (MedChem Express, Monmouth Junction, NJ, USA); pregabalin methyl ester, an antagonist of the voltage-dependent calcium channel α2δ subunit (Cayman Chemical, Ann Arbor, MI, USA and FUJIFILM Wako Pure Chemical Corporation), abrocitinib, the JAK1 selective inhibitor (Selleckchem, Houston, TX, USA) and AZ960, the JAK2 selective inhibitor (Selleckchem) were used as antipruritics. Yohimbine hydrochloride, an α1/α2 adrenergic receptors antagonist (Cayman Chemical) and methysergide maleate, a 5-HT1 receptor agonist (5-HT1 -agonist of the voltage-dependent calcium channel α2δ subunit (Cayman Chemical, Anc Arbor, MI, USA and FUJIFILM Wako Pure Chemical Corporation), abrocitinib, the JAK1 selective inhibitor (Selleckchem, Houston, TX, USA) and AZ960, the JAK2 selective inhibitor (Selleckchem) were used as antipruritics.

Preparation and administration of drugs and inhibitors
Antipruritics except for neurotropin were administered orally. Each concentration (bepotastine 3 mg/kg, baricitinib 3 mg/kg, pregabalin 0.3 g/kg, abrocitinib 15 mg/kg, and AZ960 1.58 mg/kg) was based on previous reports and interview forms (13, 14), ensuring equally effective concentrations as indicated by their IC50 values. These drugs were dissolved in dimethyl sulfoxide (DMSO, Sigma-Aldrich, St Louis, MO, USA) and suspended with 0.5 w/v% methylcellulose (FUJIFILM Wako Pure Chemical Corporation) to 5%. As the vehicle control of oral administration, DMSO in methylcellulose and adjusted 5% concentration (bepotastine according to the pharmacokinetic information), or 30 min after the administration of baricitinib, pregabalin, abrocitinib, or AZ960. Neurotropin was dissolved in saline to a concentration of 2/9 neurotropin units (NU)/kg (15). Neurotropin or vehicle control was administered and mechanical alloknesis assays were performed.

Intrathecal administration was performed to inhibit the descending inhibitory pathways with modification of the previous procedure (16). Briefly, a 33-gauge needle (Nipro, Osaka, Japan) and a 25 µl Hamilton microsyringe (Hamilton Company, Reno, NV, USA) were connected using approximately 30–40 cm of polyethylene guide cannula (PE10; Fisher Scientific, Pittsburgh, PA, USA) half-filled with water. The needle tip was dipped into the drug solution, and 5 µl of the solution containing either yohimbine or methysergide was drawn into the cannula by withdrawing the same volume of water using the Hamilton microsyringe. Under sevoflurane anaesthesia, 5 µl of the drug solution or vehicle control saline was injected into the shaved lumbar region, specifically in the groove between the lumbar vertebrae, by expelling water from the microsyringe. Five min after the intrathecal injection, an antipruritic was administered and mechanical alloknesis assays were performed.

Mechanical alloknesis assay: Mechanical stimulation was performed using von Frey filaments ( Biosèb, Chaville, France). Unless stated otherwise, we utilized filaments with bending forces of 0.07 g and 0.16 g. Only data from the 0.16 g filament were presented. The experimental procedure has previously been described (17). Briefly, the rostral back of each mouse was shaved at least 2 days prior to the assay. On the day of the assay each mouse was placed in a new cage to acclimatize for at least 1 h. Subsequently, they received 3 innocuous mechanical stimuli to the rostral back using the aforementioned filaments at approximately equivalent durations and time intervals (about 20 s, at least 5 s or more). This sequence was repeated 10 times, totalling 30 stimulations. Mechanical alloknesis scores were determined as the total number of scratching responses to these stimuli.

Statistical analysis
Data were expressed as mean and standard error of the mean (SEM) and analysed by 2-tailed Student t-test for two group comparisons and by ANOVA for multiple comparisons followed by Dunnett post hoc test with p<0.05 deemed statistically significant. Analysis was performed using GraphPad Prism 9 (GraphPad Software Inc, San Diego, CA, USA).

RESULTS
Characterization of itch in aged mice
We initially assessed the level of mechanical alloknesis and spontaneous itch in aged mice. To quantify mechanical alloknesis, we counted the frequency of hindlimb scratching evoked by von Frey filaments with varying bending forces (0.008 g to 1.0 g), which was recorded as the mechanical alloknesis score (Fig. 1a). Mechanical alloknesis scores elicited by 0.04 g to 0.16 g and 1.0 g von Frey filaments were significantly higher in aged mice than young mice (Fig. 1b). Similar trends were observed at other bending forces tested (0.008 g, 0.02 g, 0.4 g and 0.6 g), albeit differences were not statistically significant (Fig. 1b). In contrast, spontaneous scratching without mechanical stimulation was significantly less in aged mice (Fig. 1c), suggesting it was not spontaneous itch but mechanical alloknesis that was exacerbated with ageing.

We next investigated the skin condition in mice. Increased stratum corneum (SC) hydration (Fig. S1a) and transepithelial water loss (TEWL) (Fig. S1b) were observed in aged mice, suggesting that although ageing was associated with a little decline in skin barrier function, it was not associated with a decline in skin moisture content. The locomotor activity of aged mice was significantly less than that of young mice (Fig. S2).

Evaluation of the effectiveness of antipruritic drugs for pruritus in the elderly
We next examined the therapeutic effect on mechanical alloknesis in aged mice using various antipruritics:
bepotastine, neurotropin, pregabalin, and baricitinib (Fig. 2). After appropriate time intervals for each agent, the mice were subjected to mechanical alloknesis assay (Fig. 2a). Pregabalin, neurotropin, and baricitinib significantly suppressed mechanical alloknesis in aged mice (Fig. 2c–e), whereas bepotastine did not (Fig. 2b). Since the JAK1/2 inhibitor baricitinib suppressed mechanical alloknesis, we investigated the therapeutic effects of JAK1 selective inhibitor abrocitinib and JAK2 selective inhibitor AZ960 in aged mice by oral injection (Fig. 3a). Abrocitinib but not AZ960 significantly suppressed mechanical alloknesis (Fig. 3b).

**Involvement of the descending inhibitory system in age-related mechanical alloknesis**

It was reported that pregabalin and neurotropin exert analgesic effects via the descending inhibitory system (18, 19), which includes noradrenergic and serotonergic pathways. We hypothesized that the antipruritics which showed therapeutic effects in this assay (Figs 2, 3) inhibit mechanical alloknesis by activating these descending inhibitory pathways. To demonstrate this, we investigated the inhibitory effect of noradrenergic and serotonergic descending inhibitory pathways on the anti-mechanical alloknesis activity of these antipruritics on age-related mechanical alloknesis by administering either yohimbine, an α2-adrenergic receptor antagonist, or methysergide, a mixed 5-HT receptor agonist/5-HT2 and 5-HT3 receptor antagonists. Yohimbine or methysergide was administered intrathecally to aged mice 5 min before administration of the antipruritic. After the appropriate time interval for each antipruritic, mechanical alloknesis assays were performed (Fig. 4a). We found yohimbine (Fig. 4b–e) and methysergide (Fig. 4f–i), respectively, inhibited the anti-mechanical alloknesis activity of pregabalin, neurotropin, baricitinib, and abrocitinib, whereas pre-treatment by vehicle solution with antipruritics significantly inhibited mechanical alloknesis in aged mice. Furthermore, when yohimbine or methysergide was administered intrathecally to aged mice in a single administration, no significant difference was observed compared with the vehicle-administered group, whereas a slight increase in mechanical alloknesis score was observed (Fig. 5).
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Fig. 2. Effects of various antipruritic drugs on mechanical alloknesis in aged mice. (a) Schematic procedure for assessing the efficacies of antipruritics on mechanical alloknesis in aged mice. Bepotastine, pregabalin, and baricitinib were administered orally, while neurotropin was delivered via intraperitoneal injection in mice. (b) to (d) Effects of bepotastine (b, n = 15 for vehicle control, 14 for bepotastine), pregabalin (c, n = 16, 16), neurotropin (d, n = 8, 8) and baricitinib (e, n = 8, 8) on mechanical alloknesis scores. Each dot represents an individual value, and the horizontal line in the middle of scattergrams indicates the mean value. Error bars indicate SEM. p.o.: per os, i.p.: intra-abdominal, *p < 0.05, **p < 0.01, ns: not significant (2-tailed t-test).

Fig. 3. Effects of JAK1 and JAK2 selective inhibitors on mechanical alloknesis in aged mice. (a) Schematic procedure for the mechanical alloknesis assay following 2 oral JAK inhibitors: JAK1 selective inhibitor (abrocitinib) and JAK2 selective inhibitor (AZ960) in aged mice. (b) Effects of abrocitinib (n = 7) and AZ960 (n = 7) on mechanical alloknesis in aged mice. DMSO was used as a vehicle control (n = 13). Each dot represents an individual value, and the horizontal line in the middle of scattergrams indicates the mean value. Error bars indicate SEM. **p < 0.01, ns: not significant (Dunnett’s multiple comparison test).
DISCUSSION

In the present study, we re-characterized two types of itch and skin condition of aged mice, and we clarified several antipruritics that are thought to be effective against pruritus in the elderly, and their mechanism of action, at least in part, using the age-related mechanical alloknesis model.

There have been several reports regarding itch in aged mice (11, 12). Similar to these reports our study demonstrated that mechanical alloknesis is more frequent in aged mice than in young mice (Fig. 1b). Conversely, spontaneous scratching behaviour under non-mechanical stimulus was significantly lower in aged mice than in young mice (Fig. 1c). This result contrasts with the fin-
In this study, we selected bepotastine, pregabalin, neurotropin, and baricitinib as test drugs to evaluate their effectiveness for pruritus in the elderly using the age-related mechanical alloknesis model. These drugs are commonly used to treat various types of pruritus in Japan (7). Among them, bepotastine, a selective H<sub>1</sub>-receptor antagonist and one of the most well-known antipruritics, had no effect on the mechanical alloknesis in aged mice (Fig. 2b). It is known that dermal pruritus with no skin lesions is generally less responsive to antihistamines (7), which may explain these results.

In terms of pharmacological behaviour, we also found that neurotropin and pregabalin improved mechanical alloknesis in aged mice (Fig. 2c, d). Pregabalin is a drug officially indicated for neuropathic pain and pain associated with fibromyalgia, but not for itching in Japan (7). This agent is thought to act on the voltage-dependent calcium channels in the central and peripheral nervous system, inhibiting the secretion of neurotransmitters such as glutamate, substance P, and CGRP in a state of neural hyperexcitability. This, in turn, raises the threshold for itch (23). Neurotropin is a drug commonly used in Japan and China. Its injectable form is officially prescribed to treat conditions like low back pain, cervicobrachial syndrome, symptomatic neuralgia, and itching associated with skin diseases (such as eczema, dermatitis, urticaria). It is also employed for allergic rhinitis and to manage after-effect SMON symptoms like cold sensations, abnormal sensations, and pain (24). It was reported that neurotropin suppresses itch-related behaviour in NC/Nga mice with atopic dermatitis (AD)-like symptoms (15). Moreover, clinical studies have reported that the administration of pregabalin and neurotropin successfully alleviated itching in conditions like dialysis-induced pruritus and neuropathic itch, such as post-herpetic pruritus (25–27). Therefore, our data strongly suggest that these drugs are promising candidates to treat pruritus in the elderly.

Notably, baricitinib, an orally administered selective JAK1/2 inhibitor, improved mechanical alloknesis in aged mice (Fig. 2e). Further analysis revealed that the JAK1-specific inhibitor abrocitinib, but not AZ960, the JAK2-specific inhibitor, also had a significant effect on age-related mechanical alloknesis (Fig. 3b). Baricitinib and abrocitinib are approved in many countries for the treatment of moderate-to-severe AD in adults (28, 29), but not for dermal pruritus without skin lesions, at least in Japan (7). It is believed that JAK inhibitors are effective for AD primarily by suppressing Th2 inflammation via inhibiting JAK-STAT signalling (30). However, it was reported that JAK1 is enriched with pruriceptive neurons, and the specific deletion of JAK1 in sensory neurons reduces itch in AD models (31). This indicates that JAK1 signalling in sensory nerves plays a pivotal role in itch transduction. This report supports our findings that the anti-mechanical alloknesis activity of baricitinib...
and abrocitinib are suppressed by inhibiting descending inhibitory pathways, which are among the most well-known neural pathways (Fig. 4d, e, h, and i).

Not only baricitinib and abrocitinib, but also pregabalin and neurotropin demonstrated a reduction in their therapeutic effects on mechanical alloknesis by inhibiting the descending inhibitory system (Fig. 4b, c, f, and g). The descending inhibitory system can be categorized into 2 main pathways: the noradrenergic pathway, which involves the projection of noradrenaline from the locus coeruleus to the dorsal horn of the spinal cord, and the serotonergic pathway, which projects serotonin from the hypothalamus through the raphe nuclei to the dorsal horn of the spinal cord (32). Within these pathways, there are various receptor types that play a role in their function. Adrenaline receptors (which respond to noradrenaline) are broadly classified into 3 types (α1, α2 and β), while 5-HT receptors (which respond to serotonin) are classified into 7 types (HT1 to HT7), each with subtypes. It was reported that α1 and α2 receptors (33) and 5-HT1A, 5-HT1B, 5-HT1D, 5-HT2A, 5-HT3, and 5-HT4 receptors (34) are expressed in the spinal cord and contribute to the operation of the descending inhibitory system. Historically, the noradrenergic and serotonergic pathways were known to alleviate thermal, mechanical, and neuropathic pain in animal models (32). However, recent research has indicated their involvement in acute (35) and chronic itch (36).

A study by Koga et al. (37) suggested that descending noradrenergic signalling inhibits acute and chronic itch, supporting our findings that yohimbine, an α1/α2 receptor antagonist, suppressed the anti-alloknesis activity of antipruritics. Regarding 5-HT receptors, we found that the anti-mechanical alloknesis effect of our antipruritic candidates was suppressed by methysergide, a chemical drug with both 5-HT1 receptor agonist and nonselective 5-HT1-5-HT2 receptors (38, 39). It has also been reported that depletion of spinal 5-HT1A-expressing neurons has been shown to reduce itch, and intraperitoneal injection of 5-HT precursor facilitates itch, suggesting that descending 5-HT1A spinal neurons work as a facilitative pathway rather than an inhibitory pathway (40). This activation of the 5HT1A-mediated itch-facilitative pathway may be responsible for the slight increase in scratching behaviour observed in aged mice when methysergide was administered alone (Fig. 5). However, as no significant difference was observed in this case, there is a strong possibility that methysergide also acts on the descending inhibitory system mediated by a 5HT1 receptor (or possibly a 5HT7 receptor whose expression has not been confirmed) in the spinal cord, which may be the action points of these antipruritics. The fact that 2 descending inhibitory pathways may be involved in the mechanism of the anti-mechanical alloknesis activity of antipruritics strongly suggests that age-related mechanical alloknesis may be primarily influenced by the modulation of the central nervous system rather than epidermal barrier disorders or changes in the immune system.

In conclusion, the findings from this study suggest that mechanical alloknesis in elderly mice serves as a suitable animal model for evaluating pruritus in the elderly, particularly in cases without initial rashes or xerosis. Pregabalim, neurotropin, baricitinib, and abrocitinib appear to have potential as effective antipruritic agents in the elderly, as they may activate both the noradrenergic and serotonergic descending inhibitory pathways, either directly or indirectly (Fig. 6). These results provide valuable insights that can inform the selection and development of antipruritics for managing pruritus in the elderly population, particularly when skin lesions and dryness are not present.

Fig. 6. A model of the descending inhibitory system and the possible points of action for anti-mechanical alloknesis. When a mechanical stimulus is received at the periphery, primary afferents are excited. This excitement reaches the cerebral cortex where it is perceived as itch, through the dorsal horn of the spinal cord, secondary afferents, and thalamus (itch-transaction pathway: red line). The descending inhibitory system is a system that descends from the periaqueductal gray (PAG) to the dorsal horn of the spinal cord, and inhibits synaptic transmission between primary and secondary afferents in the itch-transaction pathway. This system consists of noradrenaline (green line) and serotonin (blue line) pathways, which release noradrenaline or serotonin in the spinal cord via the locus coeruleus and raphe magnus, respectively. As yohimbine and methysergide inhibited the anti-mechanical alloknesis effect of pregabalim, neurotropin, baricitinib, and abrocitinib at the spinal cord, these antipruritics may suppress mechanical alloknesis by activating both of the descending inhibitory pathways directly, or indirectly by activating their upstream pathway (such as hypothalamus or thalamus, possible points of action: orange lines).
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