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The cell type-specific ER membrane protein UGS148 is not essential in mice

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Abstract

Genomes of higher eukaryotes encode many uncharacterized proteins, and the functions of these proteins cannot be predicted from the primary sequences due to a lack of conserved functional domains. In this study, we focused on a poorly characterized protein UGS148 that is highly expressed in a specialized cell type called tanycytes that line the ventral wall of the third ventricle in the hypothalamus. Immunostaining of UGS148 revealed the fine morphology of tanycytes with highly branched apical ER membranes. Immunoprecipitation revealed that UGS148 associated with mitochondrial ATPase at least *in vitro*, and ER and mitochondrial signals occasionally overlapped in tanycytes. Mutant mice lacking UGS148 did not exhibit overt phenotypes, suggesting that UGS148 was not essential in mice reared under normal laboratory conditions. We also found that RNA probes that were predicted to uniquely detect UGS148 mRNA cross-reacted with uncharacterized RNAs, highlighting the importance of experimental validation of the specificity of probes during the hybridization-based study of RNA localization.

Introduction

Molecular functions of proteins can be generally predicted from combinations of known functional domains, of which primary sequences are highly conserved between species. However, functional annotations of conserved domains remain far from complete, and there are hundreds of "DUFs", domain of unknown functions, of which molecular function has not been demonstrated experimentally (El-Gebali et al. 2019). As such, even 20 years after the first draft sequences of the human and mouse genomes were published, nearly a thousand proteins remain functionally uncharacterized (Frankish et al. 2021). In addition, the development of ribosome profiling technology has revealed that many of the transcripts that were initially described as long noncoding RNAs (lncRNAs) indeed associate with ribosomes (Ingolia 2014). These transcripts are believed to be translated into peptides, which are shorter than the arbitrarily set thresholds for the lack of protein coding capacity of lncRNAs. A series of studies using mutant animals have revealed that novel proteins or peptides encoded by "lncRNAs" play important physiological roles, including the regulation of transcription via inhibition of P-TEF-b (Hanyu-Nakamura et al. 2008), activation of G protein-coupled receptors (Chng et al. 2013; Pauli et al. 2014), proteolytic processing of transcription factors (Kondo et al. 2007), modulation of Ca²⁺ pump activity (Anderson et al. 2015; Nelson et al. 2016), regulation of mTORC1 (Matsumoto et al. 2017), and regulation of mRNA de-capping complexes (D'Lima et al. 2017). These newly discovered peptides lack known conserved domains and thus have long been overlooked by functional predictions based on primary sequence analyses. Studies have also shown that a group of novel proteins, entirely consisting of intrinsically disordered regions, possesses outstanding molecular properties to inhibit the formation of disease-related molecular aggregates or protect DNA from irradiation (Hashimoto et al. 2016; Tsuboyama et al. 2020). Interestingly, at least two of these "superdisordered" proteins maintain their molecular activity even after the random shuffling of their primary amino acid sequences (Tsuboyama et al. 2020), suggesting that the function of these superdisordered proteins is dependent on the composition of the amino acids rather than their primary sequences. The functional independence of the amino acid sequences makes it difficult to predict the function of these proteins by conventional bioinformatics analyses. Taken together, these unpredictable proteins constitute the "dark matter" of the genome, which may regulate important processes in higher organisms.

Tanycytes are specialized cell types that align the ventral region of the third ventricle around the medial eminence and have a long process extending from the ventricular regions to the deep hypothalamus (reviewed in Langlet 2014; Rodriguez *et al.* 2019). Tanycytes express molecular markers specific to neural precursor cells (reviewed in Recabal *et al.* 2017), and have been shown to contribute to the adult neurogenesis in the hypothalamic regions (Lee *et al.* 2012). Tanycytes are also proposed to be involved in the transport of small molecules and peptides from the cerebrospinal fluid to neurons in the arcuate nucleus and vice versa (reviewed in Bolborea & Dale 2013; Langlet 2014; Rodriguez *et al.* 2019).

In this study, we performed functional analyses of the poorly characterized protein UGS148 (US patent US20020155119A1), which was previously reported as a novel ER protein (Bennetts *et al.* 2006) or a specific molecular marker for tanycytes (Ma *et al.* 2015). Immunoprecipitation analyses revealed that UGS148 associated with mitochondrial ATP synthetase at least *in vitro*, suggesting that UGS148 might be involved in the interaction of the ER and the mitochondrial membrane in this specialized cell type. UGS148 knockout (KO) mice were viable and exhibited no overt phenotype, suggesting that UGS148 was not required for the mice, at least under normal laboratory conditions.

Results

Incidental identification of UGS148 while identifying novel transcripts that exhibit unique subcellular distribution

Transcripts longer than 200 nucleotides that lack protein coding capacity are called long noncoding RNAs (lncRNAs), the threshold of which was arbitrarily established during early studies conducted in the 2000s (reviewed in Ulitsky 2016). To identify novel functional IncRNAs, we initially investigated the subcellular distribution of transcripts of uncharacterized genes by in situ hybridization, especially focusing on genes with the suffix -Rik, originally annotated as non-protein-coding transcripts, in the FANTOM project (Carninci et al. 2005). We expected that if the transcripts are highly enriched in the nucleus, they might be good candidates for a novel functional lncRNA because essentially no translation occurs in the nucleus (Dahlberg & Lund 2012). We used cultured hippocampal neurons for this study because the nervous system generally expresses a broader range of lncRNAs than other tissues and cell types (reviewed in Ng et al. 2013). We arbitrarily selected 24 genes with the suffix "Rik" from the list of highly expressed genes identified by microarray analyses of cultured hippocampal neurons (Supplemental Table S1), and intense signals were confirmed for 16 of them by fluorescent in situ hybridization using cultured hippocampal neurons (Fig. 1A). Among them, transcripts of 2 genes, 2610002J23Rik and 6330403K07Rik, were highly enriched in the nucleus (Fig. 1A). We focused on 6330403K07Rik in subsequent studies because of its intense and specific expression in the nervous system.

To investigate the expression pattern of 6330403K07Rik in the adult brain by *in situ* hybridization, we initially used a full-length cDNA clone, BC069976, for preparation of the probe (UGS148 #1, Fig. 1B). The clone did not contain any homologous sequence in the genome when searched by NCBI BLAST and RepeatMasker, except for a short 73-bp fragment 72% identical to a simple repeat, (TCTCCA)_n (Fig. 1B, C). *In situ* hybridization using a stringent hybridization condition revealed that 6330403K07Rik was broadly expressed throughout the entire brain regions (probe #1, Fig. 1D), which was consistent with the data in the Allen mouse brain atlas (Fig. 1E), a public genome-wide database for the expression of genes in the mouse brain (Sunkin *et al.* 2013). However, during the course of this study, a paper reported that 6330403K07Rik indeed encodes putative ER protein UGS148 that is highly enriched in tanycytes, a group of ependymal cells that line ventral regions of the 3rd ventricle in the hypothalamus (Ma *et al.* 2015). Thus, we speculated that the probe used in our initial

screening and the Allen Brain Atlas cross-reacted with additional transcripts via the region homologous to (TCTCCA)_n, considering that protein coding mRNAs usually localize in the cytoplasm. To test this hypothesis, we prepared probes #2 and #3 that do and do not contain the repeat-homologous region (Fig. 1B) and performed Northern blot analyses using brain RNAs (Fig. 1F). As expected, the probe #1, which we used in the in situ hybridization experiment, detected multiple transcripts on the Northern blot (arrowheads in Fig. 1F) in addition to the transcript at the expected size (asterisk in Fig. 1F). To further confirm the crossreactivity of the probes, we performed Northern blot analyses using brain RNAs derived from mutant mice lacking the 6330403K07Rik/UGS148 transcripts, which were generated during this study. Extra bands were also detected in mice lacking the 6330403K07Rik/UGS148 transcript, suggesting that #1 and the probe used in the Allen Brain Atlas were not specific to 6330403K07Rik/UGS148. Conversely, probe #3, which was designed against the 3' untranslated region of 6330403K07Rik/UGS148, detected a single band of the expected size, which was not detected in the RNA samples derived from the mutant mouse lacking this gene (Fig. 1F). Unexpectedly, these extra bands were not detected with probe #2, which targeted the region homologous to the repeat (TCTCCA)_n (Fig. 1F). These results suggested that probe #1 cross-reacted with unknown transcripts via 5' regions of these genes containing the ORF, even though the sequences were unique in the genome, as shown by conventional bioinformatic searches such as BLAST (Altschul et al. 1990), BLAT (Kent 2002), or RepeatMasker (Smit 1996). We also confirmed the specificity of the probes by *in situ* hybridization. Probe #3 mainly detected the signals enriched in the tanycytes, as described in a previous study (Ma et al. 2015), which were not observed in the brains of the mutant mice (Fig. 1D). These results suggested that RNA probes predicted to be unique by conventional sequence analyses possibly detected other transcripts when used in hybridization-based assays. The mRNAs of UGS148 detected with probe #3 were mainly localized in the cytoplasm, as expected for protein-coding genes (Fig. 1H). To detect the protein product of UGS148, we raised a monoclonal antibody that specifically detected UGS148 at the predicted size of 13 kDa (Fig. 1G). Both UGS148 mRNA and protein were specifically expressed in the brain tissue but not in other tissues in our investigation (Fig. 1I, J).

UGS148 is expressed in tanycytes and neurons in the hypothalamus

While a previous study reported highly enriched expression of UGS148 in tanycytes (Ma *et al.* 2015), the molecular properties and physiological functions of this protein remain largely unknown. To obtain further insights into the functions of this protein without known

functional domains, we first examined the expression pattern of UGS148 in the hypothalamus in detail. As previously reported, UGS148 was highly expressed in tanycytes located at the ventral region of the 3rd ventricle in the hypothalamus (Tn in Fig. 2A). Distinct signals were also observed in neurons with a large cell body in the lateral hypothalamic nuclei (LH in Fig. 2A). There are at least two groups of tanycytes, α - and β -tanycytes, which are located in dorsal and ventral regions of the 3rd ventricle in the hypothalamus (Langlet 2014; Rodriguez et al. 2019). Each group of tanycytes is further categorized into 2 cell types: dorsally located type 1 and ventrally located type 2, and four types of tanycytes, $\alpha 1$ -, $\alpha 2$ -, $\beta 1$ -, and $\beta 2$ -tanycytes, have been described and are aligned from dorsal to ventral in order (Langlet 2014; Rodriguez et al. 2019). UGS148 was highly expressed in all of these tanycytes, except for β 2-tanycytes located in the ventral medial eminence that weakly expressed UGS148 (Fig. 2B). UGS148-expressing tanycytes were distinct from ependymal cells identified by the expression of Gfap (glial fibrillary acidic protein), which were dorsally located along the 3rd ventricle (SA, Fig. 2B, D). Superresolution confocal microscopy of UGS148 revealed highly branched mesh-like apical branches (AB), elongated oval cell bodies (CB), and basal processes (BP) extending perpendicular to the ventricular surface (Fig. 2C), reminiscent of the highly developed ER membrane network found in this cell type (Ma et al. 2015; Rodriguez et al. 2019). In the lateral hypothalamic region that contains UGS148-positive neurons, there are two types of neurons with large cell bodies, hypocretin/orexin-expressing neurons and melanin-concentrating hormone-expressing neurons, which can be identified by the expression of Hert and Pmch, respectively (reviewed in Adamantidis & de Lecea 2008). Simultaneous detection of the mRNA expression of these marker genes and UGS148 revealed expression of UGS148 in all Hcrt-positive hypocretin/orexin-expressing neurons but not in the Pmch neurons in the lateral hypothalamus (Fig. 2E).

To further confirm the expression of UGS148 in tanycytes at the single-cell level, we reanalyzed RNA sequencing data prepared from hypothalamic cells (Chen *et al.* 2017). Hierarchical clustering analyses revealed that UGS148 clustered into genes that are known to be pan-tanycyte marker genes (Fig. 2F), suggesting that UGS148 is ubiquitously expressed in tanycytes at least at the mRNA level.

UGS148 is an ER membrane protein with an intrinsically disordered domain that protrudes into the cytoplasm

To study the subcellular localization of UGS148, we prepared HeLa cells that overexpressed UGS148 and compared the signals with subcellular organelle markers. Analyses of the primary amino acid sequences using TMHMM (Krogh et al. 2001) revealed that UGS148 is a single transmembrane protein consisting of an N-terminal disordered region that contains alternate stretches of basic and acidic amino acids, with a short C-terminal region consisting of the tripeptide QLA (Fig. 3A, B). ClustalW analyses (Thompson et al. 1994) revealed that the transmembrane domain is highly conserved among mice, rats, and humans, while the N-terminal disordered region was rather variable (Fig. 3A). As previously reported (Bennetts et al. 2006; Ma et al. 2015), UGS148 was localized in the cytoplasm when overexpressed, forming a mesh-like structure, which resembled the distribution of ER membranes (Fig. 3C). ER localization was also confirmed based on the co-localization of UGS148 with GFP-calreticulin, a marker for the ER membrane (Michalak et al. 1999). The two signals completely overlapped (Fig. 3C, D), suggesting that UGS148 was localized to the ER membrane, as previously reported (Bennetts et al. 2006; Ma et al. 2015). The overall morphology of the ER membrane network was not affected by the overexpression of UGS148 in HeLa cells, and a similar mesh-like pattern was observed in both the UGS148-expressing and nonexpressing cells (Fig. 3E). We also performed wndchrm analysis (Shamir et al. 2008), a machine-learning-based image recognition analysis, and found that the ER membrane pattern of the parental HeLa cells was indistinguishable from that of the HeLa cells that ectopically expressed UGS148 (Fig. 3F). The ER localization of endogenous UGS148 in tanycytes was confirmed by simultaneous detection of UGS148 and another authentic ER membrane marker, Hspa5/BiP (Munro & Pelham 1987), in the adult brain, although the signals were obscure due to the harsh antigen retrieval procedure required for the detection of Hspa5/BiP signals in mouse tissue samples (Fig. 3G, H).

To further investigate the topological organization of UGS148 in the ER membrane, we introduced N-terminal or C-terminal tagged UGS148 into HeLa cells and detected the signals after permeabilization using digitonin, which can permeabilize plasma membranes while leaving ER membranes intact (Plutner *et al.* 1992; Wilson *et al.* 1995). An N-terminal tagged FLAG epitope was detected in the digitonin-treated cells, whereas a C-terminal tagged FLAG epitope could not be detected in the cells (Fig. 3I). However, both epitopes were detected in the cells treated with Triton X-100 (Fig. 3I), which could permeabilize both plasma and ER membranes. These results suggested that the N-terminal disordered region of UGS148 faced the cytoplasmic regions, whereas the short tripeptide faced the ER lumen (Fig. 3J).

UGS148 associates with the mitochondrial ATPase subunit, at least in the cell lysate

To gain further insight into the molecular function of UGS148, we attempted to identify molecules that associate with UGS148 in the cells. We established HeLa cells that stably expressed N-terminal FLAG-tagged UGS148 and performed immunoprecipitation using cell lysates prepared from the stable cell line. Discrete bands were specifically immunoprecipitated with N-terminal FLAG-tagged UGS148 (Fig. 4A) and were identified as subunits of mitochondrial ATPase by subsequent MALDI-TOF analyses. To confirm that UGS148 formed a complex with mitochondrial ATPase in the lysate, we performed Western blot analyses of the immunoprecipitated complex using specific antibodies against the α -, β -, and γ -subunits of ATPase. All three subunits were specifically immunoprecipitated with N-terminal FLAG UGS148 (Fig. 4B), suggesting that UGS148 could associate with the ATPase complex, at least in the cell lysate. Interestingly, multiple bands were detected for FLAG-UGS148 (arrowheads in the FLAG panel in Fig. 4B), suggesting that FLAG-UGS148 formed SDS-resistant multimers during immunoprecipitation.

Because UGS148 was localized in the ER membrane, the association with mitochondrial ATPases that are localized in the inner membrane of mitochondria was unexpected. Thus, we thus simultaneously detected UGS148 and the α -subunit of the ATPase subunit in HeLa cells that stably expressing UGS148, and the two signals basically did not overlap, as expected (Fig. 4C, D). We also examined the localization of endogenous UGS148 and the α -subunit of ATPase in tanycytes. Although the UGS148 signals did not coincide with the signals of the α -subunit of ATPase in most cases, we occasionally observed UGS148 signals aligning along the mitochondrial signals (Fig. 4E, F).

UGS148 is not essential for mice reared under laboratory conditions

Finally, we examined the physiological function of UGS148 by creating KO mice that lack the expression of this molecule (Fig. 5A-C). Although the number of adult (8 weeks) KO mice obtained by crossing the heterozygous mice was slightly lower than the expected number, the difference was not significant (male: P = 0.088, female: P = 0.055, chi-square test) (Fig. 5D). Both female and male adult KO mice exhibited similar body weights compared with their wild-type (WT) littermates (Fig. 5E). No overt morphological abnormality of the tanycytecontaining hypothalamus regions was observed on hematoxylin-eosin-stained paraffin sections (Fig. 5F). We then examined the morphology of tanycytes using the ER membrane marker Hspa5/BiP, and mesh-like structures of the AB and BP in KO mice were comparable to those in wild-type mice (Fig. 5G). We also examined the differentiation of tanycyte- and hypocretin/orexin-expressing neurons using molecular markers. All of these tanycyte and hypocretin neuron markers were normally expressed in the KO mouse brain (Fig. 5H), suggesting that UGS148 was not involved in the differentiation of these cell types, at least in the animals reared under the laboratory conditions.

Discussion

We have thus characterized UGS148 as an ER membrane protein specifically expressed in brain tissue, which is not essential for the mice reared under normal laboratory conditions. Consistent with a previous report (Ma et al. 2015), the strongest expression of UGS148 was observed in tanycytes that align the ventral region of the 3rd ventricle in the hypothalamus, and antibody staining of UGS148 revealed the fine morphology of tanycytes with well-developed ER networks located apically. Given that tanycytes regulate molecular transport between the cerebrospinal fluid and hormone-secreting neurons in the ventral hypothalamus (reviewed in Langlet 2014; Rodriguez et al. 2019), UGS148 may regulate molecular trafficking across the tanycyte through the characteristic ER membranes. However, KO mice lacking UGS148 expression did not exhibit obvious changes in the ER membrane morphology of tanycytes, and tanycyte-specific molecular markers were normally expressed in the mutant mice. In addition, the body weights of the UGS148 KO mice were indistinguishable from those of the WT mice, suggesting that the proposed functions of tanycytes, e.g., regulation of the energy balance as a component of the hypothalamic network, were maintained in the UGS148 KO mice. While UGS148 is not essential in mice reared under normal laboratory conditions, this finding does not exclude the possibility that UGS148 becomes functional when animals are placed under certain stressed conditions. Furthermore, certain single-transmembrane ER proteins may compensate for the lack of UGS148 in tanycytes and function redundantly in the tanycytes of UGS148 KO mice. Notably, the N-terminal region of UGS148 is composed of amino acid sequences that are predicted to be intrinsically disordered. Moreover, the functions of certain disordered proteins are maintained even after random shuffling of the primary amino acid sequence (Tsuboyama et al. 2020), suggesting that the composition of the amino acid, rather than the primary sequence, is important for the function of disordered proteins that do not have rigid 3-dimensional structures. A conventional homology search based on the conservation of the primary amino acid sequences may thus fail to find family proteins that function redundantly with UGS148.

Recently, much attention has been given to the properties of proteins containing intrinsically disordered regions, which mediate multivalent weak molecular interactions (reviewed in Shin & Brangwynne 2017; Alberti *et al.* 2019; McSwiggen *et al.* 2019; Peng & Weber 2019; Roden & Gladfelter 2021). Intrinsically disordered regions undergo liquid-liquid phase separation *in vitro* and are suggested to be involved in the formation of various molecular

condensates in the cells, including nonmembranous organelles, transcriptional complexes, organizing sites of autophagosomes, and disease-related insoluble aggregates (reviewed in Shin & Brangwynne 2017; Alberti *et al.* 2019; Roden & Gladfelter 2021). The N-terminal region of UGS148 may thus be involved in the formation of certain molecular condensates at the surface of the ER membrane and promote efficient membrane trafficking or molecular transport. Understanding the molecular rules that specify the function of disordered regions of proteins (Wang *et al.* 2018) or lncRNAs (Lubelsky & Ulitsky 2018; Pyfrom *et al.* 2019) may lead to the discovery of hidden gene families that escape classic bioinformatics approaches based on primary sequence homologies.

Many studies have revealed functional interactions between mitochondrial and ER membranes, especially through the contact site called the mitochondria-associated ER membrane (MAM). The MAM regulates the fission and fusion of mitochondria as well as molecular changes between the two organelles through a set of proteins that localizes to the contact sites (reviewed in van der Bliek et al. 2013; Petrungaro & Kornmann 2019). We have shown that UGS148 interacts with all three subunits of mitochondrial ATP synthase at least in vitro, suggesting that UGS148 also functions as an MAM protein to regulate mitochondrial functions. ATP synthetase is normally found in the inner membranes of mitochondria and is topologically separated from ER membranes by the outer membranes of mitochondria in general. However, early electron microscopic studies have shown that the ER membrane occasionally exhibits continuity with tubules protruding from the mitochondrial outer membranes (Franke & Kartenbeck 1971; Spacek & Lieberman 1980), which are commonly found in all tanycyte subtypes (Rodriguez et al. 2019). Indeed, we occasionally observed colocalization of UGS148 and mitochondrial markers in tanycytes, although the frequency was low compared with that of other MAM proteins. In addition, UGS148 has been shown to be distributed around mitochondria and the base of ciliary processes in tanycytes (Ma et al. 2015). These previously reported signals were not clearly observed with the monoclonal antibody we used, raising the possibility that the epitope recognized by the monoclonal antibody is masked in molecular complexes outside of the ER membrane and can only be detected with the polyclonal antibodies used in a previous study (Ma et al. 2015). To clarify the possible physiological importance of the interaction of UGS148 with ATP synthetase, researchers should further dissect the precise molecular mechanism and cellular conditions that lead to the interaction of topologically separated molecules.

The probes we initially used to detect UGS148 mRNA strongly cross-reacted with certain RNAs that could not be identified by BLAST search or other bioinformatics methods

available to date to the best of our knowledge. Hybridization-based detection of target RNAs is a fundamental tool for the study of RNA localization within cells or tissues, and databases of systematic genome-wide analyses of RNA localization (Lecuyer et al. 2007; Yokoyama et al. 2009; Sunkin et al. 2013) provide highly useful platforms to investigate the expression pattern of certain RNAs of interest in silico. However, the specificity of RNA probes during hybridization-based detection might be regulated by various factors in addition to conventional base pairing, including secondary structures, the formation of non-Watson-Crick base pairs, and possibly by interactions with particular proteins. Indeed, we previously found that probes targeting a specific mRNA strongly cross-reacted through stretches of 22 nucleotides (nt) with 2 nt mismatches even under stringent hybridization conditions (Ishida et al. 2015). Therefore, the specificity of the RNA probes used in localization studies, such as in situ hybridization, must be experimentally validated using multiple approaches. Northern hybridization is a classic method for the quantification of specific RNA transcripts (Alwine et al. 1977) but still serves as the only experimental approach that can experimentally estimate the size of the target transcripts, which is particularly useful to confirm the specificity of the probes. The importance of multiple approaches to confirm the specificity of hybridization probes must not be underestimated even in the era of genome-wide high-throughput studies.

Material and Methods

Animals

All experiments were approved by the safety division of Hokkaido University (#2015-079). C57BL6/N mice were used for all experiments except if otherwise mentioned. For anesthetization, medetomidine-midazolam-butorphanol (Kawai *et al.* 2011) was intraperitoneally injected at a volume of 10 μ l/g of body weight.

cDNA cloning, plasmid construction and cell lines

The coding sequence of UGS148 was amplified by PCR using a BAC clone (RP23-479A13) as a template and cloned into pT2K XIG Δ in (Urasaki *et al.* 2006). The sequence of calreticulin was obtained by RT-PCR using a cDNA template prepared from SK-N-SH cells. For generation of pT2KXIG Δ in avGFP_2A_FLAG-UGS148 and pT2KXIG Δ in avGFP_2A_UGS148-FLAG, the synthesized ORF region was purchased from gBlocks Gene Fragments (IDT) and cloned into pT2K XIG Δ in, which was used to establish stable Neuro2A cell lines used in the coimmunoprecipitation experiments. All constructs were assembled using Gibson Assembly (#E2611L, NEB), and the sequences of inserted cDNA were confirmed by Sanger sequencing. For plasmid transfection, expression vectors were mixed with FuGENE HD transfection reagent (#E2311, Promega) at a ratio of 1:3 and incubated with target cells, following the manufacturer's instructions.

Immunohistochemistry

For immunohistochemistry of the cultured cells, cells were cultured on coverslips coated with 0.5 mg/ml poly-L-lysine (Sigma). Cells were fixed with 4% paraformaldehyde (PFA) in HEPES-buffered saline [HBSS: 10 mM HEPES (pH 7.4), 137 mM NaCl, 5.4 mM KCl, 0.34 mM Na₂HPO₄, 5.6 mM glucose, 1 mM CaCl₂, 1 mM MgCl₂] for 20 minutes at room temperature and permeabilized with 0.5% Triton X-100 in PBS for 5 minutes at room temperature. To selectively permeabilize the plasma membrane, we treated the cells in digitonin buffer [10 mM PIPES (pH 6.8), 1 mM EDTA, 0.1 M KCl, 25 mM MgCl₂, 0.3 M sucrose, 0.01% digitonin] for 5 minutes on ice. To prepare sections of mouse brains, we perfused anesthetized mice with 4% PFA in HBSS. Dissected brains were washed once with HBSS, embedded in OCT compound (Sakura Finetech), and frozen on dry ice-ethanol. Frozen samples were sectioned at a thickness of 8 µm using a cryostat (Zeiss, HM520), collected on

PLL-coated slide glasses (Matsunami), and fixed in 4% PFA in HBSS for 1 hour at room temperature. For permeabilization, slices were washed three times in HBSS and incubated in 100% methanol for 5 minutes at -20°C. Nonspecific binding was blocked in blocking buffer containing 4% skim milk (Difco) in TBST (50 mM Tris-HCl, 150 mM NaCl, 0.01% Tween 20) for 5 minutes. The samples were incubated in primary and secondary antibodies diluted in blocking buffer for 1 hour each, followed by 3 washes in TBST. After the final washing, samples were mounted with 97% 2,2-thiodiethanol mounting media and PVA-based mounting media when using Cy2 and Alexa 488, respectively, as described previously (Mito *et al.* 2016). The list of antibodies used in this study is shown in Supplemental Table S1. Fluorescence and bright field images were obtained using an epifluorescence microscope (BX51; Olympus) equipped with a CCD camera (DP70). For wndchrm analyses, 50 equivalently sized TIFF images (256 X 256 pixels) were obtained using a 40X objective lens and analyzed using wndchrm (Shamir *et al.* 2008) following the manuals provided in GitHub (https://github.com/wnd-charm/wnd-charm).

Reanalyzes of single-cell RNA-Seq data

Single-cell RNA-Seq data of hypothalamus samples (GSE87544) were downloaded, and 3,319 cells with <2,000 detected genes were selected from 1,443,737 cells using Seurat (Stuart *et al.* 2019) in R. Hierarchical clustering was performed using hclust in R, and a heatmap was generated using "heatmap" function in R using a set of genes described as tanycyte markers in the original paper (Chen *et al.* 2017).

Northern blotting

Total RNA was extracted from cultured cells using TRIzol reagent (#15596026, Thermo Fisher Scientific) according to the manufacturer's instructions. For purification of RNAs from tissues, three volumes of TRIzol LS (#10296028, Thermo Fisher Scientific) were added to 1 volume of tissue samples, and tissues were homogenized using Omni THQ (Omni International). Ten micrograms of total RNA was mixed with the same volume of Gel Loading Buffer II (AM8546G, Thermo Fisher Scientific) and heated at 65°C for 10 minutes before loading. The RNA samples were separated on a 1% agarose gel (1% agarose gel, 1X MOPS, 10% formaldehyde), washed in distilled water, and treated with 0.05 N NaCl for 20 minutes to enhance the transfer efficiency. After equilibration in 20X SSC, RNAs were transferred to a nylon membrane (#11209299001, Sigma-Aldrich) using a standard capillary method and UV-crosslinked using a FUNA UV crosslinker (Funakoshi). Membranes were hybridized with

appropriate probes diluted in DIG Easy Hyb (#11603558001, Sigma-Aldrich) at 68°C for 16 hours, washed in 2× SSC at 68°C for 30 minutes, and washed twice in 0.2× SSC at 68°C. Hybridized probes were detected with AP-conjugated anti-DIG antibody (#11093274910, Sigma-Aldrich) and CDP-star (#11685627001, Sigma-Aldrich) following the manufacturer's instructions. Chemiluminescence signals were detected using a ChemiDoc Touch Imaging System (Bio-Rad).

Identification of UGS148-interacting proteins using MALDI-TOF/MS

For extraction of total cell lysates, stable transfectants expressing FLAG-tagged UGS148 were lysed in HBST buffer [0.5% Triton X-100, 150 mM NaCl, 10 mM HEPES buffered with KOH (pH 7.4)] for 30 minutes at 4°C and centrifuged at 17,400 g for 5 minutes to remove cell debris. Anti-FLAG M2 antibody affinity gels (#A2220, Sigma-Aldrich) were washed with HBST, added to the supernatant, and incubated for 1 hour at 4°C. After extensive washing with HBST, the proteins bound to the beads were eluted by the addition of FLAG peptide (#F3290, Sigma-Aldrich). Proteins in the eluate were separated by SDS-PAGE to detect UGS148-interacting proteins by Western blotting or by silver staining. To analyze the protein sample for MALDI-TOF/MS, a Silver Stain MS kit (#299-58901, Wako Pure Chemicals) was used, and staining and destaining were performed according to the manufacturers' instructions. After silver staining of the gel, visible bands were cut out, destained and chopped into small pieces. The gel pieces were dehydrated by soaking in acetonitrile and dried using a centrifugal evaporator (CVE-2100, EYELA). The gel pieces were further reacted with reducing solution [50 mM TCEP (Thermo Fisher Scientific), 25 mM NH₄HCO₃] at 60°C for ten minutes, followed by washing in washing solution (25 mM NH₄HCO₃). The gel pieces were further soaked for 30 minutes in alkylation solution (55 mM 2-iodoacetamide, 25 mM NH₄HCO₃) to alkylate the proteins. After alkylation, the gel pieces were heavily dehydrated by soaking in dehydrating solution (50% acetonitrile, 25 mM NH₄HCO₃) 3 times and once in acetonitrile. For digestion of the proteins in gels, dehydrated gel pieces were immersed in protease solution (50 mM Tris-HCl (pH 8.5), 10 ng/µl trypsin (#90057, Thermo Fisher Scientific), and 10 ng/µl lysyl endopeptidase (#121-05063, Wako Chemicals)] for 45 minutes on ice. Digested protein fragments were recovered using the extraction solution (50% acetonitrile, 5% TFA) in an ultrasonic bath for 30 minutes. The extracted protein solution was concentrated using a centrifugal evaporator. Extracted protein fragments were spotted onto the standard CHCA matrix for MALDI-TOF/MS on the target plate (MTP AnchorChip TM var/384 TF, Bruker Daltonics), and then, peptides were detected using UltraflexII-18 TOF/TOF (Bruker Daltonics). Identification of proteins was performed by peptide mass fingerprinting using a Mascot search (Matrix Science).

Western blotting

To prepare samples for Western blotting, mouse tissues were homogenized in CSK buffer [10 mM PIPES (pH 6.8), 100 mM NaCl, 300 mM sucrose, 3 mM MgCl₂] and mixed with an equal volume of 2× SDS buffer [0.1 M Tris-HCl (pH 6.8), 4% SDS, 10% 2-mercaptoethanol, 20% glycerol, and 0.1 mM PMSF]. For cultured cells, cells were lysed in RIPA buffer [10 mM Tris-HCl (pH 7.4), 1% NP40, 0.1% sodium deoxycholate, 0.1% SDS, 250 mM NaCl, 1 mM EDTA], and the soluble fraction was mixed with a one-sixth volume of 6x Laemmli buffer [30 mM Tris-HCl (pH 6.8), 6% SDS, 12% 2-mercaptoethanol, 50% glycerol]. Boiled samples were subjected to SDS-PAGE, and separated proteins were transferred to PVDF membranes. Membranes were soaked in blocking solution (4% skim milk in PBST) for 30 minutes and then reacted with appropriate antibodies diluted in PBST. Alexa Fluor 680-conjugated or IRDye800-conjugated antibodies were used as secondary antibodies to detect the bands by an Odyssey Infrared Imaging System (LI-COR).

Preparation of RNA probes

The primers used to prepare RNA probes are provided in Supplemental Table S1. The target sequence was amplified by PCR and cloned into a plasmid using a TOPO-TA cloning kit (#452640, Thermo Fisher Scientific). DNA fragments containing the T7 promoter sequence were further amplified by PCR using M13 forward and reverse primers and purified using a Wizard SV Gel and PCR clean-up system (#A9281, Promega) to prepare templates for *in vitro* transcription. DIG/FITC-labeled probes were prepared using DIG- (#11277073910, Sigma-Aldrich) or FITC- (#11685619910, Sigma-Aldrich) RNA labeling mix according to the manufacturer's instructions. Synthesized RNA probes were purified using Centri-Sep Spin Columns (Princeton Separations) or Centri Pure Mini (emp BIOTECH) to remove unincorporated labels and stored at -20°C with an equal volume of formamide.

In situ hybridization of tissue sections and brain slices

In situ hybridization was performed as previously described (Mito *et al.* 2016). All reactions were performed at room temperature unless otherwise mentioned. Briefly, fresh-frozen tissue sections were fixed in 4% paraformaldehyde in HBSS overnight at 4°C and treated with 0.2 N HCl for 20 minutes and subsequently with 3 μ g/ml Proteinase K (#3115887001, Sigma-

Aldrich) for 7 minutes at 37°C. The proteinase reaction was stopped in 0.2% glycine in PBS for 10 minutes, and sections were postfixed in 4% paraformaldehyde in HBSS for 20 minutes. After acetylation in acetylation solution (1.5% triethanolamine, 0.25% concentrated HCl, 0.25% acetic anhydride), sections were processed for hybridization. Hybridization was performed in hybridization solution (50% formamide, 1X Denhardt's solution, $2\times$ SSC, 10 mM EDTA, 100 µg/ml yeast tRNA, 0.01% Tween-20) for <16 hours at 55°C. Probes were washed twice with $2\times$ SSC containing 50% formamide at 55°C and treated with 10 µg/ml RNase A in RNase A buffer [10 mM Tris-HCl (pH 8.0), 500 mM NaCl, 1 mM EDTA, 0.01% Tween-20] for 1 hour at 37°C. After washing with $2\times$ SSC and $0.2\times$ SSC for 30 minutes at 55°C, the hybridized probes were detected with anti-DIG or anti-FITC antibodies.

To prepare brain slices, anesthetized mice were perfused with 4% paraformaldehyde, and dissected brains were embedded in 3% agarose. Brain slices at a thickness of 150 µm were prepared using a microslicer (DTK-100N, Dosaka EM), dehydrated in 100% methanol, bleached in a 1:5 mixture of 30% H2O2/methanol for 5 hours, and stored in 100% methanol at -20°C before use. *In situ* hybridization using brain slices was performed according to the whole-mount *in situ* hybridization protocol as described previously (Nakagawa & Takeichi 1995).

Production of recombinant proteins and anti-UGS148 monoclonal antibodies

To produce the His-tagged recombinant protein His-UGS148, the corresponding cDNA was subcloned into the pET-28a vector (Millipore). The recombinant protein was produced in *Escherichia coli* BL21 (DE3) and purified with TALON resin (Clontech). For immunization, 10 µg of His-UGS148 was mixed with the adjuvant TiterMax Gold (TiterMax) to produce antigen-adjuvant emulsions and injected intraperitoneally into four BALB/c female mice every two weeks. The lymphocytes from the immunized mice were fused with myeloma P3U1 cells at a ratio of 3:1 ratio by mixing in 50% polyethylene glycol (Roche). The fused cells were dispersed in 80 ml of GIT medium (Wako, Japan) supplemented with 1 ng/ml IL-6 (PeproTech) and 1×HAT (Kohjin-Bio). The cells were seeded in four 96-well plates at 0.2 ml/well and grown for 10 days at 37°C. The first screening was performed by enzyme-linked immunosorbent assay (ELISA) with 50 ng/well His-UGS148 and subsequently screened by Western blotting and immunofluorescence.

Generation of UGS148 KO mice

UGS148 KO mice (Accession No. CDB1164K: http://www2.clst.riken.jp/arg/mutant%20mice%20list.html) were generated following previously described protocols (Murata et al. 2004). Briefly, DNA fragments were amplified by PCR using a BAC clone that covering the UGS148 genomic region and subcloned into pDT-ApA/LacZ/NeO to generate the targeting vector. The linearized targeting vector was electroporated into TT2 ES cells (Yagi et al. 1993), and G418-resistant clones were screened by PCR followed by Southern blot analysis for homologous recombination. Chimeric mice were generated with the recombinant ES clone and mated with C57Bl/6 females to generate heterozygous animals. They were then mated with Gt (ROSA)26Sortm1(FLP1)Dym (Jackson Laboratory) to flip out the frt-flanked PGK-Neo cassette, and the resultant heterozygous mice were maintained on the C57BL/6 genetic background. PCR-mediated genotyping was performed using DNAs obtained from adult or embryonic tails with the following conditions: predenaturation at 96°C for 1 minute, followed by 30 cycles of denaturation at 94°C for 30 seconds, annealing at 62°C for 30 seconds, and extension at 72°C for 30 seconds. All animal protocols were approved the Institutional Animal Care and Use Committee of the RIKEN Kobe branch. Primers used to generate the UGS148 KO mice and used for genotyping the mice are provided in Supplemental Table S1 (Fig. 5A).

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Figure legends

Figure 1. Cross-reactivity of RNA probes and identification of UGS148 using a specific probe

(A) Localization of abundantly expressed transcripts of uncharacterized genes with the -Rik suffix (green). Nuclei were simultaneously detected with DAPI (magenta). (B) Schematic drawing of 6330403K07Rik and positions of RNA probes. (C) Sequence alignment of the region that exhibits homology to (TCTCCA)n revealed by the RepeatMasker search. (D) Coronal sections at the hypothalamus level showing the expression of 6330403K07Rik in the adult brain detected with the RNA probes shown in (B). Note that probe #1 detected signals in knockout (KO) mice lacking the expression of this gene. Intense signals in the ventral region of the hypothalamus (inset) were observed only in the brains of wild-type (WT) mice but not in those of KO mice. (E) Sagittal section of the brain in the Allen Brain Atlas database showing the expression of 6330403K07Rik. The ubiquitous expression was similar to the pattern obtained with probe #1 shown in (B). (F) Northern blot analyses of transcripts detected with probes #1, #2, and #3 shown in (B). Detection of 7SK was performed to confirm equal loading of the RNA samples in each lane. The asterisk shows the band corresponding to the size of 6330403K07Rik transcript. Arrowheads indicate bands detected in the samples obtained from the WT mice but not the KO mice. (G) Western blot analyses of UGS148 in the adult mouse brain. Specific bands were detected in the samples prepared from WT adult brains. (H) Localization of UGS148 mRNAs in brain cells. It should be noted that the transcripts are mostly localized in the cytoplasm but not in the nucleus. (I, J) Tissue distribution of UGS148 mRNA (I) and protein (J) detected by Northern blot and Western blotting, respectively. Note that both UGS148 mRNA and protein are specifically expressed in the brain. SG: salivary gland, BAT: brown adipose tissue, WAT: white adipose tissue. Scale bars, 10 µm in A and 1 mm in E and D.

Figure 2. UGS148 is expressed in tanycytes and orexin neurons in the hypothalamus

(A) Coronal section of brain at the level of the hypothalamus stained with an antibody against UGS148 (green). Nuclei were counterstained with DAPI and pseudocolored in magenta. UGS148 is expressed in tanycytes (Tn) and neurons with large cell bodies in the lateral hypothalamus (LH). (B) Simultaneous fluorescent detection of UGS148 (green) and GFAP (magenta) in serial coronal sections at the level of the hypothalamus flanking the medial

eminence (ME). White boxes indicate the regions shown at higher magnification in (D). UGS148 is strongly expressed in α1-, α2-, and β1-tanycytes (Tn), and weak expression is observed in β2-tanycytes located in the ME. (C) Fine morphology of tanycytes visualized with UGS148 antibody and observed with the Airyscan superresolution confocal microscope. Note the highly branched mesh-like structures in the apical branches (AB). CB: cell bodies, BP: basal processes. (D) Higher magnification of the ventricular region shown in (B). Ventricular astrocytes strongly expressing GFAP (magenta, arrowheads) did not express UGS148 (green). (E) Simultaneous detection of UGS148 (green) and Hcrt or Pmch (magenta) in the lateral hypothalamus. UGS148 is expressed in Hcrt-expressing neurons but not in Pmch-expressing neurons. Asterisks show autofluorescent signals derived from lipofuscin. (F) Hierarchical clustering analyses and heatmap of single-cell RNA sequencing data of the hypothalamus. UGS148 is clustered into a branch containing two pan-tanycyte markers, Slc16a2 and Col23a1. Red-yellow represents relative expression levels, and green lines represent count values.

Figure 3. UGS148 is an ER membrane protein with disordered regions

(A) Multiple sequence alignment of mouse, rat, and human UGS148 generated by ClustalW. Asterisks indicate conserved amino acids, and colons marks indicate similar amino acids. Acidic and basic amino acids are indicated in blue and magenta, respectively. The bold bar indicates the transmembrane domain predicted by TMHMM. (B) Analysis of the disorder propensity of UGS148 using IUPred2A. The blue line indicates an arbitrary threshold for the intrinsically disordered regions. Note that most of the N-terminally located nontransmembrane regions are intrinsically disordered. (C) Simultaneous detection of exogenously introduced UGS148 (green) and calreticulin-GFP (magenta) in HeLa cells. The dotted line indicates the region analyzed in (D). (D) Intensity profile plot of UGS148 (green) and calreticulin-GFP (magenta) shown in (C). Note that the peaks of these two signals completely coincide. (E) Morphology of ER membranes shown by calretinin-EGFP (magenta) in UGS148-expressing (green) and nonexpressing cells. Note that no overt morphological changes were induced upon expression of UGS148. (F) Wndchrm analyses of ER morphology of cells that do (plus) and do not (minus) express UGS148. The average class probability matrix index indicates the probability of the images classified into each category. The images were equally classified into either category, suggesting that they were not distinguishable by machine-learning-based image recognition. (G) Simultaneous detection of endogenous UGS148 (green) and the ER marker Hspa5 (magenta) using an Airyscan superresolution confocal microscope. Dotted lines

indicate the region analyzed in (H). (H) Intensity profile plot of UGS148 (green) and Hspa5 (magenta) shown in (G). (I) Detection of N-terminal and C-terminal-tagged FLAG epitopes in HeLa cells permeabilized with digitonin or Triton X-100. Note that C-terminal tagged FLAG was detected only when the cells were permeabilized with Triton X-100. (J) Schematic drawing of the topology of UGS148 in relation to the ER membrane. The N-terminal intrinsically disordered region is located in the cytoplasmic compartment.

Figure 4. UGS148 associates with the mitochondrial ATPase complex

(A) Silver staining of immunoprecipitated samples run on SDS-PAGE. UGS148 was specifically coimmunoprecipitated with multiple bands at approximately 50 kDa, which were determined to be ATPase complexes by MALDI-TOF analyses. (B) Western blot analyses of proteins coimmunoprecipitated with FLAG-tagged UGS148 detected with specific antibodies against the α - (ATPA), β - (ATPB), and γ - (ATPG) subunits of ATPase. Arrowheads indicate coimmunoprecipitated proteins detected by Western blotting. Note that multiple bands were detected for FLAG-UGS148. (C) Simultaneous detection of UGS148 and ATPA in HeLa cells that exogenously expressing UGS148. The signals were observed under a conventional fluorescence microscope. Note that the two signals did not largely overlap. The white box indicates the regions shown at higher magnification in the lower panels. The dotted line indicates the region analyzed in (D). (D) Intensity profile plot of UGS148 (green) and ATPA (magenta). Note that the peaks of these two signals largely do not coincide. (E) Simultaneous detection of UGS148 and ATPA in tanycytes. The signals were observed with a structural illumination superresolution microscope. The white box indicates the regions shown at higher magnification in the lower panels. The dotted line indicates the region analyzed in (F). (F) Intensity profile plot of UGS148 (green) and ATPA (magenta). Lines below the profile plots represent areas shown in the high magnification images. Arrows indicate overlapping peaks of UGS148 and ATPA signals.

Figure 5. Genetic deletion of UGS148 does not cause overt phenotypic changes in tanycytes

(A) Targeting strategy for disruption of UGS148. The open reading frame of UGS148 was replaced with the LacZ gene followed by polyadenylation signals. The founder heterozygous mice were crossed with mice expressing Flippase recombinase to flip out the frt-flanked Neo cassette, resulting in the generation of LacZ knock-in (KI) UGS148 KO mice, which were used

for subsequent analysis. SpeI restriction sites used for the Southern blot analyses are indicated in the short lines. (B) Southern blot analysis of SpeI-digested DNA from ES cells that undergo homologous recombination. The positions of the 5'- and 3'-probes and primers used for genotyping are indicated by black bars and magenta arrows, respectively, in (A). For the 5' probe, two probes were mixed for sensitive detection. (C) Agarose gel electrophoresis of PCR fragments amplified using primers for genotyping. The molecular markers are pUC19/HinfIdigested DNA. (D) The number of littermates obtained by crossing heterozygous UGS148 mutant mice. Wild-type (+/+), heterozygous (+/-), and homozygous (-/-) mice were obtained at nearly the expected rate. (E) Body weight of 8- to 22-week-old male and female littermates. Each dot represents individual animals, and the vertical line represents standard deviations. P values were calculated by Welch's t-test. (F) Coronal paraffin sections of adult brain at the level of tanycytes stained with hematoxylin-eosin. Note that no overt phenotype was observed in the KO mice. (G) Simultaneous detection of UGS148 and Hspa5 in tanycytes of WT and KO mice. Brain slices were stained with the antibodies, and the signals were detected using an Airyscan superresolution confocal microscope. (H) Expression patterns of tanycyte markers (Col25a1, Slc16a2, Vcan, Frzb) and hypocretin/orexin-expressing neurons (Hcrt) in WT and KO mice.









