The core circadian component, Bmal1, is maintained in the pineal gland of old killifish brain

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Highlights
The amplitude of free-running circadian rhythms decreases during aging in killifish
Core clock genes are highly conserved in the turquoise killifish genome
Bmal1 protein expression decreases in whole brain tissue with aging
Bmal1 expression is relatively well maintained in the pineal gland with aging
The core circadian component, Bmal1, is maintained in the pineal gland of old killifish brain

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Summary
Circadian rhythm is altered during aging, although the underlying molecular mechanisms remain largely unknown. Here, we used the turquoise killifish as a short-lived vertebrate model to examine the effects of aging on the major circadian network comprising the four mammalian clock protein homologs, Bmal1, Clockb, Cry1b, and Per3, which are highly conserved in the killifish with 50%–85% amino acid sequence identity to their human counterparts. The amplitude of circadian rhythm was smaller in old fish (14 weeks) than in young fish (6 weeks). In old fish brain, the Bmal1 protein level was significantly downregulated. However, the Bmal1 interaction with Clockb and chromatin binding of Bmal1 to its downstream target promoters were retained. Furthermore, Bmal1 was relatively well maintained in the pineal gland compared with other regions of the old fish brain. The results suggest that the circadian clock system in the killifish becomes spatially confined to the pineal gland upon aging.

Introduction
Most organisms have a circadian system characterized by an endogenous biological rhythm with a period of approximately 24 h, which is critical for physiological, metabolic, and developmental adaptation to daily environmental changes. This rhythm evolved in response to regular repetitive changes in environmental factors such as light and temperature resulting from the Earth’s rotation, but it can also be maintained under a constant environment as an internal clock (Panda et al., 2002b). The circadian clock controls a wide range of biological responses, including transcriptional and translational regulation, hormone secretion, metabolic cycling, and the sleep-wake cycle (Parhud and Aryan, 2018). In mammals, approximately 20% of transcripts undergo daily oscillations in expression (Du et al., 2014; Koike et al., 2012; Menet et al., 2012; Mure et al., 2018; Panda et al., 2002a; Ueda et al., 2002; Zhang et al., 2014), implying that a significant portion of physiological processes is under the control of the circadian clock. The circadian clock system consists of three main components: the input pathway that transfers the external environmental changes into the core system, the core oscillator that generates the 24 h rhythm, and the output pathway that leads to rhythmic activities (Lowrey and Takahashi, 2004; Quintero et al., 2003). In animals, the core oscillator consists of transcription/translation feedback loops formed by multiple components that modulate the abundance, interactions, and localization of various proteins during the entire day (Bell-Pedersen et al., 2005; Hardin, 2006; Panda et al., 2002b; Partch et al., 2014).

Four groups of genes encode the proteins that form the core oscillatory feedback loop in mammalian circadian systems: brain and muscle Aryl Hydrocarbon Receptor Nuclear Translocator-like 1/Brain and Muscle ARNT-Like 1 (ARNTL/BMAL1), circadian locomotor output cycles kaput (CLOCK), cryptochromes (CRYs), and periods (PERs). In the morning, BMAL1 and CLOCK form heterodimers that bind to the E-boxes, which are abundant in the promoters of PERs and CRYs, and lead to a gradual increase in the expression of PERs and CRYs during the day (Bunger et al., 2000; Gekakis et al., 1998; King et al., 1997; Kume et al., 1999; Yoo et al., 2005; Zheng et al., 2001). This accumulation is attenuated by casein-kinase-dependent protein phosphorylation and SCF ubiquitin ligase-dependent degradation (Ca-Nach et al., 2001; Eide et al., 2005; Shirogane et al., 2005; Vanselow et al., 2006). As night falls, PER and CRY form a complex that is protected from further degradation and transported into the nucleus. The nuclear PER-CRY complex then negatively regulates the gene expression of PERs and CRYs by disrupting the BMAL1/CLOCK dimer (Griffin et al., 1999; Kume et al., 1999; Lee et al., 2001; Sato et al., 2006).
In the mammalian circadian system, the suprachiasmatic nucleus (SCN) of the brain is the center of the master clock oscillator and is the main site that controls circadian physiology and behavior (Moore and Eichler, 1972; Stephan and Zucker, 1972; Yamazaki et al., 2000). The core circadian oscillator in SCN generates approximately 24-h cycling rhythms by receiving light input from eye photoreceptors. The circadian signals from the SCN are delivered to other tissues, such as the pineal gland and liver. Unlike mammals, teleost fish tissues and cells can generate their own rhythms by directly sensing external light (Whitmore et al., 2000); therefore, defining a site of the master circadian clock in teleost fish has been difficult.

As an organism ages, the circadian clock system and the rhythmic physiological cycles undergo changes (Banks et al., 2015; Dijk et al., 1999; Espiritu, 2008; Farajnia et al., 2012; Pandi-Perumal et al., 2005; Roenneberg et al., 2007; Van Cauter et al., 1998). However, the mechanisms underlying age-dependent changes of circadian clock system and physiological rhythms have yet to be clearly defined. For example, it is still unclear which accessory and/or core clock components undergo age-dependent changes in function and which are responsible for the age-dependent changes in physiological rhythms (Banks et al., 2016; Bonacorsa et al., 2014; Weinert et al., 2001). Conversely, it is known that defects in circadian components can affect longevity and age-related phenotypes (Kondratov et al., 2006, 2009; Sun et al., 2006). Knockout of the key clock components, CLOCK and/or BMAL1, reduces the lifespan of mice (Dubrovsky et al., 2010; Kondratov et al., 2006, 2009; Sun et al., 2006). Knockout of BMAL1 in mice causes neurodegeneration, sarcopenia, and early mortality (Kondratov et al., 2006). Thus, aging and the circadian clock system greatly affect each other, although the molecular mechanisms underlying their interactions remain unknown. Here, we investigated age-dependent alterations of the circadian clock system in Nothobranchius furzeri (the turquoise killifish), a short-lived vertebrate model organism, and suggest a possible molecular mechanism that explains age-associated alterations in circadian physiology.

Results

Changes in free-running circadian rhythms in old fish under LL and DD conditions

To monitor the behavior of turquoise killifish under free-running conditions, we first developed a circadian rhythm measurement system using a culture facility equipped with continuous flow-through water circulation. Monitoring locomotor activity is a typical non-invasive assay for measuring circadian rhythms. As the fish move three-dimensionally in water, monitoring the locomotor activity of fish is conventionally performed using laser sensors installed at the bottom of the tank and/or near the water surface in separate behavior rooms or facilities, and the circadian rhythm is assessed by measuring the number of times the fish touch the sensors over time (Hurd et al., 1998; Lucas-Sanchez et al., 2011). However, this assay, despite its convenience, generates a square-shaped output, so it is intuitively difficult to find the key parameters (period, phase, and amplitude) of the circadian rhythm. To improve the non-invasive characterization of circadian rhythmicity in a conventional locomotor activity measurement system, we developed a cost-effective video-based LL (continuous light) and DD (continuous darkness) free-running rhythm monitoring system with a continuous water flow in a culture facility (Figure 1A, see Transparent methods for details).

Using the custom-built video-based free-running monitoring system, we examined for potential changes in circadian rhythms during fish aging. GRZ-AD, the shortest-lived strain used in this study, had a median lifespan of approximately 16 weeks under our culture conditions. Thus, we measured endogenous circadian locomotor activity of young (6 weeks old, after sexual maturation) and old (14 weeks old, approximate median lifespan) fish under free-running conditions. Fish activity was first measured under LL by determining how many times the fish entered a specific sector during 15 minutes every hour, which was reported previously to represent locomotor activity (Lucas-Sanchez et al., 2011). We also analyzed movement duration, velocity, and total distance covered by the moving fish (Figure S1).

The raw video and heatmap of movement revealed differences in the general movement of young fish between day and night compared with that of older fish (Figure 1B and Video S1). The overall movement throughout the free-running cycles was higher in young fish than in old fish. Similar results were obtained by measuring mean velocity, cumulative movement duration, and total distance of movement, so mean velocity was chosen for further analysis (Figure S1).

The mean velocity of each fish was calculated to analyze endogenous rhythmicity under LL and DD free-running conditions. In the LL condition, we first noticed that the proportion of fish with rhythmic circadian activity was far smaller among old fish than among young fish; six of nine young fish and five of fourteen old
Fish showed significant rhythmic behavior under the LL condition (Table S1A). The circadian periods of the young and old fish with rhythmic circadian activity (adjusted p < 0.1) were 24.3 and 25.6 h under the LL condition, respectively, but the difference was not statistically significant (Figure 1C and Table S1A). In addition, old fish with rhythmic circadian activity showed more variation in circadian periods than young fish (Figure 1D). Under the DD condition, circadian rhythmicity was measurable in all young and old individuals, but neither their circadian periods (23.75 and 23.88 h for young and old fish, respectively) nor their phases

Figure 1. Changes in free-running circadian rhythms in old fish under LL and DD conditions

(A) Experimental setup of a video-based circadian rhythm measurement system under free-running conditions. (B) Representative images of fish movement during the subjective day and night. Cumulative fish movements are presented as a heatmap. (C–F) (C and E) The mean velocity of young (6 weeks old) and old (14 weeks old) fish, respectively, under LL (C) and DD (E) free-running conditions. Blue and red dots represent individual young and old fish, respectively. The gray (C) and the dark gray backgrounds (E) denote subjective night. CT denotes “Circadian time.” (D and F) Characterization of circadian rhythmicity of young and old fish under LL (D) and DD (F). The rhythmic behavior of individual fish was analyzed with JTK_CYCLE (Adj. p < 0.1, see Table S1). For amplitude, data are expressed as the mean ± SD, *p < 0.05.

See also Figure S1, Video S1 and Table S1.
differed significantly (Figure 1E and Table S1B). However, the amplitude of circadian rhythm was significantly smaller in old fish than in young fish under DD conditions (Figure 1F). These results suggested that free-running circadian rhythmicity was altered in old fish.

Core circadian clock components are highly conserved in the turquoise killifish

Having confirmed that the turquoise killifish exhibits typical circadian behavior with a free running rhythm in swimming behavior, we searched for homologs of the core circadian clock components, BMAL1, CLOCK, CRY1, and PER3, in the turquoise killifish genome. The turquoise killifish Bmal1, Clockb, Cry1b, and Per3 showed 85%, 65%, 81%, and 50% overall amino acid sequence identity with the corresponding human proteins, respectively, and they showed the highest amino acid sequence homology with human counterparts (Figures 2 and S2 and Table S1C); the amino acid identity is notably higher for Bmal1 and Cry1b than for Clockb and Per3. Multiple ancient paralogs of the circadian clock genes have been discovered based on amino acid comparisons. In the turquoise killifish genome there are three bmal genes (bmal1, bmal2a, and bmal2b), two clock genes (clocka and clockb), five cry genes (cry1a, cry1b, cry2a, cry2b, and cry5), and four per genes (per1b, per2a, per2b, and per3) (Table S1C). The core clock proteins with the highest homology to their human counterparts were further compared with those of mice and zebrafish, and their relationships were visualized as a phylogenetic tree (Figures 2 and S2). The turquoise killifish homologs of the four core clock proteins contained all the conserved functional domains of core clock proteins (Figure 2). Together, these results indicated overall high conservation of core clock proteins in the turquoise killifish.
**The mRNA expression of core circadian clock genes changes slightly during aging**

To determine whether the four core clock genes show rhythmic expression in the fish brain and, if so, whether their rhythmic expression is altered in old fish, their mRNA expression in brains of young and old fish was measured throughout the day. Brain aging was first observed by staining brain sections with Fluoro-Jade B, and the level of neurodegeneration was greater in old fish than in young fish (Figure S3). In the LD (light-dark cycles) condition, expression of bmal1 and clockb showed peaks around 10 h after light on (zeitgeber time [ZT] 10 h), whereas that of cry1b and per3 exhibited a relatively higher peak around ZT 22 h (Figures 3A and S4A and Table S1D), as expected for turquoise killifish, a diurnal animal, whose expression of these genes is anti-phasic with respect to the corresponding genes in nocturnal animals such as mice. The circadian control of turquoise killifish clock gene expression was robust under the LD condition, as indicated by the >3-fold difference between peak and trough expression of all genes tested. bmal1/clockb and cry1b/per3 were controlled by antagonistic cycles, suggesting that they can form a transcriptional/translational feedback loop similar to that in other model organisms (Figure S4A). The circadian amplitude in the expression of the four clock genes was not significantly different between young and old fish brains. These results indicated that aging has a minor effect on the rhythmic mRNA expression of the core clock genes under the LD condition.

Core clock gene expression was also assessed 3 days after transferring the fish to a DD free-running condition (Figure 3B). bmal1, clockb, and per3 cycled throughout the day, similar to the pattern observed under the LD condition; however, cyclic cry1b expression was almost completely abolished in both young and old fish under the DD condition. The phase of bmal1 and per3 expression was more advanced in old fish that in young fish under the DD condition (Figures 3B and S4B and Table S1E). However, the expression patterns of the four core genes under the DD condition did not show a marked change in the brain of old fish. These results show that the rhythmic expression of cry1b is more critically affected by the DD condition than by the LL condition, but the amplitude and period of expression of the four core clock genes in the brain of old fish were similar between the DD condition and LD condition.

**The Bmal1 protein level decreases dramatically in the brain of old turquoise killifish**

Next, we examined potential post-transcriptional changes associated with organismal age (Figure 4). For this purpose, we used human BMAL1 and CLOCK antibodies to detect killifish Bmal1 and Clock proteins, because antibodies against killifish Bmal1 and Clockb were not available and Bmal1 and Clockb proteins show high amino acid identities to their human homologs (85% and 65%, respectively). First, we tested the specificity of human BMAL1 and CLOCK antibodies for the detection of synthetic Bmal1 and Clockb proteins of the turquoise killifish. Both anti-BMAL1 and anti-CLOCK antibodies detected ectopic Bmal1 and Clock proteins of the turquoise killifish and had sufficient specificity to detect endogenous Bmal1 and Clockb in the killifish brain (Figure S5).

Assessment of Bmal1 and Clockb protein expression in killifish brain tissue under LD and DD conditions showed that Bmal1 protein expression cycled during the day in both young and old brains. The expression of the Bmal1 protein peaked at approximately ZT13 under the LD condition, and a 3-fold difference in the abundance of Bmal1 protein was observed between its peak and the trough levels. In old fish brain, the abundance and amplitude of Bmal1 protein expression were lower than in young brain, whereas the phase and period were not different under the LD condition (Figures 4A and S6A and Table S1F). Clockb protein was expressed at a constant level during the day, and its abundance was not significantly different between old and young brains (Figure 4B). Similar patterns of Bmal1 and Clockb protein expression were observed under the DD condition (Figure 4C). Bmal1 protein abundance and its amplitude were significantly lower in old fish brain than in young fish brain, but there were no changes in the phase and period of its expression. Clockb protein expression did not differ significantly between young and old fish under the DD condition (Figures 4C and S6B and Table S1G).

We also examined the phosphorylation of Bmal1 and Clockb in young and old fish brains by measuring inter-gel differences in the mobility of phosphorylated and non-phosphorylated forms of these proteins (See Transparent methods for details). Bmal1 from young fish brain was detected as four separate bands, and the relative intensities of the four bands varied throughout the day, indicating that Bmal1 underwent diurnal changes in phosphorylation. The abundance of phosphorylated Bmal1 was lower in old fish brain (Figure 4A), and the relative intensities of the four bands were different from those of young fish brain during the day, indicating that Bmal1 in old fish brain was subject to different types of phosphorylation throughout the day. Clockb protein from young and old turquoise killifish brains migrated as a single band suggesting that it might be uniformly phosphorylated or not phosphorylated (Figure 4B).
The amount and function of the Bmal1/Clockb heterodimer are retained in the old killifish brain

In circadian clock system, Bmal1 and Clockb form a complex to control downstream genes including the evening clock genes, cry1b and per3. As fish age, the expression of Bmal1 dramatically decreased; thus...
we tested if this age-dependent change in Bmal1 expression affects the Bmal1/Clockb complex pathway. We first confirmed that Bmal1 and Clockb proteins interact in the killifish brain by performing bidirectional immunoprecipitation using anti-BMAL1 or anti-CLOCK antibodies (Figure S7). The Bmal1/Clockb interaction was monitored throughout the day in young and old fish brains (Figure 5A). After normalizing the Bmal1/Clockb interaction using immunoprecipitated Bmal1, we noticed that the relative Bmal1/Clockb interaction was not markedly different between old fish brain and young fish brain (Figure 5A). Although
the Bmal1 protein abundance decreased during aging, the Bmal1/Clockb interaction was maintained or protected in the aging brain.

Next, we examined the chromatin binding of Bmal1 to the E-box regions of circadian clock-regulated gene promoters. The region upstream of the start codon in the cry1b and per3 genes was searched to see if it contained canonical E-box (CACGTG) and non-canonical E-box (CACGTT and CAGCTT) sequences (Nakahata et al., 2008; Salero et al., 2003; Zhang et al., 2012). To analyze for age-dependent differences in Bmal1-chromatin association in the killifish brain, five E-box-containing regions in each of the cry1b and per3 promoter regions were targeted. Among the five amplicons tested, three cry1b and four per3 E-box-containing genomic regions were identified as Bmal1-binding sites, but not in the GFP control. Only one target sequence in the per3 promoter (amplicon 2 in Figure 5C, −4,720 to −4,917 before the start codon) showed a significant age-dependent increase in Bmal1 chromatin association. These results indicate that despite the large downregulation of Bmal1 in old fish brain, Bmal1/Clockb heterodimer formation and Bmal1 chromatin binding to E-boxes were relatively well maintained in old fish brain. This was consistent with the subtle changes in the age-dependent expression of cry1b and per3 mRNA expression.

**Bmal1 is relatively well maintained in the pineal gland of old turquoise killifish brain**

It was curious that the Bmal1/Clockb interaction and Bmal1-chromatin association were retained in old killifish brain despite the significant decrease in the Bmal1 protein level. One possibility was that Bmal1/Clockb heterodimer formation occurs in a specific location in the brain where the local expression of Bmal1 for Bmal1/Clockb heterodimer formation is relatively well sustained during aging. To test this idea, we used whole brain and detected Bmal1 and Clockb proteins in the killifish brain after making the brain transparent.

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**Figure 5. The amount and function of the Bmal1/Clockb heterodimer are retained in old killifish brain**

(A) Bmal1/Clockb heterodimer formation throughout the day in the brain of young and old fish. Co-immunoprecipitated (co-IPed) Clockb was normalized against IPed-Bmal1 for each trial. Data are presented as the mean ± SD from three replicates.

(B and C) Binding affinity of Bmal1 to the promoter regions of cry1b (B) and per3 (C) in the brains of young and old fish. Schematic diagrams of the promoter regions of cry1b and per3 are shown along with the amplicon positions. E-box, red bar; gray box, 5′-UTR; black box, protein coding sequence; solid arrow, start codon; pale blue bar, amplicon position. Data are presented as the mean ± SE from three replicates. Asterisks indicate *p < 0.05, **p < 0.01, and ****p < 0.0001 after two-way ANOVA followed by the Tukey’s multiple comparison test.

See also Figure S7.
To get an insight into the function of Bmal1 and Clockb, we first examined their intracellular and tissue localizations in young fish brain. Bmal1 protein was localized to both the nucleus and cytoplasm, whereas Clockb protein was preferentially localized to the cytoplasm (Figure 6A). The two proteins were detected throughout the brain but their spatial distribution differed slightly (Figure 6B and Videos S2, S3, and S4). For example, both Bmal1 and Clockb were detected on the surface of the brain, but Clockb protein was detected in a deeper layer than Bmal1 in the optic tectum (Figure 6B and Video S3).

Figure 6. Bmal1 is relatively well maintained in the pineal gland of old turquoise killifish brain
(A) Subcellular localization of Bmal1 and Clockb in the turquoise killifish brain. Green, red, and blue colors represent Clockb, Bmal1, and DAPI (nuclear marker) staining, respectively.
(B) Co-localization of Bmal1 and Clockb in the telencephalon, optic tectum, rhombencephalon, and SCN of the turquoise killifish brain.
(C) Whole brain immunostaining of Bmal1 and Clockb in young and old turquoise killifish brains.
(D) Bmal1 and Clockb localization in the pineal gland of young and old fish brains. Abbreviations “A” and “P” in the figure represent directions toward the anterior and posterior sides of the brain, respectively.
See also Figure S8 and Videos S2, S3, S4.
We then analyzed the effect of aging on the whole brain localization of these two proteins. Fluorescence detection of Bmal1 revealed that its level decreased more drastically than that of Clockb in old fish brain (Figures 6C and S8), which was consistent with the analysis of their expression at the protein level (Figure 4). However, Bmal1 fluorescence was better maintained in the pineal gland than in other parts of the brain. Bmal1 and Clockb co-localized in an internal area of the pineal gland, whereas Bmal1 alone was detected in the epidermal region of the pineal gland (Figures 6C and 6D). These data indicate that Bmal1/Clockb complex formation might be specifically maintained in the pineal gland of old killifish brain.

Discussion

The circadian rhythm is weakened in old killifish

A custom-built non-invasive and cost-effective endogenous circadian locomotor monitoring system was used to obtain a robust and clear picture of the locomotor circadian rhythms of the short-lived turquoise killifish Nothobranchius furzeri, strain GRZ-AD, which includes mean velocity, cumulative movement duration, and total distance of movement under LL and DD conditions. Continuous monitoring of the circadian activity using this system revealed a typical wave-like circadian oscillatory pattern.

The turquoise killifish were active during the day and calm at late night, as well expected for a diurnal animal. In the free-running LL condition, the proportion of fish with significant rhythmic behavior decreased, and the circadian period tended to be longer in old fish. An age-dependent decline in circadian rhythmicity has been reported for other model organisms, including teleost fishes such as Danio rerio (zebrafish) and Nothobranchius korthausae (Lucas-Sanchez et al., 2011; Zhdanova et al., 2008). However, the lifespans of these organisms are relatively long; the median lifespans of zebrafish and N. korthausae are 42 months and 72 weeks (Gerhard et al., 2002; Lucas-Sanchez et al., 2011), respectively. Investigation of age-associated changes in circadian activity using these model organisms requires 4 years for zebrafish and 72 weeks for N. korthausae (Lucas-Sanchez et al., 2011; Zhdanova et al., 2008). The turquoise killifish strain used in this study, GRZ-AD, is short-lived vertebrate with a median lifespan of only 16 weeks under laboratory conditions, and a decrease in circadian locomotor activity was observed even in 14-week-old fish, demonstrating that the killifish model is a reliable and useful system for the study of age-dependent changes in circadian rhythm and physiology. It should also be noted that the killifish is a diurnal animal, like human but unlike mouse, which makes it additionally advantageous as a vertebrate circadian model system.

Free-running circadian locomotor activity was different between LL and DD conditions. Under the DD condition, all young and old fish showed significant circadian behavior unlike under the LL condition. Interestingly, a phase difference between circadian rhythms under LL and DD conditions was also observed and the circadian phase was advanced in LL and delayed in DD (Figures 1D and 1F). The circadian period in DD was not significantly different in old fish; however, we speculate that, with advancing age, the circadian period will get longer in DD as well. As individual variability in circadian locomotor behavior was lower under DD, we propose that the DD condition allows a better estimation of the internal circadian rhythm in turquoise killifish.

The turquoise killifish has highly conserved circadian components

The core circadian system in animals is highly conserved across species, including flies, zebrafish, and mammals (Hardin, 2011; Lowrey and Takahashi, 2004; Vatine et al., 2011). The core circadian oscillatory component complexes in these animals, BMAL1/CLOCK and CRY1/PER3, form an interlocking feedback loop and are well conserved in the turquoise killifish genome. This suggests that the basic molecular mechanisms controlling the circadian clock are comparable between the turquoise killifish and other vertebrate models. There are multiple paralogs of these circadian core genes in the turquoise killifish genome, reflecting their early emergence during evolution and speciation from other teleost fishes (Table S1C). For example, eight ancient paralogs of bmal1 have been identified in the killifish genome and one of them is clockb. All the bmal1 paralogs showed <50% sequence identity with bmal1, and they evolved from bilateral animals, namely, arthropods and nematodes (Ecdysozoa). The clockb gene has one paralog (clocka) and seven ancient paralogs in the killifish genome that are also present in bilateral animals. The clocka gene has the highest amino acid sequence identity (62.26%) with Clockb and is duplicated in teleost fish (Clupeocephala). cry1b has one paralog (cry1α), which is a duplicate from Osteoglossocephalai, and five ancient paralogs that are similar to those found in Saccharomyces cerevisiae. per3 has three ancient paralogs that are similar to those found in arthropods and nematodes (Ecdysozoa). It would be interesting to investigate the
key features of the turquoise killifish circadian clock network that distinguish it from the clock regulatory pathways of other vertebrate and non-vertebrate organisms.

Organismal age has little impact on the Bmal1 and Clockb pathway in the brain of the turquoise killifish

The turquoise killifish core clock components show the following conserved properties across species: (1) mRNA cycling of bmal1, clockb, cry1b, and per3 with an opposite phase between bmal1/clockb and cry1b/per3; (2) cycling of the Bmal1 protein; (3) phosphorylation of Bmal1 protein; (4) Bmal1 and Clockb heterodimer formation; and (5) chromatin binding of Bmal1 to the E-Box sequences of promoters of circadian clock-controlled genes. The turquoise killifish Bmal1 and Clockb proteins form a heterodimer that binds to cry1b and per3 promoters, inducing the expression of these genes. This indicates that the negative-positive feedback loop in animal core clocks system is also operational in turquoise killifish.

The age-dependent changes of circadian core gene expression in mammals and flies still remain unclear (Asai et al., 2001; Rakshit et al., 2012). In the other fish model, zebrafish, bmal1a and per1a expression is lower in 4-year-old fish than in 1-year-old fish, whereas clocka expression does not change with age (Zhdanova et al., 2008). In our study using turquoise killifish, bmal1, clockb, cry1b, and per3 showed only minor differences in their rhythmic expression between young (6 weeks old, after sexual maturation) and old (14 weeks old, approximate median lifespan) fish brain (Figure 3). However, overall Bmal1 protein abundance was dramatically lower in old fish brain than in young fish brain throughout the day (Figure 4). This observation provides a clue to understanding the age-dependent change in the fish circadian clock; the post-transcriptional regulation of Bmal1, a core clock protein, occurs before a change occurs in its transcript level. Another important finding is that aging, despite its negative effect on Bmal1 protein abundance, had a minimal effect on Bmal1/Clockb heterodimer formation and E-box binding affinity of Bmal1 (Figure 5). These results suggest that the circadian core oscillator system involving Bmal1 remained robust in old fish, especially in the pineal gland of the brain as explained below.

Bmal1 is preferentially maintained in the pineal gland of old killifish brain

The master circadian clock oscillator is located in the SCN in the mammalian brain to control circadian physiology (Yamazaki et al., 2000; Yoo et al., 2004). However, in teleost fishes including zebrafish, the location of the master clock is unclear, because the light sensing capacity of every cell can generate endogenous rhythms (Whitmore et al., 2000). In young turquoise killifish brain, Bmal1 and Clockb proteins were widely expressed, whereas their cellular localizations were distinct. Analysis of whole brain localization of these proteins in the turquoise killifish showed that Bmal1 and Clockb localized to distinct regions of the brain (Figure 6B). Whole brain imaging of Bmal1 and Clockb suggested that the circadian network associated with Bmal1 or Clockb can form throughout the brain but can also be restricted to specific regions of the brain through the spatiotemporal co-localization of Bmal1 and Clockb.

Whole brain immunostaining provided information on the age-dependent localization of Bmal1 and Clockb in the killifish brain. The fluorescence intensity of Bmal1 was decreased in whole brain images of older fish, in agreement with the decreased overall protein level. However, the fluorescence intensity in one small area of the dorsal brain, the pineal gland, remained constant relative to that in other parts of the brain, and Bmal1 and Clockb co-localized in cells of the pineal gland (Figure 6C). This result suggests that Bmal1 acts as a critical modulator of the circadian network during aging, and that the pineal gland is a major sub-organ involved in the maintenance of circadian activity, most likely via the Bmal1/Clockb complex, in old killifish.

In this study, we identified age-dependent changes in the highly conserved core circadian clock components of the turquoise killifish. We showed that downregulation of the core clock protein Bmal1 occurs mainly at the translational level, but that its expression is maintained in the pineal gland of old killifish brain, probably to maintain circadian gene expression. The pineal gland, which secretes melatonin, is an important region for neuro-endocrine activity (Wurtman et al., 1963) and also acts as a central circadian oscillator in many non-mammalian vertebrates (Fukada and Okano, 2002). The present results suggest that the master clock of the old turquoise killifish might be located in the pineal gland. Our findings thus provide an important insight into the change that occurs in the spatiotemporal regulation of circadian clock system in the brain of aging vertebrates.
Limitations of the study
The pineal gland-specific mechanisms of Bmal1/Clockb complex regulation of circadian rhythms require further investigation. The physiological roles of age-specific and pineal gland-specific maintenance of Bmal1 with cell-type specific markers of pineal gland should help improve our understanding of circadian physiology during aging. Comparison of the spatiotemporal patterns of Bmal1 expression across species could provide further insights into the evolution of the circadian system and the role of Bmal1 in the circadian system of aging vertebrates.

Resource availability
Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Yumi Kim (yumikim@ibs.re.kr).

Materials availability
All unique reagents generated in this study are available from the Lead Contact without restriction.

Data and code availability
All data are available in the main text or Supplemental information. Raw data images can be found Mendeley Data: https://data.mendeley.com/datasets/gy57ymzh67/draft?a=b36c53c2-250b-4b92-bbe3-d1990378e123.

This study did not contain any newly generated datasets or codes.

Methods
All methods can be found in the accompanying Transparent methods supplemental file.

Supplemental information
Supplemental information can be found online at https://doi.org/10.1016/j.isci.2020.101905.

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Author contributions
Y.K. conceived the project. S.L. and Y.K. designed and performed experiments. Y.K., S.L., and H.G.N. wrote the manuscript.

Declaration of interests
The authors have no competing interests to declare.

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Supplemental Information

The core circadian component, Bmal1, is maintained in the pineal gland of old killifish brain

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Figure S1. Locomotion of circadian rhythm measurements under the free-running LL condition. Related to Figure 1. Activity counts recapitulating a conventional method for locomotion (A), mean velocity (B), total distance (C), and cumulative movement duration (D) of young and old fish were measured under the free-running LL condition. CT represent “Circadian time.” Each dot represents a single fish at each measurement (n = 4).
Figure S2. Amino acid alignment of the circadian core components BMAL1, CLOCK, PER3, and CRY1 among *Nothobranchius furzeri*, *Danio rerio*, *Mus musculus*, and *Homo sapiens*. Related to Figure 2. Compared are the translated amino acid sequences of turquoise killifish, zebrafish, mouse, and human BMAL1 [XM_015965280 (*bmal1*), NM_131577 (*bmal1a*), NM_007489, and NM_001030272, respectively], CLOCK [XM_015971720 (*clockb*), BC163244 (*clock1a*), AF000998, and AF011568, respectively], CRY1 [XM_015964931 (*cry1b*), NM_001099297 (*cry1aa*), NM_004075, and NM_004075, respectively], and PER3 (XM_015959552 (*per3*), AF254792, NM_011065, and NM_001289862, respectively). Identical amino acid sequences are highlighted in red characters and a yellow background, weakly similar amino acids are displayed in green characters, a block of similar amino acids is shown in black characters with a green background, conserved sequences are displayed in blue characters and sky-blue background, and non-similar sequences are indicated in black characters. Red box in BMAL1 consensus sequence indicate an epitope region of human BMAL1 for anti-BMAL1 antibody (ab93806, Abcam).
Figure S3. Neurodegeneration in the aged killifish brain. Related to Figure 3. Degenerated neurons were stained with Fluoro Jade B, and image sections from two independent brains are displayed.
Figure S4. Circadian characteristics of clock gene expression in LD and DD. Related to Figure 3. Period, amplitude, and phase from the cyclic expression under LD (A) and DD (B) conditions were analyzed using JTK_CYCLE. Data are presented as the mean ± SD for period and amplitude.
Figure S5. Validation of commercial BMAL1 and CLOCK antibodies. Related to Figure 4. (A) Synthesized Bmal1 and Clock proteins were detected with anti-His antibody (ab184607, Abcam). (B) Synthesized Bmal1 and Clock proteins were detected with commercial BMAL1 (ab93806, Abcam and ITM0071-100M-647, GBiosciences) and CLOCK (R1511-2, HUABIO) antibodies. (C) Specificity of antibodies in this study. Total protein extracts from the turquoise killifish brain were used to test the specificity of commercial BMAL1 and CLOCK antibodies.
Figure S6. Circadian characteristics of Bmal1 and Clockb protein abundance in LD and DD.
Related to Figure 4. Abundance cycling of Bmal1 and Clockb in LD (A) and DD (B) was analyzed using JTK_CYCLE. Data are presented as the mean ± SD for period and amplitude.
**Figure S7. In vivo interaction between Bmal1 and Clock.** Related to Figure 5. Heterodimer formation between Bmal1 and Clockb in the turquoise killifish brain was detected using BMAL1 or CLOCK antibody.
Figure S8. Biological replicates of whole brain staining with BMAL1 and CLOCK antibodies. Related to Figure 6.
Transparent Methods

Fish husbandry

The GRZ-AD strain, which is a short-lived strain of the turquoise killifish, was used and maintained as previously described (Dodzian et al., 2018). The fish were cultivated under 12 h light and 12 h dark cycle. The fish were singly housed in a 1.8 L tank and fed twice a day at 1 h and 8 h after light on. Young and old fish were sacrificed after 6 and 14 weeks after hatching, respectively.

Fish care and experiments were performed in accordance with the animal care and use protocol that is reviewed and approved by the Institutional Animal Care and Use Committee at Daegu Gyeongbuk Institute of Science and Technology, Republic of Korea (Approval number: DGIST-IACUC-17103001-00).

Development of the assay system for measuring the free-running circadian rhythm

A wide tank measured 17 cm (w) × 24 cm (l) × 18 cm (h) was designed to contain relatively shallow water. The water level was maintained at a 3 cm in depth with continuous circulation of system water in the fish culture facility. The tanks were equipped on the LED light panel (LED light pad GB4, GAOMON) for a video recording in LL and 940 nm LED illuminator (custom-built) was installed on top of the tank for a video recording in DD. A web camera [WideCam F100 (Genius) for LL measurement and C920 PRO HD WEBCAM (Logitec) for DD measurement] was installed on top of each tank. One fish per tank was placed and the fish husbandry were consistently carried out following the ordinary fish husbandry scheme. The cameras were connected to a computer, and video images were acquired automatically every 15 min for 4 days using open source software (OBS studio 23.0.1). The raw video images were concatenated into one file per fish, and analyzed and visualized with a behavior analysis program (EthoVision XT, Version 13.0, Noldus Information Technology).

Characterization of circadian rhythms

Only male fish for each age groups were used for measuring circadian rhythmicity as indicated in the main figure and Table S1. The raw data (mean velocity from EthoVision XT outputs of LL and DD free-running rhythm measurements, and normalized circadian clock gene and protein expressions) were used to analyze the circadian rhythmicity with JTK_CYCLE (Hughes et al., 2010). The outputs of every analysis are shown in Supplementary Table 1; period, amplitude, and phase were used for visualization.

Measurement of circadian gene expression

The turquoise killifish brains were dissected from young (6-week-old) and old (14-week-old) fish. Each age group contains three males and two females per time point. Total RNAs were isolated individually from five fish per age group at each time point using QIAzol Lysis Reagent (79306, QIAGEN) following the manufacturer’s instructions. cDNAs were synthesized from 2.5 μg of total RNA using SuperScript™ IV VILO™ Master Mix with ezDNase™ Enzyme (11766050, Invitrogen). Synthesized cDNAs were diluted 10-fold, and 3 μL of diluted cDNA was used as a template for qPCR (SsoAdvanced Universal SYBR Green Supermix, BioRad). The primers used are listed below. The
core gene expressions were normalized with insr expressions which have been known to be well maintained over the fish age (Hartmann et al., 2011).

| Gene name | Forward primer | Reverse primer |
|-----------|----------------|----------------|
| bmal1     | CGATGGAAAGTTTGTTTCTCGT | TGGGGCAAATACGCTAGG |
| clock     | CAGCTTTCAGCCATGCAG | TGGGTAGATTGGTTTCCATGA |
| per3      | TCATGAGGAAATAAAGATCTACAAGC | CGCTGGAGGCCATGTTGT |
| cry1b     | GGCTCTCATGCGCTCGT | TGCTGGTAATAGCGCTGATG |
| insr      | TGCCCTCTCAACCCCTGAGT | AGATGGCGATCTATCACG |

Immunoblot analysis and co-immunoprecipitation

Total proteins were isolated from young and old brains using 100 mM Tris-HCl (pH 7.5), 1 mM EDTA, 50 mM NaCl, 0.05% NP40, 3 mM DTT, 1 mM PMSF, and a protease inhibitor cocktail; approximately 10 μg of total protein was used for protein gel blot analysis. Each age group contains two males and one female per time point. The specificity of BMAL1 and CLOCK antibodies was determined using synthetic Bmal1 and Clockb proteins expressed in the turquoise killifish. XM_015965280 for Bmal1 and XM_015971720 for Clockb were used as a template to synthesize control proteins. The synthetic killifish Bmal1 and Clockb proteins were obtained using a cell-free expression system after conjugation with a 6xHIS tag for purification (Gene to Protein Synthesis service, Bioneer). The synthetic proteins were detected with an anti-6xHis antibody (ab184607, Abcam, RRID:AB_2868537). Tubulin (T5168-2mL, Sigma, RRID:AB_477579) was used as a loading control, and BMAL1 (ab93806, Abcam, RRID:AB_10675117) and CLOCK (R1511-2, HUABIO, RRID:AB_2859538) antibodies were used to detect protein expressions and for immunoprecipitation. Co-immunoprecipitation was performed with BMAL1 or CLOCK antibody conjugated with IgG-conjugated agarose beads (20423, Pierce™ Protein A/G Plus Agarose, Thermo Scientific™).

Chromatin immunoprecipitation

Each biological group contained five brains from three males and two females. Chromatin immunoprecipitation was performed using an antibody against BMAL1 (ab93806, Abcam, RRID:AB_10675117), and GFP (ab290, Abcam, RRID:AB_303395) antibody was used as the control according to a previously described method (Haring et al., 2007). Chromatin shearing was performed using a sonicator (VCX 130, Vibra-Cell™ Ultrasonic Liquid Processors, Sonics & Materials, Inc.) under the following conditions: 2 s of sonication and 2 s of rest for 16 min. Two nanograms of input and ChiPed DNA were used for qPCR. The primers targeting the E-boxes of cry1 and per3 promoter regions are listed below.

| Name        | Forward primer | Reverse primer     |
|-------------|----------------|--------------------|
| cry1b-amp1  | AATAAAGAGGGGATGGGGGCGATG | TGAATTCAGTAACGTAATGGCC |
| cry1b-amp2  | AGCAGAAAACATTACTCCTTTCT | AAGGGTGAGGAGTAGACTAAAG |
| cry1b-amp3  | CTGTGGTTTGTATATCAGCATTACCAATTAC | CCAAGTCATATAATGGAAACTTGT |
| cry1b-amp4  | AAATCAGTAATCGGTAGCTGAC | CTCAAGGTAATAGTGCAGGACC |
| cry1b-amp5  | TATAGAGCTCCGACGGCTCA | AGGTCCGTAAGAAGTCC |
| per3-amp1   | CTTACGACTCCCTCCCCTC | AGAAGTTTCACAGTGAAGAC |
| per3-amp2   | GGTGACGCTGTGAATCTGAG | GCTGAAAATCTTCAAAGGGCTA |
| per3-amp3   | TGGCCGATATGATGGCAAGA | GTCATCTGGTATACCACCTG |
| per3-amp4   | TAAACATAGCAGGATACCATC | GTGTGACTGTACATAAAC |
| per3-amp5   | CAACGTGATTCCGCCATG | CACTTGTTGCTCCTTTGT |
Whole brain immunostaining

Two young and two old male brains were dissected carefully under a microscope and fixed in 4% PFA overnight. The whole brain was cleared with a tissue immune-staining kit (Binaree Immuno Staining™ Kit for Brain, BINAREE) following the manufacturer’s protocol. Cleared brains were stained using anti-BMAL1-Alexa647 (ITM0071-100M-647, GBiosciences, RRID:AB_2868539) or anti-CLOCK (R1511-2, HUABIO, RRID:AB_2868538)/goat anti-rabbit IgG Cross-Adsorbed, Alexa Fluor488 (A11008, Invitrogen, RRID:143165) antibodies. The nucleus was labeled with DAPI. Cleared and stained brains were embedded in 2% of low melting agarose in 2-mm capillaries (inner diameter). The embedded brain was tiled by 9-30 regions to cover the whole brain, and imaged with a 20× objective lens (W Plan-APOCHROMAT 20, Zeiss) under light sheet microscopy (Lightsheet Z.1, Zeiss). Tiled images were converted (Imaris File Converter x64.9.2, Bitplane) and merged (Imaris Stitcher 9.2.1, Bitplane) into one image for visualization.

Statistics

Statistical comparison between young and old of gene and protein expression for circadian core components was performed in Graph Pad PRISM (Prism 7 for Windows, version 7.03) by putting raw data in the data tables. Statistical analysis of circadian rhythmicity was performed JTK_CYCLE (Hughes et al., 2010), and Every raw data and output after JTK_CYCLE analysis were provided in Supplemental Table 1.
Supplemental References

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