Long-term monitoring of bioluminescence circadian rhythms of cells in a transgenic Arabidopsis mesophyll protoplast culture

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Abstract The circadian system of plants is based on the cell-autonomously oscillating circadian clock. In the plant body, these cellular clocks are associated with each other, but their basic and intrinsic properties are still largely unknown. Here we report a method that enables long-term monitoring of bioluminescence circadian rhythms of a protoplast culture in a complete synthetic medium. From the leaves of Arabidopsis transgenic plants carrying the luciferase gene under a clock-gene promoter, mesophyll protoplasts were isolated and their bioluminescence was automatically measured every 20 min for more than one week. Decreasing luminescence intensities were observed in protoplasts when they were cultured in a Murashige and Skoog-based medium and also in W5 solution. This decrease was dramatically improved by adding the phytohormones auxin and cytokinin to the MS-based medium; robust circadian rhythms were successfully monitored. Interestingly, the period lengths of bioluminescence circadian rhythms of protoplasts under constant conditions were larger than those of detached leaves, suggesting that the period lengths of mesophyll cells in leaves were modulated from their intrinsic properties by the influence of other tissues/cells. The entrainability of protoplasts to light/dark signals was clearly demonstrated by using this monitoring system. By analyzing the circadian behavior of isolated protoplasts, the basic circadian system of plant cells may be better understood.

Key words: Arabidopsis thaliana, bioluminescence monitoring, circadian rhythm, entrainment, mesophyll protoplasts.

Most organisms possess a circadian rhythm that helps them adapt to environmental day-night cycles. In plants, which depend considerably on sunlight, various physiological phenomena exhibit daily rhythms that are modulated by the circadian clock (Greenham and McClung 2015). Plant circadian clocks are based on transcription-translation feedback loops of a number of clock genes (Nagel and Kay 2012). Arabidopsis CCA1 is one of these clock genes, and its gene expression oscillates with a peak around dawn. It has rhythmic expression with a period of ca. 24 h under constant conditions. Most clock components encoded by clock genes are transcription factors, and the clock cell-autonomously oscillates (Muranaka and Oyama 2018). Indeed, it was reported that cultured cells of Arabidopsis that were transformed with clock-controlled luminescence reporter genes showed robust bioluminescence circadian rhythms similar to those of intact plants (Kim et al. 2003; Nakamichi et al. 2004). By measuring rhythmicity in individual leaf cells in transgenic Arabidopsis plants with fluorescence-tagged CCA1, cell-specific differences in clock properties were demonstrated (Yakir et al. 2011). Furthermore, by developing a single-cell bioluminescence imaging system for monitoring cellular gene expression in duckweed, the circadian behavior of individual cells in an intact plant was quantitatively analyzed (Muranaka and Oyama 2016, 2018; Muranaka et al. 2013; Okada et al. 2017). Individual cellular circadian clocks oscillate with their own frequencies and respond independently to external light/dark signals. Despite the heterogeneous and unstable features of cellular clocks in the duckweed plant, partial synchronization between neighboring cells was suggested. In Arabidopsis as well, cell-to-cell interactions for synchronization of cellular clocks were suggested (Fukuda et al. 2007; Wenden et al. 2012). The circadian rhythms of vascular tissues were also reported to be robust and dominant over those of mesophyll tissues in Arabidopsis (Endo et al. 2014). Studies using Arabidopsis at an organ level suggested that the circadian clock in the shoot apex was robust and dominant over other clocks in leaves and roots (Takahashi et al. 2015).

Abbreviations: CCA1, CIRCADIAN CLOCK ASSOCIATED1; DD, constant dark; FBS, fetal bovine serum; LL, constant light; LUC, luciferase gene; MSP medium, MS-based protoplast medium; PH, phytohormones.

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Individual circadian clocks in the plant body are thus likely to be influenced in their coordinated behavior by signals of various hierarchies. Information regarding the basic features of cellular circadian rhythms without those influences should be required for better understanding of the plant circadian systems. Protoplasts isolated from plant tissues can be free from contact with other cells in the culture medium; cell-to-cell communications that influence their circadian clocks are thus unlikely. Protoplasts from *Arabidopsis* leaves transfected with luminescence reporters have been reported to exhibit bioluminescence circadian rhythms (Kim and Somers 2010). This system can be used for a convenient assay of the molecular functions of clock-related genes. Here we report that bioluminescence monitoring of protoplast-derived cells of *CCA1::LUC* transgenic *Arabidopsis* leaves can be performed long-term by improving the medium conditions. Using this monitoring system, entrainability to light/dark cycles and stable circadian rhythmicity in isolated cells was demonstrated.

We used the previously reported *CCA1::LUC* transgenic *Arabidopsis thaliana* (Col-0) (Nakamichi et al. 2004, 2005). Mesophyll protoplasts were isolated from leaves of 3–4-week-old plants aseptically grown in 0.5 × MS (Murashige and Skoog) medium (0.8% agar) with 1% sucrose under constant light conditions (32 μmol m⁻² s⁻¹ at 22°C) (Figure 1A, B). Before protoplast isolation, these plants were exposed to two 12-h dark periods to entrain (synchronize) circadian rhythms. Protoplast isolation was performed as described by Yoo et al. (2007) with minor modifications. Fifty leaves were detached from plants 3–4 h after lights-on (Figure 1A). Enzyme digestion with 20 ml enzyme solution was performed in dim light with gentle shaking for 3 h at 25°C. Luciferin was included in the enzyme solution (400 mM mannitol, 20 mM KCl, 20 mM MES-KOH (pH 5.8), 10 mM CaCl₂, 1% Cellulase R10, 0.54% Macerozyme R10, 0.1% BSA, 0.1 μM luciferin). The released protoplasts were filtered through a sterile 70 μm nylon mesh and harvested by centrifugation at 500 g for 2 min at 22°C. The protoplast pellet was resuspended and washed twice with 45 ml W5 solution [154 mM NaCl, 125 mM CaCl₂, 5 mM KCl, 1.5 mM MES-KOH (pH 5.6)]. The protoplast pellet was then resuspended with ca. 4 ml W5 solution. Cells in a 1-ml aliquot were harvested by centrifugation at 100 g for 3 min at 22°C, and the pellet was resuspended with 4 ml culture medium with 0.1 μM luciferin. The cell suspension was then transferred into a 35-mm dish for bioluminescence monitoring. Removal of undigested tissues was checked with a microscope. Protoplasts sank to the bottom and they exhibited more than 100 times less bioluminescence density than did leaf cells (Figure 1C). Luminescence intensities of protoplasts changed similarly to the leaf of the *CCA1::LUC* transgenic plant, implying that the circadian system of those protoplasts was robust. To analyze the circadian rhythms of protoplast-derived cells in detail, it was important to monitor their bioluminescence traces for a long duration. The major growth regulators, auxins and cytokinins, are known to be essential for sustained protoplast growth in plants (Davey et al. 2005). We tested an MS-based protoplast medium with or without phytohormones (MSP + PH, MSP). The MSP medium contains 1 × MS salts, 1% sucrose, 400 mM mannitol, 2 mg/L glycine, 100 mg/L myo-inositol, 3 mg/L thiamin hydrochloride, 5 mg/L nicotinic acid, 0.5 mg/L pyridoxine hydrochloride, KOH (pH 5.6). Phytohormones [1 μM NAA (1-Naphthaleneacetic Acid), 0.2 μM kinetin] were added to the MSP medium to make the MSP + PH medium (Fukazawa et al. 2014; Satoh et al. 2004).

We first monitored bioluminescence of protoplast-derived cells in the three media (MSP + PH, MSP, WS)
under constant light. Hereafter, we use “protoplast-derived cells” in our experiments for the detection of bioluminescence rhythms because protoplasts tend to begin cell-wall regeneration within 48 h after protoplast isolation (Sheahan et al. 2004; Tiew et al. 2015). Bioluminescence monitoring was performed as described previously (Muranaka et al. 2015). The bioluminescence dish-monitoring system used photomultiplier tubes for bioluminescence detection, and each dish was subjected to 30 s measurement every 20 min. Bioluminescence circadian rhythms were observed in all medium conditions under constant light and constant dark (Figure 2A, B). However, luminescence intensities in MSP medium and W5 solution gradually decreased and their rhythms were damped. Under both conditions, the period lengths of bioluminescence rhythms in MSP+PH were slightly larger than those in MSP medium and W5 solution (Figure 3). The shorter period lengths in these media might be due to the severe damping of rhythms. Meanwhile, the period lengths of protoplast-derived cells in any media were larger than those of detached leaves, especially under constant light (Figure 3). Since sustainability in luminescence intensities and its rhythms of protoplast-derived cells in MSP+PH was shown, we next examined the rhythmicity under prolonged constant dark conditions. We tested protoplast-derived cells at various densities (1×10^5–6×10^5 cells in 4 ml MSP+PH medium). At all densities, luminescence intensities were maintained for 11 days with gradual increase for ca. 5 days and gradual decrease afterwards (Figure 2C). At the three highest cell densities (2×10^5–6×10^5 cells), the circadian rhythms were maintained for

Figure 2. Bioluminescence circadian rhythms of protoplast-derived cells of the CCA1::LUC transgenic Arabidopsis under constant light and constant dark. (A, B) Circadian rhythms of luciferase activities of protoplast-derived cells in MSP+PH medium (black line), MSP medium (green line), W5 solution (magenta line) under constant light (A) or constant dark (B). Experimental schemes are shown in Figure 1A. Time of panel A is set as hours in constant light including time before protoplast isolation. Time of panel B is set as hours in constant dark starting 12 h after the last lights-on. Data for the magenta line in (B) are plotted with scale on right. (C) Long-term monitoring of bioluminescence circadian rhythms under constant dark. Circadian rhythms of protoplast-derived cells [1×10^5 cells/4 ml (light blue line), 2×10^5 cells/4 ml (black line), 4×10^5 cells/4 ml (blue line), 6×10^5 cells/4 ml (purple line)] in MSP+PH medium and a detached leaf of the CCA1::LUC transgenic Arabidopsis on NF medium (red line). The composition of NF medium was described previously (Muranaka et al. 2015).

Figure 3. Period lengths of bioluminescence circadian rhythms under constant light and constant dark in various medium conditions. Circadian rhythms of protoplast-derived cells in MSP+PH medium (n=9 in LL; n=15 in DD), MSP medium (n=9 in LL; n=18 in DD), W5 solution (n=9 in LL; n=15 in DD), and leaves on NF (n=12 in LL; n=12 in DD) were monitored as described in Figure 2. The period of each rhythm was estimated by FFT-NLLS, a multicomponent cosine fit (Zielinski et al. 2014). Luminescence data in the range between 24 h and 96 h in constant conditions were used for the analysis. Mean values with error bars (SD) are represented in the bar graph.
11 days with gradual damping that was also observed in the bioluminescence rhythm of a detached leaf. At the lowest cell density \((1 \times 10^5)\) cells, the circadian rhythm was maintained for 6 days with gradual damping and then showed severe damping, suggesting that higher cell densities were required for long-term monitoring of bioluminescence rhythms of protoplast-derived cells.

Protoplast-derived cells were only analyzed for their circadian rhythms under free-running conditions in previous studies (Hansen and Ooijen 2016; Kim and Somers 2010; Takahashi et al. 2015; Wang et al. 2013). We then tested the entrainability of the circadian rhythm in protoplast-derived cells. After protoplast isolation, cells were treated with two 12-h dark periods in phase with the light/dark cycles before protoplast isolation (Figure 4A, filled circles). The first peak occurred 2.4 h after lights-on and the next one occurred at 27.7 h under constant light. Those cells that were treated with a 12-h dark period in antiphase with the light/dark cycles showed a bioluminescence rhythm (open circles) in antiphase with that of the filled circles (Figure 4A). This indicated that individual protoplast-derived cells were capable of entraining their circadian rhythms to the 12-h dark period through light/dark responses. Without the entrainment of the dark period, the amplitude of the bioluminescence rhythm was lower than that in entrained cells, suggesting that the circadian rhythms of individual protoplasts became synchronous due to the dark entrainment (Figure 4A, crosses). Detached leaves that tended to show damped rhythms under constant light were entrained to the 12-h dark period similarly to protoplast-derived cells, suggesting that entrainment behaviors in the dark treatment were similar between them (Figure 4B). This also implied that mesophyll cells in leaves might cell-autonomously respond to the light/dark signals to entrain their circadian rhythms.

In this study, we successfully monitored the bioluminescence rhythms of protoplast-derived cells of \(CCA1::LUC\) transgenic \(Arabidopsis\) leaves. Bioluminescence rhythms of those cells were sustained for more than 10 days by improving medium conditions. The monitoring of bioluminescence rhythms of \(Arabidopsis\) protoplasts was previously reported by Kim and Somers (2010). A circadian reporter gene was transiently transfected into mesophyll protoplasts and bioluminescence of those cells in W5 solution supplemented with 5% fetal bovine serum (FBS) was monitored under constant conditions. Adding serum into W5 solution was shown to be effective in the prolonged measurement of protoplast bioluminescence though the mechanism of action was unknown. In our monitoring procedure, by using protoplasts of \(Arabidopsis\) transformants, bioluminescence monitoring is begun just after the protoplast isolation. The quick and simple steps for preparation of luminescent protoplasts is likely to guarantee the uniformity and intactness of cells that is required for reproducible results in long-term monitoring. Adding an auxin and a cytokinin into MS-based medium is critical for the prolonged measurement of protoplast bioluminescence. These phytohormones may partially induce dedifferentiation of protoplast-derived cells and lead to the survival of these cells for a long time. The MS-based medium containing the phytohormones is a complete synthetic medium, whereas the W5 solution supplemented with 5% FBS is a semisynthetic medium that enables long-term bioluminescence monitoring of protoplast-derived cells. The complete synthetic medium can be used for experiments to assess chemicals that affect cellular circadian rhythms. Circadian behavior of protoplast-derived cells in any culture media showed a tendency
for period lengths larger than those of detached leaves (Figure 3). This suggested that period lengths of individual mesophyll cells were modulated by other tissues and/or each other in leaves, and protoplast-derived cells exhibited the intrinsic period lengths. By analyzing the circadian behavior of isolated protoplast-derived cells in a culture medium, the basic circadian system, without influence from other tissues/cells, may be understood.

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