6-Year Periodicity and Variable Synchronicity in a Mass-Flowering Plant

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Abstract

Periodical organisms, such as bamboos and periodical cicadas, are very famous for their synchronous reproduction. In bamboos and other periodical plants, the synchronicity of mass-flowering and withering has been often reported indicating these species are monocarpic (semelparous) species. Therefore, synchronicity and periodicity are often suspected to be fairly tightly coupled traits in these periodical plants. We investigate the periodicity and synchronicity of Strobilanthes flexicaulis, and a closely related species S. tashiroi on Okinawa Island, Japan. The genus Strobilanthes is known for several periodical species. Based on 32-year observational data, we confirmed that S. flexicaulis is 6-year periodical mass-flowering monocarpic plant. All the flowering plants had died after flowering. In contrast, we found that S. tashiroi is a polyarpic perennial with no mass-flowering from three-year individual tracking. We also surveyed six local populations of S. flexicaulis and found variation in the synchronicity from four highly synchronized populations (>98% of plants flowering in the mass year) to two less synchronized one with 11–47% of plants flowering before and after the mass year. This result might imply that synchrony may be selected for when periodicity is established in monocarpic species. We found the selective advantages for mass-flowering in pollinator activities and predator satiation. The current results suggest that the periodical S. flexicaulis might have evolved periodicity from a non-periodical close relative. The current report should become a key finding for understanding the evolution of periodical plants.

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Introduction

Periodical organisms are very famous for their mass and synchronous reproduction, e.g., bamboos [1–3], periodical cicadas [4–6]. The characteristic features of these organisms are the periodicity and synchronicity of reproductive events, e.g., periodical mass-flowering in bamboos [1–3]. Even though periodicity and synchronicity should be considered essentially two separate phenotypic traits, periodical organisms are usually associated with both periodicity and synchronicity [5,6]. Therefore, it is particularly important to investigate the evolutionary backgrounds of periodicity and synchronicity in these periodical organisms. Specifically, in periodical mass-flowering monocarpic (semelparous) plants, synchronous flowering is the key factor for pollination and seed predation avoidance [1,7,8].

However, it is highly difficult, if not impossible, to investigate the relationship between periodicity and synchronicity in many periodical plants, because most these plants have extremely long life cycles (e.g., 15–120 years in bamboos). Based on the extensive studies, the confirmation of periodicity in periodical cicadas has been widely accepted [4–6]. However, the confirmation of periodicity in bamboos is still highly difficult [1]. It is also further difficult to evaluate whether these organisms are perfectly synchronized or not even from extensive field surveys. In bamboo species, particularly, the synchronicity of individual plants is almost impossible because we cannot identify the vegetative grown genets from other individuals [2].

Instead of studying these long-cycle periodical organisms, we here investigated a periodical plant with relatively shorter life cycles (much less than 10 years). In the genus Strobilanthes (Acanthaceae), many species are reported to have periodicity with shorter life cycles, i.e., mass-flowering with various synchronous cycles (3–16 years), while other species flower every year [1,9,10]. We investigated the periodicity and synchronicity in supposedly periodical S. flexicaulis and its closely related S. tashiroi, both occurring on Okinawa Island, in the Ryukyu Islands, Japan. We also investigate the pollinator activities and predator satiation in S. flexicaulis to examine the selective advantages of mass-flowering.

Results

Field observations were carried out at Mt. Katsuu and Mt. Yae in the Motobu Peninsula, Okinawa Island from 1980. Mass-flowering of Strobilanthes flexicaulis was recorded at a six-year cycle...
in 1980, 1986, 1992, 1998, 2004, and 2010, while the rest 12 years were either few flowers or no flowers (Fig. 1). The seedling experiments starting in 1990, 2004 and 2005 showed that 17 out of 20 seedlings flowered (and withered) at six years, and the rest three plants at seven years (Table 1). Therefore, the interval of periodic mass-flowering had been estimated six years.

Detailed examinations of the life history for *S. flexicaulis* were performed from 2006 to 2011 (Table 2). We surveyed all flowering individuals in populations at Mt. Iyu, Mt. Oppa, and Awa. In all six populations, the number of flowering individuals was highest in 2010, indicating mass-flowering. Only few flowering plants are seen in 2009 or/and 2011 in four out of six populations (Mt. Iyu, Mt. Katsuu, Mt. Yae and Mt. Oppa). The populations at Mt. Yae and Mt. Katsuu show high synchronicity, more than 99% individuals flowered in 2010 with only a few one-year displacement (Fig. 1). Mt. Nago is significantly lower than that of Mt. Katsuu and Mt. Yae \((\chi^2 = 234.46, \text{ df} = 2, P<0.01; \text{Text S1, Table S2})\). Furthermore, in Awa population, the number of flowering plants in 2011 (31 individuals) is about the same with that in 2010 (34 individuals), and 25 individuals are still not flowering at 2011 that are expected to flower next year if survived. The chi-square test shows that the synchronicity of Awa is significantly lower than that of Mt. Katsuu and Mt. Yae \((\chi^2 = 927.58, \text{ df} = 2, P<0.01; \text{Text S1, Table S3})\). Thus the flowering years are spread almost three consecutive years at Mt. Nago and Awa, showing low levels of synchronicity.

In the total of six populations, we keep track of 241 plants by individual labeling (Table S4). Out of the 241 individuals, 237 died after flowering (Table 2). The remaining four individuals, in Mt. Yae population, had flowered partially on some branches. These branches had withered after flowering. These all four plants then had flowered in the following year and died after. Thus the death after flowering is a stable phenotypic character in *S. flexicaulis* even at the level of branches.

Comparing with *S. flexicaulis*, mass-flowering has never been observed in *S. tashiroi*. In the total of three populations, we labeled 108 individuals of *S. tashiroi* (Table S4). In 2010, 76 labeled individuals were alive and the other 32 individuals were dead (Fig. 2). Out of the 76 individuals, 31 flowered again. In 2011, 36 individuals survived and the other 40 individuals died. Six individuals bloomed for three consecutive years.

We also compare pollinator activities with *S. flexicaulis* and *S. tashiroi* between the mass-flowering year and the previous/following years (Table 3). Even though these species flowered in the winter season, in the mass-flowering year, three insect pollinators are seen (1.71 individuals/hour): humming-bird hawk moths (*Macroglossum stellatarum*, 0.46 individuals/hour); honeybees (*Apis mellifera*, 1.18 individuals/hour); and butterflies (*Hypaphis alucionis laochroma*, 0.07 individuals/hour). In contrast, in the year before/after mass-flowering year, the same humming-bird hawk moth was observed to make a single visit during the total observation of 13.25 hours (0.09 individuals/hour). Thus the level of insect pollinations is extremely low.

In order to evaluate the effects of predator satiation on the selective advantages of mass-flowering, we compared the percentage of fruits predated in *S. flexicaulis* and *S. tashiroi* between the mass-flowering year and the previous/following years (Fig. 3). Fruit predation by the larva of Pierophoridae sp. was observed in Mt. Yae population. Fruit predation rates in the mass-flowering year (2010) were significantly low in both *S. flexicaulis* \((\chi^2 = 13.5, \text{ df} = 1, P<0.01; \text{Text S3, Table S5})\) and *S. tashiroi* \((\chi^2 = 59.4, \text{ df} = 1, P<0.01; \text{Text S3, Table S6})\) with the chi-square test.

### Discussion

From 32-year data, it is confirmed that *Strobilanthes flexicaulis* is a periodical plant with a six-year mass-flowering cycle (Fig. 1). Detailed observations of six populations during 2008 and 2011 revealed that all plants (branches) had died after flowering (Table 2). These results suggested that *S. flexicaulis* was a periodical plant that maintained a periodical mass-flowering on Okinawa Island. However, the synchronicity of flowering is variable among local populations (Table 2). Mt. Iyu, Mt. Yae, Mt. Katsuu and Mt. Oppa show high levels of synchrony, while there is much lower synchrony at Nago and Awa. These results indicate that the synchronicity in *S. flexicaulis* is relatively variable depending on the locality. Thus *S. flexicaulis* is confirmed a monocarpic periodical plant as in bamboos, but with some variation in the levels of synchronization.

Unlike *S. flexicaulis*, *S. tashiroi* was confirmed to be a polycarpic perennial plant; each plant produces flowers at least for several years without withering (Fig. 2). No mass-flowering had been observed in *S. tashiroi*. Both *S. flexicaulis* and *S. tashiroi* belong to Parachampionella group, a small closely related taxonomic group consisting of four species [11,12]. These two species are genetically closely related [11]. This means that two distinctively different life histories, a polycarpic perennial plant and a monocarpic periodical plant, have been occurred in a small closely related taxonomic group. From the uniqueness and rarity of monocarpic periodical life history, we suspect that perennial life history is the ancestral trait for monocarpic periodical life history. Because of its shorter life cycle, and the variation of life history in its taxonomic group, *S. flexicaulis* could be a good model organism for the study of the evolution of monocarpic periodical life history, and periodical plants in general [1].

For the evolution of periodicity and synchrony we need to evaluate two traits: (1) an internal clock, set for six years in this case, and (2) a selection to keep populations (cohorts) synchronous.
The latter synchrony comes from lower fitness of out-of-step plants (from higher selective mortality of seeds produced in off-years, or from failure to pollinate if flowering in off-years). A rarer, second out-of-step cohort growing at the same site, which would be expected to arise easily enough, would be easily eliminated due to lower fitness in the same manner, if selection favors synchrony [1].

It is important to consider the evolutionary relationships between periodicity and synchrony [1]. Even though periodicity and synchrony are separate traits, mast- or mass-flowering should be achieved by simultaneous evolution of synchrony and periodicity to some extent [7,8]. Koenig et al. [13] showed some selective advantages of simultaneous periodicity and synchrony in iteroparous plant species. Such selective advantages should be much higher in semelparous species with strict masting [7], since the latter synchrony comes from lower fitness of out-of-step flowering individuals of {Strobilanthes flexicaulis} [1]. The latter synchrony comes from lower fitness of out-of-step plants (from higher selective mortality of seeds produced in off-years, or from failure to pollinate if flowering in off-years). A rarer, second out-of-step cohort growing at the same site, which would be expected to arise easily enough, would be easily eliminated due to lower fitness in the same manner, if selection favors synchrony [1].

Table 1. Seedling experiments.

| sowing year | individuals | 6th year | 7th year |
|-------------|-------------|----------|----------|
| 1998        | 9           | 9        | 0        |
| 2004        | 8           | 5        | 3        |
| 2005        | 3           | 3        | 0        |

The seedling experiments starting in 1998, 2004 and 2005 were performed in the greenhouse in Medicinal Botanical Garden, Faculty of Pharmaceutical Sciences, Setsunan University. The numbers of flowering individuals in each year were shown. doi:10.1371/journal.pone.0028140.t001

Table 2. Numbers of flowering individuals and survived individuals.

| population | flowering | 2008 | 2009 | 2010 | 2011 |
|------------|-----------|------|------|------|------|
| Mt. Iyu    | flowering | –    | 0    | 15   | 0    |
|            | survived/labeled | –    | –    | 0/15 | –    |
| Mt. Katsu  | flowering* | 0    | 11   | 1882 | 0    |
|            | survived/labeled | –    | 0/11 | 0/21 | –    |
| Mt. Yae    | flowering* | 0    | 9    | 1043 | 1    |
|            | survived/labeled | –    | 4/9  | 0/102 | –    |
| Mt. Oppa   | flowering | –    | 1    | >50  | 0    |
|            | survived/labeled | –    | –    | 0/18 | –    |
| Mt. Nago   | flowering* | 13   | 6    | 484  | 45   |
|            | survived/labeled | 0/13 | 0/6  | 0/30 | –    |
| Awa        | flowering* | –    | 0    | 34   | 31   |
|            | survived/labeled | –    | –    | 0/16 | –    |

The numbers of flowering individuals for {Strobilanthes tashiroi} were enumerated in six populations from 2008 to 2011. We surveyed all flowering individuals at Mt. Iyu, Mt. Oppa, and Awa. The number of survived individuals/labeled individuals is shown in the second row of each locality, where some flowering plants were labeled and their survivals were examined in the following year.

*The chi-square test shows that the synchronicity of Mt. Nago and Awa was significantly lower than Mt. Katsu and Mt. Yae (Text S1). doi:10.1371/journal.pone.0028140.t002

Table 3. Pollinator observations.

| year | time (hour) | Macroglossum corythis platyxaanthum | Apis mellifera | Byasa alcinous loochoona |
|------|-------------|------------------------------------|---------------|------------------------|
| 2009 | 10.75       | 0.09                               | 0.00          | 0.00                   |
| 2010 | 15.25       | 0.46                               | 1.18          | 0.07                   |
| 2011 | 2.5         | 0.00                               | 0.00          | 0.00                   |

Pollinator activities (observed individuals/hour) in Mt. Yae population before/during/after mass-flowering year are shown. doi:10.1371/journal.pone.0028140.t003

Seed predation might be lowered significantly with masting [1,7,17,21,22]. We compare the rates of fruit predation in mass-flowering monocarpic {S. flexicaulis} and related polycarpic {S. tashiroi} before/during/after mass-flowering year at the same locality (Fig. 3). In both {S. flexicaulis} and {S. tashiroi}, predation rates are low when mass-flowering, but much higher a year before and after. This suggests that mass-flowering at least reduces the level of predation considerably. Thus the selection favoring synchronicity may be promoted by pollinator activities [7,23] and predator avoidance [1,7].
The lack of synchrony in some populations may be basically errors by individual plants, e.g., deleterious mutations and recombination. These out-of-synchronous plants should be weeded out by natural selection later. The current variation in the level of synchrony may be a result of such stochastic factors in small populations and intermittent errors in plant cycle length, that have been appearing temporally before the selection weeds out.

**Materials and Methods**

**Ethics Statement**

We were permitted to collect and observe in the Okinawakaigan Quasi-National Park from the Governor of Okinawa, and the Natural Conservation Zone of Mt. Katsuu, Mt. Awa and Mt. Yae from the Okinawa Prefectural Board of Education.

**Species and morphology**

*Strobilanthes flexicaulis* is a subshrub distributed in the Ryukyu Islands, Japan, and Taiwan [12]. *Strobilanthes tashiroi* is a perennial herb endemic to the Ryukyu Islands [12]. These species are closely related and morphologically very similar [12,24]. Around the Motobu peninsula on Okinawa Island in the Ryukyu Islands, *S. flexicaulis* and *S. tashiroi* grow sympatrically. We identified these species based on the bract shape and the difference in length between longer (anterior) and shorter (posterior) pairs of stamens, which are didynamous (Text S2, Figure S1, Table S4).

**Study sites and observation of life histories**

We recorded mass-flowering years of *S. flexicaulis* and *S. tashiroi* around the Motobu Peninsula on Okinawa Island from 1980 to 2011. Quantitative research of mass-flowering species was performed in six populations (Table S4). All flowering individuals of *S. flexicaulis* were enumerated in the definite zones along paths at Mt. Katsuu, Mt. Yae, and Mt. Nago from 2008 to 2011, and at Awa, Mt. Oppa, and Mt. Iyu from 2009 to 2011. The difference in the synchronicity among populations was checked by the chi-square test (Text S1). Some representative flowering individuals of *S. flexicaulis* were labeled in winter from 2008 to 2010 (Table S4). After flowering, life or death of labeled individuals was checked in the following summer. We labeled some flowering individuals of *S. tashiroi* at Mt. Nishime, Mt. Yae and Mt. Nago in 2008 and checked whether these labeled individuals survived and flowered, survived but did not flower, or died in the two successive years, 2009–2010 (Table S4). The seedling experiments starting in 1998, 2004 and 2005 were performed in the greenhouse in Medicinal Botanical Garden, Faculty of Pharmaceutical Sciences, Setsunan University (Hirakata, Japan). A total of 20 individuals were sown and cultivated until flowering and dying.

**Pollinator activities and fruit predation**

Pollinator observations and surveys of fruit predation rates are performed in high synchronous Mt. Yae population from 2009 to 2011. Because pollinators were not seen in nighttime in preliminary observations, we observed pollinators in daytime in mostly *S. flexicaulis* in the mass-flowering year and mostly *S. tashiroi* in the previous and following years, as these are the available flowers in these years, respectively. To evaluate the fruit predation rates, we examined all the fruits on a plant before the capsules were exploded, at the end of March or the beginning of April from 2009 to 2011. The difference in the fruit predation rates in each species between the mass-flowering year and before/after was checked by the chi-square test (Text S3).
Supporting Information

Text S1 The chi-square test was performed to examine the difference in synchronicity among populations.

Text S2 *Strobilanthes flexicaulis* and *S. tashiroi* were identified by the combination of the bract shape and the difference in length between longer and shorter pairs of stamens.

Text S3 The chi-square test was performed to examine the difference in fruit predation rates between the mass-flowering year and off years.

Figure S1 The bract shape (length/width) and the difference in length between longer and shorter pairs are shown. Blue circles are *Strobilanthes flexicaulis* individuals and red squares are *S. tashiroi*. (TIF)

Table S1 The contingency table for the chi-square test to examine the difference in synchronicity between Mt. Yae and Mt. Katsu. (XLS)

Table S2 The contingency table for the chi-square test to examine the difference in synchronicity between Mt. Yae, Mt. Katsu and Mt. Nago. (XLS)

Table S3 The contingency table for the chi-square test to examine the difference in synchronicity between Mt. Yae, Mt. Katsu and Awa. (XLS)

References

1. Janzen DH (1976) Why bamboo wait so long to flower. Annu Rev Ecol Syst 7: 347–391.
2. Makita A (1998) The significance of the mode of clonal growth in the life history of bamboos. Plant Species Biol 13: 85–92.
3. Keeley JE, Bond WJ (1999) Mass flowering and semelparity in bamboos: the bamboo fire cycle hypothesis. Am Nat 154: 383–391.
4. Alexander RD, Moore TE (1962) The evolutionary relationships of 17-year and 13-year cicadas, and three new species. (Homoptera: Cicadidae, *Mégacanthus*). University of Michigan Museum of Zoology Miscellaneous Publication 121: 1–59.
5. Williams KS, Simon C (1995) The ecology, behavior, and evolution of periodical cicadas. Annu Rev Entomol 40: 269–295.
6. Yoshimura J (1997) The evolutionary origins of periodical cicadas during ice ages. Am Nat 149: 112–124.
7. Kelly D (1994) The evolutionary ecology of mast seeding. Trends Ecol Evol 9: 465–470.
8. Kelly D, Sork VI (2002) Mast Seeding in Perennial Plants: Why, How, Where? Annu Rev Ecol Syst 33: 427–447.
9. Daniel TF (2006) Synchronous flowering and monocarpity suggest pterilesian life history for neotropical *Simmoutheus chiapensis* (Acanthaceae). Proc Calif Acad Sci 57: 1011–1018.
10. Wood JRI (1994) Notes relating to the flora of Bhutan: XXIX. Acanthaceae, with special reference to *Strobilanthes*. Edinb J Bot 51: 175–273.
11. Sook DI, Hsieh CF, Murata J (2004) A new species of *Strobilanthes* (Acanthaceae) from Lan Yu (Orchid Island), Taiwan, with special reference to the flower structure. J Jap Bot 79: 145–154.
12. Wood JRI, Scotland RW (2003) The 2-lipped species of *Strobilanthes* (Acanthaceae). Kew Bull 58: 83–129.
13. Koenig WD, Kelly D, Sork VI, Duncan RP, Elkinton JS, et al. (2003) Dissecting components of population-level variation in seed production and the evolution of masting behavior. Oikos 102: 581–591.
14. Burd M, Read J, Samson GD, Jaffre T (2006) Age-size plasticity for reproduction in monocarpic plants. Ecology 87: 2755–2764.
15. Read J, Samson GD, Burd M, Jaffre T (2008) Mass flowering and parental death in the regeneration of *Cerberiopsis candela* (Apocynaceae), a long-lived monocarpic tree in New Caledonia. Am J Bot 95: 558–567.
16. Tsuura Z, Griffi ths ME, Gunton RM, Lawes MJ (2011) Predator satiation and recruitment in a mast fruiting monocarpic forest herb. Am J Bot 107: 379–387.
17. Forget P, Kitajima K, Foster RB (1999) Pre- and postdispersal seed predation in *Tachigali rosea* (Caesalpiniaceae): effects of timing of fruitying and variation among trees. J Trop Ecol 15: 61–81.
18. Kitajima K, Aupperger CK (1989) Seed and seedling ecology of a monocarpic tropical tree, *Tachigali rosea*. Ecology 70: 1102–1114.
19. Kelly D, Hart DE, Allen RB (2001) Evaluating the wind pollination benefits of mast seeding. Ecology 82: 117–126.
20. Herrera CM, Jordano P, Guittin J, Travest E (1998) Annual variability in seed production by woody plants and the mastng concept: reassessment of principles and relationship to pollination and seed dispersal. Am Nat 152: 576–594.
21. Janzen DH (1971) Seed Predation by Animals. Annu Rev Ecol Syst 2: 465–492.
22. Silvertown JW (1980) The evolutionary ecology of mast seeding in trees. Biod J Linn Soc 14: 235–250.
23. Westphal C, Steffen-Dewenter I, Tschamke T (2003) Mass flowering crops enhance pollinator densities at a landscape scale. Ecol Lett 6: 961–965.
24. Yamazaki T (1991) On *Strobilanthes flexicaulis* Hayata, *S. tashiroi* Hayata and *S. glandulifera* Hatusima. J Jap Bot 66: 303–306.

Table S4 Localities and geographical coordinates of observed or/sampling populations, where the extent of observed areas in each population and the sampling numbers of *Strobilanthes flexicaulis* and *S. tashiroi* in pure populations are shown.

Table S5 The contingency table for the chi-square test to examine the difference in fruit predation rate between the mass-flowering year and off years in *Strobilanthes flexicaulis*.

Table S6 The contingency table for the chi-square test to examine the difference in fruit predation rate between the mass-flowering year and off years in *Strobilanthes tashiroi*.

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Author Contributions

Conceived and designed the experiments: SK JY JM. Performed the experiments: SK HM JM. Analyzed the data: SK. Wrote the paper: SK JY JM.