Effects of time delay and space on herbivore dynamics: linking inducible defenses of plants to herbivore outbreak

Gui-Quan Sun1-4*, Su-Lan Wang2*, Qian Ren2*, Zhen Jin1 & Yong-Ping Wu3

Empirical results indicate that inducible defenses of plants have effects on herbivore populations. However, little is known about how inducible defenses of plants have influences on herbivore outbreak when space effect is considered. To reveal the relationship between inducible defenses and herbivore outbreak, we present a mathematical model to describe the interaction of them. It was found that time delay plays dual effects in the persistence of herbivore populations: (i) large value of time delay may be associated with small density of herbivore populations, and thus causes the populations to run a higher risk of extinction; (ii) moderate value of time delay is beneficial for maintaining herbivore density in a determined range which may promote the persistence of herbivore populations. Additionally, we revealed that interaction of time delay and space promotes the growth of average density of herbivore populations during their outbreak period which implied that time delay may drive the resilience of herbivore populations. Our findings highlight the close relationship between inducible defenses of plants and herbivore outbreak.

Herbivores are diverse, ranging in size from microscopic zooplankton to the largest of land vertebrates from the point of view of taxonomy and ecology1. By feeding on different plant parts or materials, herbivores can affect plant growth, transfers of nutrients to the soil surface, and habitat and resource conditions for other organisms. In many plants, particularly trees, damages or stresses by herbivores populations can result in changes in the chemical, physical or other aspects of leafs, which are called as “inducible defenses”2-6. Both theoretical and experimental studies have shown that inducible defences affect stability and persistence of herbivore populations7-16.

Empirical findings suggested that populations of many herbivorous insects exhibit outbreak, in which short-lived peaks of high density and lots of fallen leaves alternate with long periods of low density17-21. As a result, the mechanisms on herbivore outbreak have been attracted considerable attention by ecologists and other experts in the relevant research area. The existing work revealed that interactions with enemies22-24, physiological stress2,9,11,25, the case that herbivore populations's parents and grandparents experienced in preceding generations26, environmental forcing27 and limited resource28,29 may be the significant factors for herbivore outbreak. Although some previous works link inducible defenses to population-level effects on herbivore2,11,25, internal connections of inducible defenses and herbivore outbreak are far from being well understood. Especially, two main questions need to be well addressed: (1) Can inducible defenses of plants induce herbivore outbreak when space is considered? (2) How do...
inducible defenses affect the persistence of herbivore populations in different aspects during their outbreak period?

It is difficult to characterize the relationship between inducible defenses and herbivore outbreak empirically due to that long time series of the density of both plant and herbivore is needed. It may provide useful information by constructing mathematical models to explain the phenomenon observed in the real world. Edelstein-Keshet posed a model to show how changes in plant quality have influence on herbivore populations\(^3\). Clark and Harvell used dynamic-optimization models to estimate the relative fitness consequences of inducible versus constitutive defenses strategies and found that inducible defenses played a more important role\(^3\). Abbott and Dwyer showed that outbreaking insects may be induced by a food limitation in the herbivore and defoliation and intraspecific competition in the host plant\(^2\). Anderson et al. presented a mathematical model on herbivore competition mediated by inducible changes in plant quality and obtained several types of competition outcomes\(^2\). Most studies to date only consider the evolution in time\(^2\). Nevertheless, it has been observed in the literature that spatial effects on plant and herbivore had been generally overlooked despite its potential ecological reality and intrinsic theoretical interest. In our paper, we will investigate the plant-herbivore interactions with time delay (it arises between herbivore damage and deployment of inducible defenses) and spatial diffusion and aim to link inducible defences to herbivore outbreak.

Results

Since overall data of herbivore is not available, we may not find out the intrinsic mechanisms on herbivore outbreak empirically. Instead, we are aim to use a simple model to reflect the interactions between inducible defenses and herbivore populations (see Method section).

Our analysis is to link inducible defenses and herbivore outbreak by three steps. Firstly, we obtain the conditions on critical value of time delay for herbivore outbreak analytically. Secondly, we revealed dual effects of time delay on herbivore outbreak: on the one hand, large value of time delay brings about herbivore density to be zero which implied that time delay may be harmful to survival of herbivore; on the other hand, moderate value of time delay promotes the persistence of herbivore during the stage of herbivore outbreak. Finally, we display that joint forces of time delay and space boost the growth of average density of herbivore populations.

Critical value of time delay for herbivore outbreak. Based on mathematical analysis, one can find the critical value of time delay to ensure the outbreak of herbivore populations (see Method section). The smallest critical value of time delay has the following form:

\[
\tau_c = \frac{1}{w_1} \left[ \arccos \left( -w_1^2 + d_1 d_2 - (d_2 a_{11} + d_1 a_{22}) + a_{11} a_{22} \right) \right].
\]

To well see the effect of time delay on herbivore outbreak, critical value of time delay is shown as a function of diffusion coefficient of herbivore populations in Fig. 1. As seen from this figure that, when the moving speed of herbivore populations is small \((d_2 < 0.45)\), larger diffusion rate of herbivore requires larger value of time delay to ensure their outbreak; when the moving speed is large enough, critical value

![Figure 1. Critical value of time delay with respect to diffusion rate of herbivore populations.](image_url)
of time delay is a decreasing function of the diffusion rate. Biological speaking, there is a balance between time delay and spatial motion of herbivore populations in the mechanisms on herbivore outbreak. When diffusion rate of herbivore populations is small, herbivore populations will consume more resources as $d$ increases, which needs larger time delay to hold back their growth and thus herbivore will periodic outbreak. When diffusion rate of herbivore populations is large enough, the remained resources are limited which may induce negative growth of herbivore populations. In this case, smaller time delay can lead to herbivore outbreak.

In Fig. 2, we show the herbivore outbreak numerically for fixed parameters sets: $\alpha = 200$, $\beta = 1$, $\delta = 0.75$, $b = 5$, $\theta = 3$, $r = 1$, $K = 10$, $m = 0.01$, $d_1 = 0.01$ and $d_2 = 0.25$. Under these circumstances, one can find that $\tau_c \approx 4.7412$. Herbivore populations are considered as a function of space (in one-dimensional space) and time. In Fig. 2(A), the solutions are stable as $\tau = 3.2 < \tau_c$; while in Fig. 2(B), periodic solutions emerge as $\tau = 4.8 > \tau_c$. In other words, time delay induces the outbreak of herbivore populations.

In order to better show the outbreak of herbivore populations, time series are shown in Fig. 3. In Fig. 3(A), herbivore populations exhibit an oscillation behavior with decreased amplitude and converge to a constant state with $\tau < \tau_c$. This figure suggests that periodic outbreak of herbivore will not appear with $\tau < \tau_c$. However, when $\tau > \tau_c$, herbivore populations will outbreak with fixed period and amplitude as evolution time is long enough showed in Fig. 3(B).

**Dual effects of time delay on the persistence of herbivore.** Synchronization is a fundamental phenomenon arising in many biological contexts, which can be an important part of the function or malfunction of a biological system. We checked that during the period of herbivore outbreak, herbivore populations and inducible defenses exhibit synchronous phenomenon (cf. Fig. 4). However, this figure also indicates that, when time delay is too large, the minimum value of herbivore density will reach zero which may cause the herbivore populations to run a high risk of extinction which can be seen from Fig. 4(A). This phenomenon can be explained in two different directions: on the one hand, the presence of synchronization may decrease the global persistence; on the other hand, it was observed that herbivore can remain persistent when inducible defenses are small and thus it may go extinct with high density of inducible defenses. In this sense, time delay plays a negative role for the persistence of herbivore populations.
As seen from Figs 2–3, we know that herbivore populations have property of periodic solutions. One may ask whether the periodic solutions are stable or not. Based on stability analysis (see Method section), we obtained that for all the parameters sets to ensure the outbreak of herbivore, $\beta < 0$ which means the periodic solutions are stable. From biological point of view, time delay plays a positive role in herbivore persistence due to that it can keep the density of herbivore in a determined range from extinction.

The period of the periodic solutions has the following expression:

$$T = -\frac{1}{\mu} \left[ \text{Im} (C) + \mu \text{Im} (\lambda' (\tau_x)) \right],$$

with $\mu = -\frac{\text{Re} (C)}{\text{Re} [\lambda (\tau_x)]}$. Figure 5 shows the period of periodic solutions as a function of time delay. It shows that period is an increasing function of time delay. At the same time, the maximum value of herbivore density is becoming larger and the minimum value is becoming smaller as time delay increases.

**Combination of time delay and space promotes the growth of average density of herbivore populations.** In the parameters sets which ensure the emergence of herbivore outbreak, it is found that average density of herbivore populations increases as time delay increases which was shown in Fig. 6. We checked that if herbivore populations do not outbreak, i.e., value of time delay is smaller than the critical value, this phenomenon can not be observed. Meanwhile, the results can not be obtained if space is not included. Accordingly, we concluded that interaction of time delay and space promotes the growth of average density of herbivore and then drive more resilience for herbivore populations. The existing results revealed that spatial scaling laws\(^{37}\), multiple scale spatial patterns\(^{38,39}\) or insects populations\(^{40}\) may increase the robustness in some biological systems. Therefore, our results enrich the findings in ecosystem functioning.
Discussion

An extensive body of scientific research on inducible defenses of plant to herbivore populations demonstrated that inducible defenses may have great influences of dynamics of herbivore populations. However, the studies on how inducible defenses exactly affect herbivore populations when space is under

Figure 4. Synchrony of the herbivore and inducible defenses with $\tau = 8$ and the other parameters are as the same in Table 1. This figure also shows that herbivore populations suffer a higher likelihood of extinction if time delay is too large.

Figure 5. Period of periodic solutions with respect to time delay. As time delay increases, the period will increase with large amplitude.
consideration, especially on the effect of herbivore outbreak, is still limited. As a result, a simple model to describe the interaction of inducible defenses and herbivore populations is investigated. It was found that time delay arising from plant defenses response to herbivore attacks can lead to periodic outbreak of herbivore populations. Furthermore, time delay has dual functions on the persistence of herbivore populations: large time delay may result in the extinction of herbivore and moderate value of time delay enlarges the possibility of herbivore persistence by keeping the periodic solutions to be stable. This finding implies that inducible defense with different intensities resolves the paradox of enrichment in the spatial sense.

Vos et al. found that inducible defenses play different roles on herbivore populations. On the one hand they promote local stability and thus persistence and on the other hand they may reduce the likelihood of herbivore persistence. In our paper, we confirm the conclusions still hold when spatial effects are included. Meanwhile, we found that interaction of time delay and space may drive the resilience of herbivore populations on account of that time delay and space increase average density of herbivore populations during their outbreak period.

It should be noted that herbivore populations not only consume resources, but they are resources for other consumers. Consequently, they have much potential roles of connecting link between upper and down trophic chains in evolution process of the whole ecosystems. In this sense, it needs to be a balance in the control of herbivore populations and thus human beings can not blindly kill or protect herbivore populations.

**Method**

**Mathematical Model.** We give four main assumptions on our model: (1) To reflect delays in the deployment of inducible defenses, we assumed that induction changes at time \( t \) dependent on herbivore densities at \( t - \tau \) time steps previously; (2) Inducible defenses is dependent on herbivore density (in saturation form) and the level of already inducible defenses; (3) In the absence of induced changes in plant quality, herbivore populations grows logistically with intrinsic rate \( r \) and carrying capacity \( K \); (4) For some plants, their weeds can move in the space caused by environmental factors such as wind. Consequently, we consider that both inducible defenses and herbivore randomly move in the space with diffusion coefficients \( d_1 \) and \( d_2 \) respectively. Based on the above assumptions, we arrive at the following reaction diffusion equation:

\[
\begin{align*}
\frac{\partial I(\gamma, t)}{\partial t} &= [\alpha - \beta I(\gamma, t)] \frac{H^0(\gamma, t - \tau)}{b^0 + H^0(\gamma, t - \tau)} - \delta I(\gamma, t) + d_1 \Delta I(\gamma, t), \\
\frac{\partial H(\gamma, t)}{\partial t} &= rH(\gamma, t) \left( 1 - \frac{H(\gamma, t)}{K} \right) - mI(\gamma, t)H(\gamma, t) + d_2 \Delta H(\gamma, t),
\end{align*}
\]

where \( I(\gamma, t) \) and \( H(\gamma, t) \) represent inducible defenses and herbivore density in both space and time. \( \alpha \) is maximum per capita induced defenses, \( \beta \) is per unit reduction in the elicitation rate due to plant self-limitation, \( \delta \) is per-unit induction decay rate, \( m \) is per unit reduction in the growth rate of herbivore caused by induction of defenses, \( \gamma \) represents space, and \( \Delta = \partial^2/\partial x^2 \) is Laplacian operator in one-dimensional space. More details can be found in Table 1.
Analysis on Critical Value. Mathematical speaking, if a system undergoes hopf bifurcation, then it will exhibit periodic solutions. For system (3), if it has hopf bifurcation, the herbivore populations will outbreak. In this sense, we need to find the critical value for herbivore outbreak.

Denote \( E^* = (I^*, H^*) \) as the positive equilibria of system (3). We deduce the eigenpolynomial associated with wavenumber \( \kappa \):

\[
\lambda^2 + [(d_1 + d_2)\kappa^2 - (a_{11} + a_{22})]\lambda + d_2 d_2 e^\kappa^2 - (d_2 a_{22} + d_2 a_{11})\kappa^2 + a_{11} a_{22} - a_{12} a_{21} e^{-\lambda \tau} = 0, \tag{4}
\]

where \( a_{11} = \frac{\partial f}{\partial I}|_{E^*}, a_{12} = \frac{\partial f}{\partial H}|_{E^*}, a_{21} = \frac{\partial g}{\partial I}|_{E^*}, a_{22} = \frac{\partial g}{\partial H}|_{E^*} \) with \( f = (\alpha - \beta II) - \frac{H^2}{b^2 + H^2} - \delta I \) and \( g = rH (1 - \frac{H}{F}) - mIH \).

Setting \( \lambda = iw (w > 0) \) is a root of the eigenpolynomial (4) and separating the real and imaginary parts, one can have:

\[
\begin{aligned}
-w^2 + d_2 d_2 e^\kappa^2 - (d_2 a_{22} + d_2 a_{11})\kappa^2 + a_{11} a_{22} &= a_{12} a_{21} \cos w \tau, \\
[(d_1 + d_2)\kappa^2 - (a_{11} + a_{22})]w &= -a_{12} a_{21} \sin w \tau.
\end{aligned}
\tag{5}
\]

Then,

\[
w_\kappa = \sqrt{\frac{B_\kappa + \sqrt{B_\kappa^2 - 4C_\kappa}}{2}},
\]

where

\[
B_\kappa = - [(d_1 + d_2)\kappa^4 + 2k^2 (d_1 a_{11} + d_2 a_{22}) - (a_{11}^2 + a_{22}^2)],
\]

\[
C_\kappa = [d_2 d_2 e^\kappa^2 - (d_2 a_{22} + d_2 a_{11})\kappa^2 + a_{11} a_{22}] - a_{12} a_{21}. \]

The corresponding critical value of time delay is:

\[
\tau_\kappa = \frac{1}{w_\kappa} \left[ \arccos \left( \frac{-w_\kappa^2 + d_2 d_2 e^\kappa^2 - (d_2 a_{22} + d_2 a_{11})\kappa^2 + a_{12} a_{21}}{a_{12} a_{21}} \right) + 2j\pi \right], j = 0, 1, 2...
\]

By calculations, the transversality condition \( \text{sign} \left\{ \text{Re} \left( \frac{d_1}{\tau} \right) \right\} \mid_{\tau = \tau_j} > 0 \) holds. As a result, system (3) undergoes a spatial Hopf bifurcation at the equilibrium \( E = (I, H) \) when \( \tau = \tau_j \) (\( j = 1, 2, \ldots \)) and periodic solutions emerge in system (3) when \( \tau > \tau_j \).

Stability of Periodic Solutions for Herbivore Populations. We can use normal form and the center manifold theory to investigate the stability of the bifurcated periodic solutions. In order to determine the properties of Hopf bifurcating periodic solutions at the critical value, we can compute the following values:

\[
\beta = 2 \text{Re} [C],
\]

with \( C = \frac{i}{2w_\kappa} \left( g_{20} g_{11} - 2 |g_{11}|^2 - \frac{1}{3} |g_{20}|^2 \right) + \frac{1}{2} g_{21} \). Since the expressions of \( g_{20}, g_{11}, g_{20}, \) and \( g_{21} \) are complex, we omit them here. The bifurcating periodic solutions are stable (unstable) if \( \beta < 0 \) (\( \beta > 0 \)).

| Parameter | Value | Comments | References |
|-----------|-------|----------|------------|
| \( \alpha \) | 200 | maximum induction rate per herbivore | 47 |
| \( \beta \) | 1 | per-unit reduction of induction rate by self-limitation | 32 |
| \( \delta \) | 0.75 | per-unit induction decay rate | 32 |
| \( b \) | 5 | half-maximum for herbivore effectiveness of damage | 47 |
| \( \theta \) | 3 | herbivore damage effectiveness shape tuning parameter | 47 |
| \( r \) | 1 | intrinsic rate of herbivore populations growth | 32 |
| \( K \) | 10 | herbivore carrying capacity | 32 |
| \( m \) | 0.01 | mortality rate by induction | 32 |
| \( d_1 \) | 0.01 | diffusion rate of plant populations | 48 |
| \( d_2 \) | 0.01 \( - 1 \) | diffusion rate of herbivore populations | 49 |

Table 1. Summary of the parameters used in system (3).
References

1. Huntly, N. Herbivores and the dynamics of communities and ecosystems. *Annu. Rev. Ecol. Syst.* **22**, 477–503 (1991).

2. Edelstein-Keshet, L. & Rausher, M. D. The effects of inducible plant defenses on herbivore populations. 1. Mobile herbivores in continuous time. *Am. Nat.* **133**, 787–810 (1989).

3. Fowler, S. V. & Lawton, J. H. Rapidly Induced Defenses and Talking Trees: The Devil’s Advocate Position. *Am. Nat.* **126**, 181–195 (1985).

4. Karban, R. & Baldwin, I. T. Induced Responses to Herbivory. University of Chicago Press, Chicago (1997).

5. Tollrian, R. & Harvell, C. D. The Ecology and Evolution of Inducible Defenses. Princeton University Press, Princeton, NJ (1999).

6. Pavia, H. & Tochgunilla B. Inducible chemical resistance to herbivory in the brown seaweed ascolyphodium nodosum. *Ecology* **81**, 3212–3225 (2000).

7. Stastny, J. V. D., Vos, M. & Mooij, W. M. Linking herbivore-induced defences to population dynamics. *Freshwater Biol.* **51**, 424–434 (2006).

8. Lin, H., Kogan, M. & Fischer, D. Induced Resistance in Soybean to the Mexican Bean Beetle (Coleoptera: Coccinellidae): Comparisons of Inducing Factors. *Environ. Entomol.* **12**, 1852–1857 (1990).

9. Lundberg, S., Jarenko, J. & Nilsson, P. Herbivory, inducible defence and population oscillations: a preliminary theoretical analysis. *Oikos* **71**, 537–539 (1994).

10. Abrams, P. A. & Walters, C. J. Invulnerable prey and the paradox of enrichment. *Ecology* **77**, 1125–1133 (1996).

11. Underwood, N. The influence of plant and herbivore characteristics on the interaction between induced resistance and herbivore population dynamics. *Am. Nat.* **133**, 282–294 (1989).

12. Ramos-Estévez, R. Population dynamics of prey exhibiting inducible defences: the role of associated costs and density-dependence. *Theor. Popul. Biol.* **64**, 221–231 (2003).

13. Vos, M., Kooi, B. W., DeAngelis, D. L. & Mooij, W. M. Inducible defences and the paradox of enrichment. *Oikos* **105**, 471–480 (2004).

14. Peacock, S. D. & Werner, E. E. Predator effects on an assemblage of consumers through induced changes in consumer foraging behavior. *Ecology* **81**, 1988–2010 (2000).

15. Raimondi, P. T., Forde, S. E., Delph, L. E. & Lively, C. M. Processes structuring communities: evidence for trait-mediated indirect effects through induced polymorphisms. *Oikos* **91**, 353–361 (2000).

16. Turner, A. M., Bernt, R. J. & Roes, C. M. Chemical cues modify species interactions: the ecological consequences of predator avoidance by freshwater snails. *Oikos* **88**, 148–158 (2000).

17. Varley, G. C., Gradwell, G. R. & Hassell, M. P. Insect Population Ecology. Blackwell Scientific, Oxford, UK (1973).

18. Crawley, M.J. Herbivory: The Dynamics of Animal-Plant Interactions. University of California Press, Los Angeles, CA, USA (1983).

19. Berryman, A. A. The theory and classification of outbreaks. In: Insect Outbreaks (eds Barbosa, P. & Schultz, J. C.), pp. 3–30. Academic Press, New York (1987).

20. Peacock, S. D. & Werner, E. E. Predator effects on an assemblage of consumers through induced changes in consumer foraging behavior. *Ecology* **81**, 1988–2010 (2000).

21. Edelstein-Keshet, L. & Rausher, M. D. The effects of inducible plant defenses on herbivore populations. 1. Mobile herbivores in continuous time. *Am. Nat.* **133**, 787–810 (1989).

22. Abbott, K. C. & Dwyer, G. Food limitation and insect outbreaks: complex dynamics in plant-herbivore models. *Oecologia* **86**, 594–602 (2005).

23. Ginzburg, L. R. & Taneyhill, D. E. Population cycles of forest lepidoptera: a maternal effect hypothesis. *J. Anim. Ecol.* **63**, 79–92 (1994).

24. Hunter, A. F. Traits that distinguish outbreaking and nonoutbreaking macrolepidoptera feeding on northern hardwood trees. *Oikos* **60**, 275–282 (1991).

25. Abbott, K. C. & Dwyer, G. Food limitation and insect outbreaks: complex dynamics in plant-herbivore models. *J. Anim. Ecol.* **76**, 1004–1014 (2007).

26. Abbott, K. C., Morris, W. F. & Gross, K. Simultaneous effects of food limitation and inducible resistance on herbivore population dynamics. *Theor. Popul. Biol.* **73**, 63–78 (2008).

27. Ritchie, M. E. & Oliff, H. Spatial scaling laws yield a synthetic theory of biodiversity. *Nature* **400**, 557–560 (1999).

28. Pringle, R. M., Doak, D. F., Brody, A. K., Jacque, R. & Palmer, T. M. Spatial pattern enhances ecosystem functioning in an african savanna. *PLoS Biol.* **8**, e1000377 (2010).

29. Liu, Q.-X. & et al. Pattern formation at multiple spatial scales drives the resilience of mussel bed ecosystems. *Nat. Commun.* **5**, 5234 (2014).

30. Bonachela, J. A., Pringle, R. M., Sheffer, E. et al. Termite mounds can increase the robustness of dryland ecosystems to climatic change. *Science* **347**, 651–655 (2015).

31. Stiling, P. & Rossi, A. M. Complex effects of genotype and environment on insect herbivores and their enemies. *Ecology* **77**, 2212–2218 (1996).

32. Underwood, N. & Rausher, M. D. The effects of host-plant genotype on herbivore population dynamics. *Ecology* **81**, 1565–1576 (2000).

33. Whitham, T. G., Bailey, J. K., Schweitzer, J. A. et al. A framework for community and ecosystem genetics: from genes to ecosystems. *Nat. Rev. Genet.* **7**, 510–523 (2006).

34. Abdala-Roberts, L. & Mooney, K. A. Environmental and plant genetic effects on tri-trophic interactions. *Oikos* **122**, 1157–1166 (2013).

35. Schmitz, O. J. Herbivory from Individuals to Ecosystems. *Annu. Rev. Ecol. Evol. Syst.* **39**, 133–152 (2008).

36. Wu, J. Theory and Applications of partial functional differential equations. Springer-Verlag, New York (1996).
47. Sun, G.-Q., Chakrabort, A., Liu, Q.-X., Jin, Z., Anderson, K. E. & Li, B.-L. Influence of time delay and nonlinear diffusion on herbivore outbreak. *Commun. Nonlinear Sci. Numer. Simulat.* **19**, 1507–1518 (2014).

48. Rietkerk, M., Boerlijst, M. C., van, L. F., HilleRisLambers, R., van, d.K.J., Kumar, L., Prins, H.H.T. & de, R.A.M. Self-organization of vegetation in arid ecosystems. *Am. Nat.* **160**, 524–530 (2002).

49. Morris, W. F. & Dwyer, G. Population consequences of constitutive and inducible plant resistance: herbivore spatial spread. *Am. Nat.* **149**, 1071–1090 (1997).

Acknowledgments
This work is supported by the National Natural Science Foundation of China under Grants 11331009, 11171314 and 11301490, and International Exchange Program of Postdoctor in Fudan University.

Author Contributions
G.S., Y.W. and Z.J. designed the study, carried out the analysis and contributed to writing the paper, G.S., S.W. and Q.R. performed numerical simulations.

Additional Information
Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Sun, G.-Q. et al. Effects of time delay and space on herbivore dynamics: linking inducible defenses of plants to herbivore outbreak. *Sci. Rep.* **5**, 11246; doi: 10.1038/srep11246 (2015).

This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/
Erratum: Effects of time delay and space on herbivore dynamics: linking inducible defenses of plants to herbivore outbreak

Gui-Quan Sun, Su-Lan Wang, Qian Ren, Zhen Jin & Yong-Ping Wu

Scientific Reports 5:11246; doi: 10.1038/srep11246; published online 18 June 2015; updated on 03 August 2015

In the original version of this Article, the affiliations for Zhen Jin and Yong-Ping Wu were incorrectly listed as ‘College of Physics Science and Technology, Yangzhou University, Yangzhou, Jiangsu Province, 225002, P.R. China’ and ‘Complex Systems Research Center, Shanxi University, Taiyuan, Shanxi 030006, P.R. China’, respectively. The correct affiliations are listed below.

Zhen Jin
Complex Systems Research Center, Shanxi University, Taiyuan, Shanxi 030006, P.R. China

Yong-Ping Wu
College of Physics Science and Technology, Yangzhou University, Yangzhou, Jiangsu Province, 225002, P.R. China

These errors have now been corrected in both the HTML and PDF versions of the Article.