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## Testing the Terminal Investment Hypothesis in California Oaks


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# Testing the Terminal Investment Hypothesis in California Oaks

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Submitted September 6, 2016; Accepted December 7, 2016; Electronically published March 1, 2017

Online enhancements: appendix. Dryad data: <http://dx.doi.org/10.5061/dryad.51ft1>.

**ABSTRACT:** The terminal investment hypothesis—which proposes that reproductive investment should increase with age-related declines in reproductive value—has garnered support in a range of animal species but has not been previously examined in long-lived plants, such as trees. We tested this hypothesis by comparing relative acorn production and radial growth among 1,000+ mature individuals of eight species of California oaks (genus *Quercus*) followed for up to 37 years, during which time 70 trees died apparently natural deaths. We found no significant differences in the radial growth, acorn production, or index of reproductive effort, taking into consideration both growth and reproduction among dying trees relative to either conspecific trees at the same site that did not die or growth and reproduction from earlier years for the focal trees that did eventually die. Furthermore, we found no consistent trade-off between growth and reproduction among trees that died, nor did dying trees significantly alter their relative investment in reproduction even as they underwent physical decline. Trees approaching the end of their lives are often in poor physical condition but do not appear to differentially invest more of their diminished resources in reproduction compared with healthy trees.

**Keywords:** demography, life history, oaks, senescence, terminal investment.

## Introduction

Among the many contributions to evolutionary biology made by the late G. C. Williams is the hypothesis that individuals should shift more of their resources into reproduction as they approach the end of their lives (Williams 1966). Known as the terminal investment hypothesis (Clutton-Brock 1984), it assumes that there are trade-offs between reproduction, growth, and survivorship and that the probability of survival decreases with age—that is, that organisms senesce. Given these assumptions, it follows that there is

likely to be a fitness advantage to invest more in reproduction as individuals approach the limits of their life span.

Thus far, support for the terminal investment hypothesis has been reported in many animals that have been examined, including species as diverse as burying beetles (Creighton et al. 2009), fishes (Bercovitch et al. 2009), birds (Velando et al. 2006), and primates (Hoffman et al. 2010), including humans (Tifferet and Kruger 2010). Several studies have not supported this hypothesis, however (Yoccoz et al. 2002), and there are few comparable studies for plants. Indeed, Hamilton (1966) suggested that actuarial senescence in plants may be negligible, a hypothesis that now has considerable support from subsequent studies (Baudisch et al. 2013; Caswell and Salguero-Gómez 2013). Nonetheless, there has been little work examining the pattern of reproductive senescence in plants, which is the relevant information for the terminal investment hypothesis. In one of the few studies of which we are aware, Shefferson and Roach (2013) found evidence countering this hypothesis in the short-lived iteroparous perennial *Plantago lanceolata*, where both size and reproduction decline with age. There appear to be no prior studies examining reproductive senescence or terminal investment in long-lived trees.

In contrast, there are many studies regarding the role of stress as indicated by growth rates and mortality of trees; these studies have typically found that growth is reduced before death (Waring 1987; Pederson 1998; Das et al. 2007). Given the possibility that this reduction is a consequence of a reallocation of resources to reproduction rather than physiological decline affecting both growth and reproduction, studies investigating the possibility that long-lived trees support the terminal investment hypothesis clearly warrant investigation.

Here we test the terminal investment hypothesis as part of a long-term study of reproduction and life-history strategies of California oaks (genus *Quercus*). We gathered data on acorn production on 1,067 individuals of eight species at a series of sites across the state over a period of 20 years. In addition, we obtained data on both acorn production and

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Am. Nat. 2017. Vol. 189, pp. 564–569. © 2017 by The University of Chicago. 0003-0147/2017/18905-5723\$15.00. All rights reserved.

DOI: 10.1086/691161

radial growth on 250 trees of five species at a site in central coastal California over an even longer time period. Over the course of the study, enough individuals died of apparently natural causes that we can now examine patterns of growth and reproduction just before death as a test of the hypothesis that the trees differentially invest in reproduction, as predicted by the terminal investment hypothesis.

Two complications of such a test are that oaks can live for hundreds of years and are typically mast fruiting, with reproduction varying greatly among both years and individuals (Kelly and Sork 2002; Koenig and Knops 2005). As such, testing for terminal investment requires a long-term study involving enough individuals that variability across years and among trees can be accommodated.

## Methods

### Data

Two related data sets were used for this analysis. The first was a sample of 250 mature trees of five species studied at Hastings Natural History Reservation in central coastal California since 1980 (Koenig et al. 1994b). These trees—for which we had data on both growth and reproduction—were used in analyses involving radial growth. The second data set, involving data on only acorn production, included 817 mature individuals of eight species from various sites across California, a complete list of which is provided in table A1 (tables A1–A3 are available online). Although some changes and additions were made throughout the course of the study, trees were generally subject to visual acorn counts each year between 1994 and 2016.

Acorn production was estimated by means of visual surveys conducted each autumn. Surveys consisted of two observers counting as many acorns as they could in 15 s; values were then added and ln-transformed in order to reduce skew in the data (Koenig et al. 1994a), to better match prior analyses and because generalized linear mixed models in which untransformed acorn count data were used with a Poisson error distribution exhibited poor convergence. To measure radial growth, trees at Hastings were fitted with dendrometers (Cattellino et al. 1986) in 1993, and radial growth was measured annually thereafter. The relationship between radial growth as measured by mean circumference increment and tree size (diameter at breast height [DBH]) was not significantly size dependent (mixed-effects model with species included as a random factor; mean growth increment =  $0.013 (\pm 0.011 \text{ SE}) \times \text{DBH}$ ;  $t = 1.17$ ,  $N = 238$  trees,  $P = .24$ ). Thus, we used mean annual circumference increment as measured by the dendrometers as our index of growth investment.

Trees were visited each year in September, when we conducted the acorn survey and assessed whether individuals

were dead or alive on the basis of the presence of new or green leaves. In all cases, trees included in the died category either fell or remained standing but exhibited no new growth; trees that were killed by fire and four individuals that may have been cut were excluded from the analyses. For each tree that died (the focal tree), we compiled data on acorn production and (when available) radial growth for the years before death along with parallel data for the mean values for all trees of the same species at the same site during the same years (comparison trees). Results were unchanged using the two trees (one the same size or larger; the other the same size or smaller) of the same species at the same site that were closest in size to the focal tree as comparison trees to control for any size-dependent scaling of reproduction. We then recorded data for the focal tree along with the mean acorn crop and mean radial growth for the comparison trees. Data were analyzed using R 3.3.1 (R Development Core Team 2016).

### Analyses

Two sets of analyses were conducted. The first set compared the population of trees that died (focal trees) to trees of the same species at the same site that lived (comparison trees). Two tests were conducted. The first involved reverse-age analyses in which trees were standardized on the basis of their year of death, similar to analyses performed previously by Martin and Festa-Bianchet (2011) on bighorn sheep (*Ovis canadensis*), Hammers et al. (2012) on Seychelles warblers (*Acrocephalus sechellensis*), and Shefferson and Roach (2013) on *Plantago*. Analyses involved linear mixed-effects models (procedure lmer in the lme4 package; Bates et al. 2015). For each focal tree and each set of comparison trees, we compiled data on the acorn crop (ln-transformed) and (when available) radial growth as dependent variables and size (DBH), year relative to time of death of the focal tree, whether the tree died (i.e., whether it was a focal tree or the comparison trees), and the interaction between these latter two variables as fixed effects. Year of death and individual within species within locality (individual within species for the analysis of growth, since these data were from only a single site) were included as random effects to account for temporal and spatial variation in conditions. Since we were interested in the differences between trees that died and trees that survived, the relevant results consisted of the interaction terms between year relative to time of death and whether the tree died (the focal tree vs. comparison trees). We analyzed and plotted the effect sizes for the interaction terms for the 6 years just before death (year  $-5$  to year  $0$ ) compared with the average of the values for the prior 5 years (year  $-10$  to year  $-6$ ).

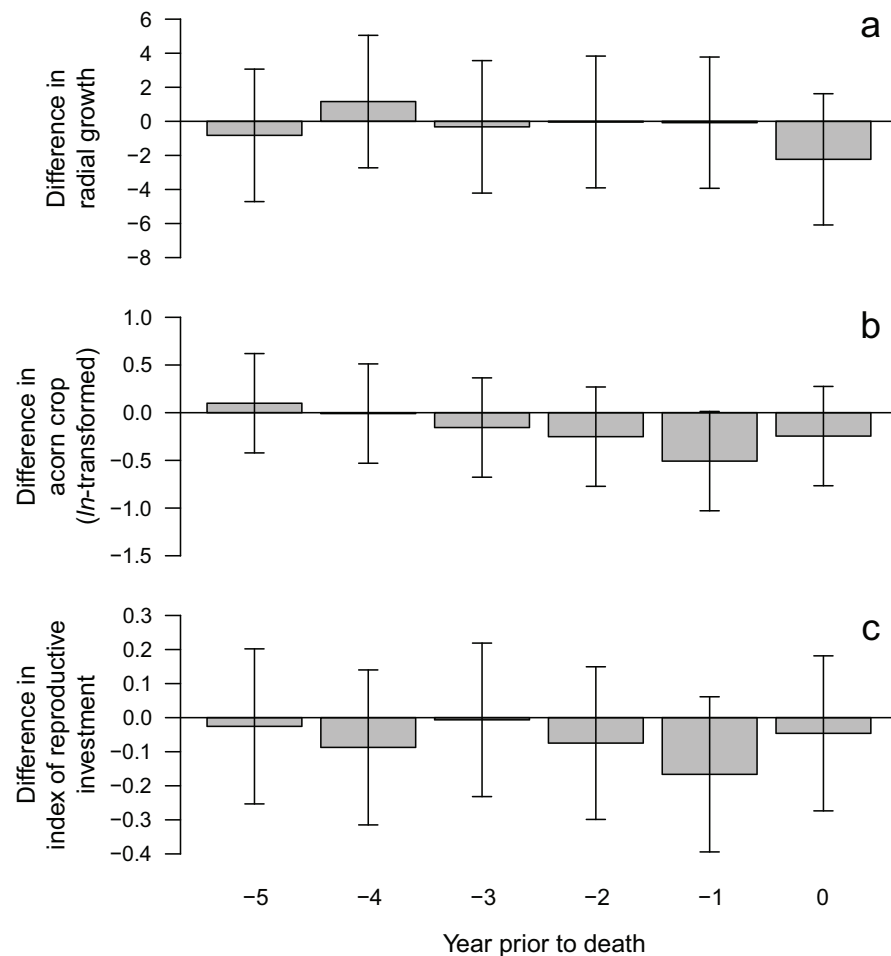
The second test that compared trees that died with those that lived involved standardizing both the acorn crop (ln-transformed) and radial growth of trees to between 0 to 1

on the basis of their maximum and minimum values over the complete length of time for which we had data. We then calculated an index of reproductive investment—defined as the standardized acorn crop divided by the sum of the standardized acorn crop plus standardized radial growth—over the last 6 years of life (year  $-5$  to year  $0$ ). The index of reproductive investment for each focal tree was then compared with the mean of all trees that did not die of the same species during the same years. (All data for this analysis were from the Hastings Natural History Reservation site, where radial growth was measured.)

We conducted a second set of analyses examining the hypothesis that reduced growth may be a cost of increased investment in acorn production within the same tree rather than on a population level. First, we performed Spearman rank correlations between acorn production and radial

growth of trees that died over the 6 years from year  $-5$  to year  $0$ , and we report the mean correlation coefficient and the proportion of trees for which the correlation was negative, as predicted by this hypothesis. (Results were unchanged using data going back as far as year  $-10$  before death.) This analysis—testing for a potential trade-off between growth and reproduction—was similar to analyses we have performed previously (Knops et al. 2007), except that here we restricted our analyses to the years just before death, thus testing the relationship between growth and reproduction during the last years of a tree's life.

Finally, we tested the possibility that dying trees may invest relatively more of their reserves into reproduction compared with radial growth, even if their relative investment in both growth and reproduction decrease. We averaged the index of reproductive investment as defined above for years



**Figure 1:** Mean ( $\pm$  95% confidence interval) difference in absolute radial growth (a), acorn production (ln-transformed; b), and index of reproductive effort (c) for trees that died compared with trees of the same species and site that lived in the years before death. Values estimated by the effect size of the interaction terms between years before death and whether the tree died derived from mixed-effects models. Values are plotted for the 6 years before death (year  $-5$  to year  $0$ ). For further details of the analyses, see table A3, available online.

–10 through –6 for each tree that died and compared this value with the index of reproductive investment for the same trees for each of years –5 to 0, using paired Wilcoxon two-sample tests. We then tested for a trend in values over the last 6 years of life with a Spearman rank correlation.

### Results

During the study, a total of 70 trees died for reasons that were apparently unrelated to fire or other anthropogenic causes (table A2). Of the trees that died, 23 were from Hastings; 17 of these provided data on both radial growth and acorn production.

Compared with the mean for the prior 5 years, both radial growth and acorn production were smaller among trees that died compared with those that lived for the last 6 years of life, with one exception (year –4 for radial growth; year –5 for acorn production); none of the differences was significant, however (fig. 1*a*, 1*b*). Comparing the index of reproductive investment by trees that died to the mean of trees that lived, the index of investment was lower among trees that died in all cases but was again not significant for any

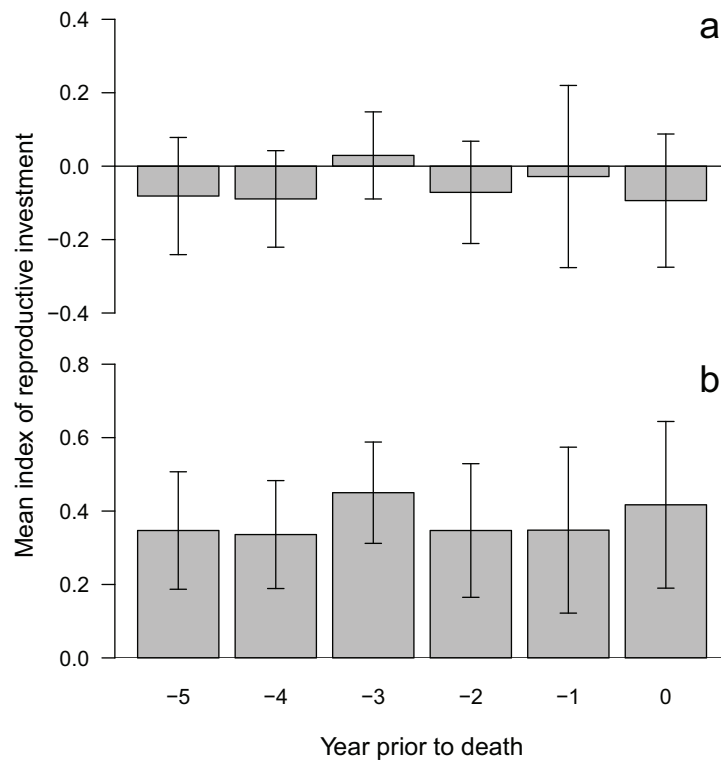
of the years (fig. 1*c*). Complete details regarding these analyses are provided in table A3.

Of the 15 trees for which we were able to calculate correlations between growth and reproduction during the last 6 years of their lives, nine were positive and six were negative. Overall, the mean  $\pm$  SD Spearman rank correlation coefficient was  $0.21 \pm 0.56$ .

Seventeen trees provided data for estimating the index of reproductive effort during the last years of their lives; nine of these trees had data for all years we analyzed (year –5 to year 0 along with a mean for years –10 to –6). Compared with the mean value of the index for years –10 to –6, there were no significant differences for any of years –5 to 0 (six pairwise tests; all  $P \geq .25$ ; fig. 2*a*). There was also no significant trend in the index including either all trees ( $r_s = 0.002$ ,  $P = .99$ ; fig. 2*b*) or only the nine trees with data for all years ( $r_s = 0.36$ ,  $P = .42$ ).

### Discussion

Since it was first proposed 50 years ago, the terminal investment hypothesis has received considerable—although by no



**Figure 2:** *a*, Mean ( $\pm$  95% confidence interval [CI]) difference in the index of reproductive effort for the 6 years before death (year –5 to year 0) compared with the mean of the index for the same trees over years –10 to –6. *b*, Mean ( $\pm$  95% CI) index of reproductive effort for the 6 years before death.

means universal (Yoccoz et al. 2002)—support in animals ranging from insects to humans. However, we know of only one prior study investigating this hypothesis in iteroparous plants (Shefferson and Roach 2013). Indeed, the only scientific reference to this phenomenon in trees of which we are aware is in Crawley and Long's (1995, p. 692) article on *Quercus robur*, in which they reported that “acorn production increased significantly with the degree of stag-headedness . . . consistent with forester's folk lore that dying trees increase their reproductive output.” Beyond this, terminal investment bears some similarities to the increased flowering and fruit production that often accompanies the girdling of trees, a procedure than typically results in death within several years (Noel 1970). It differs, however, in that the effects of girdling on productivity are due to interference with normal water transport and nutrient translocation, whereas terminal investment—if it exists—is presumably an evolved capacity of individuals that are senescing naturally.

Prior studies indicate that trees approaching the end of their lives typically exhibit inferior growth indicative of environmental stresses that are often exhibited during drought or other stressful periods (Pederson 1998, 1999). Other changes in allocation that have been found in trees undergoing stress include reductions in canopy growth, photosynthetic activity, and storage reserves (Waring 1987) and, in the case of *Acer rufinerve*, a shift in resources toward female flowers (Nanami et al. 2004)—a change that potentially is in accord with the terminal investment hypothesis.

Here we found no significant differences in growth, reproduction, or an index of reproductive investment that considered both reproduction and growth in California oaks during the last 6 years of their lives compared with either conspecific trees at the same site (fig. 1) or to growth and reproduction by the same trees during prior years (fig. 2a). Differences were, however, generally in the direction of trees investing less in reproduction during the years before their demise rather than more, thus countering the likelihood that significant support for terminal investment would be detected with additional data.

The terminal investment hypothesis is predicated on a trade-off between growth and reproduction. Such a trade-off is an assumption of several hypotheses of aging, including antagonistic pleiotropy (Williams 1957) and the disposable soma theory (Kirkwood 1977), but not of, for example, the mutation accumulation hypothesis (Medawar 1952). Such a trade-off has often not been found in prior studies of trees (Sork et al. 1993; Crawley and Long 1995), nor was it evident in our data either here or in prior analyses of oaks at this site (Knops et al. 2007). Together, these results counter the hypothesis that oaks shift resources into reproduction in response to conditions potentially culminating in their demise.

This conclusion is based both on a relatively small sample of trees and on data taken during the last several years of life. Many oak species live hundreds of years, and death may be a decades-long phenomenon (Pederson 1998). It is therefore possible that oaks and other long-lived trees shift resources into reproduction many years before they finally die but still in response to stresses that may eventually kill them. Testing this possibility will clearly be a challenge, given the timescales potentially involved. In the short-term, however, our results do not support the hypothesis that oaks suffering from stress and thus more likely to die differentially shift resources into reproduction, as proposed by the terminal investment hypothesis.

#### Acknowledgments

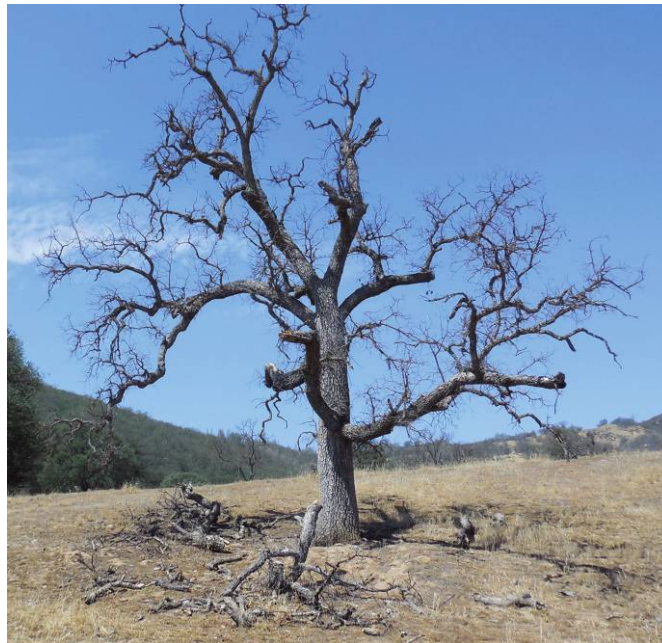
We thank M. Crawley and L. Hadj-Chikh for discussion, W. Hochachka for statistical advice, and the reviewers for their comments. The study was supported by the National Science Foundation (most recently DEB-1256394 to W.D.K.) and the Integrated Hardwoods Range Management Program (University of California).

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Natural History Editor: Mark A. McPeck



Valley oak (*Quercus lobata*) from central coastal California, one of the 70 trees that died of natural causes studied to test whether oaks invest more in producing acorns than expected in the years before their demise. Photo credit: Walter D. Koenig.

**Appendix from W. D. Koenig et al., “Testing the Terminal Investment Hypothesis in California Oaks”  
(Am. Nat., vol. 189, no. 5, p. 564)**

**Details of Sampling Sites, Trees That Died, and Linear Mixed-Effects Models**



**Table A1:** Sites and species included in the analyses

Site	First year	<i>Quercus</i> species											Notes	
		<i>Q. lobata</i>	<i>Q. douglasii</i>	<i>Q. chrysolepis</i>	<i>Q. agrifolia</i>	<i>Q. wislizenii</i>	<i>Q. kelloggii</i>	<i>Q. engelmannii</i>	<i>Q. garryana</i>					
Hastings Natural History Reservation, Monterey Co.	1980	88	57	21	63		21							
Jasper Ridge, San Mateo Co.	1989	39	29		31									
Pozo, San Luis Obispo Co.	1989	24	25		24									
Hopland Field Station, Mendocino Co.	1994	10	20	20		13	22							<i>lobata</i> added 1999; <i>wislizenii</i> added 2000
Tower House, Shasta Co.	1994	17		19			20							
Douglas City, Trinity Co.	2002												12	
Dye Creek, Tehama Co.	1994	18	22			10								<i>wislizenii</i> added 2000
Sierra Foothills, Yuba Co.	1994	9	25			12								<i>wislizenii</i> added 1999; <i>lobata</i> added 2001
Yosemite National Park, Mariposa Co.	1994			20										<i>wislizenii</i> added 1999
San Joaquin Experimental Range, Madera Co.	1994	26	25		20	12	25							Additional <i>lobata</i> added 2008
Sedgwick, Santa Barbara Co.	1994		20							19				
Santa Rosa Plateau, Riverside Co.	1994													
Palomar Mountain, San Diego Co.	1994			20			20							
Switzer's, San Gabriel Mountains, Los Angeles Co.	1994			20			20							
Liebre Mountain, Los Angeles Co.	1994	11	20											<i>lobata</i> added 1999
Kaweah Oaks, Tulare Co.	1999	10												
Kaweah River Preserve, Tulare Co.	1999		14	10				10						<i>wislizenii</i> added 2000
James Reserve, San Jacinto Mountains, Riverside Co.	2004			12										
San Jacinto Mountains, Riverside Co.	2004				12									Site abandoned 2014
Oak Grove Campground, San Diego Co.	2014				10									Site added 2014
Total trees (1,067)		252	257	142	188	57	140	19	12					
Total sites (49)		10	10	8	7	5	7	1	1					

**Table A2:** Species and locality of trees that died during the study

Site	<i>Quercus</i> species								
	<i>Q. lobata</i>	<i>Q. douglasii</i>	<i>Q. chrysolepis</i>	<i>Q. agrifolia</i>	<i>Q. wislizenii</i>	<i>Q. kelloggii</i>	<i>Q. engelmannii</i>	<i>Q. garryana</i>	
Hastings Natural History Reservation, Monterey Co.	5	2	1	5		10			
Jasper Ridge, San Mateo Co.	1	1		1					
Pozo, San Luis Obispo Co.	2	1		3					
Hopland Field Station, Mendocino Co.	0	1	0		1	0			
Tower House, Shasta Co.	1		0			3			
Douglas City, Trinity Co.								0	
Dye Creek, Tehama Co.	1	1			3				
Sierra Foothills, Yuba Co.	2	0			0				
Yosemite National Park, Mariposa Co.			0			0			
San Joaquin Experimental Range, Madera Co.		0			2				
Sedgwick Reserve, Santa Barbara Co.	0	0		0					
Santa Rosa Plateau, Riverside Co.				4			3		
Palomar Mountain, San Diego Co.			0			1			
Switzer's, San Gabriel Mountains, Los Angeles Co.			1	0					
Liebre Mountain, Los Angeles Co.	4	1				3			
Kaweah Oaks, Tulare Co.	5								
Kaweah River Preserve, Tulare Co.		0	0		1				
James Reserve, San Jacinto Mountains, Riverside Co.			0			0			
San Jacinto Mountains, Riverside Co.				0					
Oak Grove Campground, San Diego Co.				0					
Total mortality	21	7	2	13	7	17	3	0	

**Table A3:** Results of linear mixed-effects models (lmer) of radial growth, reproduction (acorn crop, ln-transformed), and index of reproductive investment on size (diameter at breast height), whether trees died, years before death (year -5 to year 0), and interaction between the latter two variables

Fixed factor	Effect size $\pm$ SE	<i>t</i>	<i>P</i>
Radial growth ( <i>N</i> = 17 trees):			
Died	-2.24 $\pm$ 1.46	-1.54	.13
Diameter at breast height	-.004 $\pm$ .014	-.29	.77
Year -5	.48 $\pm$ 1.56	.31	.76
Year -4	.95 $\pm$ 1.57	.60	.55
Year -3	1.88 $\pm$ 1.59	1.18	.24
Year -2	.20 $\pm$ 1.59	.13	.90
Year -1	.03 $\pm$ 1.60	.02	.99
Year 0	.30 $\pm$ 1.60	.19	.85
Died $\times$ year -5	-.82 $\pm$ 2.05	-.40	.69
Died $\times$ year -4	1.16 $\pm$ 2.05	.57	.57
Died $\times$ year -3	-.32 $\pm$ 2.05	-.16	.88
Died $\times$ year -2	-.04 $\pm$ 2.04	-.02	.99
Died $\times$ year -1	-.08 $\pm$ 2.03	-.04	.97
Died $\times$ year 0	-2.23 $\pm$ 2.03	-1.10	.27
Acorn crop ( <i>N</i> = 70 trees):			
Died	-.16 $\pm$ .19	-.86	.39
Diameter at breast height	.003 $\pm$ .002	1.54	.12
Year -5	-.53 $\pm$ .20	-2.65	.008
Year -4	-.20 $\pm$ .20	-1.02	.31
Year -3	-.18 $\pm$ .20	-.91	.36
Year -2	-.30 $\pm$ .20	-1.47	.14
Year -1	-.18 $\pm$ .20	-.89	.37
Year 0	-.36 $\pm$ .21	-1.76	.08
Died $\times$ year -5	.10 $\pm$ .27	.37	.71
Died $\times$ year -4	-.01 $\pm$ .27	-.04	.97
Died $\times$ year -3	-.16 $\pm$ .27	-.59	.56
Died $\times$ year -2	-.25 $\pm$ .27	-.95	.34
Died $\times$ year -1	-.51 $\pm$ .27	-1.91	.06
Died $\times$ year 0	-.25 $\pm$ .27	-.92	.36
Index of reproductive investment ( <i>N</i> = 17 trees):			
Died	-.039 $\pm$ .082	-.48	.63
Diameter at breast height	.0020 $\pm$ .0008	2.44	.02
Year -5	-.044 $\pm$ .083	-.54	.59
Year -4	.011 $\pm$ .008	.13	.31
Year -3	.048 $\pm$ .084	.57	.57
Year -2	-.016 $\pm$ .083	-.19	.85
Year -1	.048 $\pm$ .082	.58	.56
Year 0	-.033 $\pm$ .082	.40	.69
Died $\times$ year -5	-.026 $\pm$ .116	-.22	.83
Died $\times$ year -4	-.087 $\pm$ .116	-.75	.45
Died $\times$ year -3	-.006 $\pm$ .115	-.06	.96
Died $\times$ year -2	-.075 $\pm$ .114	-.65	.51
Died $\times$ year -1	-.166 $\pm$ .116	-1.43	.15
Died $\times$ year 0	-.046 $\pm$ .116	-.40	.69

Note: For radial growth, year of death and individual within species within locality were included as random effects; for acorn crop and index of reproductive investment, year of death and individual within species were random effects. Values are relative to the mean of years -10 to -6. Interactions—representing the differences in the effect sizes for trees that died versus those that did not—are plotted in figure 1.