Hypotheses accounting for ultimate causes (i.e., masting would be a selective trait that increases fitness) are based on the benefits of economies of scale, i.e., massive but random reproductive events would be more beneficial than producing regular crops of similar but smaller size (e.g., the predator-satiation hypothesis, see Curran and Webb, 2000; Espelta et al., 2008, 2017). Hypotheses accounting for proximate causes, though, are focussed on the mechanisms by which plants produce fruits (Norton and Kelly, 1988; Kelly and Sork, 2002). These hypotheses are being intensely debated,
and many studies have investigated the potential causes of the high interannual variability of fruit production: stored carbohydrates (Hoch et al., 2013; Ichie et al., 2013), reproductive costs (Sala et al., 2012), availability of nutrients (Crone et al., 2009; Fernández-Martínez et al., 2016), pollination efficiency (Koenig et al., 2015; Bogdziewicz et al., 2017a,b), or weather, either as a cue prompting reproduction (Kelly et al., 2013; Pearse et al., 2014; Bogdziewicz et al., 2017b) or by its effect on plant productivity either by enhancing photosynthesis (Fernández-Martínez et al., 2015, 2017) or by its interaction with nutrient availability (Smalil et al., 2011). Amongst proximate causes, meteorological variability may be an inseparable component of variable seed production (Pearse et al., 2016), because weather is the most evident source of interannual variability that can affect plant productivity. However, it has often been claimed that factors other than weather have to be involved in driving mast seeding behavior (Koenig and Knops, 2000; Kelly and Sork, 2002; Crone and Rapp, 2014; Pearse et al., 2016).

Two main arguments have been used to claim that weather cannot be the sole proximate cause of masting. First, fruit production is much more interannually variable (i.e., CV = SD · mean\(^{-1}\), at the population level, the standard way to characterize masting behavior; Herrera et al., 1998) than meteorological variables such as rainfall or temperature (Koenig and Knops, 2000, 2005; Kelly and Sork, 2002), two of the most commonly used meteorological variables for predicting crop size. Second, meteorological variables are rarely bimodally distributed, but fruit production sometimes is (Norton and Kelly, 1988; Herrera et al., 1998). These arguments, though, are only correct when linear relationships between seed production and weather are assumed, even though a large number of mast studies have relied on logarithmic transformations for seed production to normalize model residuals (Sork et al., 1993; Koenig et al., 1994; Kelly et al., 2013; Pearse et al., 2014). This transformation linearises the relationship between crop size and weather, but the true relationship remains unequivocally non-linear and is thus within the domain of Jensen’s inequality.

Jensen’s inequality (Jensen, 1905; Ruel and Ayres, 1999; Denny, 2017) is a mathematical property that applies to non-linear functions. Understanding this property is useful in order to better predict and understand the consequences of predictor variability on the response variables for functions of different shapes. Briefly, Jensen’s inequality implies that the variance of a predictor variable will increase the variance of the response variable in accelerating functions (i.e., convex functions where the slope increases; the 2nd derivative is positive, as in exponential-growth functions). Conversely, the variance of a predictor variable will decrease the variance of the response variable in decelerating functions (i.e., the slope decreases in concave down functions; the 2nd derivative is negative, as in logarithmic functions, see Denny (2017) for a detailed explanation).

Many of the published relationships between fruit production and weather are subject to Jensen’s inequality because fruit production is usually log-transformed to fit the models, implying that meteorological variability can increase or decrease the variability of fruit production depending on the shape of the function. Jensen’s inequality also means that response and predictor variables will likely be distributed differently in non-linear relationships (Ruel and Ayres, 1999). A Gaussian distribution for a predictor can easily lead to an exponentially distributed response variable (e.g., gamma distribution) in accelerating functions. Similarly, other types of non-linear relationships, such as sigmoidal functions, can produce a bimodal response variable, another property sometimes found in fruit-production data (Herrera et al., 1998).

The aim of this study was thus to demonstrate, using computer simulations, that common and even expected non-linear relationships between weather and fruit production could lead to the differences in distribution and temporal variability observed in the time series of weather and fruit production. Positive results would return meteorological variability to the focus of mastings research.

METHODS

We performed five simulations in which fruit production (response variable) followed a linear, exponential, logarithmic, sigmoidal, and Gaussian relationship with a meteorological variable to demonstrate that the extreme variability in fruit production data can be explained by a normally distributed, less variable predictor (e.g., weather). We simulated 48 meteorological time series of 100 years following a normal distribution with a mean of 600 (e.g., proximate annual rainfall in Barcelona) and an SD ranging from 10 to 200. Hence, the simulated interannual variability (CV) ranged from 0.017 to 0.33, which are plausible ranges of interannual meteorological variability of annual temperature (~5%) and highly variable annual rainfall in the Mediterranean regions (~30%). However, the discussion of our results apply to any meteorological variable (relative humidity, wind speed, rainy days) that could affect the biology of a given species that produces fruits (Crone and Rapp, 2014). For each simulated meteorological time series, we then simulated fruit production using linear, exponential, logarithmic, sigmoid, and Gaussian functions with a normally distributed error for different slopes of the functions (or width for Gaussian). The SD of the predicted values was 10% of the average of the predicted values for all simulations (ensuring a statistically significant relationship between the dependent and the predictor variables for most of the slopes). We simulated 100 years of fruit production 1,000 times and calculated the average proportional variability (PV) for each slope and meteorological time series. PV is calculated as: \[ PV = \frac{\sum D(z)}{C} \] where \( D(z) = 1 - \frac{\min(z_i, z_j)}{\max(z_i, z_j)} \) is each of the pairwise comparisons of all the \( z \) values of the variable and \( C \) is the number of all possible pairwise combinations within the set of values (see Heath, 2006 for further calculation details). The PV index is a robust measure of variability that overcomes some of the mathematical problems that the CV presents, especially when assessing variability in non-normally distributed data (Heath, 2006; Fernández-Martínez et al., 2016). However, because the CV index is still widely used in ecology, we also calculated it in our simulations and presented the results in Supplementary Material (Figure S1). All analyses were performed using R (R Core Team, 2015); the full code...
used to extract the results is available in Supplementary Material, section 1.

We also searched the literature for exemplar masting studies that provided relationships between fruit production and weather to summarize information about fruit production and interannual meteorological variability (CV and PV) and the shape (linear or non-linear) of their relationship. The results of these analyses should help other authors interpret their results when using non-linear relationships in their studies.

**RESULTS AND DISCUSSION**

Our simulations indicated that the variability (PV) of the response variable, in a linear relationship between two variables (i.e., \( f(\text{fruit production}) \approx \beta \text{weather} + \varepsilon \)), where \( \beta \) represents the slope of the rainfall and \( \varepsilon \) is the error term, increased slightly for a very low slope coefficient (\( \beta \)) at any variability of the predictor variable (weather) but quickly reached an asymptote at very similar values of the predictor (Figure 1A). The normal distribution of the independent variable in this case was also translated into the response variable, without being affected by the slope of the relationship (Figures 2A,B). This simulation, however, produced fruit-production data very different from those reported in masting studies (Kelly, 1994; Herrera et al., 1998; Fernández-Martínez et al., 2016). Additionally, linear relationships, as in this first scenario, have rarely been reported in masting studies (Table 1), because fruit production is usually log-transformed before fitting the models to normalize the residuals. The variability in the meteorological variables in these cases is always similar in magnitude to the variability of fruit production, as our simulation also suggested (Table 1).

Our second simulation used log-transformed data for fruit production (exponential function), the most common analysis performed in masting studies (e.g., Sork et al., 1993; Koenig et al., 1994; Kelly et al., 2013; Pearse et al., 2014), where Jensen's inequality already plays a role.

The variability of fruit production increased steeply with \( \beta \) in an exponential function of the type \( f(\text{fruit production}) \approx e^{(\beta \text{weather})} + \varepsilon \), especially at high predictor variability (Figure 1B). When this kind of relationship between response and predictor was assumed, the variability of the response variable rapidly increased above the variability of the predictor, using both PV and the CV indices (Figure S1). The shape of the distribution of the response variable varied with \( \beta \), from a normal distribution similar to that of the meteorological predictor at \( \beta = 0.001 \), to a highly left-skewed distribution at \( \beta = 0.01 \) (Figures 2C,D) similar to those reported for fruit production in masting studies. Our bibliographical search clearly supported these results. Fruit production was always more variable than the predictor meteorological variable in relationships in which fruit production was log-transformed (Table 1). The change from constant to highly variable fruit production along with the magnitude of the slope of the relationship implied that masting behavior may be a consequence of the hypersensitivity of plants to meteorological variability [e.g., high exponential slopes (\( \beta \))], providing the first mathematical support for previous hypotheses (Kelly, 1994; Kelly and Sork, 2002). This hypersensitivity to weather cues could be explained by synergistic effects boosting fruit production because of favorable weather for photosynthesis and nutrient availability. For instance, acorn production in Mediterranean oaks has been reported to be positively associated to spring rainfall, which, a part from drought, ameliorates nutrient limitation by enhancing nitrogen mineralization (Smaill et al., 2011; Fernández-Martínez et al., 2015; Bogdziewicz et al., 2017a). Overall, this phenomenon reflects the high costs of reproduction of some species (Sala et al., 2012).

The third simulation, using the logarithmic function \( f(\text{fruit production}) \approx \beta \ln(\text{weather}) + \varepsilon \), draws the typical shape of a function describing a process of diminishing returns, in which the response variable increases steeply at low values of the predictor but saturates at high values (Figure 1C). In this case, the behavior of the variability is completely opposite to the behavior reported in the two previous simulations. Because of the concave-down shape of the function, variability of the response was rather insensitive to variability in the predictor. For high values of variability of the predictor, the logarithmic function even shrinks variability of the response. Like in linear functions, the slope of the relationship did not affect variability of the response. Consequently, the shape of the distribution of the response variable barely varied \( \beta \), presenting right-skewed Gaussian-like distributions for low and high slopes (Figures 2E,F).

Our fourth simulation used a sigmoidal function of the type \( f(\text{fruit production}) \approx a + \frac{b-a}{1+e^{-(\text{weather}-d})} + \varepsilon \), where \( a \) and \( b \) are the minimum and maximum values in the function, respectively, \( d \) is the value of the predictor variable at the inflection point, and \( c \) is the degree of curvature of the function, which is linear at low values of \( c \) and predicts only \( a + \varepsilon \) and \( b + \varepsilon \) values for high values of \( c \). The variability of fruit production rapidly increased with \( c \) until it approached an asymptote near \( PV = 0.7 \) (Figure 1D) or \( CV = 1 \) (Figure S1). The increase, though, was even steeper for highly variable weather. This simulation further indicated that phenomena produced by sigmoidal relationships could potentially increase the variability of response variables much more than that of the predictors. Sigmoidal relationships between weather and fruit production produced a distribution of fruit production similar to a Gaussian distribution at low values of \( c \), but the distribution became bimodal at high values of \( c \) (Figures 2G,H). This kind of relationship could account for cases like the one reported for Chionochloa sp. in which flowering only occurred when average temperature for January and February was \( >10^\circ C \) (McKone et al., 1998; Rees et al., 2002). However, we could not find any sigmoidal relationships in masting studies (Table 1).

Our last simulation used a Gaussian function of the type \( f(\text{fruit production}) \approx ae^{-\frac{(\text{weather}-b)^2}{2c^2}} + \varepsilon \), where \( a \) is maximum \( y \) value of the function, \( b \) is the value of the predictor variable at the center of the bell, and \( c \) is the standard deviation (or width) of the bell, determining the curvature of the function, which defines a flat function at high values of \( c \) and a narrow bell at low values. The variability of the response variable increased with the variability
FIGURE 1 | Summary of theoretical responses of an ecosystem, or organism, variable as a function of an environmental predictor. Shown responses are, from narrow to wide environmental range of the environmental predictor, linear (A), exponential (B), logarithmic (C), sigmoidal (D), and Gaussian (E). Coloured lines represent the different types of responses within functions (blue, exponential growth; black, linear; red, logarithmic). Gray horizontal bars show the distribution of the response variable for each case. Boxes on the right side show the variation of the response variable (e.g., fruit production, color gradient) as a function of the variability of the predictor (e.g., weather) and the strength of the relationship between both variables for each of the theoretical responses. The variability of the response variable was estimated using the proportional variability index (PV, see Heath, 2006). Simulations performed with the coefficient of variation are shown in Figure S1. Blue colors indicate low values of variability (i.e., PV) and red colors indicate high values.
FIGURE 2 | Histograms of the change in the distribution of fruit production for different slopes and functions [linear (A,B), exponential (C,D), logarithmic (E,F), sigmoidal (G,H), and Gaussian (I,J)] assuming that fruit production depends on a Gaussian meteorological variable with a mean of 600 and an SD of 150 (CV = 0.25). Distributions for low slopes—or c parameters for sigmoidal and Gaussian functions—a (A,C,E,G,I) and distributions for high slopes (B,D,F,H,J). The code for this simulation is provided in Supplementary Material, section 1.
of the predictor but decreased with increasing $c$ (Figure 1E), being even lower than that of the predictor for high values of $c$. Simulated distributions were Gaussian-like for high values of $c$ but almost uniform for narrow (low $c$) Gaussian relationships, except for a high frequency of low values (Figures 2I,J).

Overall, our results point out that the claim that weather cannot be the main proximate driver of fruit production in masting species was based on the erroneous assumption that the relationship between weather and fruit production must be linear. Most relationships in nature, though, are theoretically non-linear (Denny, 2017). Operative values for environmental gradients for life on Earth are optimal, so most of the relationships between organisms and the environment should be Gaussian-like or similar (Figure 1E), increasing from low values of the independent variable until they reach the optimal value at which the response peaks. The relationship then becomes negative at high values of the independent variable. Linear and exponential relationships, such as those usually reported in masting studies (Table 1), should appear within the mid-low or the mid-high range of the independent variable in which the response occurs (Figure 1). To the best of our knowledge, however, second-order polynomial or Gaussian-like relationships between any environmental variable and fruit production have not yet been reported, indicating that studies with larger ranges of environmental variability are required to better understand the link between weather and fruit production. Second-order polynomial or Gaussian-like relationships are very common in other fields of ecology (e.g., species distributions) but are very rare in many others such as masting.

Observational and experimental studies in ecology are still very limited in space and time, so determining the complete range of responses for a dependent variable remains difficult (see Figure 1). The biosphere has evolved to operate nearest to its optimal values, which may be another reason for the lack of more Gaussian-like relationships. In other words, we may not be able to find Gaussian-like relationships between, for example, precipitation and fruit production in a focal population, because when the given species is pushed far away from its optimum (e.g., by climate), it will be replaced by a different one that will operate better (or will be more competitive). This hypothesis could easily be tested experimentally, but we predict that observational studies will provide little evidence.

Masting studies should reconsider the role of meteorological variability as the most likely predictor of ecosystemic functioning, because weather is the main source of interannual variability to which plants are subjected. Comprehending the role of weather as a proximate cause for masting is especially important after the recent discoveries that resources invested in reproduction are being used as a proximate cause for masting. Understanding the role of weather in plant reproduction can be facilitated by observing how weather affects both the acquisition of resources by plants, by conditioning their photosynthetic rates, and pollination

### TABLE 1 | Examples extracted from the literature showing the differences in the CVs and PVs of weather and fruit production for linear and exponential relationships.

| Species          | Fruit CV | Fruit PV | Meteorological variable       | Weather CV | Weather PV | Shape | References       |
|------------------|----------|----------|--------------------------------|------------|------------|-------|------------------|
| Quercus ilex     | 1.18     |          | Summer water deficit           | 0.14       | EXP        |       | 1                |
| Q. ilex          | 1.72     | 0.83     | Spring torrential rainy days   | 0.46       | EXP        |       |                  |
| Q. humilis       | 1.08     | 0.64     | Spring water deficit           | 0.28       | EXP        |       |                  |
| Q. kelloggi      | 0.77     |          | Mean max April T               | 0.12       | EXP        |       |                  |
| Q. canariensis   | 1.40     |          | Spring water deficit           | 0.93       | EXP        |       |                  |
| Q. suber         | 0.42     |          | Spring water deficit           | 0.93       | LIN        |       |                  |
| Q. robur         | 1.81     | 0.79     | Summer P                       | 0.27       | EXP        |       |                  |
| Q. petraea       | 1.60     | 0.78     | Autumn T                       | 0.19       | EXP        |       |                  |
| Fagus sylvatica  | 1.78     | 0.81     | Summer P                       | 0.30       | EXP        |       |                  |
| Pinus abies      | 1.18     | 0.63     | Spring P                       | 0.31       | EXP        |       |                  |
| Pseudotsuga menziesii | 1.17 | 0.70 | Summer T | 0.18 | EXP | 5 |
| Abies alba       | 0.99     | 0.64     | Winter P                       | 0.36       | EXP        |       |                  |

All reported relationships were significant at the 0.05 level. Shape indicates the shape of the relationship (LIN, linear; EXP, exponential) and Ref is the reference. Note that the variability of fruit production was always higher than the variability of the meteorological variables for all exponential relationships. $T$, temperature; $P$, precipitation; max, maximum; min, minimum.

The literature search was not exhaustive, because our aim was to present a general overview of the relationship between seed production and weather and not to review all studies.

1 Pérez-Ramos et al. (2010); 2 Fernández-Martínez et al. (2012); 3 Köning et al. (2016); 4 Pérez-Ramos et al. (2013); 5 Fernández-Martínez et al. (2016).
efficiency (Fernández-Martínez et al., 2012; Koenig et al., 2015; Pesendorfer et al., 2016; Bogdziewicz et al., 2017a,b). We propose that rejecting weather as the most parsimonious driver of high interannual variability of seed production in mastling plants was premature and should be revised. We hypothesize that more knowledge could be gained from studying plant reproduction if we combined observational studies with experiments that modify the amount of interannual meteorological variability to which plants are subjected, amongst other factors (e.g., pollen dispersal). These kinds of experiments may more easily identify non-linear relationships and would certainly provide new insights into the response of the reproductive behavior of plants to meteorological variability (Figure 1).

AUTHOR CONTRIBUTIONS

MF, MB, JE, and JP conceived the paper. MF performed the statistical analyses and MB provided data. All authors contributed substantially to the writing of the paper.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fevo.2017.00134/full#supplementary-material

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