Global mean temperature is expected to significantly increase by the end of the twenty-first century and could have dramatic impacts on a plant's growth, physiology, and ecosystem processes. Temperature manipulative experiments have been conducted to understand the responsive pattern of plant ecophysiology to climate warming. However, it remains unknown how different methodology used in these experiments will affect plants' ecophysiological responses to warming. We conducted a comprehensive meta-analysis of the warming manipulative studies to synthesize the ecophysiological traits responses to warming treatment of different intensities, durations, and conducted for different species and under different experimental settings. The results indicated that warming enhanced leaf dark respiration ($R_d$) and specific leaf area (SLA) but decreased net photosynthetic rate ($A_{net}$) and leaf nitrogen content (LN). The positive and negative effects of warming on $R_d$ and $A_{net}$ were greater for C$_4$ species than C$_3$ species, respectively. The negative effect of warming treatment on $A_{net}$ and LN and the positive effect on $R_d$ were more evident under >1 year warming treatment. Negative effects of warming were more evident for plants grown at <10 L pots when experiment duration was longer than 1 year. The magnitude of warming treatment had a significant impact on most of the parameters that were investigated in the study. Overall, the results showed that warming effects on plant ecophysiological traits varied among different response variables and PFTs and affected by the magnitude of temperature change and experimental methodology. The results highlight the need for cautiously selecting the values of plant ecophysiological parameters in forecasting ecosystem function changes in future climate regimes and designing controlled experiments to realistically reflecting ecosystems responses to future global warming.

**Keywords:** photosynthesis, respiration, controlled experiment, climate warming, meta-analysis
INTRODUCTION

Based on the current trends in fossil energy production and use, deforestation, and population growth, it is expected that the increase of global mean surface temperatures for 2081–2100 relative to 1986–2005 is projected to be in the ranges of 0.3 to 1.7°C (RCP2.6), 1.1 to 2.6°C (RCP4.5), 1.4 to 3.1°C (RCP6.0), and 2.6 to 4.8°C (RCP8.5), which will have dramatic effects on economics, agriculture, and environment (AR5, IPCC, 2013). Plant traits are sensitive to climate warming and ecologists use plant trait-climate relationships to simulate plant physiology and growth in current and future climate situations (Farquhar and Sharkey, 1982; Wang et al., 2012; Jing et al., 2016). Therefore, understanding the patterns of plant physiological and morphological responses to global warming is of great importance in simulating and predicting the impact of global change on natural systems and agriculture.

Predictions of response to global warming may be derived from experimental and observational studies (Tilman, 1989; Wang et al., 2008, 2018; Knapp et al., 2012). While both types of study are common, relatively few authors have investigated whether they produce similar predictions or reflect reality (Dunne et al., 2004; Knapp et al., 2018). Experimental global change studies are typically limited in scope both spatially and temporally (Rustad et al., 2001). Observational studies often have broader spatial and temporal scales but suffer from a lack of control over covariates in biophysical and biochemical parameters of weather and soil. To minimize the weaknesses of each approach, it has been suggested that more research should explicitly unite observational and experimental work, perhaps by nesting experiments at multiple sites within a larger observational context or through summarized meta-analysis (Dunne et al., 2004; Jing et al., 2016).

Many manipulative experiments controlling physical and environmental factors have been conducted around the world to investigate the potential effects of global change on plants and terrestrial ecosystems (Sage and Kubien, 2007; Rustad, 2008; Wang et al., 2018). However, the methodology used in these experiments was different in their research settings, treatment intensities and durations and targeted species. The impact of short-term vs. long-term warming on plants traits would probably be different due to plants' acclimation capacity in photosynthesis, respiration and other physiological processes and these impacts would vary among different plant functional types (PFTs) under natural or controlled settings (Smith and Dukes, 2017). Plants' physiological and morphological responses to short-term warming treatment, however, are often used to parameterize the sub-models of photosynthesis, stomatal conductance, and respiration in plant growth and terrestrial ecosystem models, which would likely unrealistically simulate plant energy, carbon, and water fluxes in the long term. Indoor or outdoor settings and pot sizes could also affect the magnitude of ecophysiological responses to temperature increase by implicating root growth and plant above-ground and below-ground tissue interactions (Arp, 1991). To accurately predict the impacts of climatic change and develop proper adaptive agricultural management practices, it is imperative to understand how temperature changes of different intensities and duration and changes manipulated under different experimental settings affect photosynthetic carbon gain, loss and allocation through a comprehensive analysis of relevant studies.

Previous research and meta-analyses have indicated that global warming will promote plant photosynthesis, dark respiration, leaf nitrogen content, specific leaf area, and other metabolisms (Poorter et al., 2012). It has been reported that the modulation of leaf traits and trait relationships by site climatic properties was modest (Wright et al., 2005). However, the modulation of leaf traits by warming treatment of different intensities and duration has not been extensively analyzed. Understanding how these processes vary among different species and plant functional types is a major goal for plant ecology and crucial for modeling how nutrient fluxes and vegetation boundaries will shift under global warming. The effect of the intensities and the treatment duration of global warming manipulative experiments on the plant physiology and growth among different plant functional groups, however, remain unclear. Therefore, the main objective of this study was to investigate the effects of global warming treatment with different magnitudes and durations on plant response in ecophysiological traits. Specifically, we aim to: (1) assess the impact of global warming of different magnitudes and durations on plant ecophysiological traits at leaf level; (2) detect the variations of ecophysiological traits response of different plant functional types to warming treatment of different durations; (3) explore the effect of different experimental settings on the response of a plant's traits to global warming. Accordingly, we propose: (1) due to plant acclimation capacity, short-term vs. long-term warming has different impacts on plant traits, with short-term warming having a more stimulating effect on the physiological functions of plants; (2) different experimental facilities may change the response of plants traits to warming treatment. To test these hypotheses, we conducted a comprehensive meta-analysis of the warming manipulating studies published from 1980 to 2018.

MATERIALS AND METHODS

Data Collection

Journal articles were searched on the Web of Science database with the keyword “leaf traits & warming,” “leaf traits & temperature increase” and etc. The articles were later cross-checked with review articles and book chapters. The articles were imported into EndNote software and formed a database. All articles about warming effects on leaf traits were screened to ensure that all the articles available were included for the analysis. The articles published from 1980 to 2018 and meeting the following two conditions were included in the analysis: (1) the control group in the experiment was treated at ambient temperature situation; (2) physiological and morphological measurements were performed on both ambient and manipulated groups. Articles were rejected if: (1) plant physiological changes under warming treatments led to death of or severe damage to the plant; (2) there were other stressing factors impacting the warming treatments. Finally, 80 papers meeting the requirements were included in
the database (Supplementary Material S1). Data was obtained directly from the table or was extracted using the GetData Graph Digitizer software from the selected articles. In these studies, the magnitude of warming treatment ranged between 0.3 and 25°C, with only two studies showing a warming treatment above 20°C. The magnitude of warming treatment ranged between 0.3 and 25°C, and the warming treatments were evaluated by a second-degree polynomial or linear regression analysis with the R statistical programming language (R 3.2.2 for Windows GUI front-end).

Categorization of the Studies
Temperature treatment was divided into two categories: AT (ambient temperature) and ET (elevated temperature). Plant species were classified into different photosynthetic pathways (C3, C4, or CAM), growth forms (herb or wood) and economic values (crop or non-crop). Experimental facilities were categorized into indoor (growth chambers or greenhouses) and outdoor (open top chambers or fully-open) settings and <10L and >10L growing pots. In our dataset, exposure time (i.e., how long plants were exposed to warming) ranged from <10 days to >10 years. To analyze the possible different responses under various warming durations, we banded the temperature treatment into two categories: short-term (<1 year) and long-term (>1 year). Warming treatments that were applied through air warming were included in the analyses. We listed the species, PFTs information and relevant experimental methodology used in this study (Supplementary Material S1).

Meta-Analysis Methods
To avoid the adverse effects of different units, we used the response ratio \( r = \frac{X_t}{X_c} \) to estimate the magnitude of the effect of warming treatment, where \( X_t \) is the treatment mean and \( X_c \) is the control mean. For ease of comparison, we calculated the natural logarithm of the response ratio (lnr). The standard deviation (SD) and the sample size (n) for each observation were collected to calculate the variance of the effect size.

The lnr was calculated without and with being standardized by warming magnitude (Equations 1, 2).

\[
\log_{e} r = \log_{e} \left( \frac{X_t}{X_c} \right) = \log_{e}(X_t) - \log_{e}(X_c)
\]

\[
\log_{e} r = \frac{\log_{e} \left( \frac{X_t}{X_c} \right)}{T_t - T_c} = \frac{\log_{e}(X_t) - \log_{e}(X_c)}{T_t - T_c}
\]

where \( T_t \) and \( T_c \) are the temperature in the warming and control treatments, respectively.

Using METAWIN software 2.1 (Sinauer Associates, Inc. Sunderland, MA, USA), we calculated the effect size of the target variables and used a weighted fixed-effect model to assess the effect of plant functional types, experimental settings, and treatment duration. If the 95% confidence interval (CI) of the effect size produced by the fixed-effect model overlaps with 0, no significant effect was detected on the response variables. If the upper limit of 95% CI is less than 0, the effect is considered significantly negative. In contrast, if the lower limit of 95% CI is greater than 0, the effect is considered significantly positive. If the 95% CI of the effect size among different species, pot size, and treatment duration does not overlap, their response is considered significantly different. Unless otherwise indicated, significance level was set at \( p < 0.05 \). The publication bias for effect size (lnr) in this meta-analysis was also calculated. We calculated Spearman’s rank order correlation (rs) which indicates the relationship between the effect size (lnr) and the sample size (Begg and Mazumdar, 1994), and Rosenthal’s fail-safe number which represents the number of additional studies with a mean effect size of zero needed to eliminate the significance of a significant effect (Rosenthal, 1979). Publication bias was significant if \( p \)-value of rs was smaller than 0.05. However, the publication bias may be safely ignored if the fail-safe number is larger than a critical value of 5n+10 where n is the number of studies (Rosenberg, 2005).

Statistical Analysis
Original data collected from these two studies were arranged into a database in which the value of response variables was lnr. The effect of warming duration on lnr was considered significant if the 95% confidence interval (CI) of lnr does not overlap with 0. And when the 95% confidence intervals (CI) of lnr of different PFTs, facilities or pot size did not overlap with each other, the response was considered significantly different among different categories, the means of the ratio of the Rd/A_{net} and A_{net}/Gs in the control and warming treatments were compared using paired t-test. The relationship between lnr of all the variables and the magnitude of warming treatments were evaluated by a second-degree polynomial or linear regression analysis with the R statistical programming language (R 3.2.2 for Windows GUI front-end).
RESULTS

Effects of the Duration of Warming Treatment on Plant Ecophysiological Traits Across Plant Functional Types (PFTs) and Growth Forms

Warming treatment increased dark respiration ($R_d$) and specific leaf area (SLA) and decreased net photosynthetic rate ($A_{net}$) and leaf N concentration (LN) across all the experiments (Figure 1). The response of standardized (triangle symbols) or unstandardized (circle symbols) rate of $A_{net}$, $G_s$, $R_d$, LN, and SLA to warming treatment differed with different warming durations (Figure 2). Long-term warming treatment (>1 year) had a greater positive effect on $R_d$ than short-term (<1 year), regardless of whether the effect was standardized or unstandardized. LN was decreased by long-term warming but was increased or not changed by short-term warming treatment for unstandardized and standardized effect, respectively. Long-term warming treatment increased SLA, while short-term treatment had no effect on SLA. For standardized response of SLA, there was no difference between long and short-term treatments. For $G_s$, long-term treatment had a positive but short-term treatment had a negative effect on the standardized effect size. However, for the unstandardized effect size, short-term treatment did not have a significant but long-term had a negative effect on $G_s$. Short-term had a positive and long-term a negative effect on $A_{net}$ for the unstandardized form of the effect. And for standardized effect of $A_{net}$, long-term treatment had a more negative effect than short-term treatment (Figure 2).

The response of $A_{net}$, $G_s$, $R_d$, LN, and SLA to warming treatment differed among PFTs with different photosynthetic pathways (Figure 3). Warming had a more positive effect on $R_d$ for $C_4$ species than for $C_3$ species, regardless of whether the effect size was standardized. Warming had a negative effect for $C_3$ but a positive effect for $C_4$ species on LN, SLA, and $G_s$. In contrast, warming had a negative effect for $C_4$ but near-zero effect for $C_3$ species on $A_{net}$ (Figure 3).

Warming duration had a significant effect on the response of $A_{net}$, $G_s$, $R_d$, LN, and SLA for PFTs with different photosynthetic pathways (Figure 4). Long-term warming treatment had a more positive effect than short-term on $R_d$ for both $C_3$ and $C_4$ species, regardless of whether the effect was standardized. For LN, long-term treatment had a negative effect but short-term treatment had a positive effect for $C_3$ and $C_4$ species. For $C_3$ species, short-term warming treatment had a positive and long-term had a negative effect on $A_{net}$; for $C_4$ species, long-term warming treatment had a positive but short term a negative effect on $A_{net}$. Similar trend was found for standardized $A_{net}$, even though the magnitude of the effect differed.

Effects of Warming Duration on Plant Traits Across Different Experimental Settings

The responses of $A_{net}$, $G_s$, $R_d$, LN, and SLA to warming treatment differed among in-door and outdoor experimental settings (Figure 5). Warming had a more positive impact on $R_d$ in the in-door than the out-door settings for unstandardized effect size. Warming had a positive effect on LN for in-door, but a negative effect for outdoor settings. Being standardized with temperature treatment, warming had no impact on LN for the in-door but negative impact on outdoor experimental settings. Warming had a positive effect on SLA for in-door settings but tended to have a negative effect for outdoor settings. $G_s$ responded positively to warming under in-door but negatively under outdoor settings. Warming treatment had a positive effect on unstandardized $A_{net}$ under in-door settings but a negative effect under outdoor settings. For standardized $A_{net}$, warming had a negative effect for both in-door and outdoor settings (Figure 5).

The response of $A_{net}$, $G_s$, $R_d$, LN, and SLA to warming treatment under indoor and outdoor experiment settings also differed with different treatment durations (Figure 6). Short-term warming had a positive effect but long-term had a negative effect on $R_d$ for indoor experimental settings. Long-term warming had a more positive impact on $R_d$ than short-term for outdoor experimental settings for both standardized and unstandardized effect size. Short-term warming treatment had a more positive impact than long-term treatment on $A_{net}$ for unstandardized effect size but had no difference on standardized effect size. Short-term had a positive impact on $A_{net}$ for outdoor settings, but long-term treatment had a negative impact on $A_{net}$ for unstandardized effect. Long-term warming treatment had a more negative effect on standardized $A_{net}$ than short term for outdoor settings (Figure 6).

Pot size had a significant impact on the responses of $A_{net}$, $G_s$, $R_d$, and LN to warming treatment (Figure 7). Warming had a positive impact on $R_d$ for plants grown in pots larger than 10 L, while a negative effect for plants grown in pots smaller than 10 L. $G_s$ responded positively to warming when grown at <10 L plots but negatively at >10 L plots. $A_{net}$ of plants grown at >10 L plots responded negatively to warming. Warming had no impacts on unstandardized $A_{net}$ but a negative effect on standardized $A_{net}$ of plants grown at <10 L pots.

The response of $A_{net}$, $G_s$, LN, and SLA to warming treatment differed among different treatment durations when plants were grown in pots of different volumes (Figure 8). Short-term warming had a positive effect but long-term, a positive effect on LN for plants grown at both <10 L and >10 L pots. Short-term warming had a negative effect on SLA, but long-term a positive effect for both <10 L and >10 L pots. $G_s$ responded positively with both short and long-term warming treatments at <10 L pots but negatively at >10 L pots. $A_{net}$ responded positively to long-term warming treatment at <10 L pots but negatively at >10 L pots (Figure 8).

Effects of Warming Magnitude on Plant Traits Across Different Experimental Settings

$A_{net}$, $R_d$, LN, and SLA formed a quadratic relationship to warming treatment (Figure 9). The effect size of $A_{net}$, $R_d$, LN, and SLA to warming was highest or lowest when temperature change was 6.6, 2.5, 6.6, and 5.2°C above ambient temperature, respectively (Figure 9).
DISCUSSION

Several meta-analyses have investigated the general tendency of warming impacts on plant physiology and production (Rustad et al., 2001; Jing et al., 2016). However, it remains unclear how the experimental methodology of warming treatment affects the responses of plant ecophysiological traits to warming at leaf level. In this study, we collected data from warming manipulative studies and analyzed changes in the ecophysiological responses in the leaf traits. Overall, we found that (1) the direction and degree of the effect of warming treatment of different durations and settings on plant ecophysiological traits varied...
Warming Methodology Effects on Plants Traits

FIGURE 3 | Standardized (triangle symbols) and unstandardized (circle symbols) of net photosynthetic rate ($A_{\text{net}}$), stomatal conductance ($G_s$), leaf nitrogen content (LN), specific leaf area (SLA), and leaf dark respiration rate ($R_d$) of $C_3$ (closed symbols) and $C_4$ (open symbols) species to increased temperatures. Each data point represents the mean±95% CI. The number of observations for each variable is given on the right of the graph.

FIGURE 4 | Responses of net photosynthetic rate ($A_{\text{net}}$), stomatal conductance ($G_s$), leaf nitrogen content (LN), specific leaf area (SLA) and leaf dark respiration rate ($R_d$) of $C_3$ (closed symbols) and $C_4$ (open symbols) species to $<1$ year (circle symbols) and $>1$ year (triangle symbols) temperature treatment. Each data point represents the mean±95% CI. The number of observations for each variable is given on the right of the graph.

significantly; (2) there were significant variations among plant functional types in response to warming treatment of different methodology.

Consistent with previous findings from other studies, this meta-analysis confirmed that $R_d$ and SLA were stimulated by warming treatment (Rustad et al., 2001; Jing et al., 2016). Increasing, decreasing or neutral impacts of experimental warming have been observed for net photosynthetic rates (Bruhn et al., 2007; Bronson and Gower, 2010; Li et al., 2013). The net photosynthetic rate in this analysis was significantly decreased by warming treatment. The decrease in plant photosynthetic capacity may be attributed to the decreased LN under warmed conditions. Many studies showed that plant photosynthetic capacity was positively related to leaf N concentrations (Kattge et al., 2009; Reich et al., 2009). Compared with the negative effect of warming for non-legumes, there was
Warming Methodology Effects on Plants Traits

FIGURE 5 | Standardized (triangle symbols) and unstandardized (circle symbols) responses of net photosynthetic rate ($A_{\text{net}}$), stomatal conductance ($G_s$), leaf nitrogen content (LN), specific leaf area (SLA), and leaf dark respiration rate ($R_d$) to increased temperatures at in-door (closed symbols) and out-door (open symbols) experimental settings. Each data point represents the mean±95% CI. The number of observations for each variable is given on the right of the graph.

FIGURE 6 | Responses of net photosynthetic rate ($A_{\text{net}}$) and leaf dark respiration rate ($R_d$) to <1 year (circle symbols) and >1 year (triangle symbols) temperature treatment at in-door (closed symbols) and out-door (open symbols) experimental settings. Each data point represents the mean±95% CI. The number of observations for each variable is given on the right of the graph.

A positive or neutral effect on LN and $A_{\text{net}}$ for legume species (Supplementary Material S3). Contrary to the expectations, stomatal conductance remained unchanged under warming, thus highlighting the key roles of biochemical and nutritional limitations on the negative responses of net photosynthesis to warming treatment. The response of $G_s$ to global warming is critical for modeling ecosystem and landscape-scale water fluxes and CO$_2$ exchange. The ratio of respiration to photosynthesis ($R/P$) has been used to express the proportion of consumed to fixed C of plants (Atkin et al., 2007; Campbell et al., 2007) and shown to be enhanced (Danby and Hik, 2007; Wan et al., 2009), suppressed (Jochum et al., 2007), or maintained (He et al., 2005) by experimental warming. The ratio of $R_d/A_{\text{net}}$ was increased at warming conditions (effect size is 0.3623, $n = 275$) in this study, suggesting that the respiration was more affected and a greater proportion of fixed C was consumed, implying a decline of the net amount of C fixed by leaves by warming, at least in the controlled experiments.
Ecophysiological traits responses of terrestrial plants to increased temperature varied among plant functional types with different photosynthetic pathways (PFTs; Wang et al., 2012; Jing et al., 2016). Previous studies indicated that global warming had stronger effects on $A_{\text{net}}$ of $C_3$ species than $C_4$ species (Wahid et al., 2007). In this study, the positive and negative effects of warming on $R_d$ and $A_{\text{net}}$ were greater for $C_4$ species than $C_3$ species, in spite of positive or neutral effects of warming on LN, SLA, and $G_s$ for $C_4$ and $C_3$ species, respectively. The contradictory findings posed great challenges for projecting the responses and feedbacks of terrestrial ecosystems to global warming. The more disadvantaged situation for $C_4$ species under warming might be associated with higher growth and treatment temperature applied in the experiment (Supplementary Material S1). The metabolic balance of the photosynthetic and respiratory processes under climate warming plays a critical role in regulating ecosystem carbon storage and cycling (Schimel, 1995; King et al., 2006).

Warming stimulated $A_{\text{net}}$ in woody but suppressed it in herbaceous plants (Supplementary Material S4).
Effect of warming on $A_{\text{net}}$ for woody species was unrelated to either $G_s$ or LN, as $G_s$ and LN both were decreased under warming treatments (Supplementary Material S4). The results from this study were similar to the trend reported for trees showing a lower percentage decrease in $G_s$ compared to herbaceous species (Wang et al., 2012). Warming had a positive effect on $G_s$ and LN for crops, while a negative effect for non-crops (Supplementary Material S5). The changes in $G_s$ at warming treatment may alter leaf temperature and result in a change in latent heat loss through evaporation, which may further affect net carbon balance (Warren et al., 2011). Warming could influence vegetation dynamics and ecosystem structure through shifting competitive interactions among different functional groups in natural or agricultural systems. Therefore, knowledge of photosynthetic and stomatal responses to increased temperature of different PFTs instead of species will facilitate the prediction of terrestrial C- and water-cycle feedback to climate warming.

Ecophysiological trait responses of terrestrial plants to increased temperature varied among warming treatments of differing durations. The physiological acclimation can lead to smaller enhancements of plant photosynthesis and respiration under long term warmer conditions than predicted with photosynthesis/respiration-temperature relationships (Medlyn et al., 2002; Dwyer et al., 2007; Tjoelker and Zhou, 2007; Gunderson et al., 2010). The thermal acclimation of $R_d$ could minimize the effects of climate warming on C loss via plant respiration (Gifford, 1995; Ziska and Bunce, 1998; Loveys et al., 2002) and mitigate the positive feedback between climate change and atmospheric CO$_2$ (King et al., 2006; Atkin et al., 2008). The findings in this meta-analysis indicated that the negative effect of warming treatment on $A_{\text{net}}$ and LN and the positive effect on $R_d$ were more evident under >1 year warming treatment and the trend was confirmed for both C$_3$ and C$_4$ species (Figure 4), which contrasted to other studies showing significant declines in the photosynthetic and/or respiratory response with increasing exposure time, a thermal acclimation to warming (Hikosaka et al., 2006; Gunderson et al., 2010).

Potential confounding factors must be accounted in the meta-analysis because many studies were conducted under variable conditions and targeted on different species. In this analysis, studies in which plants were grown under other environmental stresses such as drought, low nutrients, light deficiency or elevated ozone were excluded. In addition to the variation caused by plant functional types and treatment duration, different experimental facilities could be responsible for the responses of different PFTs (Cheesman and Klaus, 2013; Rehmani et al., 2014). This study mainly focused on the effects of pot size (<10 L vs. >10 L) and experimental settings (in-door vs. out-door) on plant ecophysiological responses. Warming had a negative and positive effect on LN and $G_s$ when plants were grown at outdoor and in-door settings, respectively. Pot size significantly altered the responses of $R_d$, LN and SLA to warming treatments. Warming had a negative effect on $R_d$ for plants grown at <10 L pots, while a positive effect at >10 L pot. For both LN and $G_s$, warming had a negative effect for plants grown at >10 L pots, while a neutral effect at <10 L pot. We were expecting that warming would have a more negative effect on LN and $G_s$ in smaller pots or in-door settings considering that below-ground growth would be more constrained and thus limited the nutrients and water supply to the aboveground growth (Walters and Reich, 1989; Climent et al., 2011), the analysis indicated that this was true only when...
experiment duration was longer than 1 year when negative effects of warming were more evident for plants grown at <10 L pots.

Warming treatment duration had a significant interactive effect with experimental settings (in-door vs. outdoor) on R_d and A_net. Long-term warming had a negative effect on R_d for in-door and on A_net for outdoor experimental settings. The findings in this meta-analysis indicated that the negative effect of warming treatment on A_net and LN and the positive effect on R_d were more evident under >1 year warming treatment and the trend was confirmed for both C_3 and C_4 species (Figure 4). The negative effect of warming on R_d could be related to the higher treatment temperature applied at the in-door settings (Supplementary Material S2). Temperature conditions in which plants live may be another possible reason for the contradictory findings (Rustad et al., 2001). The discrepancy of the response of A_net and R_d to warming treatment under different experimental settings provided difficulty in parameterizing ecosystem models and raised concerns in proper experimental designs when dealing with climate change questions.

The intensities of temperature treatment also had a significant impact on most of the parameters that were investigated in the study. The effect size of A_net, R_d, LN, and SLA responded to temperature increase in a quadratic relationship. Consistent with the results discussed before, the peak value of the ecophysiological traits of A_net, R_d and LN occurred at temperatures higher than the ambient. Plant physiological responses to warming may also depend on the temperature regime they are grown at. Studies often report a positive response to warming in Rubisco carboxylation, photosynthesis, and growth in cool-climate species but reduced growth and carbon gain in species that exist in warm low-latitude climates (Way and Oren, 2010; Crous et al., 2018).

CONCLUSION

Overall, we found that warming treatment of different durations and settings had different impacts on plant ecophysiological traits and the responses varied significantly among different plant functional types. Warming stimulated R_d and SLA but suppressed A_net and LN and the effect varied among different PFTs and experimental designs. The positive and negative effects of warming on R_d and A_net, were greater for C_4 than C_3 species, in spite of the positive or neutral effects of warming on LN, SLA, and G_S for C_4 and C_3 species, respectively. The findings in this meta-analysis also indicated that the negative effect of warming treatment on A_net and LN and the positive effect on R_d were more evident under >1 year warming treatment and the trend was confirmed for both C_3 and C_4 species. Negative effect of warming was more evident for plants grown at <10 L pots only when experiment duration was longer than 1 year. The magnitude of temperature treatment did have an impact on most of the parameters that were investigated in the study. The functional type specific response patterns of plant traits to warming are critical for obtaining credible predictions of the changes in food production, carbon sequestration and climate regulation. This result also highlights the need for cautiously selecting parameter values in forecasting ecosystem function changes in future climate regimes, evaluating much more broadly what can and cannot be learned from experimental studies and designing controlled experiments to realistically reflecting ecosystems responses to future global warming.

DATA AVAILABILITY

All datasets for this study are included in the manuscript and the Supplementary Files.

AUTHOR CONTRIBUTIONS

DW and ZY conceived and wrote the paper. The rest of the authors helped collection data and ran data analysis.

ACKNOWLEDGMENTS

Funding for this research was provided by The National Natural Science Foundation of China (31500503 & 31770485), Nanjing University of Information Science and Technology (2013r115), Jiangsu Distinguished Professor Scholarship, Jiangsu six talent peaks (R2016L15), Jiangsu Natural Science Foundation (BK20150894), and the Jiangsu Overseas Research & Training Program for University Prominent Young & Middle-aged Teachers and Presidents through DW.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2019.00957/full#supplementary-material

REFERENCES

An, Y. A., Wan, S. Q., Zhou, X. H., Subedar, A. A., Wallace, L. L., and Luo, Y. (2005). Plant nitrogen concentration, use efficiency, and contents in a tallgrass prairie ecosystem under experimental warming. Glob. Chang. Biol. 11, 1733–1744. doi: 10.1111/j.1365-2486.2005.01030.x

Anderson, L., and Cipollini, J. (2013). Gas exchange, growth, and defense responses of invasive Allaria Petiolata (Brassicaceae) and native Geum Vurnum (Rosaceae) to elevated atmospheric CO_2 and warm spring temperatures. Am. J. Bot. 100, 1544–1554. doi: 10.3732/ajb.1300014

Arp, W. J. (1991). Effects of source-sink relations on photosynthetic acclimation to elevated CO_2. Plant Cell Environ. 14, 869–875. doi: 10.1111/j.1365-3040.1991.tb01450.x

Atkin, O. K., Atkinson, L. J., Fisher, R. A., Campbell, C. D., Zaragoza-Castells, J., Pitchford, J. W., et al. (2008). Using temperaturedependent changes in leaf scaling relationships to quantitatively account for thermal acclimation of respiration in a coupled global climate-vegetation model. Glob. Chang. Biol. 14, 2709–2726. doi: 10.1111/j.1365-2486.2008.01664.x

Atkin, O. K., Scheurwater, I., and Pons, T. L. (2007). Respiration as a percentage of daily photosynthesis in whole plants is homeostatic at
moderate, but not high, growth temperatures. *New Phytol.* 197, 367–380. doi: 10.1111/j.1469-8137.2007.02011.x

Begg, C. B., and Mazumdar, M. (1994). Operating characteristic of a rand correlation test for publication bias. *Biometrics* 50, 1088–1101. doi: 10.2307/2534446

Bronson, D. R., and Gower, S. T. (2010). Ecosystem warming does not affect photosynthesis or aboveground autotrophic respiration for boreal black spruce. *Tree Physiol.* 30, 441–449. doi: 10.1093/treephys/tqp001

Bruhn, D., Egerton, J. J. G., Lovesey, R. B., and Ball, M. C. (2007). Evergreen leaf respiration acclimates to long-term nocturnal warming under field conditions. *Glob. Chang. Biol.* 13, 1216–1223. doi: 10.1111/j.1365-2486.2007.01351.x

Campbell, C., Atkinson, L., Zaragoza-Castells, J., Lundmark, M., Atkin, O., and Hurr, V. (2007). Acclimation of photosynthesis and respiration is asynchronous in response to changes in temperature regardless of plant functional group. *New Phytol.* 176, 375–389. doi: 10.1111/j.1469-8137.2007.02183.x

Cheesman, A. W., and Klaus, W. (2013). Elevated night-time temperatures increase growth in seedlings of two tropical pioneer tree species. *New Phytol.* 197, 1185–1192. doi: 10.1111/nph.12098

Climent, J., Chambl, M. R., Pardos, M., Lario, F., and Villar-Salvador, P. (2011). Biomass allocation and foliage heteroblasty in hard pine species respond differentially to reduction in rooting volume. *Eur. J. For. Res.* 130, 841–850. doi: 10.1007/s10342-010-0476-y

Crous, K. Y., Drake, J. E., Aspinwall, M. J., Sharwood, R. E., Tjoelker, M. G., and Ghannoum, O. (2018). Photosynthetic capacity and leaf nitrogen decline along a controlled climate gradient in provenances of two widely distributed *Eucalyptus* species. *Glob. Change Biol.* 24, 4626–4644. doi: 10.1111/gcb.14330

Danby, R. K., and Hik, D. S. (2007). Responses of white spruce (*Picea glauca*) to experimental warming at a subarctic alpine treeline. *Glob. Chang. Biol.* 13, 437–451. doi: 10.1111/j.1365-2486.2006.01302.x

Drake, J. E., Aspinwall, M. J., Pfuttsch, S., Rymer, P. D., Reich, P. B., Smith, R. B., et al. (2015). The capacity to cope with climate warming declines from experimental and gradient methods in ecological climate change research. *Tree Physiol.* 35, 53–66. doi: 10.1093/treephys/20.2.87

Duan, B. L., Zhang, X. L., Li, Y. P., Li, L., Korpelainen, H., and Li, C. (2013). Temperature acclimation of photosynthesis: mechanisms involved in the changes in temperature dependence of photosynthetic rate. *J. Exp. Bot.* 57, 291–302. doi: 10.1093/jxb/erj049

IPCC (2013). *Climate Change 2013 AR5: The Physical Science Basis*. Intergovernmental Panel on Climate Change. Cambridge: Cambridge University Press, 548.

Jacques, M., Lapointe, L., Rice, K., and Montgomery, R. A., Stefanski, A., and Reich, P. B. (2015). Responses of two understory herbs, *Maianthemum canadense* and *Eurybia macrophylla*, to experimental forest warming: early emergence is the key to enhanced reproductive output. *Am. J. Bot.* 102, 1610–1624. doi: 10.3732/ajb.1500046

Jing, P., Wang, D., Zhu, C., and Chen, J. (2016). Plant physiological, morphological and yield-related responses to night temperature changes across different species and plant functional types. *Front. Plant Sci.* 7.1774.

Jochum, G. M., Mudge, K. W., and Thomas, R. B. (2007). Elevated temperatures increase leaf senescence and root secondary metabolite concentrations in the understory herb *Panax quinquefolius* (*Araliaceae*). *Am. J. Bot.* 94,819–826. doi: 10.3732/ajb.94.5.819

Kattinge, J., Knoor, W., Raddatz, T., and Wirth, C. (2009). Quantifying photosynthetic capacity and its relationship to leaf nitrogen content for global-scale terrestrial biosphere models. *Glob. Chang. Biol.* 15, 976–991. doi: 10.1111/j.1365-2486.2008.01744.x

King, A. W., Gunderson, C. A., Post, W. M., Weston, D. J., and Wullschleger, S. D. (2006). Atmosphere: plant respiration in a warmer world. *Science* 312, 536–537. doi: 10.1126/science.1114166

Knapp, A. K., Carroll, C. J. W., Griffin-Nolan, R. J., Suttle, I. J., Chaves, F. A., Baur, L. E., et al. (2018). A reality check for climate change experiments: do they reflect the real world? *Ecology* 99, 2145–2151. doi: 10.1002/ecy.2474

Kroner, Y., and Way, D. A. (2016). Carbon fluxes acclimate more strongly to elevated growth temperatures than to elevated CO$_2$ concentrations in a northern conifer. *Glob. Chang. Biol.* 22, 2931–2928. doi: 10.1111/gcb.13215

Levin-Sanchez, L., Nicolas, E., Nortes, P. A., Maestre, F. T., and Querejeta, J. I., et al. (2016). Photosynthesis and growth reduction with warming are driven by nonstomatal limitations in a Mediterranean semi-arid shrub. *Ecol. Evol.* 6, 2725–2738. doi: 10.1002/ece3.2074

Liu, Y., Zhang, Y., Zhang, X., Korpelainen, H., Berninger, F., and Li, C. (2013). Effects of elevated CO$_2$ and temperature on photosynthesis and leaf traits of an understory dwarf bamboo in subalpine forest zone, China. *Physiol. Plant* 148, 261–272. doi: 10.1111/j.1399-3054.2012.01705.x

Lovesey, B. R., Scheurwater, I., Pons, T. L., Fitter, A. H., and Atkin, O. K. (2002). Growth temperature influences the underlying components of relative growth rate: an investigation using inherently fast- and slow-growing plant species. *Plant Cell Environ.* 25, 975–987. doi: 10.1046/j.1365-3040.2002.00879.x

Maenpää, M., Riihokon, J., Konttinen-Soppela, S., Rouss, M., and Oksanen, E. (2011). Vertical profiles reveal impact of ozone and temperature on carbon assimilation of *Betula pendula* and *Populus tremula*. *Tree Physiol.* 31, 808–818. doi: 10.1093/treephys/trp675

Medlynn, B. E., Dreyer, E., Ellsworth, D., Forstreuter, M., Harley, P. C., Kirschbaum, M. U. F. et al. (2002). Temperature response of parameters of a biochemically based model of photosynthesis. II. A review of experimental data. *Plant Cell Environ.* 25, 1167–1179. doi: 10.1046/j.1365-3040.2002.00891.x

Niu, S., Li, Z., Xia, J., Han, Y., Wu, M., and Wan, S. (2008). Climatic warming changes plant photosynthesis and its temperature dependence in a temperate steppe of northern China. *Environ. Exp. Bot.* 63, 91–101. doi: 10.1016/j.envexpbot.2007.10.016

Poorter, H., Niklas, K. J., Reich, P. B., Oleksyn, J., Poot, M., and Pommersheim, L. (2012). Biomass allocation to leaves, stems and roots: meta-analyses of interspecific variation and environmental control. *New Phytol.* 193, 30–50. doi: 10.1111/j.1469-8137.2011.03952.x
Quesada, J. A., Sinosic, T., and Wright, I. J. (2009). Leaf photosynthesis influences the photosynthesis-nitrogen relation: a cross-biome analysis of 314 species. Oecologia 160, 207–212. doi: 10.1007/s00442-009-1291-3

Rudiger, R. G., and Baker, J. E. (2013). Carbon transfer in aquatic ecosystems. In *Aquatic ecosystem processes: a synthesis*. Eds. R. T. P. and R. P. (Boca Raton, FL: CRC Press), 103–136. doi: 10.1201/b16687-7

Rustad, L. E. (2008). The response of terrestrial ecosystems to global climate warming. *Ecol. Evol.* 9, 531–541. doi: 10.1093/jpe/rtw010

Quesada, J. A., Sinosic, T., and Wright, I. J. (2009). Leaf photosynthesis influences the photosynthesis-nitrogen relation: a cross-biome analysis of 314 species. Oecologia 160, 207–212. doi: 10.1007/s00442-009-1291-3

Rudiger, R. G., and Baker, J. E. (2013). Carbon transfer in aquatic ecosystems. In *Aquatic ecosystem processes: a synthesis*. Eds. R. T. P. and R. P. (Boca Raton, FL: CRC Press), 103–136. doi: 10.1201/b16687-7

Rustad, L. E. (2008). The response of terrestrial ecosystems to global climate change: towards an integrated approach. *Sci. Total Environ.* 404, 222–235. doi: 10.1016/j.scitotenv.2008.04.050

Sage, R. F., and Kubien, D. S. (2007). The temperature response of C3 and C4 photosynthesis. *Plant Cell Environ.* 30, 1086–1106. doi: 10.1111/j.1365-3040.2007.01682.x

Schimel, D. S. (1995). Terrestrial ecosystems and the carbon-cycle. *Glob. Chang. Biol.* 1, 77–91. doi: 10.1038/1365-2486.1995.800008.x

Shen, H., Klein, J. A., Zhao, X., and Tang, Y. (2009). Leaf photosynthesis and simulated carbon budget of Gentiana straminea from a decade-long warming experiment. *J. Plant Ecol.* 2, 207–216. doi: 10.1093/jpe/rtp025

Smith, N. G., and Dukes, J. S. (2017). Short-term acclimation to warmer temperatures accelerates leaf carbon exchange processes across plant types. *Glob. Chang. Biol.* 23, 4840–4853. doi: 10.1111/gcb.13735

Song, B., Niu, S., and Wan, S. (2016). Precipitation regulates plant gas exchange and its long-term response to climate change in a temperate grassland. *J. Plant Ecol.* 9, 531–541. doi: 10.1093/jpe/rtw010

Song, X., Wang, Y., and Lv, X. (2016). Responses of plant biomass, photosynthesis and lipid peroxidation to warming and precipitation change in two dominant species (*Stipa grandis* and * Leymus chinensis*) from North China Grasslands. *Ecol. Evol.* 6, 1871–1882. doi: 10.1002/eee.31982

Tian, Y., Chen, J., Chen, C., Deng, A., Song, Z., Zheng, C., et al. (2012). Warming impacts on winter wheat phenophase and grain yield under field conditions in Yangtze Delta Plain, China. *Field Crops Res.* 134, 193–199. doi: 10.1016/j.fcr.2012.05.013

Tilman, D. (1989). “Ecological experimentation: strengths and conceptual problems,” in *Long-Term Studies in Ecology*, ed G. E. Likens (New York, NY: Springer), 136–157. doi: 10.1007/978-1-4615-7358-6_6

Tjoelker, M. G., and Zhou, X. (2007). The many faces of climate warming. *New Phytol.* 176, 739–742. doi: 10.1111/j.1469-8137.2007.02279.x

Tjoelker, M. G., Oleksyn, J., Lorenz-Plucinska, G., and Reich, P. B. (2009). Acclimation of respiratory temperature responses in northern and southern populations of * Pinus banksiana*. *New Phytol.* 181, 218–229. doi: 10.1111/j.1469-8137.2008.02624.x

Valencia, E., Quero, J. L., and Maestre, F. T. (2016). Functional leaf and size traits determine the photosynthetic response of 10 dryland species to warming. *J. Plant Ecol.* 9, 773–783. doi: 10.1093/jpe/rtv081

Wahid, A., Gelani, S., Ashraf, M., and Foolad, M. R. (2007). Heat tolerance in plants: an overview. *Environ. Exp. Bot.* 61, 199–223. doi: 10.1016/j.envexpbot.2007.05.011

Walters, M. B., and Reich, P. B. (1989). Response of Ulmus americana seedlings to varying nitrogen and water status. 1 Photosynthesis and growth. *Tree Physiol.* 5, 159–172. doi: 10.1093/treephs/5.2.159

Wan, S., Xia, J., Liu, W., and Niu, S. (2009). Photosynthetic overcompensation under nocturnal warming enhances grassland carbon sequestration. *Ecology* 90, 2700–2710. doi: 10.1890/08-2261

Wang, D., Bariua, D., Joshi, P., LaCroix, J., Hamilton, E. W., and Heckathorn, S. A. (2008). Effects of elevated CO2 on the tolerance of photosynthesis to acute heat stress in 11 (C3, C4, and CAM) species. *Am. J. Bot.* 95, 1–13. doi: 10.3732/ajb.95.2.165

Wang, D., Heckathorn, S. A., Wang, X., and Philpot, S. M. (2012). A meta-analysis of plant physiological and growth responses to temperature and elevated CO2. *Oecologia* 169, 1–13. doi: 10.1007/s00442-011-2172-0

Wang, D., Ling, T., Wang, P., JIng, P., Fan, J., Wang, H., et al. (2018). Effects of 8-year nitrogen and phosphorus treatments on the ecophysiological traits of two key species on Tibetan Plateau. *Front. Plant Sci.* 9:1290. doi: 10.3389/fpls.2018.01290

Warren, J. M., Norby, R. J., and Wullschleger, S. D. (2011). Elevated CO2 enhances leaf senescence during extreme drought in a temperate forest. *Tree Physiol.* 31, 117–130. doi: 10.1093/treephs/tpr002

Way, D. A., and Oren, R. (2010). Differential responses to changes in growth temperature between trees from different functional groups and biomes: a review and synthesis of data. *Tree Physiol.* 30, 669–688. doi: 10.1093/treephs/tqp015

Wright, I. J., Reich, P. B., Cornelissen, J. H. C., Falster, D. S., Groom, P. K., Hikosaka, K., et al. (2005). Modulation of leaf economic traits and trait relationships by climate. *Glob. Ecol. Biogeogr.* 14, 411–421. doi: 10.1111/j.1466-822x.2005.00172.x

Xiong, F. S., Mueller, E. C., and Day, T. A. (2000). Photosynthetic and respiratory acclimation and growth response of Antarctic vascular plants to contrasting temperature regimes. *Am. J. Bot.* 87, 700–710. doi: 10.2307/2656856

Xu, X., Peng, G., Wu, C., and Han, Q. (2010). Global warming induces female cuttings of * Populus cathayana* to allocate more biomass, C and N to aboveground organs than do male cuttings. *Asiat. J. Bot.* 58, 519–526. doi: 10.1071/BT10108

Xu, X., Yang, F., Xiao, X. W., Zhang, S., Korpeleim, H., and Li, C. (2008). Sex-specific responses of *Populus cathayana* to drought and elevated temperatures. *Plant Cell Environ.* 31, 850–860. doi: 10.1111/j.1365-3040.2008.01799.x

Ziska, L. H., and Bunce, J. A. (1998). The influence of increasing growth temperature and CO2 concentration on the ratio of respiration to photosynthesis in soybean seedlings. *Glob. Chang. Biol.* 4, 637–643. doi: 10.1046/j.1365-2486.1998.00179.x

Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2019 Wang, Wang, Wang, Ling, Tao and Yang. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.