

Impact of elevated CO₂ in Casuarina equisetifolia rooted stem cuttings inoculated with Frankia

Arumugam Karthikeyan¹

Received: 19 April 2016 /Accepted: 4 September 2016 / Published online: 10 September 2016 \oslash Springer Science+Business Media Dordrecht 2016

Abstract Impact of different levels of elevated $CO₂$ on the activity of Frankia (Nitrogen-fixing actinomycete) in Casuarina equisetifolia rooted stem cuttings has been studied to understand the relationship between C. equisetifolia, *Frankia* and $CO₂$. The stem cuttings of *C. equietifolia* were collected and treated with 2000 ppm of Indole Butyric Acid (IBA) for rooting. Thus vegetative propagated rooted stem cuttings of C. equisetifolia were inoculated with Frankia and placed in the Open top chambers (OTC) with elevated $CO₂$ facilities. These planting stocks were maintained in the OTC for 12 months under different levels of elevated $CO₂$ (ambient control, 600 ppm, 900 ppm). After 12 months, the nodule numbers, bio mass, growth, and photosynthesis of C. equisetifolia rooted stem cuttings inoculated with Frankia were improved under 600 ppm of $CO₂$. The rooted stem cuttings of C. equisetifolia inoculated with Frankia showed a higher number of nodules under 900 ppm of $CO₂$ and cuttings without Frankia inoculation exhibited poor growth. Tissue Nitrogen (N) content was also higher under 900 ppm of $CO₂$ than ambient control and 600 ppm levels. The photosynthetic rate was higher (17.8 μ mol CO₂ m⁻² s⁻¹) in 900 ppm of CO₂ than in 600 ppm (13.2 μ mol CO₂ m⁻² s⁻¹) and ambient control (8.3 μ mol CO₂ m⁻² s⁻¹). This study showed that Frankia can improve growth, N fixation and photosynthesis of C. equietifolia rooted stem cuttings under extreme elevated $CO₂$ level conditions (900 ppm).

Keywords Casuarina equisetifolia \cdot Frankia \cdot CO₂ \cdot Nodulation . N fixation

1 Introduction

Increase of carbon dioxide $(CO₂)$ and other green house gases in atmosphere due to burning of fossil fuels, clearing forests and converting lands for industrial purpose results in global warming and climate change. It was predicted that the amount of $CO₂$ in the atmosphere is rising by approximately 3 Pg carbon per year (UNESCO/UNEP [2011\)](#page-5-0). The recent report of NOAA ([2016\)](#page-5-0) stated that at present (May 2016) the $CO₂$ concentration in the atmosphere is 407.70 ppm. To mitigate the global warming through carbon sequestration, studies are being undertaken worldwide particularly on afforestation, reforestation and reclamation of waste lands with suitable tree species. However, studies on microorganisms are equally important to reduce the $CO₂$ levels as the soil microorganisms contribute significantly in the consumption of greenhouse gases such as $CO₂$, methane (CH₄), nitrous oxide (N₂O), and nitric oxide (NO) (Wiley et al. [2009\)](#page-5-0). For e.g. it was reported that a bacteria Methylokorus infermorum consuming methane about 11 kg/year for their energy and multiplication (Jenkinson et al. [1991\)](#page-5-0). Similarly, mycorrhizal fungi consumes 10–20 % of photosynthetically fixed carbon from plants for their survival in plant roots (Staddon et al. [1999](#page-5-0)) particularly under elevated $CO₂$ conditions (Quoreshi et al. [2003\)](#page-5-0). These microbial symbionts associated with plants contribute to carbon sequestration by increasing nutrient uptake in plants (Garcia et al. [2011\)](#page-5-0). Plants rely upon microbial symbionts like mycorrhizal fungi and symbiotic nitrogen fixing bacteria to acquire nutrients such as phosphorus (P) and nitrogen (N) for their growth and metabolism. These microbial symbionts scavenge nutrients from soils and transfer to the host

 \boxtimes Arumugam Karthikeyan karthika@icfre.org; karthikarumugam13@gmail.com

¹ Institute of Forest Genetics and Tree Breeding, Coimbatore 641 002, India

plant and in turn the symbionts obtain carbohydrates from the host plant (Hodge [1996\)](#page-5-0). In an experimental work total biomass, root biomass and mycorrhizal colonization of Quercus alba and Pinus echinata seedlings were increased under elevated $CO₂$ (O'Neill et al. [1987\)](#page-5-0). It was also reported that N fixing plants respond positively to elevated $CO₂$ than other plants due to their high nutrient demand (Temperton et al. [2003\)](#page-5-0) and the N- fixing plants improved their nutrient supply through N fixing bacteria under elevated $CO₂$ (Arnone and Gordon [1990;](#page-5-0) Vogel et al. [1997\)](#page-5-0). Trees under elevated CO₂ also showed increased growth and photosynthesis due to high nutrient supply through microbial symbionts (Ceulemans et al. [1999](#page-5-0)). Hence it was understood that the microbial symbionts facilitate to sequestrate the carbon in plants. Microbial symbionts also stimulate host plant photosynthesis to a greater extent at elevated $CO₂$ than at ambient $CO₂$ (Staddon et al. [1999\)](#page-5-0). This was also confirmed by Tissue et al. [\(1997\)](#page-5-0) as they found increased photosynthetic rates and carbon storage in Gliricidia sepium inoculated with Rhizobium sp. under elevated CO2. Based on these informations, a study has undertaken to determine the inoculation effect of Frankia in Casuarina equisetifolia rooted stem cuttings under elevated $CO₂$ to find out the response of N fixation and biomass improvement in C. equisetifolia.

Frankia is a symbiotic actinomycete which associates with C. equisetifolia and form N fixing root nodules. As part of the symbiotic relationship with Frankia, C. equisetifolia can fix N up to 300 kg ha^{-1} year^{-1} (Wheeler and Miller [1990](#page-5-0)) and in return for the fixed N, the tree supply carbon to the symbiotic bacteria (Santi et al. [2013](#page-5-0)). This tree is used in agro forestry system along with vegetable and pulse crops in India. It grows up to 50 m height with 50 cm girth and the final yield is within 3.5 to 4 years. The annual production of pulp wood alone from C. equiseitifolia is 10 million tonnes that worth of \$ 300,000 (Karthikeyan et al. [2009\)](#page-5-0). At present the poles of C. equisetifolia costs \$ 100–120 /tonne in India. It is also used asfuel wood, poles for services like shelterbelts, windbreaks, rehabilitating mine spoils and nutrient poor areas (Diagne et al. [2013\)](#page-5-0). This tree was also recorded for good nutrient turnover through litter decomposition (Uma et al. [2014\)](#page-5-0). However, there are no earlier reports on effect of $CO₂$ on Frankia and C. equisetifolia association. The relationship between elevated $CO₂$ and C. equisetifolia in the presence and absence of Frankia will be helpful to understand the impact of elevated $CO₂$ on the growth and photosynthesis of C. equisetifolia.

2 Materials and methods

2.1 Culture of Frankia

Root nodules of C. equisetifolia were collected from the mature trees at farm fields of Coimbatore, India. The nodules

were transported in an ice box and stored at −4 °C and surface sterilized with 30 % H_2O_2 . Later, the nodules were kept at room temperature for 30–40 min. Under aseptic conditions the nodules were rinsed in sterile distilled water and 0.2 g of nodule was ground manually in a sterile mortar and pestle. The nodule solution was centrifuged at 1000 rpm for 20 min and the supernatant was filtered through Whatman No.1 filter paper. The suspension was then spread on P media* (Shipton and Burgraff [1983](#page-5-0)) plates and incubated at 25 °C for 3– 4 weeks. *(One litre of P medium contained: 10 g $CaCl₂, 2H₂O$, 20 g MgSO₄, 0.46 g propionic acid, 0.15 g H3BO3, 0.15 g ZnSO4.7H2O, 0.45 g MnSO4. H2O, 0.004 g $CuSO₄.5H₂O$, 0.028 g Na₂MoO₄.2H₂O, 0.009 g CaCl₂.6H₂O, 0.04 g Biotin, 100 g K_2HPO_4 , 67 g NaH₂PO.2H₂O, 0.1 g FeNa EDTA, and 8.g agar; pH of the medium is 6.8). After 25 days of incubation, the Frankia growth appeared as fluffy white cloudy colonies. These colonies were transferred in to P media broth for scaling up the inoculum.

2.2 Propagation of rooted stem cuttings

The stem cuttings of *C. equisetifolia* were obtained from the Casuarina germplasm bank at Model Nursery of Institute of Forest Genetics and Tree Breeding, Coimbatore, India. Uniform sized (5 cm length: 1 mm girth) stem cuttings with 10 g (± 0.8) of total biomass were treated with 0.1 % carbendazim fungicide for 3 min. The cuttings were later treated with 2000 ppm of IBA $(40 \text{ mg of IBA} + 20 \text{ g of } 1$ powder) by immersing the basal end of the cuttings in the hormonal solution for 0.5 min. The treated cuttings were then placed in 100 cm^3 root trainers containing the inert vermiculite. The rooted stem cuttings were thereafter placed in polytunnels made of polythene sheets (180 cm \times 90 cm) and maintained under a temperature range of 32–35 °C and 60– 65 % relative humidity for 30 days for the development of roots. Previously the stem cuttings of C. equisetifolia were analysed for major reserved tissue nutrients according Jackson ([1973](#page-5-0)).

2.3 Inoculation of Frankia

After the development of adventitious roots, Frankia was inoculated at the rate of 10 ml /rooted stem cutting. The rooted stem cuttings of C. equisetifolia grown in 100 cm^3 root trainers with or without Frankia inoculation were placed in Open Top Chambers (OTC) and maintained for 12 months from April 2014 to March 2015. These OTC are cubical structures of $3 \times 3 \times 3$ m dimension fabricated with galvanized iron pipe frame and covered with polyviny chloride sheet. The upper part of the chamber was uncovered to maintain the atmospheric conditions. A software facility called supervisory control and data acquisition (SCADA) was used to control the $CO₂$ supply.

The control and *Frankia* inoculated rooted stem cuttings were replicated at 10 times consists of 5 rooted stem cuttings/ replicate (Totally 50 rooted stem cuttings/treatment). The rooted stem cuttings of C. equisetifolia were watered daily however, no fertilizers were added. Three OTC were used for this study viz., (i) OTC with 600 ppm $CO₂$ supply /day (ii) OTC with 900 ppm $CO₂$ supply/day (iii) and an ambient $CO₂$ controlled chamber. 598 (\pm 2.2) ppm of CO₂ was provided throughout the day in 600 ppm chamber and 899 (± 1.7) ppm of $CO₂$ was provided in 900 ppm chamber. These $CO₂$ levels were supplied using $CO₂$ cylinder in the chambers for the entire study period and monitored through SCADA. The ambient CO_2 chamber showed 380 (\pm 1.1) ppm of CO_2 . All the chambers were built in the premises of Institute with an espacement of 4×4 m. The average temperature in the chambers was $36.8 \ (\pm 1.00)$ and the average relative humidity was 65 % (± 1.2) . The mean annual rainfall was recorded in Coimbatore; India during the period of study was 796.8 mm.

2.4 Harvest and analyses

The rooted stem cuttings of C. equisetifolia were harvested after 12 months from the OTC chambers and measured for their growth characteristics like shoot length, root length, number of nodules, root collar diameter and biomass. The tissue N content of rooted stem cuttings was analyzed according to Jackson ([1973](#page-5-0)).

2.5 Photosynthetic rate

At the end of the study period the light saturated photosynthetic rate (A_{sat} , μ mol CO₂ m⁻² s⁻¹) was measured on the 15 days old needle leaves of C. equisetifolia rooted stem cuttings from the top of the stem using photosynthetic meter (Li 6400 XT, Licor linc, USA). These needle leaves are usually will be matured after 10 days as they emerged from the matured rooted stem cuttings. The leaf chamber of photosynthetic meter was set at 380 ppm of $CO₂$ concentration, 24 °C temperature and saturating photosynthetic rate of 1500 μ mol CO₂ m⁻² s⁻¹. All the rooted stem cuttings of C. equisetifolia with/without inoculation of Frankia placed in OTC were measured for determination of photosynthetic rates under ambient, 600 ppm and 900 ppm $CO₂$ conditions.

2.6 Statistical analyses

Each measured variable in the OTC experiments were statistically analyzed using Duncan's multiple range test (SPSS ver. 17). Standard error (±SE) was also applied on the data of photosynthetic rate and tissue N content.

3 Results

3.1 C. equisetifolia Rooted stem cuttings

At the end of 12 months (Mar 2015), the effect of elevated CO2 on Frankia inoculated rooted stem cuttings of C. equisetifolia showed that the growth and biomass were improved under 900 ppm of elevated $CO₂$ conditions. The shoot biomass includes needle leaves and stem (65.3 g plant $^{-1}$), root biomass (44.5 g plant $^{-1}$) and number of nodules (24.3 plant⁻¹) were significantly ($P = 0.05$) increased in C. equisetifolia rooted stem cuttings inoculated with Frankia under 900 ppm of elevated $CO₂$ conditions than 600 ppm and ambient $CO₂$ conditions (Table [1](#page-3-0)). Root nodules were observed in the rooted stem cutting of C. equisetifolia inoculated with Frankia and grown in the inert media (vermiculite) under elevated $CO₂$ conditions (Fig. [1\)](#page-3-0) Nodule numbers were significantly ($P = 0.05$) higher under 600 ppm of elevated $CO₂$ conditions due to inoculation of *Frankia* than ambient $CO₂$ conditions. However, the uninouclated control plants grown under 900 ppm and 600 ppm of elevated $CO₂$ had poor growth, biomass than ambient elevated $CO₂$ conditions. *Frankia* inoculation significantly $(P = 0.05)$ increased the collar diameter under 600 ppm and 900 ppm of elevated $CO₂$ conditions. Under ambient $CO₂$ conditions, seedlings inoculated with *Frankia* showed significantly ($P = 0.05$) higher growth and biomass and number of nodules than control plants (Table [1\)](#page-3-0).

In overall, the results showed that the rooted stem cuttings inoculated with Frankia had improved growth and biomass under elevated $CO₂$ conditions, whereas, the uninoculated control plants had poor performance under elevated $CO₂$ conditions particularly under 900 ppm.

3.2 Photosynthetic activity

C. equisetifolia rooted stem cuttings showed increased photosynthetic rates in 600 ppm and 900 ppm of elevated $CO₂$ conditions in the presence of Frankia. The photosynthetic rate was significantly ($P = 0.05$) increased in 900 ppm level (17.8 μ mol CO₂ m⁻² s⁻¹) of elevated CO₂ conditions than 600 ppm (13.2 μ mol CO₂ m⁻² s⁻¹) and ambient control (8.3 μ mol CO₂ m⁻² s⁻¹) conditions. The control plants had poor photosynthetic rates compared to Frankia inoculated seedlings particularly under 900 ppm of elevated $CO₂$ conditions (Fig. [2\)](#page-3-0).

3.3 Tissue nutrient content

Low major tissue nutrients (N, P, K) were showed in the stem cuttings of C. equisetifolia that considered as reserved food material (Fig. [3](#page-4-0)). However, the tissue N content (mg/g) was significantly ($P = 0.05$) higher for C. equisetifolia rooted stem

Means followed by same letters are not significantly different at 5 % level of DMRT

Means followed by same letters are not significantly different at 5 % level of DMRT

Ĭ.

Fig. 1 C. equisetifolia rooted stem cuttings inoculated with Frankia showed root nodules under 900 ppm of elevated CO₂ conditions (White arrow indicate root nodules)

cuttings inoculated with Frankia at 600 and 900 ppm of elevated $CO₂$ conditions. Further, Frankia inoculated C. equisetifolia rooted stem cuttings showed significantly $(P = 0.05)$ higher N content (3.2 mg g⁻¹) under 900 ppm of elevated CO ² conditions than ambient and 600 ppm of elevat-ed CO₂ conditions (Fig. [4\)](#page-4-0).

4 Discussion

Global CO ² levels are rising and it is anticipated that by the year 2100 these levels could reach 815 ppm (UKCIP [2011\)](#page-5-0). The microbial symbionts like *Frankia* can contribute to carbon sequestration by increasing nutrient uptake by plants (Garcia et al. [2011\)](#page-5-0) as found in this study. In this study elevated $CO₂$ greatly influenced the growth, biomass, nutrient content and photosynthesis in C. equisetifolia inoculated with Frankia. The rooted stem cuttings of C. equisetifolia grown in soilless media (vermiculite) without any fertilization the plants have responded well in growth and biomass under elevated $CO₂$ due to inoculation of *Frankia*. It was also confirmed that the inoculation of Frankia has only promoted the growth of

Fig. 2 Photosynthetic rates of C. equisetifolia rooted stem cuttings under elevated CO ² conditions (mean of 10 replicates). Bars indicating same letters are not significantly different according to DMRT ($p < 0.05$). Error bard indicating $SE(\pm)$ of mean

Fig. 3 Major tissue nutrients (N, P, K) content in stem cuttings of C. equisetifolia (mean of 10 replicates). Error bard indicating SE (\pm) of mean

C. equisetifolia rooted stem cuttings through N fixing root nodules as the other microbes were absent in the inert media. In earlier studies, the N_2 fixing microbes have been attempted in legume or actinorhizal plants at the seedling stage with inoculation of N fixing bacteria to find out the response under elevated $CO₂$ conditions (Arnone and Gordon [1990](#page-5-0); Vogel and Curtis [1995](#page-5-0); Ryle et al. [1992;](#page-5-0) Tissue et al. [1997\)](#page-5-0). It was reported that the symbiotic N fixers (Frankia, Rhizobium) promoted the growth and biomass of N fixing trees under elevated $CO₂$ conditions (Norby [1987\)](#page-5-0). Inoculation of Frankia mitigate the temperature and nutrient stress of the *C. equisetifolia* under the elevated $CO₂$ (AbdElgawad et al. [2015\)](#page-5-0) which may be the reasons of growth improvement in C. equisetifolia. The rooted stem cuttings of C. equisetifolia in the present study responded positively to elevated $CO₂$ in growth, bio mass, photosynthetic rates and nutrient accumulation which is in accordance with an earlier study (Xu et al. [2014\)](#page-5-0). The supply of carbon to nodules was used in the nitrogenase enzyme system as source of energy to fix N and development of root nodules (Hartwig and Nosberger [1994](#page-5-0)). An increase in the number of root nodules might have increased the nitrogenase activity of Frankia / nodule biomass that led to higher fixation of N in C. equisetifolia rooted stem cuttings. The inoculated Frankia with P medium contains N free and

Fig. 4 Tissue N content in rooted stem cuttings of C. equisetifolia inoculated with Frankia under elevated $CO₂$ conditions (mean of 10 replicates). Bars indicating same letters are not significantly different according to DMRT ($p < 0.05$). Error bard indicating SE (\pm) of mean

low amount of and also contains propionic acid which is the main carbon source for Frankia growth. Though N or P fertilizers were not applied in control and Frankia inoculated C. equisetifolia, root nodules still developed on roots of inoculated cuttings which could be attributed to the reserve food material present in the rooted stem cuttings of C. equisetifolia. This may be the reason in extreme $CO₂$ elevated conditions (900 ppm) the rooted stem cuttings of C. equisetifolia in the absence of Frankia had poor growth due to deficient N supply. Nigom et al. [\(2016\)](#page-5-0) also reported the successful tolerance of casuarinas to environmental stress in the presence of Frankia through N fixation. Some of earlier studies have shown that the inoculation of microbial symbionts under elevated $CO₂$ conditions could improve the efficiency of nutrient uptake by plants (Tang et al. [2012;](#page-5-0) Song et al. [2013\)](#page-5-0). Song et al. ([2014\)](#page-5-0) found enhanced growth and biomass in Lolium *perenne* inoculated with *Trichoderma* under ambient $CO₂$ conditions. Elevated $CO₂$ conditions (900 ppm) increased collar diameter compared to ambient $CO₂$ which is in agreement with the earlier studies (Yazaki et al. [2004](#page-5-0)). Higher elevated $CO₂$ (900 ppm) increased nodulation by *Frankia* which indicates an increased availability of carbon in form of carbohydrate to the bacterial symbiont. Similar results were also reported for Alnus hirsuta inoculated with Frankia under ele-vated CO₂ conditions (Tobita et al. [2005](#page-5-0)). It was also reported in earlier studies that $CO₂$ positively correlated with the amount of N acquired through N fixing bacteria in plants (Vogel et al. [1997\)](#page-5-0). Nasser et al. ([2007](#page-5-0)) found an increased leaf area index in Lentil under 700 ppm of elevated $CO₂$ level. They also found higher nodule numbers in response to rhizobial inoculation at 700 ppm of elevated $CO₂$ conditions. Tissue et al. [\(1997\)](#page-5-0) reported an increased photosynthetic rates and carbon storage in Gliricidia sepium inoculated with $Rhizobium$ under elevated $CO₂$ conditions which are coherent with the present study. Increased photosynthetic rates observed in the present study might have enhanced the rate of N fixation as evidenced by higher concentration of N in Frankia inoculated plants (Thomas et al. [1991\)](#page-5-0). In overall this study showed that C. equisertifolia along with Frankia responded positively at 900 ppm of elevated CO₂ conditions.

5 Conclusion

Nitrogen fixing microbial symbiont, Frankia plays important roles in improvement of Casuarinas. In this present study under high atmospheric $CO₂$ conditions the Frankia facilitate the C. equisetfolia rooted stem cuttings for growth and biomass improvement. Universally, C. equisetifolia is propagating through rooted stem cuttings from genetically superior clones for establishing commercial plantations to make paper and pulp. These commercial plantations of C. equisetifolia may be inoculated with *Frankia* to improve the growth and

biomass and to mitigate the increasing $CO₂$ levels by carbon sequestration.

Acknowledgments The author thanks Indian Council of Forestry Research and Education, Dehra Dun, India for financial assistance for this study.

References

- AbdElgawad H, Vignola EAR, de Vos D, Asard H (2015) Elevated CO₂ mitigates drought and temperature, induce oxidative stress differently in grasses and legumes. Pl Sci 23:1–10
- Arnone J, Gordon JC (1990) Effect of nodulation, nitrogen fixation and CO2 enrichment on the physiology, growth and drymass allocation of seedlings of Alnus rubra bong. New Phytol 116:55–66
- Ceulemans R, Janssens IA, Jach ME (1999) Effects of $CO₂$ enrichment on trees and forests, lesions to be learned in view of future eco system studies. Ann Bot 84:577–590
- Diagne N, Karthikeyan A, Ngom M, Mathish NV, Franche C, Krishnakumar N, Laplaze, L (2013) Use of Frankia and actinorhizal plants for degraded lands reclamation. Bio Med Res Int 948258 9 p
- Garcia NS, Fu FX, Breene CL, Berhandt PW, Mulholland MR, Sohm JA, Hutchins DA (2011) Interactive effects of irradiance and $CO₂$ on $CO₂$ fixation and N2 fixation in the Diazotroph Trichodesmium erythraeum (cyanobacteria). J Physiol 47:1292–1303
- Hartwig UA, Nosberger J (1994) What triggers the regulation of nitrogenase activity in forage legume nodules after defoliation? Plant Soil 161:109–114
- Hodge A (1996) Impact of elevated $CO₂$ on mycorrhizal association and implications for plant growth. Biol Fertil Soils 23:388–398
- Jackson ML (1973) Soil chemical analysis. Prentice Hall, New Delhi India, pp. 183–192
- Jenkinson DS, Adams DE, Wild A (1991) Model estimate of CO2 emissions from soil in response to global warming. Nature 351:304–306
- Karthikeyan A, Deeparaj B, Nepolean P (2009) Reforestation in bauxite mine spoils with Casuarina equisetifolia Frost. and beneficial microbes. For Trees Live 19:153–165
- Nasser RR, Fuller MP, Jellings AJ (2007) Effect of elevated and nitrogen levels in lentil growth and nodulation. Agron Sustain Dev 28:1–6
- Nigom M, Oshone R, Diagne N, Cissoka M, Svistoonoff S, Tisa LS, Laplaze L, Quereysy M, Champion A (2016) Tolerance to environmental stress by the nitrogen fixing actinobacterium *Frankia a*nd its role in actinorhizal plants adaptation. Symbiosis. doi:[10.1007](http://dx.doi.org/10.1007/s13199-016-0396-9) [/s13199-016-0396-9](http://dx.doi.org/10.1007/s13199-016-0396-9)
- NOAA (2016). National Oceanic and Atmospheric Administration report. U.S. Department of Commerce, U.S (www.noaa.gov).
- Norby RJ (1987) Nodulation and nitrogenase activity in nitrogen fixing woody plants stimulated by $CO₂$ enrichment of the atmosphere. Physiol Plant 71:77–82
- O'Neill EG, Luxmore RJ, Norby RJ (1987) Increases in mycorrhizal colonization and seedling growth in Pinus echinata and Quercus alba in an enriched $CO₂$ atmosphere. Can J For Res 17:878-883
- Quoreshi AM, Maruyama Y, Koike T (2003) The role of mycorrhiza in forest eco systems under $CO₂$ semiarid atmosphere. Eurasian J For Res 6:171–176
- Ryle GJA, Powell CE, Davidson JA (1992) Growth of white clover, dependent on N_2 fixation, in elevated CO_2 and temperature. Ann Bot 70:221–228
- Santi C, Bogsuz D, Franchie C (2013) Biological nitrogen fixation in non legume plants. Ann Bot. doi[:10.1093/aob/mct048](http://dx.doi.org/10.1093/aob/mct048)
- Shipton WA, Burgraff AJP (1983) Aspects of the cultural behaviour of Frankia and possible ecological implication. Can J Bot 61:2783– 2792
- Song HN, Tang SR, Wanf FL, Zhang C, De Geo JK, Ju XH, Smith DC (2013) Fungal inoculation and elevation of $CO₂$ mediated growth of Lilum moniliforme and Phytolacca americana metal uptake and metal bio availability in metal contaminated soil evidence from diffusing gradient in thin film measurement. Int Phytorem 15:268–282
- Song N, Ma Y, Zhao Y, Tang S (2014) Elevated ambient carbon dioxide and Trichoderma inoculums could enhance cadmium uptake of Lolium perenne explained by changes of soil pH, cadmium availability and microbial bio mass. Appl Soil Ecol 85:56–64
- Staddon PL, Fitter AH, Robinson D (1999) Effects of mycorrhizal colonization and elevated atmospheric carbon dioxide on carbon fixation and below ground carbon partitioning in Plantago lanceolata. J Exp Bot 50:853–860
- Tang SR, Liao SQ, Guo JK, Song ZS, Wang RG, Zhou XM (2012) Growth and cesium uptake responses Phytolacca americana Linn and Aaranthhus curentis L grown on cesium contaminated soil to elevated $CO₂$ on inoculation with a plant growth promoting Rhizobacterium Burkholdeia sp. D54 or in combination. J Hasand meter 198:188–197
- Temperton VM, Grayston SJ, Jackson G, Barton CVM, Millard P, Jarvis PG (2003) Effects of elevated carbon dioxide concentration on growth and nitrogen fixation in Alnus glutinosa in a long term field experiment. Tree Physiol 23:1051–1059
- Thomas RB, Richter DD, Ye H, Heine PR, Strain BR (1991) Nitrogen dynamics and growth of seedlings of an N fixing tree (Gliricidia sepium (Jacq.) Walp) exposed elevated atmospheric carbon dioxide. Oecologia 88:415–421
- Tissue DT, Megonigal JP, Thomas RB (1997) Nitrogenase activity and N_2 fixation are stimulated by elevated $CO₂$ in a tropical N2 fixing tree. Oecologia 109:28–33
- Tobita H, Kituo M, Koika T, Maryuma Y (2005) Effects of elevated CO₂ and nitrogen availability on nodulation of Alnus hirsuta. Phyton 45: 125–131
- UKCIP (2011) Making progress. UKCIP and adaptation in the UK. UK climate impacts programme, Oxford UK, pp. 23–26
- Uma M, Saravanan TS, Rajendran K (2014) Growth, litterfall and litter decomposition of Casuarina equisetifolia in a semi arid zone. J Trop For Sci 26:125–133
- UNESCO/UNEP (2011) Climate change starter's guide book: an issues guide for educating planners and practitioners. United Nations Educational, Scientific and Cultural organization and the United Nations Environment Programme, Paris
- Vogel CS, Curtis PS (1995) Leaf gas exchange and nitrogen dynamics of N2-fixing, field-grown Alnus glutinosa under elevated atmospheric CO2. Global Change Biology 1(1):55–61
- Vogel CS, Curtis PS, Thros RB (1997) Growth and nitrogen accretion of di nitrogen fixing. Alnus glutinosa (L). Gertn. Under elevated carbon dioxde. Plant Ecol 130:63–70
- Wheeler CT, Miller TM (1990) Current potential uses of actinorhizal plants in Europe. In: Schwintzer RC, Tjepkema JD (eds) The biology of Frankia and actinorhizal plants. Academic Press, San Diego, CA, pp. 365–389
- Wiley IM, Sherwood LM, Woolverton CJ (2009) Prescott's principals of microbiology. Mc Graw-Hill, New York, NY
- Xu LI, Ahmad G, Zhang Y, Shang G, Sum Z, Shou J, Zhou Y, Xiav Yu J, Hi K (2014) Carbon diozide enrichment alleviates root stress by improving cellular redox homesteads through and ABA- independent pres in tomato plants. Plant Soil. doi:[10.1111/pid.12.11](http://dx.doi.org/10.1111/pid.12.11)
- Yazaki K, Ishida S, Kawagish T, Fukatsu E, Maruyama Y, Kitao M, Tobita HT, Koike T, Funada R (2004) Effects of elevated $CO₂$ concentration on growth, annual ring structure and photosynthesis in Larix kaempferi seedlings. Tree Physiol 24:941–949