

International Journal of Environment and Climate Change

Volume 14, Issue 12, Page 846-866, 2024; Article no.IJECC.129234 ISSN: 2581-8627 (Past name: British Journal of Environment & Climate Change, Past ISSN: 2231–4784)

Resilience of C⁴ Crops to Climate Vagaries

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Authors' contributions

This work was carried out in collaboration among all authors. All authors read and approved the final manuscript.

Article Information

DOI: <https://doi.org/10.9734/ijecc/2024/v14i124667>

Open Peer Review History:

This journal follows the Advanced Open Peer Review policy. Identity of the Reviewers, Editor(s) and additional Reviewers, peer review comments, different versions of the manuscript, comments of the editors, etc are available here: <https://www.sdiarticle5.com/review-history/129234>

Review Article

Received: 28/10/2024 Accepted: 30/12/2024 Published: 30/12/2024

ABSTRACT

Climate change poses significant challenges to agriculture worldwide, affecting productivity and threatening food security. Key drivers of climate change like altered water availability, temperature fluctuations and increased carbon dioxide concentrations, influence crop performance and

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Cite as: Kumar, Swathy Anija Hari, Usha Chacko Thomas, P Shalini Pillai, Roy Stephen, G. Rajasree, and B. Aparna. 2024. "Resilience of C4 Crops to Climate Vagaries". International Journal of Environment and Climate Change 14 (12):846-66. https://doi.org/10.9734/ijecc/2024/v14i124667.

ecosystem stability. The World has committed to eradicate extreme poverty and hunger by 2030. But climate change is undermining the livelihoods and food security of the rural poor, who constitute almost 80% of the world's poor. If plant genotypes that can withhold climate extremes are insufficient, food scarcity occurs, and the price of existing food resources would increase enormously, making it practically hard for the rural poor to obtain enough food. To avoid this happening, we need to develop plant genotypes that are climate resilient. Photosynthetic pathways in plants C_3 , C_4 , and CAM play a critical role in determining their adaptability to changing climatic conditions. Understanding the physiological responses of C⁴ crops to various environmental stresses like water and temperature stress, highlights their potential for future climate resilience. Due to the efficient carbon concentrating mechanism, low stomatal conductance and high water use efficiency in C_4 plants, they are expected to show higher drought tolerance relative to C_3 plants. Studying the responses of C_4 crops to climate change is essential as they play a vital role in global food production, especially in tropical and subtropical regions prone to climate extremes. While C⁴ crops are more resilient to heat and drought than their C_3 counterparts, their yield potential is still constrained by the increasing severity of abiotic stresses, such as prolonged droughts, heatwaves, and soil salinity. Understanding how C_4 crops respond to these challenges can provide insights into optimizing their growth and productivity in future climate scenarios. These insights are important for breeding stress-resilient varieties and improving crop management practices, to ensure global food security amidst escalating climate challenges.

Keywords: Climate change; drought; heat stress; C⁴ photosynthesis; carbon concentrating mechanism.

1. INTRODUCTION

The climate of Earth has undergone alterations throughout history. There have been eight cycles of ice ages and warmer periods during the past 800,000 years, with the end of the last ice age approximately 11,700 years ago marking the beginning of the current climate era and the rise of human civilisation (NASA, 2024). Climate change occurs due to various factors, including atmospheric changes like El Niño, driven by winds and ocean currents. External forces, such as Earth's axial tilt and orbital shape, likely influence ice age cycles. Additionally, greenhouse gases like carbon dioxide $(CO₂)$ trap heat, raising Earth's surface temperature (National Geographic Society, n.d.). Human activities, such as burning fossil fuels for energy and transportation or using technology to boost meat production, release greenhouse gases (Garnett, 2009; Stavi & Lal, 2013). Deforestation for timber or industrial development reduces the carbon dioxide absorbed by trees, while factories contribute to the greenhouse effect by emitting pollutants into the atmosphere (Rykowski, 2000; Aju et al., 2015; National Geographic Society, n.d). Global warming is linked to burning fossil fuels and tropical deforestation, with human activities increasing atmospheric $CO₂$ by about 30% over the past 150 years (National Geographic Society, n.d.). Greenhouse gases like methane and chemicals such as chlorofluorocarbons, hydrofluorocarbons, and

hydrochlorofluorocarbons contribute significantly to climate change (Sovacool et al., 2021). Methane levels are rising due to agriculture, industrial activities, and decomposing waste in landfills (Karakurt et al., 2012). Similarly, these gases, used in refrigeration and aerosol sprays, trap heat in the atmosphere. While many countries are phasing them out through laws and regulations, their impact remains a concern (National Geographic Society, 2024). The effects of climate change are becoming increasingly evident. Global temperatures are rising, and oceans are getting warmer, with much of the heat being absorbed by the top layers of the ocean (Levitus et al., 2017; NASA, 2024). Ice sheets are shrinking, particularly in Greenland and Antarctica, while glaciers are retreating in many regions around the world (Velicogna et al., 2020). Snow cover is decreasing, and sea levels are rising because of melting ice and thermal expansion of seawater. Global sea level rose about 8 inches in the last century (Nerem et al., 2018). Arctic sea ice is declining, and ocean acidification is increasing. The ocean has absorbed between 20% and 30% of total anthropogenic CO₂ emissions in recent decades (Sabine et al., 2004). Carbon dioxide in the atmosphere warms the planet, causing climate change. Human activities have raised the atmosphere's $CO₂$ content by 50% in less than 200 years, with the current level of 424 ppm (NASA, 2024). C⁴ crops, with their efficient carbon concentrating mechanism and higher

water-use efficiency, present a promising solution to mitigate the impacts of these human-induced stresses. Understanding the interplay between anthropogenic climate change and the resilience mechanisms of C⁴ crops is crucial for developing adaptive strategies that ensure sustainable production in the face of escalating environmental challenges.

Climate change impacts on agriculture are being witnessed all over the world in the recent years affecting farm level productivity and adversely impacting stability in food grain availability at the national level. Temperature, water, and $CO₂$ are three of the main causes of climate change. Adopting sustainable methods that lower greenhouse gas emissions, encourage reforestation, and boost the use of renewable energy sources is crucial to addressing the issues caused by climate change, particularly in agriculture. Under drought, high temperatures, and nitrogen or CO₂ limitation, C₄ plants are more competitive than C_3 plants. (Watson-Lazowski and Ghannoum, 2021). C₄ plants possess the carbon concentrating mechanism, and can have increased photosynthetic rate, yield, water-use efficiency (WUE), and nitrogenuse efficiency (NUE) compared with ecologically similar C_3 plants. Sorghum is a promising C_4 crop for tackling extreme food insecurity because of its resistance to drought and capacity to flourish in a variety of soil types. Sorghum's ability to withstand water further increases its resilience (Mwamahonje et al., 2024). Implementing agricultural practices that favor C⁴ plants could increase crop resilience to climate stressors, leading to higher productivity and better food security.

2. IMPACT OF CLIMATE VAGARIES ON CROPS

Climate change may adversely impact the yield of irrigated crops throughout the world, attributable to rising temperatures and alterations in water supply (Lal, 2000; Mall et al., 2017). Crops may experience severe weather events such as drought, flooding, excessive heat, and cold during their life cycle, leading to significant yield reductions (Raza et al., 2019). The effects of these factors may differ based on area, crop, cropping systems, soil types, and management practices. Sub-Saharan Africa frequently endures prolonged droughts and unpredictable weather patterns that threaten crop yields (Lombe et al.,

2024). South Asia is projected to experience more frequent extreme weather events, such as heatwaves and intense rainfall, along with greater interannual variability in daily precipitation during the Asian summer monsoon (Sivakumar & Stefanski, 2011). India is among the most drought-prone nations globally, with around 53% of its geographical area classified as arid and semi-arid. Climate change affects crop production both directly and indirectly (Samuel et al., 2021). The immediate consequences primarily result from alterations in crop length and influence reproductive processes, including pollination and fertilisation. The indirect effects of climate change are mostly attributable to variations in water availability, as well as changes in insect, disease, and weed dynamics. Weather aberrations can impact numerous factors, particularly in rainfed regions where over 80 percent of farmers are smallholders, hence possessing limited capacity to mitigate adverse effects (Maheswari et al., 2015). Rainfed drylands, covering over 41% of Earth's surface, are vital for global food and fiber production while supporting biodiversity and ecosystem services. As temperatures increase, evapotranspiration increases, and precipitation decreases, 25–30% of drylands are already degraded, mostly because of soil erosion from wind and water in regions like Australia, East and Central Asia, and Sub-Saharan Africa (Chary et al., 2022).

3. CLIMATE RESILIENCE

The capacity to foresee, plan for, and react to potentially dangerous occurrences, patterns, or disruptions associated with the climate is known as climate resilience. Assessing how climate change will increase or change existing climate-related risks and acting to better manage them are key components of increasing climate resilience (C2ES, 2021).

3.1 Climate Smart Agriculture

Climate Smart Agriculture (CSA) is characterised as a methodology that directs the necessary actions to transform and realign agricultural systems to effectively promote development and guarantee food security in a changing climate. The initiative seeks to achieve three primary objectives: sustainably enhancing agricultural output and income; adapting to and fostering resilience against climate change; and minimising and/or eliminating greenhouse gas emissions, where feasible (Cordaid, 2016). CSA is frequently used interchangeably with climateresilient agriculture (CRA); however, CRA is a subset of CSA that specifically addresses the impacts of climate change (Viswanathan et al*.,* 2020).

An on-farm trial in Karnal, India, assessed the greenhouse gas mitigation potential of CSA practices (CSAPs) in rice–wheat systems across six scenarios: Sc1 (conventional tillage without residue), Sc2 (conventional tillage with residue), Sc3 (reduced tillage + residue + fertilizer), Sc4 (reduced/zero tillage + residue + fertilizer), Sc5 (zero tillage + residue + fertilizer + GreenSeeker + Tensiometer), and Sc6 (Sc5 + nutrient-expert tool). CSAPs (Sc4, Sc5, Sc6) reduced global warming potential by 33–40% and emission intensity by 36–44% compared to conventional practices (Sc1). Additionally, they improved nitrogen productivity by 32–57%, eco-efficiency by 70–105%, and wheat yields by 0.62–0.84 Mg/ha, showing resilience to climate extremes. These results highlight CSAPs as effective for mitigating GHG emissions and enhancing sustainability in rice–wheat systems (Kakraliya et al., 2021).

Climate-smart agriculture has been promoted by the Department of Agricultural Extension in Bangladesh through climate field schools since 2010 to improve food security in the context of climate change. A study involving 118 farmers from Kalapara, Patuakhali, identified 17 CSA practices, including saline- and flood-tolerant crop varieties, floating-bed vegetables, the sorjan method, urea deep placement, and rainwater harvesting. On average, farmers adopted seven practices, with 32% of households classified as food secure, 51% mildly to moderately food insecure, and 17% severely food insecure. CSA adoption positively influenced food security (Hasan et al., 2018). Small-scale coconut farmers in Philippines frequently adopted CSA practices such as early harvesting and weather forecasting, driven by the recurring natural disasters like typhoons. However, practices like coconut-banana intercropping were rarely implemented due to the high initial investment and a lengthy payback period of approximately four years, challenges that were even greater for adopting typhoon-tolerant coconut varieties (Ruales et al., 2020).

In water-scarce regions, advanced irrigation strategies such as deficit irrigation and the use of

non-conventional water resources, including treated wastewater and brackish groundwater, have emerged as effective measures to combat water scarcity. Protected cultivation systems like greenhouses equipped with artificial intelligence further enhance water productivity by creating controlled environments that reduce water loss (Nikolaou et al., 2020). These approaches collectively provide sustainable solutions for improving water management and mitigating the impacts of climate change on agriculture, especially in vulnerable areas like the Mediterranean region. Modernizing irrigation methods significantly improves water application efficiency, with systems like trickle irrigation achieving up to 90% water savings (Mesa-Jurado et al., 2012; Nikolaou et al., 2020). Agricultural water management plays a crucial role in mitigating unsustainable water use and improving water resilience and adaptation to climate change. Alternate wetting and drying (AWD) is another recommended irrigation alternative for irrigated rice systems. In AWD, rice fields experience alternating periods of saturation and desaturation, during which irrigation is paused and water is permitted to recede until the ponded water is eliminated and the soil attains a specific moisture content (Carrijo et al., 2017).AWD can decrease irrigation water inputs by 23%–43% and can lower the global warming potential by 45%–90% in comparison to continuously flooded rice (Sikka et al., 2022). A 25%–30% decrease in global warming potential through the implementation of intermittent flooding compared to continuous flooding was observed in rice cultivation (Pathak et al., 2011).

4. CLIMATE CHANGE AND PHOTOSYNTHESIS

Through the process of photosynthesis, all plants absorb $CO₂$ from the atmosphere and transform it into sugars and starches, but they do it in different ways (Paul & Foyer, 2001; Long et al., 2004). Each class of plants uses a different type of photosynthesis, known as a pathway, which is a variant on a series of chemical reactions known as the Calvin Cycle. A plant's capacity to tolerate low carbon atmospheres, higher temperatures, and decreased water and nitrogen is crucial for understanding climate change (Dusenge et al., 2019). These reactions also affect the quantity and kind of carbon molecules a plant produces, the locations where those molecules are stored, and more.

The processes of photosynthesis designated by botanists as C_3 , C_4 , and CAM, are directly relevant to global climate change studies because C_3 and C_4 plants respond differently to changes in atmospheric $CO₂$ concentration and changes in temperature and water availability. Scientists have started looking into how plants might be able to adapt to the changing climate as the world continues to warm. Modifying the processes involved in photosynthesis could be one approach to do it (Lara and Andreo, 2011). A promising approach is the introduction of a C⁴ carbon concentration mechanism (CCM) into C3 crops, such as rice. The C⁴ Rice project aims to achieve this, with predictions suggesting that incorporating a C⁴ photosynthetic pathway into rice could increase yields by as much as 50 percent (Sheehy et al., 2008). The C⁴ Rice Consortium is employing strategies like metabolic C⁴ engineering and the identification of leaf anatomy determinants through mutant screens to develop C_4 rice (Kajala et al., 2011).

The Crassulacean Acid Metabolism (CAM) cycle is a $CO₂$ fixation process occurring during the dark phase of photosynthesis in Crassulaceae plants, with malic acid as the first product (Black & Osmond, 2003; Osmond, 2007). Most CAM plants are succulents with fleshy leaves, abundant chloroplasts in mesophyll cells, and vascular bundles lacking well-defined bundle sheath cells. Their stomata open at night and close during the day. While less efficient in photosynthesis than C⁴ plants, CAM plants are better adapted to extreme desiccation (Black & Osmond, 2003; Osmond, 2007; Schiller & Bräutigam, 2021). Poplar (*Populus* spp.) and willow (*Salix* spp.) which are fast-growing, shortrotation forestry bioenergy crops being susceptible to hydraulic failure following drought stress due to their relatively high stomatal conductance and isohydric nature. Improving water-use efficiency (WUE) by engineering crassulacean acid metabolism (CAM) into C³ crops represents a viable strategy for sustaining plant productivity. CAM enhances water-use efficiency by enabling nighttime CO2 uptake when vapor pressure deficit is low. CAM in tree species like *Clusia* demonstrates its role in conserving water and maintaining carbon uptake during droughts. Bioengineering CAM into bioenergy trees offers a promising strategy for sustaining agroforestry amid climate change (Sekhar et al., 2021).

While the C_3 pathway is the most common, it is also inefficient due to the photorespiration, a process that wastes assimilated carbon when RuBisCO reacts not only with $CO₂$ but also $O₂$.
Under prevailing atmospheric conditions, Under prevailing atmospheric conditions, potential photosynthesis in C_3 plants is suppressed by $O₂$ as much as 40 %. The extent of that suppression increases under stress conditions such as drought, high temperatures and high light (Hirst, 2021). The C_4 plants are more efficient in photosynthesis than the C_3 plants (Schmitt & Edwards, 1981; Gowik & Westhoff, 2011). In C_4 plants, the photorespiration is suppressed by increasing the CO² concentration at the Rubisco site, hence suppressing the enzyme's oxygenase activity. C⁴ photosynthesis relies on a biochemical CO₂ pump with spatial separation of $CO₂$ fixation and assimilation, typically involving Kranz anatomy where mesophyll and bundle sheath cells cooperate (Edwards et al., 2004; Lara and Andreo, 2011). The enzyme, phosphoenol pyruvate carboxylase of the C⁴ cycle is found to have more affinity for CO₂ than the ribulose diphosphate carboxylase of the C_3 cycle in fixing the molecular $CO₂$ in organic compound during Carboxylation (Hatch, 1987; Lara and Andreo, 2011).

5. CARBON CONCENTRATING MECHANISM IN C⁴ PLANTS

C⁴ plants achieve high productivity through a carbon concentrating mechanism (CCM) that increases CO₂ concentration around RuBisCO (ribulose-1,5-bisphosphate

carboxylase/oxygenase), the primary $CO₂$ -fixing enzyme in plants, algae, and cyanobacteria. This CCM operates between two cell types: mesophyll cells, where $CO₂$ is initially fixed, and bundle sheath cells, where it is ultimately fixed (Watson-Lazowski & Ghannoum, 2021). In C₄ plants, PEP carboxylation occurs in specialized mesophyll cells, which also perform the full light-dependent reactions of photosynthesis (Schlüter & Weber, 2020; Romanowska & Wasilewska-Dębowska, 2022). Malate or aspartate is then transported to bundle sheath cells, characteristic of Kranz anatomy, where pre-fixed carbon is released for final $CO₂$ fixation by RuBisCO. This process raises the CO₂ concentration in bundle sheath cells to about 10 times that of ambient air, significantly reducing photorespiration. Notably, most bundle sheath cells do not complete the full light-dependent reactions of photosynthesis (Zabaleta et al., 2012; Watson-Lazowski & Ghannoum, 2021).

Table 1. C3 vs C4 cycle

(Source: Ehleringer & Cerling, 2002; Lara and Andreo, 2011; Hirst, 2021).

6. ADVANTAGES OF THE C4 PHOTOSYNTHETIC PATHWAY

The C_4 pathway increases CO_2 concentration in bundle sheath cells to 70 μM, compared to 4 μM in mesophyll cells, reducing RuBisCO's oxygenase activity by over 80%, though this effect varies with temperature. Consequently, C⁴ plants exhibit up to double the photosynthetic rate and yield, 1.5 to 3 times greater WUE, and 2.5 times higher NUE compared to ecologically similar C_3 plants (Lin et al., 2019). In addition, elevating the concentration of $CO₂$ within the bundle sheath allows RuBisCO to increase its in vivo catalytic activity two- to fivefold in warm climates. As a result, C_4 plants have only $50-$ 80% of the RuBisCO content found in C_3 plants, allowing them to sustain a higher leaf area production rate with lower leaf nitrogen levels compared to C3 species (Sage and Zhu, 2011b; Lin et al., 2019). WUE is enhanced in C_4 plants because $CO₂$ is fixed more effectively and stomata remain less open, reducing transpiration rates (Way et al., 2014; Leakey et al., 2019). This improved WUE allows C⁴ plants to have longer growing seasons and more adaptable allocation patterns, such as directing more biomass to shoots in moist conditions or to roots in dry conditions (Lopes et al., 2011; Leakey et al., 2019). The C⁴ pathway is particularly advantageous in hot, high-light environments that encourage high photorespiration rates (Long, 1983; Sage & Kubien, 2007; Lara & Andreo, 2011).

7. SUBTYPES OF C⁴ PHOTOSYNTHESIS

C⁴ species not only exhibit superior physiological traits compared to C_3 species but also display considerable diversity within themselves. C⁴ photosynthesis is broadly categorized into three subtypes: NADP-malic enzyme (NADP-ME), NAD-malic enzyme (NAD-ME), and phosphoenolpyruvate carboxykinase (PEP-CK), each adapted to specific environmental conditions (Ghannoum et al*.,* 2011; Wang et al., 2014; Borghi, 2021).

In NADP-ME plants, malate serves as the main C⁴ acid transported between mesophyll and bundle sheath cells, whereas aspartate is the primary transport acid in NAD-ME and PCK C⁴ grasses. The C⁴ subtypes are closely linked to specific grass subfamilies. Species with NADP-ME type anatomy are found in the Panicoideae subfamily within the Andropogoneae, Arundinelleae, and Paniceae tribes (Morrone et al., 2012; Venter, 2015). The NAD-ME and PEP-CK types are primarily associated with the Chloridoideae subfamily and have evolved only once in Panicoideae (Voznesenskaya et al., 2006; Koteyeva et al., 2023). Among major C⁴ crops, sorghum exclusively uses NADP-ME, maize primarily employs NADP-ME with PEP-CK as a secondary decarboxylase, and millets exhibit a mix of NADP-ME, NAD-ME, and some PEP-CK species (Sonawane et al*.,* 2018).

Research on C⁴ grasses, categorized into three biochemical subtypes—NAD-ME, PCK, and NADP-ME—grown under ambient (400 μL L^{-1}) and interglacial (280 μ L L⁻¹) CO₂ conditions revealed that the Chloridoideae/NAD-ME group had higher leaf mass per area and leaf nitrogen content. In contrast, NADP-ME and PCK grasses exhibited enhanced photosynthetic nitrogen use efficiency (Pinto et al*.,* 2016).

Characteristics	NADP-ME	NAD-ME	PEP-CK			
Decarboxylation	NADP-ME	NAD-ME	PEP-CK			
enzymes						
Main C4 acid exported	Malate	Aspartate	Aspartate			
to bundle sheath cells						
Main 3C acid returned	Pyruvate	Alanine	Alanine/pyruvate			
to mesophyll cells						
Site of	Chloroplast	Mitochondrion	Cytosol			
decarboxylation						
Chloroplast position in	Reduced grana chloroplast with	Granal chloroplast	Centrifugal			
the BSC	centrifugal arrangement	with centripetal				
		arrangement				
(Sonawane, 2016)						

Table 2. Differences between the C4 subtypes in grasses

8. C⁴ CROPS

Approximately 60% of C⁴ species are grasses, with around 40% of grasses utilizing the C⁴ photosynthetic pathway. Most C⁴ grasses belong to warm-origin taxa, particularly the PACMAD clade, and dominate warm-climate grasslands. These grasses include ecologically and economically significant species such as major staple food, fodder, and biofuel crops, as well as numerous prominent weeds (Sage et al., 2011a; Watson-Lazowski & Ghannoum, 2021). C₄ crops are particularly prevalent in warm (Korres et al., 2016), drought-prone climates (Lopes et al., 2011; Korres et al., 2016) and are becoming increasingly crucial for food and
bioenergy security (Watson-Lazowski & security (Watson-Lazowski & Ghannoum, 2021).

Maize, sorghum, and sugarcane are major C⁴ crops. Maize, the most produced cereal crop globally, is mainly grown in North and South America, as well as Eastern and Southern Africa, with over 60% of production in temperate regions (Watson-Lazowski & Ghannoum, 2021). Maize is primarily used in feed, starch, and biofuel industries, with 83% of its production allocated for industrial purposes. Among 125 developing countries, 75 consider maize a staple crop, accounting for 70% of global maize production (Burlakoti et al., 2024). Sorghum, more droughttolerant than maize, thrives in dry climates and is valuable in areas with limited rainfall and resources for fertilizers (Watson-Lazowski & Ghannoum, 2021; Khalifa & Eltahir, 2023; Liaqat et al., 2024. Sorghum is a grain crop used for fodder, food, and bioenergy production. Its high photosynthesis rate makes it excellent for silage, while its stalks produce juice for jaggery, sugar, and ethanol (Wasi et al., 2023). The 2001–2020 global average harvested area for sorghum is 40.90 mha (Khalifa & Eltahir, 2023). All parts of sorghum like grain, juice, and bagasse are utilized for fodder, food, ethanol, and electricity. Sorghum thrives on waterlogged, saline-alkaline soils and is drought-resistant (Wasi et al., 2023). The drought resistance in sorghum is attributed to its root system, leaf rolling, osmotic adjustment, and ability to delay reproductive development (Nadew et al., 2021). Sugarcane, a key industrial crop for sugar and bioenergy, grows in tropical and subtropical regions (Raza et al., 2019). Worldwide, sugarcane ranks as the fifth largest crop in terms of production value and acreage, and it is the second largest bioenergy crop (Luo et al., 2023). It benefits from high CO²

levels, showing improvements in biomass, photosynthesis, and overall growth, and is capable of coping with rising $CO₂$ concentrations due to its low $CO₂$ compensation point and carbon sequestration abilities (Watson-Lazowski & Ghannoum, 2021). Marin et al.
(2013) observed improved sugarcane sugarcane WUE and yield in parts of Brazil due to climate change using crop simulation models. As a C⁴ plant with a $CO₂$ compensation point of 0-10 ppm, sugarcane can deplete atmospheric CO² under certain conditions. High $CO₂$ levels partially close stomata, reducing transpiration and sap flow, enhancing xylem potential and water status. Sugarcane also sequesters carbon naturally, mitigating CO₂ emissions and global warming by forming phytoliths (PhytOC), storing \sim 300 Mt of CO₂ annually in soil for thousands of years (Misra et al., 2019).

Millets, including pearl millet (*Pennisetum glaucum*) and foxtail millet (*Setaria italica*), are C⁴ crops vital for food and fodder, with over 95% produced in developing countries. Their drought and heat tolerance make them suitable for harsh climates, and their short life cycle (12-14 weeks) helps escape stress. Traits like small leaves, thickened cell walls, and dense roots enhance stress resilience. The C₄ mechanism concentrates CO² around RuBisCO, reducing photorespiration (~80%) and boosting photosynthesis, WUE, and NUE. This also improves growth, biomass allocation, and ecological performance in warm conditions (Lenka et al.,2020).

9. CLIMATE RESILIENCE IN C⁴ GRASSES

Grass species are divided into two distinct clades: BOP (Bambusoideae, Oryzoideae, Pooideae) and PACMAD (Panicoideae, Arundinoideae, Chloridoideae, Micrairoideae, Aristidoideae, Danthonioideae) (Hodkinson, 2018; Pardo and VanBuren, 2021; Gallaher et al., 2022). BOP grasses, primarily cool-season species found in temperate climates, utilize C3 photosynthesis, which outperforms C4 in these regions. Frost tolerance has independently evolved in many Pooideae grasses. PACMAD grasses, mainly warm-temperate and tropical species, include agriculturally significant crops like sugarcane, maize, sorghum, and various millets (Panicoideae) and underutilized grains like finger millet (Chloridoideae) (Pardo and VanBuren, 2021).

9.1 Climate Resilience among PACMAD Grasses

Grass stomatal anatomy enhances resilience with unique elongated, dumbbell-shaped guard cells and two subsidiary cells, enabling faster responses and higher WUE compared to kidneyshaped guard cells in eudicots and most nongrass monocots (Nunes et al., 2020; Pardo and VanBuren, 2021; Zahedi et al., 2024). Stomatal arrangement and density also influence drought tolerance (Mehri et al., 2009; Huang et al., 2020). Grasses typically have hypostomatic leaves with pores on the abaxial surface or amphistomatic leaves with pores on both surfaces, the latter promoting efficient CO₂ diffusion and higher photosynthetic rates. Unlike eudicots with dorsoventral leaves, grasses have isobilateral leaves oriented parallel to light, deeper veins, and vertical angles, minimizing WUE costs of amphistomaty (Pardo and VanBuren, 2021).

10. RESPONSE OF C⁴ PLANTS TO CLIMATE VAGARIES

10.1 Effect of Water Stress on C⁴ Photosynthesis

Water stress significantly limits global plant productivity (Kijne, 2006). A study on the effects of water stress on winter wheat's photosynthesis, growth, yield, WUE, and irrigation water productivity (IWP) identified four stress levels based on field water capacity: severe (30–40%), moderate (40–50%), mild (50–60%), and wellwatered (60–80%). Results showed that moderate and severe stress significantly reduced photosynthetic parameters (net photosynthetic rate, intercellular carbon concentration, stomatal conductance and transpiration), height, biomass, and grain size, leading to lower WUE and IWP. In contrast, mild stress only slightly decreased net photosynthetic rate compared to the wellwatered group, with improvements in dry biomass and 1000-grain weight by 2.07% and 1.95%, respectively, during flowering and grainfilling (Zhao et al., 2020). Severe stress causes metabolic inhibition, including photoinhibition and damage to photosynthetic enzymes, leading to irreversible loss of photosynthetic activity (Goh et al., 2012) .C₄ plants, with high WUE and low stomatal conductance, exhibit greater drought tolerance, reducing water stress development. However, high WUE often prioritizes biomass production over water conservation in areas with

high evapotranspiration (Ghannoum, 2016; Watson-Lazowski & Ghannoum, 2021).

The C_4 pathway enhances WUE, enabling C_4 grasses to thrive in drier, more exposed habitats. The high substrate affinity of PEPcase and the carbon-concentrating mechanism allow C⁴ plants to function at lower mesophyll $CO₂$ levels and stomatal conductance, achieving higher $instantaneous WUE than C₃ plants (Ghannoum)$ et al., 2011; Pardo & VanBuren, 2021). Under drought stress, leaf-level WUE increases as reduced water loss from stomatal closure outweighs the decline in $CO₂$ assimilation. WUE also varies by C⁴ subtype, with NAD-me grasses showing higher WUE than NADP-me grasses under drought condition (Ghannoum et al., 2002; Pardo & VanBuren, 2021).

Sorghum genotypes exhibit varied responses and tolerance to drought, influenced by the interaction between genotype and water stress levels. Research by Tingting (2010) found that sweet sorghum exhibited the highest WUE under moderate drought stress during early and middle growth stages, and under severe drought stress in the late growth stage. Increasing drought stress raised the light compensation point but reduced the light saturation point, apparent quantum yield, and dark respiration rate. Severe drought stress caused photoinhibition, lowering WUE and stem biomass. In contrast, normal water conditions avoided photoinhibition and increased stem biomass but reduced WUE. Overall, moderate drought stress conditions maximized both WUE and stem biomass. Jabereldar et al. (2017) identified sorghum genotype Tagat 10 as the most drought-tolerant, followed by Tagat 14, while Tagat 9 and cv. Gadambalea were the most drought-sensitive. Withholding irrigation at the 3-leaf stage improved crop WUE, reflecting the crop's ability to convert water into grain. Tagat 10 demonstrated superior WUE due to its higher seed yield compared to other genotypes.

10.2 Effect of High Temperature on C⁴ Photosynthesis

Understanding temperature effects on C⁴ plants is crucial for predicting their performance in future climates. As temperature rises, the oxygenation reaction increases, reducing RuBisCO's CO² specificity and limiting carbon gain. C⁴ plants overcome photorespiration by concentrating CO₂ around RuBisCO in bundle sheath cells, maximizing carboxylation (Watson-Lazowski and Ghannoum, 2021).

Location	Ambient temperature warming	Continual	Heatwave (°C / hours/ growth stage)	Photosynthesis Yield		Reference
Yucheng, China	~13.1	-2	N.A.	Increase	N.A.	Zheng et al., (2018)
Illinois, USA	-22.7	-2.64	6/72/Vegetative	Decrease	ns	Siebers et al. (2017)
Illinois, USA	-22.7	N.A.	6/72/ Reproductive	Decrease		Decrease Ruiz-Vera et al., (2015)

Table 3. Results of warming and heat stress studies in Maize

C4 crops like maize and sorghum show varying responses to temperature, with sorghum having higher photosynthetic temperature optima and greater heat and drought tolerance (Watson-Lazowski and Ghannoum, 2021). In a study by Correia et al. (2021), two maize genotypes, B73 and P0023, with contrasting drought and heat tolerance levels, were acclimatized to high temperatures (38°C vs. 25°C) under wellwatered and water deficit (WD) conditions. Both genotypes successfully acclimatized to high temperatures, employing different mechanisms: B73 maintained photosynthetic rates by increasing stomatal conductance (gs), while P0023 preserved g^s and exhibited limited transpiration. The study concluded that key traits for drought and heat tolerance in maize include limited transpiration rates and synchronized regulation of carbon assimilation metabolism.

10.3 CO2 Levels

Elevated CO₂ concentration can influence the growth of C⁴ plants through several mechanisms. One effect is the increase in intercellular $CO₂$ partial pressure, which enhances $CO₂$ partial pressure, which enhances CO₂ assimilation rate. Another is the reduction in stomatal conductance, leading to lower leaf transpiration rate. This decrease in leaf transpiration rate can boost leaf $CO₂$ assimilation rate and growth by conserving soil water, improving shoot water relations, and raising leaf temperature. Additionally, elevated CO₂ may lower mitochondrial respiration, which reduces overall plant respiratory losses and contributes to increased biomass (Ghannoum et al*.,* 2000).

As CO² concentrations rise, some regions will also experience increased frequency and severity of droughts (Lara & Andreo, 2011). The potential for enhanced growth and yield of C⁴ plants at elevated CO2 concentrations is primarily attributed to reduced water use and decreased drought stress, rather than a direct increase in photosynthesis (Lara & Andreo, 2011; Pignon and Long, 2020). Pignon and Long (2020) found that C_4 species with CO_2 concentration in bundle sheath cells showed an indirect stimulation of photosynthesis when atmospheric $CO₂$ increased from 400 µmol mol⁻¹ to 550 µmol mol–1 . However, no yield gains were observed under elevated $CO₂$ without drought stress.

Elevated CO² reduced midday stomatal conductance of FACE-grown sorghum by 32% with irrigation and by 37% under drought stress (Wall et al., 2001). As atmospheric CO₂ continues to rise, sorghum yield is expected to increase in areas with limited water availability (Ottman et al., 2001). Some C_4 plants grown under Free-Air Carbon Dioxide Enrichment (FACE) showed enhanced photosynthetic rates only during drought or under conditions of high atmospheric vapor pressure deficits (Leakey et al., 2009). Additionally, cultivating sorghum under elevated CO₂ mitigated the loss in grain quality caused by drought during the grainfilling stage by delaying physiological and metabolic responses to drought (De Souza et al., 2015).

In a future high-CO₂ environment, water requirements for irrigated sorghum will decrease, while dry-land productivity is expected to rise, assuming minimal global warming (Conley et al., 2001). Elevated $CO₂$ in controlled environments has been shown to increase sugarcane photosynthesis, WUE, biomass, and productivity. The improved WUE of sugarcane under elevated CO² is mainly due to reduced stomatal conductance. Sugarcane grown in elevated CO² had lower leaf stomatal conductance and transpiration, leading to greater leaf WUE. This helped delay the adverse effects of drought, allowing the plants to continue photosynthesis for at least an additional day during episodic drought cycles (Vu and Allen, 2009).

10.3.1 General effects of elevated CO² on photosynthetic heat tolerance

In C_3 species, elevated CO_2 generally enhances heat tolerance for photosynthesis, although at supra-optimal temperatures, this benefit may be diminished or even result in a decrease in photosynthesis. In contrast, C⁴ species often experience reduced photosynthetic thermotolerance at both near-optimal and supraoptimal growing temperatures with elevated CO2. While both C_3 and C_4 plants show similar reductions in stomatal conductance with increasing CO2, C⁴ plants have lower stomatal conductance at any given $CO₂$ level. This leads to reduced transpiration and higher leaf temperatures in C4 plants, which could make them more susceptible to heat-related damage compared to C_3 plants in the same environment (Lara and Andreo, 2011).

The growth of maize and pearl millet under elevated CO² and temperature improved their cellular tolerance to osmotic stress and heat shock. However, maize appeared to benefit more from increased CO2, while pearl millet seemed to benefit more from higher temperatures. The effects of current and anticipated global climate changes are likely to vary between these two species and may similarly impact other C⁴ plant species across different ecosystems, whether natural or managed (Bordignon et al., 2019).

Elevated $CO₂$ is expected to enhance carbon uptake and water-use efficiency, leading to increased productivity of broomcorn millet in semi-arid regions under future high-CO₂ climates (Zhang et al., 2021). Similarly, elevated CO² significantly boosted grain yield and the accumulation of Zn, K, and Mn over three years, as well as enhancing the concentration and accumulation of P in foxtail millet (Gong et al., 2021).

11. ROLE OF C⁴ CROPS IN THE FUTURE

Breeding crop varieties that can better withstand higher temperatures and extreme conditions is crucial for adapting to future climate challenges. Advances in technologies, particularly CRISPR-Cas9 gene editing, are significantly improving our ability to enhance germplasm. This technology allows for efficient overexpression, knockouts, and base pair edits within genetic sequences, making genetic improvements faster and more precise than traditional methods (WatsonLazowski & Ghannoum, 2021). Conventional CRISPR technologies generally comprise two elements: a Cas protein that cleaves nucleic acids and a single guide RNA that associates with the Cas protein, directing it to a specific nucleic acid sequence, which is invariably adjacent to a conserved and compatible protospacer adjacent motif or protospacer flanking site for cleavage (Gaj, 2021). Studies have demonstrated its effectiveness in enhancing disease resistance in rice by correcting specific point mutations without causing double-strand breaks (Gupta et al., 2023; Chen et al., 2024). CRISPR/Cas technology allows precise genetic modifications to improve drought tolerance by targeting genes that regulate water use efficiency and osmotic balance (Shelake et al., 2022). Recent research has shown that modifying the ZmHDT103 gene in maize enhances drought tolerance by improving the plant's ability to withstand water scarcity while maintaining growth and yield under non-stress conditions (Chen et al., 2024). In wheat, engineering the TaRPK1 gene using CRISPR has been found to improve water absorption (Chen et al., 2024; Rahim et al., 2024). Additionally, CRISPR/Cas technology has been used to develop wheat with deeper root systems, improving water access from deeper soil layers. Another breakthrough involves manipulating Sal1 genes to increase osmoprotectant production, such as proline, which enhances drought resistance in wheat by helping plants endure dry periods (Mohr et al., 2022; Chen et al., 2024).

12. CLIMATE RESILIENT MAIZE FOR ASIA (CRMA)

The "Climate Resilient Maize for Asia" project is a collaborative initiative aimed at addressing the challenges faced by resource-poor maize farming communities in South and Southeast Asia, especially considering the anticipated impacts of climate change. Supported by Germany's development agency GIZ and implemented through a public-private partnership, the project focuses on enhancing the resilience of maize crops by developing and distributing abiotic stress-tolerant maize hybrids. These hybrids are specifically designed to thrive in rain-fed, stress-prone production systems, thereby promoting crop diversification, intensification, and higher yields. Building on the successes of the GIZ-funded "Abiotic Stress-Tolerant Maize for Increasing Income and Food Security among the Poor in South and Southeast Asia" project, this initiative addresses critical challenges related to improving maize production, enhancing food security, and building economic stability for smallholder farmers in the region (CIMMYT, 2020). Systematic breeding for heat stress-tolerant maize began at CIMMYT under the Heat Tolerant Maize for Asia project, funded by the United States Agency for International Development (USAID). The project focuses on breeding heat-tolerant maize through collaboration with public research institutions and private seed companies in Bangladesh, Bhutan, India, Nepal, and Pakistan (Zaidi et al., 2023). The Heat Tolerant Maize for Asia (HTMA-II) project, funded by CIMMYT with a budget of ₹73.50 lakhs, focused on developing climate-resilient maize through advanced breeding techniques and collaborations. The project successfully identified and deployed stress-tolerant maize hybrids, including ZH191085, which performed well under heat stress, and ZH182082, which yielded over 8.0 t/ha in heat-stressed conditions (IIMR, n.d.).

13. CLIMATE RESILIENCE IN C4 WEEDS

Climate change is expected to cause shifts in weed community composition, impacting their population dynamics, life cycles, phenology, and infestation levels. Some weed species may go extinct, while others may become more aggressive and invasive. While elevated $CO₂$ levels are likely to boost the productivity of major C³ crops, many of the troublesome agricultural weeds are expected to respond more positively

to the increase in $CO₂$ than the crops themselves, potentially leading to their dominance in agro-ecosystems. Rising temperatures will likely favor the growth of C⁴ weeds, which could result in significant crop yield losses. As climatic factors shift, crop-weed interactions may change, with weeds gaining an advantage and some previously non-threatening species taking over cultivated land. Under conditions of elevated temperature and drought, C_4 weeds are expected to dominate over C_3 crops, while C_3 weeds may prevail under higher $CO₂$ concentrations. However, when both $CO₂$ and temperature levels rise, C_4 weeds are predicted to dominate, further impacting crop production (Anwar et al., 2021). Barnyardgrass (*Echinochloa crus-galli* (L.) Beauv.) a C⁴ weed (Elmore & Paul, 1983) is a strong competitor of rice and can cause severe losses in grain yield across all rice cultures (Awan et al., 2024). A study conducted in China observed that grain yield losses ranged from 12.7% to 42.6% due to competition with *Echinochloa crus-galli* var. *mitis*, 22.3% to 55.2% with *Echinochloa crus-galli* var. *zelayensis*, and 1.5% to 12.1% with junglerice (*Echinochloa colona* (L.) Link) (Zhang et al., 2017; Damalas & Koutroubas, 2023). Another study demonstrated that an increase in the density of the C⁴ weed pigweed redroot (*Amaranthus retoflexus*) led to a reduction in grain and biomass yield components, including ear length, ear diameter, the number of grains per row, the number of rows per ear, total grain number per ear, grain yield, and biological yield in corn (*Zea mays*) (Vazin, 2012).

⁽Lenka et al., 2020)

Climate change is expected to significantly impact weed demographics, leading to shifts in weed species within agroecosystems (Peters et al., 2014; Ramesh et al., 2017). These shifts are crucial for weed management strategies and agricultural productivity. For species to persist in a particular habitat, they must adapt to environmental changes that can result in the alteration of weed flora, range expansion, and migration to new areas. Climate change will likely create opportunities for weeds to invade new ecosystems (Clements & Ditommaso, 2011; Peters et al., 2014). In fact, climate change is predicted to enhance the ability of introduced weed species to adapt to new environments, increasing their potential for invasion in both native and managed ecosystems. Weeds that are well-suited to altered environmental conditions, particularly with higher CO² concentrations, are expected to be more successful in utilizing available resources (Anwar et al., 2021). While C_3 crops may have a competitive edge over C4 weeds under elevated CO² conditions alone, the simultaneous rise in both CO2 and temperature could favor the growth of C⁴ weeds (Alberto et al., 1996). For instance, soybean yields were reduced from 45% to 30% when grown alongside *Amaranthus* retroflexus under elevated CO₂ compared to ambient levels (Ziska, 2003).

14. CONCLUSION

The changing climate has the potential to exert considerable adverse effects on plant physiology, soil fertility, carbon sequestration, and microbial activity, hence inhibiting plant growth and productivity, which would ultimately influence food production. C4 plants are affected by significant global change variables in ways that contrast with C³ plants. Comprehending the responses of C⁴ plants to factors such as temperature, CO2, nutrients, and water is essential for forecasting the adaptability of agricultural and wild C_4 populations to climatic variations, particularly those projected with global climate change. Understanding the mechanisms of stress responses in C4 crops can aid in developing climate-resilient genotypes, thereby maintaining productivity amid fluctuating climatic conditions. This understanding should guide the formulation of targeted policies and programs that promote sustainable farming practices, strengthen climate resilience, and guarantee food security. Policymakers, researchers, and stakeholders must unite to prioritise investments in climate-resilient crops, enhance resource

management, and alleviate the effects of climate change on global food systems.

DISCLAIMER (ARTIFICIAL INTELLIGENCE)

Author(s) hereby declare that NO generative AI technologies such as Large Language Models (ChatGPT, COPILOT, etc) and text-to-image generators have been used during writing or editing of this manuscript.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

REFERENCES

- Aju, P. C., Iwuchukwu, J. J., & Ibe, C. C. (2015). Our forests, our environment, our sustainable livelihoods. *European Journal of Academic Essays, 2*(4), 6–19.
- Alberto, A. M., Ziska, L. H., Cervancia, C. R., & Manalo, P. A. (1996). The influence of increasing carbon dioxide and temperature on competitive interactions between a C_3 crop, rice (*Oryza sativa*), and a C⁴ weed (*Echinochloa glabrescens*). *Functional Plant Biology, 23*(6), 795–802. https://doi.org/10.1071/PP9960795
- Anwar, M., Islam, A. K. M., Yeasmin, S., Rashid, M., Juraimi, A. S., Ahmed, S., & Shrestha, A. (2021). Weeds and their responses to management efforts in a changing climate. *Agronomy, 11*(10), 1921. https://doi.org/10.3390/agronomy11101921
- Awan, T. H., Sta Cruz, P. C., & Chauhan, B. S. (2024). Influence of *Echinochloa crus-galli* density and emergence time on growth, productivity and critical period of competition with dry-seeded rice. *International Journal of Pest Management*, *70*(2),167-179. https://doi.org/10.1080/09670874.2021.196
- 9469 Black, C. C., & Osmond, C. B. (2003). Crassulacean acid metabolism photosynthesis: Working the night shift. *Photosynthesis Research, 76*, 329–341. https://link.springer.com/article/10.1023/a:1 024978220193
- Borghi, G. L. (2021). *Evolution and diversity of photosynthetic metabolism in C3,* C_3 - C_4 *intermediate,* [Doctoral dissertation, Universität Potsdam]. Universität Potsdam. https://doi.org/10.25932/publishup-52220
- Bordignon, L., Faria, A. P., França, M. G., & Fernandes, G. W. (2019). Osmotic stress at the membrane level and photosystem II activity in two C⁴ plants after growth in elevated CO₂ and temperature. Annals of **Applied** Biology, 174(2), 113-122. *Applied Biology, 174*(2), 113–122. https://doi.org/10.1111/aab.12483
- Borrell, A. K., Mullet, J. E., George-Jaeggli, B., van Oosterom, E. J., Hammer, G. L., Klein, P. E., & Jordan, D. R. (2014). Drought adaptation of stay-green sorghum is associated with canopy development, leaf anatomy, root growth, and water uptake. *Journal of Experimental Botany, 65*(21), 6251–6263.

https://doi.org/10.1093/jxb/eru232

Burlakoti, S., Devkota, A. R., Poudyal, S., & Kaundal, A. (2024). Beneficial plant– microbe interactions and stress tolerance in maize. *Applied Microbiology*, *4*(3), 1000- 1015.

> https://doi.org/10.3390/applmicrobiol40300 68

Carrijo, D. R., Lundy, M. E., & Linquist, B. A. (2017). Rice yields and water use under alternate wetting and drying irrigation: A meta-analysis. *Field Crops Research*, *203*, 173-180.

https://doi.org/10.1016/j.fcr.2016.12.002

- Center for Climate and Energy Solutions (C2ES). (2021). *Climate resilience overview*. Center for Climate and Energy Solutions. https://www.c2es.org/building-climateresilience/climate-resilience/
- Chary, G. R., Bhaskar, S., Gopinath, K., Prabhakar, M., Prasad, J., Rao, C. R., & Rao, K. (2022). Climate resilient rainfed agriculture: Experiences from India. In *Climate Change Adaptations in Dryland Agriculture in Semi-Arid Areas* (pp. 3-18). https://link.springer.com/chapter/10.1007/9 78-981-16-7861-5_1
- Chen, F., Chen, L., Yan, Z., Xu, J., Feng, L., He, N., Guo, M., Zhao, J., Chen, Z., & Chen, H. (2024). Recent advances of CRISPRbased genome editing for enhancing staple crops. *Frontiers in plant science*, *15*, 1478398.

https://doi.org/10.3389/fpls.2024.1478398

- CIMMYT. (2020). *Climate-resilient maize for Asia (CRMA)*. International Maize and Wheat Improvement Center. https://www.cimmyt.org/projects/climateresilient-maize-for-asia-crma/
- Clements, D., & Ditommaso, A. (2011). Climate change and weed adaptation: Can evolution of invasive plants lead to greater

range expansion than forecasted? *Weed Research, 51*(3), 227–240. https://doi.org/10.1111/j.1365- 3180.2011.00850.x

- Conley, M. M., Kimball, B. A., Brooks, T. J., Pinter Jr, P. J., Hunsaker, D. J., Wall, G. W., Adam, N. R., LaMorte, R. L., Matthias, A. D., Thompson, T. L., & Leavitt, S. W. (2001). CO² enrichment increases water use efficiency in sorghum. *New Phytologist, 151*(2), 407–412. https://doi.org/10.1046/j.1469- 8137.2001.00184.x
- Cordaid. (2016). *Promoting climate-resilient agriculture for sustainable livelihoods*. https://www.cordaid.org/en/wpcontent/uploads/sites/3/2016/11/2016-11- Cordaid-4P-lowres-Climate-Resilient-Agriculture.pdf
- Correia, P. M., da Silva, A. B., Vaz, M., Carmo-Silva, E., & da Silva, J. M. (2021). Efficient regulation of CO² assimilation enables greater resilience to high temperature and drought in maize. *Frontiers in Plant Science, 12.* https://doi.org/10.3389/fpls.2021.675546
- Damalas, C. A., & Koutroubas, S. D. (2023). Herbicide-resistant barnyardgrass (*Echinochloa crus*‐*galli*) in global rice production. *Weed Biology and Management*, *23*(1), 23-33. https://doi.org/10.1111/wbm.12262
- De Souza, A. P., Cocuron, J. C., Garcia, A. C., Alonso, A. P., & Buckeridge, M. S. (2015). Changes in whole-plant metabolism during the grain-filling stage in sorghum grown under elevated CO² and drought. *Plant Physiology, 169*(3), 1755–1765. https://doi.org/10.1104/pp.15.01054
- Dusenge, M. E., Duarte, A. G., & Way, D. A. (2019). Plant carbon metabolism and climate change: Elevated CO₂ and temperature impacts on photosynthesis, photorespiration, and respiration. *New Phytologist, 221*(1), 32–49. https://doi.org/10.1111/nph.15283
- Edwards, G. E., Franceschi, V. R., & Voznesenskaya, E. V. (2004). Single-cell C⁴ photosynthesis versus the dual-cell (Kranz) paradigm. *Annual Review of Plant Biology, 55*(1), 173–196. https://doi.org/10.1146/annurev.arplant.55. 031903.141725
- Ehleringer, J. R., & Cerling, T. E. (2002). C_3 and C⁴ photosynthesis. *Encyclopedia of global environmental change*, *2*(4), 186-190.

Elmore, C. D., & Paul, R. N. (1983). Composite list of C⁴ weeds. *Weed Science*, *31*(5), 686-692. https://doi.org/10.1017/S00431745000701 93

- Gaj, T. (2021). Next-generation CRISPR technologies and their applications in gene and cell therapy. *Trends in biotechnology*, *39*(7), 692-705. https://www.cell.com/trends/biotechnology/ abstract/S0167-7799(20)30287-0
- Gallaher, T. J., Peterson, P. M., Soreng, R. J., Zuloaga, F. O., Li, D. Z., Clark, L. G., Tyrrell, C. D., Welker, C. A., Kellogg, E. A., & Teisher, J. K. (2022). Grasses through space and time: An overview of the biogeographical and macroevolutionary history of Poaceae. *Journal of Systematics and Evolution, 60*(3), 522–569. https://doi.org/10.1111/jse.12857
- Garnett, T. (2009). Livestock-related greenhouse gas emissions: Impacts and options for policymakers. *Environmental Science & Policy, 12*(4), 491–503. https://doi.org/10.1016/j.envsci.2009.01.00 6
- Ghannoum, O. (2016). How can we breed for more water use-efficient sugarcane? *Journal of Experimental Botany, 67*(3), 557–559.

https://doi.org/10.1093/jxb/erw009

- Ghannoum, O., Caemmerer, S. V., & Conroy, J. P. (2002). The effect of drought on plant water use efficiency of nine NAD-ME and nine NADP-ME Australian C₄ grasses. *Functional Plant Biology, 29*(11), 1337– 1348. https://doi.org/10.1071/FP02056
- Ghannoum, O., Caemmerer, S. V., Ziska, L. H., & Conroy, J. P. (2000). The growth response of C4 plants to rising atmospheric CO2 partial pressure: A reassessment. *Plant, Cell & Environment, 23*(9), 931–942. https://doi.org/10.1046/j.1365- 3040.2000.00609.x
- Ghannoum, O., Evans, J. R., & Caemmerer, S. (2011). Nitrogen and water use efficiency of C⁴ plants. In A. S. Raghavendra & R. F. Sage (Eds.), *C⁴ photosynthesis and related CO² concentrating mechanisms* (pp. 129–146). https://www.researchgate.net/profile/JohnE vans11/publication/306203835_Nitrogen_a nd_water_use_efficiency_of_C4_plants/lin ks/5a7781bfaca2722e4df105fd/Nitrogenand-water-use-efficiency-of-C4-plants.pdf
- Goh, C.-H., Ko, S.-M., Koh, S., Kim, Y.-J., & Bae, H.-J. (2012). Photosynthesis and

environments: photoinhibition and repair mechanisms in plants. *Journal of Plant Biology*, *55*, 93-101. https://link.springer.com/article/10.1007/s1 2374-011-9195-2

- Gong, Z., Dong, L., Lam, S., Zhang, D., Zong, Y., Hao, X., & Li, P. (2021). Nutritional quality in response to elevated $CO₂$ concentration in foxtail millet (*Setaria italica*). *Journal of Cereal Science, 103*, 318. https://doi.org/10.1016/j.jcs.2021.103318
- Gowik, U., & Westhoff, P. (2011). The path from C³ to C⁴ photosynthesis. *Plant Physiology*, *155*(1), 56-63. https://doi.org/10.1104/pp.110.165308
- Gupta, A., Liu, B., Chen, Q. J., & Yang, B. (2023). High‐efficiency prime editing enables new strategies for broad‐spectrum resistance to bacterial blight of rice. *Plant biotechnology journal*, *21*(7), 1454-1464. https://doi.org/10.1111/pbi.14049
- Hasan, M. K., Desiere, S., D'Haese, M., & Kumar, L. (2018). Impact of climate-smart agriculture adoption on the food security of coastal farmers in Bangladesh. *Food Security*, *10*, 1073-1088. https://doi.org/10.1007/s12571-018-0824-1
- Hatch, M. D. (1987). C4 photosynthesis: A unique blend of modified biochemistry, anatomy, and ultrastructure. *Biochimica et Biophysica Acta (BBA) - Reviews on Bioenergetics, 895*(2), 81–106. https://doi.org/10.1016/S0304- 4173(87)80009-5
- Hirst, D. (2021). Aviation, decarbonisation and climate change. *Aviation*. https://nvvl.eu/wpcontent/uploads/2022/07/Hearst-CBP-8826-kopie.pdf
- Hodkinson, T. R. (2018). Evolution and taxonomy of the grasses (Poaceae): A model family for the study of species-rich groups. *Annual Plant Reviews Online*, 255–294. doi: 10.1002/9781119312994.apr0622
- Huang, S., Knight, C. A., Hoover, B. K., & Ritter, M. (2020). Leaf functional traits as predictors of drought tolerance in urban trees. *Urban Forestry & Urban Greening, 48,* 126577.

https://doi.org/10.1016/j.ufug.2019.126577

- Indian Institute of Maize Research (IIMR). (n.d.). *Heat tolerant maize for Asia (HTMA-II)*. Indian Institute of Maize Research. https://iimr.icar.gov.in/?p=367
- Jabereldar, A. A., El Naim, A. M., Abdalla, A. A., & Dagash, Y. M. (2017). Effect of water stress on yield and water use efficiency of

sorghum (*Sorghum bicolor* L. Moench) in a semi-arid environment. *International Journal of Agriculture and Forestry, 7*(1), 1–6. 1 https://0.5923/j.ijaf.20170701.01

- Jin, K., Chen, G., Yang, Y., Zhang, Z., & Lu, T. (2023). Strategies for manipulating Rubisco and creating photorespiratory bypass to boost C3 photosynthesis: Prospects on modern crop improvement. *Plant, Cell & Environment*, *46*(2), 363-378. https://doi.org/10.1111/pce.14500
- Kajala, K., Covshoff, S., Karki, S., Woodfield, H., Tolley, B. J., Dionora, M. J. A., Mogul, R. T., Mabilangan, A. E., Danila, F. R., Hibberd, J. M., & Quick, W. P. (2011). Strategies for engineering a two-celled C⁴ photosynthetic pathway into rice. *Journal of Experimental Botany, 62*(9), 3001–3010. https://doi.org/10.1093/jxb/err022
- Kakraliya, S. K., Jat, H. S., Sapkota, T. B., Singh, I., Kakraliya, M., Gora, M. K., Sharma, P. C., & Jat, M. L. (2021). Effect of climatesmart agriculture practices on climate change adaptation, greenhouse gas mitigation and economic efficiency of ricewheat system in India. *Agriculture*, *11*(12), 1269.

https://doi.org/10.3390/agriculture1112126 9

- Karakurt, I., Aydin, G., & Aydiner, K. (2012). Sources and mitigation of methane emissions by sectors: A critical review. *Renewable Energy, 39*(1), 40–48. https://doi.org/10.1016/j.renene.2011.09.00 6
- Khalifa, M., & Eltahir, E. A. (2023). Assessment of global sorghum production, tolerance, and climate risk. *Frontiers in Sustainable Food Systems, 7,* 1184373. https://doi.org/10.3389/fsufs.2023.1184373
- Kijne, J. W. (2006). Abiotic stress and water scarcity: Identifying and resolving conflicts from plant level to global level. *Field Crops Research, 97*(1), 3–18. https://doi.org/10.1016/j.fcr.2005.08.011
- Korres, N. E., Norsworthy, J. K., Tehranchian, P., Gitsopoulos, T. K., Loka, D. A., Oosterhuis, D. M., Gealy, D. R., Moss, S. R., Burgos, N. R., & Miller, M. R. (2016). Cultivars to face climate change effects on crops and weeds: A review. *Agronomy for Sustainable Development, 36,* 1–22. https://doi.org/10.2307/3565625
- Koteyeva, N. K., Voznesenskaya, E. V., Pathare, V. S., Borisenko, T. A., Zhurbenko, P. M., Morozov, G. A., & Edwards, G. E. (2023). Biochemical and structural diversification

of C⁴ photosynthesis in Tribe Zoysieae (Poaceae). *Plants, 12*(23), 4049. https://doi.org/10.3390/plants12234049

- Lal, M. (2000). Climatic change: Implications for India's water resources. *Journal of Social and Economic Development, 3*(1), 57–87. https://web.archive.org/web/201907130121 20id_/http://www.isec.ac.in:80/JSED/JSED _V3_I1_57-87.pdf
- Lara, M. V., & Andreo, C. S. (2011). C4 plants adaptation to high levels of $CO₂$ and drought environments. In P. Ahmad & M. N. V. Prasad (Eds.), *Abiotic stress in plants: Mechanisms and adaptations* (pp. 415–428).
- Leakey, A. D., Ferguson, J. N., Pignon, C. P., Wu, A., Jin, Z., Hammer, G. L., & Lobell, D. B. (2019). Water use efficiency as a constraint and target for improving the resilience and productivity of C_3 and C_4 crops. *Annual review of plant biology*, *70*, 781-808. https://doi.org/10.1146/annurevarplant-042817-040305
- Leegood, R. C. (2013). Strategies for engineering C⁴ photosynthesis. *Journal of plant physiology*, *170*(4), 378-388. https://doi.org/10.1016/j.jplph.2012.10.011
- Lenka, B., Kulkarni, G. U., Moharana, A., Singh, A. P., Pradhan, G. S., & Muduli, L. (2020). Millets: Promising crops for climate-smart agriculture. *International Journal of Current Microbiology and Applied Sciences, 9*(11), 656-668.

https://doi.org/10.20546/ijcmas.2020.911.0 81

- Levitus, S., Antonov, J., Boyer, T., Baranova, O., Garcia, H., Locarnini, R., Mishonov, A., Reagan, J., Seidov, D., & Yarosh, E. (2017). NCEI ocean heat content, temperature anomalies, salinity anomalies, thermosteric sea level anomalies, halosteric sea level anomalies, and total steric sea level anomalies from 1955 to present calculated from in situ oceanographic subsurface profile data (NCEI Accession 0164586). *NOAA National Centers for Environmental Information*, *10*, v53f54mvp.
- Liaqat, W., Altaf, M. T., Barutçular, C., Mohamed, H. I., Ahmad, H., Jan, M. F., & Khan, E. H. (2024). Sorghum: a Star Crop to Combat Abiotic Stresses, Food Insecurity, and Hunger Under a Changing Climate: a Review. *Journal of Soil Science and Plant Nutrition*,1-28.

https://doi.org/10.1007/s42729-023-01607- 7

- Lin, H. C., Coe, R. A., Quick, W. P., & Bandyopadhyay, A. (2019). Climateresilient future crop: Development of C⁴ Rice. *Sustainable solutions for food Security: Combating climate change by adaptation*,111-124. https://doi.org/10.1007/978-3-319-77878- 5_6
- Lombe, P., Carvalho, E., & Rosa-Santos, P. (2024). Drought Dynamics in Sub-Saharan Africa: Impacts and Adaptation Strategies. *Sustainability*, *16*(22), 9902. https://doi.org/10.3390/su16229902
- Long, S. (1983). C₄ photosynthesis at low temperatures. *Plant, Cell & Environment*, *6*(4),345-363. https://doi.org/10.1111/1365- 3040.ep11612141
- Long, S. P., Ainsworth, E. A., Rogers, A., & Ort, D. R. (2004). Rising atmospheric carbon dioxide: Plants FACE the future. *Annual Review of Plant Biology, 55*(1), 591-628. https://doi.org/10.1146/annurev.arplant.55. 031903.141610
- Lopes, M. S., Araus, J. L., Van Heerden, P. D., & Foyer, C. H. (2011). Enhancing drought tolerance in C⁴ crops. *Journal of Experimental Botany, 62*(9), 3135-3153. https://doi.org/10.1093/jxb/err105
- Luo, T., Liu, X., & Lakshmanan, P. (2023). A combined genomics and phenomics approach is needed to boost breeding in sugarcane. *Plant Phenomics*, *5*, 0074. https://doi.org/10.34133/plantphenomics.0 074
- Maheswari, M., Sarkar, B., Vanaja, M., Rao, S. M., Prasad, J. V. N. S., Prabhakar, M., Chary, R. G., Venkateswarlu, B., Choudhury, R. P., Yadava, D. K., and Bhaskar, S. 2015. *Climate resilient crop varieties for sustainable food production under aberrant weather conditions*. NICRA Bulletin No.4. ICAR-Central Research
Institute for Dryland Agriculture. Institute for Dryland Agriculture, Hyderabad. 64p.https://www.researchgate. net/profile/BasudebSarkar/publication/3317 31142_Climate_Resilient_Crop_Varieties_f or_Sustainable_Food_Production_under_ Aberrant_Weather_Conditions/links/61a1d ae707be5f31b7ba875b/Climate-Resilient-Crop-Varieties-for-Sustainable-Food-Production-under-Aberrant-Weather-Conditions.pdf
- Mall, R. K., Gupta, A., & Sonkar, G. (2017). Effect of climate change on agricultural crops. In *Current developments in biotechnology and bioengineering* (pp. 23-

46). https://doi.org/10.1016/B978-0-444- 63661-4.00002-5

- Marin, F. R., Jones, J. W., Singels, A., Royce, F., Assad, E. D., Pellegrino, G. Q., & Justino, F. (2013). Climate change impacts on sugarcane attainable yield in southern Brazil. *Climate Change, 117*(1), 227-239. https://doi.org/10.1007/s10584-012-0561-y
- Mehri, N., Fotovat, R., Saba, J., & Jabbari, F. (2009). Variation of stomata dimensions and densities in tolerant and susceptible wheat cultivars under drought stress. *Journal of Food Agriculture and Environment*, *7*(1), 167-170.
- Mesa-Jurado, M. A., Martin-Ortega, J., Ruto, E., & Berbel, J. (2012). The economic value of guaranteed water supply for irrigation under scarcity conditions. *Agricultural water management*, *113*, 10-18. https://doi.org/10.1016/j.agwat.2012.06.00 9
- Misra, V., Shrivastava, A. K., Mall, A. K., Solomon, S., Singh, A. K., & Ansari, M. I. (2019). Can sugarcane cope with
increasing atmospheric $CO₂$ atmospheric CO₂ concentration? *Australian Journal of Crop Science, 13*(5), 780-784.
- Morrone, O., Aagesen, L., Scataglini, M. A., Salariato, D. L., Denham, S. S., Chemisquy, M. A., Sede, S. M., Giussani, L. M., Kellogg, E. A., & Zuloaga, F. O. (2012). Phylogeny of the Paniceae (Poaceae: Panicoideae): integrating plastid DNA sequences and morphology into a new classification. *Cladistics*, *28*(4), 333- 356. https://doi.org/10.1111/j.1096- 0031.2011.00384.x
- Mohr, T., Horstman, J., Gu, Y. Q., Elarabi, N. I., Abdallah, N. A., & Thilmony, R. (2022). CRISPR-Cas9 gene editing of the Sal1 gene family in wheat. *Plants*, *11*(17), 2259. https://doi.org/10.3390/plants11172259
- Mwamahonje, A., Mdindikasi, Z., Mchau, D., Mwenda, E., Sanga, D., Garcia-Oliveira, A. L., & Ojiewo, C. O. (2024). Advances in Sorghum Improvement for Climate Resilience in the Global Arid and Semi-Arid Tropics: A Review. *Agronomy*, *14*(12), 3025.

https://doi.org/10.3390/agronomy14123025

Nadew, D., Bejiga, T., & Teressa, T. (2021). Breeding of sorghum crop for resistance and tolerance to drought. *International Research Journal of Plant Crops Science, 188*,188-195.

https://www.advancedscholarsjournals.org/

- NASA [National Aeronautics and Space Administration]. (2024). NASA home page [Online]. Available at: https://climate.nasa.gov/ [Accessed 12 December 2024].
- National Geographic Society, n.d. *Earth's Changing Climate*. National Geographic Education. Available at: https://education.nationalgeographic.org/re source/earths-changing-climate/ (Accessed: 23 December 2024).
- Nerem, R. S., Beckley, B. D., Fasullo, J. T., Hamlington, B. D., Masters, D., & Mitchum, G. T. (2018). Climate-change–driven accelerated sea-level rise detected in the altimeter era. *Proceedings of the National Academy of Sciences*, *115*(9), 2022-2025. https://doi.org/10.1073/pnas.1717312115
- Nikolaou, G., Neocleous, D., Christou, A., Kitta, E., & Katsoulas, N. (2020). Implementing sustainable irrigation in water-scarce regions under the impact of climate change. *Agronomy*, *10*(8), 1120. https://doi.org/10.3390/agronomy10081120
- Nunes, T. D., Zhang, D., & Raissig, M. T. (2020). Form, development and function of grass stomata. *The Plant Journal*, *101*(4), 780- 799. https://doi.org/10.1111/tpj.14552
- Osmond, C. B. (2007). Crassulacean acid metabolism: now and then. In *Progress in Botany* (pp. 3-32). https://link.springer.com/content/pdf/10.100 7/978-3-540-36832- 8_1?pdf=chapter%20toc
- Ottman, M. J., Kimball, B. A., Pinter, P. J., Wall, G. W., Vanderlip, R. L., Leavitt, S. W., LaMorte, R. L., Matthias, A. D., & Brooks, T. J. (2001) . Elevated $CO₂$ increases sorghum biomass under drought conditions. *New Phytologist, 150*(2), 261- 273. https://doi.org/10.1046/j.1469- 8137.2001.00110.x
- Pardo, J., & VanBuren, R. (2021). Evolutionary innovations driving abiotic stress tolerance in C⁴ grasses and cereals. *Plant Cell, 33*(11), 3391-3401. https://doi.org/10.1093/plcell/koab205
- Pathak, H., Tewari, A., Sankhyan, S., Dubey, D., Mina, U., Singh, V. K., & Jain, N. (2011). Direct-seeded rice: potential, performance and problems-Areview. *Current Advances in Agricultural Sciences (An International Journal)*, *3*(2), 77-88.
- Paul, M. J., & Foyer, C. H. (2001). Sink regulation of photosynthesis. *Journal of Experimental Botany, 52*(360), 1383-1400. https://doi.org/10.1093/jexbot/52.360.1383
- Peters, K., Breitsameter, L., & Gerowitt, B. (2014). Impact of climate change on weeds in agriculture: a review. *Agronomy for Sustainable Development*, *34*, 707-721. https://doi.org/10.1007/s13593-014-0245-2
- Pignon, C. P., & Long, S. P. (2020). Retrospective analysis of biochemical limitations to photosynthesis in 49 species: C⁴ crops appear still adapted to pre‐industrial atmospheric [CO2]. *Plant, Cell & Environment, 43*(11), 2606-2622. https://doi.org/10.1111/pce.13863
- Pinto, H., Powell, J. R., Sharwood, R. E., Tissue, D. T., & Ghannoum, O. (2016). Variations in nitrogen use efficiency reflect the biochemical subtype while variations in water use efficiency reflect the evolutionary lineage of C4 grasses at interglacial CO2. *Plant, Cell & Environment, 39*(3), 514-526. https://doi.org/10.1111/pce.12636
- Rahim, A. A., Uzair, M., Rehman, N., Fiaz, S., Attia, K. A., Abushady, A. M., Yang, S. H., & Khan, M. R. (2024). CRISPR/Cas9 mediated TaRPK1 root architecture gene mutagenesis confers enhanced wheat yield. *Journal of King Saud University-Science*, *36*(2), 103063. https://doi.org/10.1016/j.jksus.2023.103063
- Ramesh, K., Matloob, A., Aslam, F., Florentine, S. K., & Chauhan, B. S. (2017). Weeds in a changing climate: vulnerabilities, consequences, and implications for future weed management. *Frontiers in Slant science*, *8*, 95. https://doi.org/10.3389/fpls.2017.00095
- Raza, A., Razzaq, A., Mehmood, S. S., Zou, X., Zhang, X., Lv, Y., & Xu, J. (2019). Impact of climate change on crops adaptation and strategies to tackle its outcome: A review. *Plants*, *8*(2), 34. https://doi.org/10.3390/plants8020034
- Raza, G., Ali, K., Hassan, M. A., Ashraf, M., Khan, M. T., & Khan, I. A. (2019). Sugarcane as a bioenergy source. *Sugarcane biofuels: status, potential, and prospects of the sweet crop to fuel the world*, 3-19. https://doi.org/10.1007/978-3- 030-18597-8_1
- Romanowska, E., & Wasilewska-Dębowska, W. (2022). Light-Dependent Reactions of Photosynthesis in Mesophyll and Bundle Sheath Chloroplasts of C⁴ Plant Maize. How Our Views Have Changed in Recent
Years. Acta Societatis Botanicorum Years. *Acta Societatis Botanicorum Poloniae*,*91*.

https://doi.org/10.5586/asbp.9112

Ruales, J. H., Seriño, M. N. V., Ratilla, T. C., Cuizon, J. G., & Enerlan, W. C. (2020). Investment appraisal of selected climate smart agricultural (CSA) practices among small scale coconut farmers in Leyte, Philippines.

https://managementjournal.usamv.ro/pdf/v ol.20_3/Art54.pdf

Ruiz‐Vera, U. M., Siebers, M. H., Drag, D. W., Ort, D. R., & Bernacchi, C. J. (2015). Canopy warming caused photosynthetic acclimation and reduced seed yield in maize grown at ambient and elevated [CO₂]. *Global Change Biology, 21*(11), 4237-4249.

https://doi.org/10.1111/gcb.13013

- Rykowski, K. (2000). The role of forest ecosystems and wood in controlling the absorption and emission of carbon dioxide. *Geographia Polonica*, *73*(2), 65-88.
- Sabine, C. L., Feely, R. A., Gruber, N., Key, R. M., Lee, K., Bullister, J. L., Wanninkhof, R., Wong, C., Wallace, D. W., & Tilbrook, B. (2004). The oceanic sink for anthropogenic CO₂. *Science, 305*(5682), 367-371 https://doi.org/10.1126/science.1097403.
- Sage, R. F., & Kubien, D. S. (2007). The temperature response of C_3 and C_4 photosynthesis. *Plant, Cell & Environment*, *30*(9), 1086-1106. https://doi.org/10.1111/j.1365- 3040.2007.01682.x
- Sage, R. F., Christin, P. A., & Edwards, E. J. (2011a). The C⁴ plant lineages of planet Earth. *Journal of Experimental Botany, 62*, 3155–3169.

https://doi.org/10.1093/jxb/err048

- Sage, R. F., & Zhu, X. G. (2011b). Exploiting the engine of C⁴ photosynthesis. *Journal of Experimental Botany, 62*(9), 2989-3000. https://doi.org/10.1093/jxb/err179
- Samuel, J., Rao, C. A. R., Raju, B. M. K., Reddy, A. A., Pushpanjali, Reddy, A. G. K., Kumar, R. N., Osman, M., Singh, V. K., & Prasad, J. V. N. S. (2021). Assessing the impact of climate resilient technologies in minimizing drought impacts on farm incomes in drylands. *Sustainability*, *14*(1), 382. https://doi.org/10.3390/su14010382
- Schiller, K., & Bräutigam, A. (2021). Engineering of crassulacean acid metabolism. *Annual Review of Plant Biology, 72*(1), 77-103. https://doi.org/10.1146/annurev-arplant-071720-104814
- Schlüter, U., & Weber, A. P. (2020). Regulation and evolution of C⁴ photosynthesis. *Annual Review of Plant Biology, 71*(1), 183-215.

https://doi.org/10.1146/annurev-arplant-042916-040915

- Schmitt, M. R., & Edwards, G. E. (1981). Photosynthetic capacity and nitrogen use efficiency of maize, wheat, and rice: A comparison between C_3 and C_4 comparison between C_3 and C_4 photosynthesis. *Journal of Experimental Botany, 32*(3), 459-466. https://doi.org/10.1093/jxb/32.3.459
- Sekhar, K. M., Kota, V. R., Reddy, T. P., Rao, K., & Reddy, A. R. (2021). Amelioration of plant responses to drought under elevated CO 2 by rejuvenating photosynthesis and nitrogen use efficiency: Implications for future climate-resilient crops. *Photosynthesis Research*, *150*, 21-40. https://doi.org/10.1007/s11120-020-00772- 5
- Sheehy, J. E., Ferrer, A. B., Mitchell, P. L., Elmido-Mabilangan, A., Pablico, P., & Dionora, M. J. A. (2008). How the rice crop works and why it needs a new engine. In *Charting new pathways to C4 rice* (pp. 3- 26). https://doi.org/10.1142/9789812709523_00

 $^{\circ}$

- Siebers, M. H., Slattery, R. A., Yendrek, C. R., Locke, A. M., Drag, D., Ainsworth, E. A., Bernacchi, C. J., & Ort, D. R. (2017). Simulated heat waves during maize reproductive stages alter reproductive growth but have no lasting effect when applied during vegetative stages. *Agricultural Ecosystems & Environment, 240*, 162-170. https://doi.org/10.1016/j.agee.2016.11.008
- Sikka, A. K., Alam, M. F., & Mandave, V. (2022). Agricultural water management practices to improve the climate resilience of irrigated agriculture in India. *Irrigation and Drainage*, 71, 7-26. https://doi.org/10.1002/ird.2696
- Sivakumar, M. V., & Stefanski, R. (2011). Climate change in South Asia. *Climate change and food security in South Asia*, 13-30. https://doi.org/10.1007/978-90-481- 9516-9_2
- Sonawane, B. V. (2016). *Environmental regulation of CO² concentrating mechanisms in C⁴ grasses with different biochemical subtypes* (Doctoral dissertation). Western Sydney University, Australia.
- Sonawane, B. V., Sharwood, R. E., Whitney, S., & Ghannoum, O. (2018). Shade compromises the photosynthetic efficiency of NADP-ME less than that of PEP-CK and

NAD-ME C⁴ grasses. *Journal of Experimental Botany, 69*, 3053–3068. https://doi.org/10.1093/jxb/ery129

Sovacool, B. K., Griffiths, S., Kim, J., & Bazilian, M. (2021). Climate change and industrial F-gases: A critical and systematic review of developments, sociotechnical systems and policy options for reducing synthetic greenhouse gas emissions. *Renewable and Sustainable Energy Reviews*, *141*, 110759.

https://doi.org/10.1016/j.rser.2021.110759

- Stavi, I., & Lal, R. (2013). Agriculture and greenhouse gases, a common tragedy. A review. *Agronomy for Sustainable Development*, *33*, 275-289. https://doi.org/10.1007/s13593-012-0110-0
- Tingting, X., Peixi, S. U., & Lishan, S. (2010). Photosynthetic characteristics and water use efficiency of sweet sorghum under different watering regimes. *Pakistan Journal of Botany, 42*(6), 3981–3994. https://www.pakbs.org/pjbot/PDFs/42(6)/PJ B42(6)3981.pdf
- Vazin, F. (2012). The effects of pigweed redroot (Amaranthus retoflexus) weed competition and its economic thresholds in corn (Zea mays). *Planta Daninha*, *30*, 477-485. https://www.scielo.br/j/pd/a/sYjyV4t3vp3Z MK6LnnK7vbr/?lang=en
- Velicogna, I., Mohajerani, Y., Landerer, F., Mouginot, J., Noel, B., Rignot, E., Sutterley, T., van den Broeke, M., van Wessem, M., & Wiese, D. (2020). Continuity of ice sheet mass loss in Greenland and Antarctica from the GRACE and GRACE Follow‐On missions. *Geophysical Research Letters*, *47*(8), e2020GL087291.

https://doi.org/10.1029/2020GL087291

- Venter, N. (2015). Drought responses of selected C4 photosynthetic NADP-ME and NAD-ME Panicoideae and Aristidoideae grasses (Doctoral dissertation). Rhodes University. https://core.ac.uk/download/pdf/145040434 .pdf
- Viswanathan, P. K., Kavya, K., & Bahinipati, C. S. (2020). Global patterns of climateresilient agriculture: A review of studies and imperatives for empirical research in India. *Review of Development and Change, 25*(2), 169–192. https://doi.org/10.1177/0972266120966211
- Vu, J. C., & Allen Jr., L. H. (2009). Growth at elevated CO2 delays the adverse effects of drought stress on leaf photosynthesis of the C4 sugarcane. *Journal of Plant*

Physiology, 166(2), 107–116. https://doi.org/10.1016/j.jplph.2008.02.009

- Voznesenskaya, E. V., Franceschi, V. R., Chuong, S. D., & Edwards, G. E. (2006). Functional characterization of phosphoenolpyruvate carboxykinase-type C4 leaf anatomy: immuno-, cytochemical and ultrastructural analyses. *Annals of Botany*, *98*(1), 77-91. https://doi.org/10.1093/aob/mcl096
- Wall, G. W., Brooks, T. J., Adam, N. R., Cousins, A. B., Kimball, P. J., Pinter Jr, R. L., La Morte, R. L., & Matthias, A. D. (2001). Elevated atmospheric CO2 improved sorghum plant water status by ameliorating the adverse effects of drought. *New Phytologist, 152(2),* https://doi.org/10.1046/j.0028- 646X.2001.00260.x
- Wang, Y., Bräutigam, A., Weber, A. P., & Zhu, X.-G. (2014). Three distinct biochemical subtypes of C₄ photosynthesis? A modelling analysis. *Journal of Experimental Botany, 65*(13), 3567–3578. https://doi.org/10.1093/jxb/eru058
- Wasi, A., Tahseen, S., Arishakausar, Bhatt, A. Y., & Shahzad, A. (2023). Current Status and Future Prospectus of Bioenergy Crops. In *Biotechnology and Omics Approaches for Bioenergy Crops* (pp. 271- 288). Springer. https://doi.org/10.1007/978- 981-99-4954-0_13
- Watson-Lazowski, A., & Ghannoum, O. (2021). The outlook for C₄ crops in future climate scenarios. In K. M. Becklin, J. K. Ward, & D. A. Way (Eds.), *Photosynthesis, respiration, and climate change* (pp. 251– 281). Springer, Cham. https://doi.org/10.1007/978-3-030-64926- 5_9
- Way, D. A., Katul, G. G., Manzoni, S., & Vico, G. (2014). Increasing water use efficiency along the C_3 to C_4 evolutionary pathway: A stomatal optimization perspective. *Journal of Experimental Botany, 65*(13), 3683– 3693. https://doi.org/10.1093/jxb/eru205
- Zabaleta, E., Martin, M. V., & Braun, H. P. (2012). A basal carbon concentrating mechanism in plants. *Plant Science, 187*, 97–104. https://doi.org/10.1016/j.plantsci.2012.02.0 01
- Zahedi, S. M., Karimi, M., Venditti, A., Zahra, N., Siddique, K. H., & Farooq, M. (2024). Plant Adaptation to Drought Stress: The Role of Anatomical and Morphological Characteristics in Maintaining the Water

Status. *Journal of Soil Science and Plant Nutrition*, 1-19. https://doi.org/10.1007/s42729-024-02141 w

- Zaidi, P. H., Vinayan, M. T., Nair, S. K., Kuchanur, P. H., Kumar, R., Singh, S. B., & Prasanna, B. M. (2023). Heat-tolerant maize for rainfed hot, dry environments in the lowland tropics: From breeding to improved seed delivery. *The Crop Journal, 11*(4), 986-1000. https://doi.org/10.1016/j.cj.2023.06.008
- Zhang, Z., Gu, T., Zhao, B., Yang, X., Peng, Q., Li, Y., & Bai, L. (2017). Effects of common *Echinochloa* varieties on grain yield and grain quality of rice. *Field Crops Research*, *203*, 163-172. https://doi.org/10.1016/j.fcr.2016.12.003
- Zhang, D., Li, A., Lam, S. K., Li, P., Zong, Y., Gao, Z., & Hao, X. (2021). Increased carbon uptake under elevated $CO₂$ concentration enhances water-use efficiency of C4 broomcorn millet under

drought. *Agricultural Water Management, 245*, 106631. https://doi.org/10.1016/j.agwat.2020.10663 1

- Zhao, W., Liu, L., Shen, Q., Yang, J., Han, X., Tian, F., & Wu, J. (2020). Effects of water stress on photosynthesis, yield, and water use efficiency in winter wheat. *Water*, *12*(8), 2127. https://doi.org/10.3390/w12082127
- Zheng, Y. P., Li, R. Q., Guo, L. L., Hao, L. H., Zhou, H. R., Li, F., Peng, Z. P., Cheng, D. J., & Xu, M. (2018). Temperature responses of photosynthesis and respiration of maize (*Zea mays*) plants to experimental warming. *Russian Journal of Plant Physiology, 65*(4), 524–531. https://doi.org/10.1134/S10214437180401 92
- Ziska, L. W. (2003). Evaluation of yield losses in field sorghum from a C_3 and C_4 weeds with increasing CO2. *Weed Science, 51*, 914– 918. https://doi.org/10.1614/WS-03-002R

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