Terrestrial plants are regularly subjected to strong temperature variations. These variations can reach an amplitude of 40°C or even more, both in polar regions and in hot desert areas. Being rooted, they have reduced mobility and must cope with changes in their environment. The assimilation of CO$_2$ by plants via photosynthesis is the gateway to carbon in the biosphere. What is the thermal amplitude that allows it to function? How does photosynthesis react to rapid and slow temperature variations? What is the diversity of responses? What are the physiological processes that limit it? Crucial questions tare o be considered in the context of global warming.

1. Plant production and climate change

The current increase in greenhouse gas emissions will cause an increase in atmospheric temperature of 2 to 3°C in the next 50 years (see A carbon cycle disrupted by human activities). At the same time, heat waves and extreme heat periods will be more frequent and of longer duration [1]. Agricultural production and the functioning of forests will therefore be greatly affected. Models based on large-scale observations indicate that, in the absence of agronomic adaptation, the decrease in crop yields can reach 17% for each 1°C increase in the temperature of the growing season [2].

The production of higher plants depends in particular (but not only [3]) on leaf photosynthesis (see Shedding light on photosynthesis & The The path of carbon in photosynthesis). CO$_2$ enters the leaf where its reduction in the chloroplasts is accompanied by O$_2$ production. Its entry is almost exclusively through the stomata (Figure 1). For each molecule of CO$_2$ absorbed, 50 to 300 molecules of water are transpired from the leaves, depending on the plant. This water allows, among other things, the cooling of the leaf (see Focus Leaf transpiration and heat protection).
The leaf is a converter of solar energy into chemical energy and, like any energy converter, requires a permanent cooling system.

The climate changes that are currently occurring make it necessary to understand the effects of temperature on photosynthesis.

2. The thermal optimum of photosynthesis

2.1. Diagram of the thermal response

Photosynthetic CO₂ uptake varies with temperature. In most cases its response to temperature is rapidly reversible between about 10 and 34°C. In this range of temperatures it presents a maximum value: a thermal optimum.

Below 10°C and above 34°C plants start to set up protective mechanisms. For these extreme values, CO₂ assimilation is often unstable and can be cancelled more or less quickly: the leaf is then under stress (Figure 2).

2.2. A thermal optimum based on the average temperature of the environment
Plants in cold environments or with a cold growing season have a higher photosynthesis at low temperatures. Plants in warm environments, or growing during the warm season, have a higher photosynthesis at high temperatures.

Figure 3. Deschampsia antarctica is one of two flowering plants found in Antarctica. It is often subjected to negative temperatures. The snow that frequently covers it protects it from extreme temperatures. [Source: Lomvi2, CC BY-SA 3.0, via Wikimedia Commons]

For example, the thermal optimum for CO₂ assimilation [41] in Deschampsia antarctica (Figure 3) and Colobanthus quitensis, the only two Antarctic flowering plants, is between 8 and 15°C, while it is around 45°C in Tridestomia oblongifolia, a warm desert plant from Central America. The latter species probably holds the world record for flowering plants in this respect.

2.3. Acclimatization to the thermal conditions of the environment

Figure 4. Variations in CO₂ assimilation as a function of leaf temperature, in a plant grown at 10°C (red) or 25°C. Measurements made on Pea, under a light close to saturation. CO₂ content in ambient air: 390 ppm. [Source: Author’s diagram]

Differences in the thermal response of photosynthesis are also found in individuals of the same species growing at different
temperatures. Figure 4 shows CO₂ assimilation in pea grown at 10 or 25°C.

In the first case (cultivation at 10°C) the thermal optimum is about 16°C, while it is higher than 25°C in the second (cultivation at 25°C). At low temperatures, CO₂ assimilation is higher in plants grown at 10°C.

In this case the adjustment to cool conditions is a gain for the plant.

2.4. Acclimatization can be rapid

For example, the photosynthesis of *Hammada scoparia*, a bush in the deserts of the Middle East (Negev, Wadi Rum) follows the seasonal variations in temperature: its thermal optimum varies from 29°C in early spring to 41°C in summer and then to 28°C in autumn.

Changes in the thermal optimum can be even more rapid and of great amplitude. For example, in a seaside clone of *Encelia californica*, a change in growth temperature from 30°C (constant day and night temperature) to 15°C during the day and 2°C during the night for three days is sufficient to lower the thermal optimum by about ten degrees.

In general, these changes can be measured in both growing and mature leaves, with the response being of greater amplitude in growing leaves.

2.5. Heat-sensitive versus cold-sensitive species
Warm acclimation of cool-adapted species (or ecotypes [6]) occurs with an increase in thermal optimum but a general decrease in photosynthesis.

This is, for example, the case of *Atriplex sabulosa*. One can then wonder about the interest of this change. The opposite may be true for plants strictly adapted to warm conditions, such as *Tridestomia oblongifolia*. Figure 7 illustrates the case of *Atripex lentiformis*. [7] a perennial leafy plant, which occurs in California in both Death Valley and in cool, wet coastal habitats:

The assimilation of the desert ecotype (Figure 7A) and the coastal ecotype (Figure 7B) show almost the same response to temperature when grown under 23°C during the day and 18°C at night (in red in Figure 7).

Under the alternating 43°C day and 30°C night (blue in Figure 7), only the desert ecotype shows plasticity, maintaining high CO₂ assimilation under these new conditions. The activity of the coastal ecotype is low at all temperatures. Only the displacement of the thermal optimum remains of its acclimatization capabilities [8].

### 2.6. C3 plants versus C4 plants

![Image of Atriplex lentiformis ecotypes](https://example.com/image.png)
**C3 plants** were the first to appear and constitute about 85% of current plant species. They mainly colonize **cool and humid** environments (or seasons). Trees, for example, with rare exceptions, are C3 plants (Read *The path of carbon in photosynthesis*) (Figure 8).

**C4 plants**, of which there are traces only from the end of the Tertiary Era, constitute only 5% of the species. They tend to colonize **hot and dry** environments (or seasons) (See Restoring savannas and tropical herbaceous ecosystems). Maize and sugarcane are examples.

On average, the **thermal optimum of C4 plants** is located at **higher temperatures** than that of C3 plants.

However, **C3 plants** are the **most plastic**. In fact, their thermal optimum varies from around 7 to 35 °C, while that of C4 plants oscillates, with a few exceptions, between 30 and 40 °C. In addition, when the temperature is below 20 °C, the **photosynthesis** of C4 plants is **on average lower** than that of C3 plants.

### 3. CO₂ assimilation results from the interaction of processes whose response to temperature is different

The absorption of light at the **collecting antennae** (Figure 9) and the transfer of its energy to the PSII reaction centres are **not** temperature sensitive.

Are temperature sensitive:

- **The diffusion of CO₂** from the ambient air to the chloroplasts: its speed increases with temperature.

- **The fixation of CO₂** on Ribulose 1,5-bisphosphate (RuBP), a sugar whose skeleton is formed by 5 carbon atoms (Read Focus Deciphering the Benson-Bassham-Calvin cycle)

- **The transfer of electrons** from PSII to PSI.
Figure 9. Diagram of the interacting processes during photosynthetic CO₂ fixation (case of a C₃ plant). PSI and PSII: respectively photosystem I and II. They are included in the thylakoid membrane, which is made up of two lipid layers forming "sacs" in the chloroplast. The interior of the thylakoid is the lumen. Rubisco: enzyme that catalyzes the fixation of CO₂ on a sugar with 5 carbon atoms (Ribulose 1,5-bisphosphate: C₅). Benson-Calvin cycle: allows the regeneration of C₅, and at the same time gives the plant the necessary carbon. ATP is synthesized when protons from the lumen return to the stroma through an ATPase using inorganic phosphate, Pi. The lumen protons have two origins: (1) oxidation of water in the lumen by PSII which also provides electrons, e⁻ and (2) operation of a proton pump in the thylakoid that passes protons from the stroma into the lumen. [Source: Author’s diagram]

The regeneration of RuBP occurs via the operation of the Benson-Calvin cycle (This is the "biochemistry" of the process) which uses reducing power (in the form of NADPH) provided by electron transfer to function. The necessary ATP is synthesized when protons accumulated in the lumen pass into the stroma through an ATPase (Figure 9).

The formation of reducing power and the synthesis of ATP have a thermal sensitivity close to that of electron transfer.

4. What are the processes at work in setting the thermal optimum for CO₂ assimilation in C₃ and C₄ plants?

4.1. Photosystem activity and the resulting electron transfer are not involved

Measured in vitro on isolated thylakoids (see legend Figure 9), in the presence of artificial acceptors, electron transfer increases with temperature and shows a clear thermal optimum. It is located around 30°C and corresponds to that of CO₂ assimilation when the latter is saturating [9]. The activity of PSII has a thermal optimum identical to that of the electron transfer chain.

PSI activity is not inhibited at high temperatures (above 30°C, up to 45°C) where it remains stable or even increases: it is the activity of PSII that limits the activity of the electron chain.

Moreover, PSII is very sensitive to high temperatures which damage the protein complex that allows the oxidation of water (see Figure 9).

The thermal response of electron transfer is similar in C₃ and C₄ plants. However, there are organizational differences between these two types of plants (see The path of carbon in photosynthesis).

The supply of energy cannot therefore explain the differences in thermal optimum. It is the way in which the energy produced is used that makes the difference.

4.2. An answer? Comparison of the effect of atmospheric O₂ on CO₂ assimilation of C₃ and C₄ plants

In normal air [10], 21% O₂ (+ N₂) + 360 ppm CO₂: the thermal optimum is 27°C in Maize (C₄ plant), while it is only 22°C in Pea (C₃ plant) (Figure 10): the thermal optimum of the C₄ plant is higher than that of the C₃ plant (see also section 2.6).

In an oxygen-deficient atmosphere, 1% O₂ (+ N₂) + 360 ppm CO₂: the CO₂ uptake of Maize is not affected, while that of Pea is stimulated above about 17°C, with a shift in its thermal optimum to near that of Maize.
In C3 plants, atmospheric oxygen inhibits CO₂ uptake when the leaf temperature is sufficiently high, whereas it has no effect (or negligible effect) in C3 plants.

![Image: Pea (A; Pisum sativum) and Maize (B; Zea mays) leaves.]

Figure 10. Variation of CO₂ assimilation measured in leaves of Pea (A; Pisum sativum) and Maize (B; Zea mays) as a function of leaf temperature. The plants were grown in natural light at a temperature of 20 ± 2°C. [Source: Author’s diagram - royalty-free image / Pixabay]

Note that the variation in electron transfer estimated in vivo, by measuring chlorophyll fluorescence emission as a function of temperature, is very similar in 1% and 21% O₂ in Pea: the variation in thermal optimum is therefore not due to a change in photochemistry.

4.3. Rubisco properties explain the difference in response

Case of C3 plants

CO₂ and O₂ compete to occupy the active sites of Rubisco: This enzyme has a carboxylase function and an oxygenase function. CO₂ enters the Benson-Calvin cycle and the photosynthetic fixation of O₂ is at the origin of a metabolic pathway responsible for photorespiration (Figure 11; see also The path of carbon in photosynthesis).

CO₂ occupies a high number of active sites on the Rubisco when the O₂ content of the ambient air is low (1% for example) or that of CO₂ is high.

O₂ is mainly fixed if its content increases or if that of CO₂ decreases (the latter then releases active sites which are then occupied by O₂).

In normal air, there are two reasons why O₂ fixation increases (and consequently CO₂ fixation decreases) when the temperature increases [11].

The affinity of Rubisco for CO₂ decreases more than that for O₂; a factor that favours the assimilation of O₂.
The water solubility coefficient of \( \text{CO}_2 \) decreases more than that of \( \text{O}_2 \), leading to a more rapid decrease in the amount of \( \text{CO}_2 \) than \( \text{O}_2 \) in the chloroplast; this is a factor that favours \( \text{O}_2 \) fixation.

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The water solubility coefficient of \( \text{CO}_2 \) decreases more than that of \( \text{O}_2 \), leading to a more rapid decrease in the amount of \( \text{CO}_2 \) than \( \text{O}_2 \) in the chloroplast; this is a factor that favours \( \text{O}_2 \) fixation.

In an \textit{O}_2-poor atmosphere (Figure 10), \textit{competition between \textit{O}_2 and \textit{CO}_2} is very reduced. Energy is then used mainly for CO
In normal air, the effect of O₂ on photosynthetic CO₂ fixation (Figure 11) is very low (or even nil) when the temperature is low: competition on the carboxylation sites is in favour of CO₂.

On the other hand, when the temperature increases, the competition on these sites favours the fixation of O₂ which then consumes an increasing part of the energy produced by the activity of the photosystems. This energy is therefore no longer available for CO₂ fixation, which reaches its maximum value around 22°C.

Case of C4 plants.

CO₂ is concentrated at the Rubisco by a mechanism that is insensitive to oxygen. Its content can reach 800 to 2000 ppm depending on the plant in C4: that is to say contents from 2 to 5 times higher than its current atmospheric content.

Under these conditions, photosynthetic O₂ fixation is weak or even non-existent because the active sites of the Rubisco are all occupied by CO₂. The energy supplied by the activity of the photosystems is therefore used only in the fixation of CO₂ when the leaf temperature increases, explaining the higher thermal optimum in this type of plant.

C4 plants evolved from C3 plants during the global decrease in atmospheric CO₂ content at the end of the Tertiary Era [12].

This decrease would then have "released" the oxygenase function of the Rubisco of C3 plants, resulting in a loss of fixed carbon via photorespiration.

The establishment of a CO₂ concentration mechanism is an advantage because it prevents this carbon loss. We currently find species that are "intermediates" between C3 and C4.

5. The thermal optimum of C3 photosynthesis is modulated by certain environmental parameters

5.1. The CO₂ content in the atmosphere

The thermal optimum increases with increasing ambient CO₂ content. In the case shown in Figure 12, it increases from about 10°C when the content is 100 ppm to more than 30°C when it is 800 ppm.
This effect is explained by the competition between CO\(_2\) and O\(_2\) for the occupation of the active sites of the Rubisco: at 800 ppm CO\(_2\) the active sites are occupied mainly by CO\(_2\); at 100 ppm CO\(_2\) the occupation of these sites by atmospheric O\(_2\) is in majority.

In a world with steadily increasing atmospheric CO\(_2\) (Figure 13), the thermal optimum of C3 plants is expected to increase. This does not mean, however, that plant production will then be higher (see note 3 section 1): episodes of high heat will, like droughts, certainly be more frequent.

**5.2. Lack of water**

The photosynthetic apparatus is resistant to drought. It retains all its capacity to absorb CO\(_2\) on the Rubisco, and to produce
CO₂ uptake decreases in this range of water loss, because the stomata close (see Focus Leaf transpiration and heat protection). This closure slows down the entry of CO₂ into the leaf and consequently leads to a decrease of the CO₂ content in the mesophyll.

However, the O₂ content in the chloroplasts remains high. Indeed, its content in the atmosphere (21% or 210,000 ppm) is, compared to that of CO₂ (@ 400 ppm), very high and in any case sufficient for a very substantial quantity to pass through the epidermis even when the stomata are closed.

The competition between CO₂ and O₂ for the occupation of the active sites of the Rubisco is thus in favour of O₂.

Figure 14. A, Variations in CO₂ assimilation as a function of leaf temperature. Leaves with different amounts of water loss found in air with an ambient CO₂ content of 400 ppm. B, The electron transfer rate estimated on the same leaves by measuring the chlorophyll fluorescence emission. [Source: Author’s diagram, after Cornic et al. ref. 14]

Therefore, the thermal optimum for photosynthesis must lower in C3 plants that dry out.

This is shown in Figure 14A, in which the thermal optimum drops from about 23°C, in a Pea leaf at maximum turgor, to 17°C when it has lost 20% of its water.

Electron transfer in the thylakoid membrane is not affected by water loss in the range shown (Figure 14B). When water loss is 20%, the energy produced by photosystem activity is primarily used to bind atmospheric oxygen to RuBP [14], resulting in increased photorespiration.

6. Why, from its thermal optimum, CO₂ assimilation decreases as temperature decreases or increases?

6.1. When the temperature lowers

Several reasons probably all contribute, to varying degrees, to this decrease:

The rate of RuBP turnover decreases: there is a slowdown in the activity of some enzymes controlling this turnover, notably
Sequestration of phosphorylated compounds in chloroplasts. The triose phosphate is no longer (or less) exported when sucrose synthesis is inhibited. The inorganic phosphate in the chloroplast is no longer renewed leading to a decrease in ATP synthesis.

Inhibition of the electron transfer chain (see section 4.1), resulting in reduced energy production (reducing power and ATP).

In C4 plants it is the activity of the Rubisco that appears to be preponderant, although the cold sensitivity of enzymes involved in CO₂ accumulation at the Rubisco is well known.

6.2. As the temperature increases

In C₃ plants the increase in photorespiration decreases the fraction of electrons produced by PSII and used to assimilate CO₂. However, other factors are at play since CO₂ assimilation measured (1) in an atmosphere with little or no photorespiration (ambient O₂ content of 1%), and (2) measured in a normal atmosphere in a C₄ plant decreases in both cases (Figure 10).

Several reasons can be given:

The slowing down of PSII activity leading to that of the electron transfer chain from PSII to PSI.

To perform its role Rubisco must be activated by an enzyme called Rubisco activase, the activity of which decreases when the temperature is higher than about 33°C (incidentally, high-temperature resistant activases appear in some plants subjected to periods of high heat [15]). However, since activase must itself be activated by an electron transfer-dependent process, it cannot be ruled out that the latter is also involved in limiting [15].

The "catalytic misfiring" of Rubisco increases with temperature and increasing amounts of an inhibitor of the enzyme (Xylulose-1,4-bisphosphate), which is structurally close to RuBP (see Figures 9 and 11), are synthesized.

In C₄ plants (case of Maize) the activation and activity of enzymes that participate in the CO₂ concentration system at the Rubisco are not very sensitive to high temperatures. The same reasons as above may explain the decrease in CO₂ assimilation when the temperature increases beyond that of the thermal optimum.

7. Hardening after plant exposure to cool (≤ about 10°C) and high (≥ about 37°C) temperatures

Maintaining plants at cool or high temperatures causes, along with the changes in photosynthesis described above, increase in their resistance to otherwise lethal temperatures (frost and high temperature). This is hardening.

In this process, temperature and light interact and the metabolic changes induced are sometimes very rapid (from minutes to hours).

Thus, cold hardening can be achieved at ordinary temperature by modulating the length of the light period or its spectral composition in the red [16]. However, cold is still required to achieve full hardening. Also the lack of light in the cold prevents hardening to varying degrees.

At elevated temperatures : the transmitted signals activate the synthesis of chaperone proteins (HSPs: Heat Schock Proteins) that repair denaturing proteins, also prevent their coagulation or even help mark them for degradation.

At cool temperatures : the synthesis of chaperone proteins is also activated. It is accompanied by (i) the synthesis of "antifreeze" proteins that interfere with ice crystal formation and (ii) an increase in sugar synthesis tending to increase osmotic pressure in the cells.

Note that the signaling pathways and their interactions inducing the genome response are only partially known. The references given in "Learn More" and an attached Focus allow for further exploration of this evolving point.

8. Effects of temperature on photosynthesis: summary diagram

The summary diagram (Figure 15) classifies the effects of temperature on photosynthesis according to the speed of temperature change and the extent of its variation. Note that hardening allows leaf maintenance in perennial leaf plants and therefore minimizes energy loss under extreme temperature conditions.
The rapidity of current climate change makes it necessary to delve deeper into the responses of plants to their environment: the hope is to be able to maintain sufficient primary production to keep the biosphere functioning.

9. Messages to remember

The uptake of CO$_2$ by a leaf has a **thermal optimum** close to the average temperature of its growth environment.

This thermal optimum can **change rapidly** when the conditions of the environment are durably modified: this is a process **acclimatization**.

This thermal optimum is on average **less in C3 plants** than in C4 plants: this is mainly due to photosynthetic fixation of atmospheric O$_2$ via Rubisco activity in C3 plants.

This optimum depends on the **CO$_2$ content of the ambient air** in C3 plants: at high content it becomes identical to that in C4 plants.

This optimum depends on **the hydration state of the leaf**.

Subjected to cool or hot temperatures plants bring into play **processes hardening** to otherwise lethal temperatures. These processes involve protein syntheses and changes in the fluidity of chloroplast and cell membranes.

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**Notes and references**

**Cover image.** Sunset over the Sonora Arizona desert. [Source: royalty free / Pixabay]


[3] For example, when growing plants are subjected to drought, the amount of carbon they assimilate decreases initially because leaf growth is inhibited. Boyer JS (1970) Plant Physiol. 46, 233-235

[4] The values of thermal optima given here, are from measurements made in "normal air", containing **21% O$_2$ and about 400**
When this is not the case the O$_2$ and CO$_2$ contents are shown. The CO$_2$ uptake in air containing 21% O$_2$ is saturated from about 1200 ppm CO$_2$ when light is close to saturation. The evaporative power of the air is also regulated in most cases during the measurements. It is estimated by the saturation deficit of the partial pressure of water vapor in the ambient air around the leaves.

Plants from the same individual by vegetative reproduction. They are genetically identical.

Ecotype: Plants of the same species from different environments, which, grown from seed to flower under identical conditions show different physiological characteristics.

It fetches water from as far as the water table, hence its name of phreatophyte plant.


See note #4, section 2.2


