Article

Do Bio-Ethanol and Synthetic Ethanol Produced from Air-Captured CO₂ Have the Same Degree of “Greenness” and Relevance to “Fossil C”?

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Abstract: This paper discusses the epochal change in the reputation of carbon dioxide, which is now considered as a raw material alternative to fossil C for the synthesis of chemicals, materials and fuels, as opposed to a waste material that must be confined underground. In particular, its use as renewable C is compared to biomass. In this paper, a specific point is discussed: is ethanol (or any fuel) produced via the catalytic conversion of atmospheric CO₂ different from the relevant biomass-sourced product(s)? The answer to this question is very important because it ultimately determines whether or not fuels derived from atmospheric CO₂ (either e-fuels or solar fuels) have the right to be subsidized in the same way that biofuels are. Conclusions are drawn demonstrating that ethanol derived from atmospheric CO₂ deserves the same benefits as bio-ethanol, with the additional advantage that its synthesis can be less pollutant than its production via the fermentation of sugars. The same concept can be applied to any fuel derived from atmospheric CO₂.

Keywords: carbon dioxide capture and utilization; bio-ethanol; e-fuels; renewable fuels of non-bio-origin

1. Introduction

Nowadays, we notice a revolution concerning the reputation of the tiny CO₂ molecule: from “waste” to “resource” [1]. I am one of those people who, for decades, has disseminated the concept that “CO₂ is a resource” [2,3]. As a matter of fact, CO₂ is at the origin of life, it was the original source of carbon for constructing organisms and life on our planet and still is the most abundant form of easily accessible carbon that we can dispose of (Table 1).

Table 1. Distribution of carbon among different environments on Earth (as GtC).

| Carbon reservoir       | Total (GtC) |
|------------------------|-------------|
| Atmospheric CO₂        | 750         |
| Biosphere [4]          | 550–560     |
| Plants                 | 450–460     |
| Animals                | 2 (Humans 0.06) |
| Bacteria               | 70          |
| Archaea                | 7           |
| Fungi                  | 12          |
| Protists               | 4           |
| Viruses                | 0.2         |
| Carbonate rocks a (sediments) [5] | 1.8 × 10⁹ |
| Fossil carbon [1]      |             |
| Coal                   | 607         |
| Oil                    | 167         |
| Natural Gas            | 167         |
| CO₂ deposits (degas rate 0.1 GtC/y) | 30,000     |
| Ocean deep floor [6]   | 37,000      |

a During the Phanerozoic aeon (system of rocks deposited during the Phanerozoic era—the Paleozoic, Mesozoic and Cenozoic eras—from 541 My to present), 2100 × 10¹⁵ t of carbonate rocks were deposited with a mass cycle of 8.6 × 10¹⁴ tC/My and a decay constant of 0.0025 My⁻¹; 45,500 GtC above surface equal to 10–15% of all rocks.
However, the atmosphere is the most abundant source of immediately available CO$_2$, even if it is quite diluted (415 ppm). Subsurface deposits of CO$_2$, less easily reachable but more concentrated (up to 99% pure), are usually close to volcanic areas and spread all over the planet. The amount of carbonate rocks is estimated to be ca. $10^9$ GtC and varies over time because of the cycle of carbon (see Table 1 and Figure 1). The natural carbon cycle turns carbon from the atmosphere into plants, microorganisms or animals, and then returns to the atmosphere, soil or sub-surface (fossil C). Figure 1 shows the amount of carbon cycled yearly (ca. 200 GtC/y).

![Human respiration: ca 1 kg/pax d, or ca 2Gt/y CO$_2$](image)

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*a During the Phanerozoic aeon (system of rocks deposited during the Phanerozoic era—the Paleozoic, Mesozoic and Cenozoic eras—from 541 My to present), 2100 $\times 10^{15}$ t of carbonate rocks were deposited with a mass cycle of $8.6 \times 10^{14}$ tCaCO$_3$/My and a decay constant of 0.0025 My$^{-1}$; 43,500 GtC above surface equal to 10–15% of all rocks.

**Figure 1.** The natural C cycle.

### 2. The Need to Substitute Fossil C

The continuous extraction of fossil C and its combustion is causing an accumulation of CO$_2$ in the atmosphere as the natural C cycle cannot buffer the excess 10 GtC of anthropogenic origin, despite representing ca. 5% of the amount cycled yearly (Figure 1). Such an accumulation is considered to cause climate change. In my opinion, the increase in atmospheric CO$_2$, which parallels the amount of burned C-based fuels by our society, is not the protagonist in the “climate change drama” played these days across the world in everyday life, but the third actor, with the inefficient use of chemical energy from fossil-fuels as the protagonist and the increase in atmospheric water vapor as the deuteragonist [1]. As a matter of fact, over 65% of the chemical energy of burned fossil C, due to the low efficiency of the conversion processes into electric, mechanical or thermal energy [7], is directly discharged into the atmosphere in the form of heat even at temperatures as high as 800–1000 °C, resulting in the direct heating of the atmosphere. This is causing an increase in atmospheric water vapor, which is a stronger greenhouse gas than CO$_2$ [8]. The combination of all of these causes the adverse environmental events that we are currently observing, and may seriously affect our planet in the future. Such a negative impact could be mitigated by increasing the efficiency of energy production and a wiser use of energy. For example, IGCC technology has increased the efficiency of the chemical-to-electric energy conversion from an average of 32–35% to 55+%, causing a reasonable decrease of ca. 60%
in the use of fossil C for electricity production, and thus CO₂ emissions. Although very promising, such technology is rarely applied, despite its low costs [7].

Overall, individual and collective responsibilities play a role in improving the use of energy.

3. Alternatives to Fossil C

The alternative to the aforementioned efficiency technologies is a fossil C substitute that uses primary sources from non-fossil-C energy. The need to limit the use of fossil C as source of energy is now very urgent. After the almost unattended COP2016 Paris Agreement, hopefully, the urgency of an inversion of trend has been re-assessed by a large number of governments at the COP2021 in Glasgow. Moving away from fossil C also meets the goals of the 2030 Agenda for Sustainable Development that includes 17 Sustainable Development Goals (SDGs), first adopted by the UN General Assembly on September 2015. [8] As a matter of fact, we need to act in the direction of returning to nature, which does not produce waste and recycles carbon through the conversion of CO₂. Although CO₂ has been industrially used as raw material for over 150 years (aspirin and urea syntheses, the latter consuming more than 160 Mt CO₂/y today) and CO₂ chemistry has been intensively investigated during the last fifty years [1,9–11], attention is mainly devoted to the use of CO₂ as a building block of chemicals, because its conversion into fuels requires energy and hydrogen, both of which are still mainly produced today from fossil C (>81% and 95%, respectively) [7,12]. Today, the large volume conversion of CO₂ makes sense as it is possible without producing more CO₂ than is used. As a matter of fact, the large-scale conversion of perennial energies (solar, wind, hydro, geothermal) into more easily usable forms of energy (electric and thermal) at affordable costs is making possible an energetic transition from fossil C to carbon-free energies for powering our lives [13].

In such an epochal transition, the utilization of CO₂ as a building block of chemicals and/or source of carbon for energy products will play a key role. Carbon dioxide capture and utilization (CCU) is an active part of the circular economy strategy; at the same level, carbon dioxide capture and Storage (CCS) is the logical end-of-pipe technology of the linear economy. However, CCS and CCU are divergent technologies in the sense that, while the former pushes the extraction of fossil C, the latter avoids the extraction of fossil C. Therefore, CCS is somehow tied up with fossil-C-fed power plants, a moderately concentrated source of CO₂ (5–14% v/v of flue gases), while CCU is attuned to direct air capture (DAC). CCS, which has received significant funding over the last thirty years, had to already be a large-scale technology for CO₂ mitigation, but this was not achieved as several scientific gaps exist in its deployment (energetic costs for exploitation, permanence in disposal sites and environmental impact). However, in a world mainly powered by perennial energy sources, its role will be less and less important. As the energetic transition would not guarantee a future for energy produced by power plants, and because capturing CO₂ from plants fired with fossil C would perpetuate the use of the latter, CCU correctly orientates towards DAC to make CO₂ available for conversion, a technology that requires an energy and economic cost reduction (see below). On the other hand, CO₂ is renewable C; biomass is made from it.

Moving to circularity represents a dramatic but necessary change in both the economy and our lives. The use of renewable carbon merges CCU and bioeconomy, with hybrid (chemo-enzymatic) catalytic systems boosting the potential of carbon recycling.

Three strategies can be envisaged for implementing such a major change in CCU: i. Large-scale non-fossil-H₂ production and its use in chemo-catalytic CO₂ reduction in fuels (e-fuels). ii. Coprocessing of CO₂ and water under solar irradiation to afford energy products (solar fuels). iii. Integration of biotechnology and catalysis. In all cases, CO₂ and water are at the core of the production of energy products; an economy based on CO₂ and water [14]!

Strategy (i) is immediately deployed by using perennial energies (SWHG) to produce cheap and abundant electrons for the electrochemical reduction of non-drinkable water (recycled, salty) to afford hydrogen, which can then be used for the chemo-catalytic reduc-
tion of CO₂ to gaseous (CH₄) or liquid fuels (CH₃OH, hydrocarbons, superior alcohols), using known technologies that operate in the Syngas-FT frame. This approach has some technological barriers that can be identified by the availability of: a. low-cost electrons, b. large-scale and long-life electrolyzers, and c. cheap and stable electrodes. It can be boosted by new technologies such as: d. the potential increase in the electrolysis temperature, e. working under pressure for immediate H₂ distribution to users, and f. the exploitation of solid-state electrolysis. The use of renewable H₂ for CO₂ reduction into energy products may have a most immediate implementation, supposing that cheap sources of CO₂ and cheap “electrons” (reducing power) are found to meet the cost target [15]. Fuels produced in this way, or fuels obtained from CO₂ using non-fossil electricity and H₂, are labeled “E-fuels”, the technology has a high TRL (7–9, depending on the targeted fuel: CH₄ or CH₃OH), and demo-plants are available in several countries. The bottleneck of this technology is its worldwide dissemination in the short term.

The second strategy (ii) is based on an advanced approach of co-processing CO₂ and water to produce fuels without intermediate H₂ production [16]. Such an approach has the great advantage of cutting the costs of producing, storing, transporting, and using H₂, resulting in a much lower CAPEX, and even a lower OPEX, in terms of the safety measures to be implemented. A more probable approach is to use the Sun to power direct photochemical, photoelectrochemical, photobiological, and photobioelectrochemical routes of CO₂ reduction and water oxidation [15], producing fuels that can be called solar fuels for simplicity [16]. The TRL of such technology is very low (2–3), but of great interest for the future. The third strategy (iii) is also at its early stage of development and combines electricity, enzymes, microorganisms and metal systems as catalysts for CO₂ conversion into energy products and chemicals [17,18]. However, fossil fuels can potentially be substituted with biofuels, E-fuels, or solar fuels, each having their own peculiar character and potential to reduce carbon dioxide emissions. Because “CO₂ avoidance” receives subscriptions from governments, unfortunately, such a large transition is increasing the discussions of how the new regulations and agreements can match their old counterparts or how older statements can be rephrased. Significantly, new non-fossil fuels enter into a confrontation with biosourced fuels or fuels derived from biomass, a class of products known for a long time and already on the market. It is worth recalling that biodiesel (produced from lipids, vegetal or animal sources) represents ca. 10% of the diesel used in several countries, while ethanol (produced via sugars fermentation, a very old practice) is added to gasoline in a 5–7% v/v ratio. All around the world, there are actions in favor of CCU: both calls for large-scale projects that support innovation actions at a high TRL (7–9) (Carbon Prize, USA–Canada; Green Deal and Innovation in the EU) and calls that address more basic research at a low TRL (EU Calls in Horizon) are funding the deployment of and search for innovative solutions that may make the knowledge we have for mimicking nature useful for building and deploying a man-made C cycle. Nowadays, at the EU level, there is a broad confrontation regarding the introduction of such new fuels into older schemes (transition from RED II to RED III), and understanding if they can be assimilated with biofuels and entitled to have similar benefits. This is not an easy task and requires agreement among different actors and lobbies (industrialists, farmers, fuel producers, policy men, etc.), because this task implements new rules and standards, avoiding the double counting of benefits as well as attributing benefits to unentitled goods.

4. Biofuels vs. Renewable Fuels of Non-Bio-Origin

Here enters the question that is the title of this paper: “Ethanol produced from biomass and ethanol produced catalytically from air-captured CO₂: do they have a different degree of “Greenness” or “Fossil C”? The answer to the question is important. First of all, it is necessary to clarify if there exists any difference between fossil CO₂ and bio CO₂, where fossil CO₂ is formed in the combustion of fossil C, and bio CO₂ is formed in the conversion (combustion/fermentation) of biomass. The use of biomass is seen as antithetic to the use of fossil C: the former avoids the extraction of fossil C, and thus avoids the production of
new CO\textsubscript{2} that will accumulate in the atmosphere. Using biomass (and its derived biofuels), is considered an action that does not overload the atmosphere with CO\textsubscript{2} of fossil-fuel origin. Burning biomass or bio-fuels is considered a zero-emission option for energy production and use, even if it is not exactly so. In fact, accurate life cycle assessment (LCA) studies show that the use of biofuels in the current production–utilization–accounting scheme is the transfer of carbon from surface subsoil to the atmosphere, similar to the use of fossil C, even if it is much less intensive \cite{19}. As a matter of fact, biomass is generated from atmospheric CO\textsubscript{2}, and when burned, it is believed to return to the atmosphere the same amount of CO\textsubscript{2}, as if the sequence was part of the natural C cycle. As a matter of fact, the cycle is not really closed as it occurs in nature. In fact, one should also consider, in addition to the carbon dioxide generated by the combustion of biomass, the amount of CO\textsubscript{2} emitted by the various human activities that accompany the production and work-up of biomass (soil cultivation, planting, agricultural practices, the use of pesticides and herbicides, harvesting, converting the products into the final goods, producing biofuels, etc.), and even the soil carbon pauperization caused by agricultural practices. The latter aspect is often not considered, causing mistakes in assessment studies.

Nonetheless, biofuels are subsidized by governments (through tax breaks, grants, loans, and loan guarantees) for their environmentally friendly qualities and for the fact that they save natural fossil resources for future generations.

However, the question of whether converting anthropogenic CO\textsubscript{2} into energy products is an action that mimics nature can be raised; therefore, such fuels can be considered as biofuels.

Let us start with clarifying what “anthropogenic CO\textsubscript{2}” is:

Anthropogenic CO\textsubscript{2}, aside from that emitted by humans during respiration (ca. 1 kg/d pax), is the amount of CO\textsubscript{2} produced by burning biomass, biofuels or fossil C (such as coal, oil, gas) to produce energy or goods used by society, or even the part that accumulates in the atmosphere because of deforestation or is generated in forest fires. “Anthropogenic CO\textsubscript{2}”, therefore, encompasses all of the non-natural CO\textsubscript{2} emitted into the atmosphere that overbalances the natural C cycle.

Let us now concentrate on two particular classes of fuels: (i) fuels derived from biomass (made from atmospheric CO\textsubscript{2}), and (ii) fuels made from CO\textsubscript{2} emitted by industrial and chemical industries, cement manufacturing, stainless-steel manufacturing, or power plants. From these two cases, a very interesting and almost “philosophical” debate has been occurring for a long time. The difference between these two classes is not clear and requires a detailed and circumstantiated set of rules.

On a general basis, both classes of fuels make fuels from CO\textsubscript{2} and avoid fossil C, and their use can be considered beneficial. However, are they equal with respect to subsidies or subventions?

Industrial CO\textsubscript{2} has its origin in fossil C that is fed to industries and fuels derived from it are, therefore, considered of “fossil” origin. Conversely, CO\textsubscript{2} produced by burning biomass and its derivatives is said to be “biogenic”, and its emission does not cause, in principle, an increase in the atmospheric concentration of CO\textsubscript{2} as the level of CO\textsubscript{2} released should be fixed in the biomass. As we have discussed above, this is not completely true, and we must also consider that there is a gap of time between the combustion of bio-carbon and its fixation into biomass; its combustion is some 1000–10,000 times faster than biomass growth \cite{1}.

Moreover, the use of biomass-derived fuels produces less-intense CO\textsubscript{2} emissions than the combustion of fossil C. However, industrial CO\textsubscript{2} (fossil C-derived) has different characteristics compared to biogenic CO\textsubscript{2}, the latter being produced by actions of microorganisms in biomass or in the process of burning bio C.

Let us go further and conduct a more detailed analysis of the relationship between the origin of CO\textsubscript{2} and its conversion into fuels.

At the end of the 1990s, at the EU level, during the very early days of the question of CO\textsubscript{2} recovery and conversion (the “Recovery and Utilization of Carbon Dioxide-RUCADI”
project was the first EU-funded project for CCUS, founded in 1998 and co-ordinated by M. Aresta), and at the beginning of the discussion about the classification of fuels and goods derived from CO$_2$, a scenario was depicted for the consideration of scientists, lawyers and policy makers.

**Scenario:** Let us suppose that we have a source of CO$_2$ from fossil fuels (power plants) and the stream follows two separate routes. **Route 1:** CO$_2$ from the power plant is emitted into the atmosphere and then a pond of algae, located somewhere, fixes the aerobic CO$_2$. **Route 2:** the same CO$_2$ flow is, instead, the only C source, and is fed directly into a pond of algae located next to the power plant. Let us now extract lipids from both batches of algae and convert them into diesel by using the same technologies, obtaining diesel1 and diesel2.

**Question:** Are the diesel1 and diesel2 produced in the two ponds different? Are they distinguishable? Could both be labeled “biodiesel”? Do the two fuels have the same rights to be subsidized?

This may appear a dull question: two different opinions commonly arise: (i) “Biomass is biomass, however it is grown”, so the two diesels are equal. (ii) If we feed fossil CO$_2$, we produce fossil fuels, so the two diesels are different.

Then, the question arises: if they are different, how can we distinguish them? This question is easy to answer: the $^{14}$C analysis will clearly show which is which: fossil CO$_2$ and atmospheric CO$_2$ have a different $^{14}$C level. Diesel1 and diesel2 are, therefore, different.

The most recent conceptualization, based on a broad consultation and averaging different points of view, arrives at the same conclusions but moves from a quite different position: using CO$_2$ captured from a fossil C-fed power plant (or industrial plant) supports the continued extensive use of fossil C, and this cannot be subsidized. CO$_2$ captured from industrial and power-generation emissions is totally different from bio CO$_2$. Only bio CO$_2$ can be subsidized, and the products derived from it can be considered of bio-origin and called bio-fuels.

Now we present a different case. CO$_2$ produced by burning fossil C is not captured but emitted into the atmosphere. Then, ethanol is made, or any other fuel, following two routes. **Route A:** Corn is grown, and then ethanol is produced by fermentation. **Route B:** CO$_2$ is taken from the atmosphere, and ethanol is produced catalytically in a chemical plant. Are ethanolA and ethanolB different? The technique mentioned above ($^{14}$C isotope abundance) will not distinguish the two: in both cases, atmospheric CO$_2$ is used. One can say that the two butches of ethanol will still have different analytical characters: (A) will most likely contain proteins and other biocomponents that will reveal its bio-origin; (B) will not have such properties. However, such differences are due to the different production technologies and not the different origins of CO$_2$. Moreover, ethanol B is cleaner than ethanol A.

Then, is the corn-sourced ethanol different from synthetic ethanol when atmospheric CO$_2$ is used for the production of both? Is this a new sophism? If one looks at the source of CO$_2$ (in both cases, this is the atmosphere), they are not different. It is not the fact that one is produced using a vegetal plant and the other using a chemical plant that can differentiate between them. The common source of carbon (atmospheric C) makes them equal: both have the same link to fossil C, and both recycle atmospheric carbon. Intriguingly, their chemical synthesis is not seasonal, and the rate of ethanol production and the volume produced per unit area of the occupied land would be higher than for the bio-process. The chemical synthesis would be more intensive and present other advantages over the biological route, as arable soil would not be required, C soil pauperization would not occur and much less water would be required per unit volume of ethanol. Developing synthetic procedures based on the utilization of atmospheric CO$_2$ (man-made C cycle) would be a win–win situation that may side with the natural C cycle and alleviate the impact of the use of fossil C by progressively reducing its demand.

The tendency is to capture CO$_2$ from the atmosphere as the only source that guarantees infinitely large volumes of immediately available CO$_2$, as stated above: fuels derived from such CO$_2$ should not be distinguished from bio-fuels; they should be considered
as “renewable fuels of non-biological origin, RF-NBO” and subsidized in the same way that biofuels are.

In a world in which primary energy will mainly be provided by perennial sources (SWHG), synthetic fuels obtained from atmospheric CO$_2$ may be more environmentally beneficial than biofuels. Using recyclable heterogeneous catalysts with airborn CO$_2$ and non-drinkable water as hydrogen sources will produce less pollution than growing biomass (this requires high-quality soil, nutrients for soil with the emission of N compounds and other pollutants, in addition to agrochemicals, such as herbicides and pesticides, and large volumes of water) and a conversion into bio-fuels by using biotechnologies based on fermentation by microorganisms.

Now, let us tell this story with numbers. To make 1 L of ethanol (789 g, 17 mol) from corn, some 10–17 L [20] of clean water is necessary (equal to 10,000–17,000 g; or 554–940 mol) solely for farming and not considering the water needed for fermentation and work-up. Each mol of ethanol is will, thus, requires 32.6–55.4 mol of water. To make ethanol from air-captured CO$_2$, according to Equation (1), assuming an efficiency in water electrolysis to H$_2$ of 80% and a chemical yield of 60%, one needs only 12.5 moles of water per mol of ethanol produced, saving from 20 to 43 mol of water per mol ethanol, or 7.8–12.9 t$_{H2O}$/t$_{ethanol}$: a huge amount considering that today the consumption of bioethanol is close to 50 Gt/y. Moreover, the watering of soil requires soft water, while hydrogen can be produced from salty water, a large difference that will save on large volumes of high-quality water:

$$2 \text{CO}_2 + 6 \text{H}_2 \rightarrow \text{CH}_3\text{CH}_2\text{OH} + 3 \text{H}_2\text{O}$$ (1)

Considering the energy necessary for producing bio-ethanol and synthetic ethanol, the latter made from DAC-CO$_2$ and H$_2$ from PV–water electrolysis, one finds that the non-optimized energy for bioethanol production is 1.316 MJ/mol [21], not accounting for soil C pauperization, while synthetic ethanol requires 2.50 MJ/mol [22,23]. The former value might be optimized to 0.92 MJ/mol by using the most well-known technologies [21], not accounting for soil restoration, which remains an ongoing debate, while the latter can be more than halved to 1.00 MJ/mol by: i. increasing PV efficiency (solar to electrons) from 20% to a perspective of 40% by 2040 [1]; ii. reducing the cost of PV electrons by using cheaper materials for PV (organic materials instead of Si-based materials); iii. improving the electrolyzer size, life and cost; iv. implementing heat recycling; and v. improving the DAC technology, that today is very expensive, both economically (ca. 200 EUR/t$_{CO2}$) and energetically (0.06–0.16 MJ/mol$_{CO2}$) [24]. Therefore, one can say that, by 2040, the energy necessary for making bio- or synthetic ethanol will be almost the same, with the great advantage that synthetic ethanol can be produced everywhere without climate constraints and will require much less water of a lower quality, a great positive. Another advantage of the synthetic route is that will require less space for producing the same amount of ethanol with respect to the bio-route, and lower quality soil. A drawback of bio-ethanol is that it will produce large volumes of waste biomass that must be used (production of thermal energy, or further working up to produce sugars to increase the amount of ethanol) for not aggravating the energy balance. In the synthetic route, heat recovery will improve the overall energy balance, and developing selective catalysts will avoid the production of side-products and loss of carbon. The industry will require a higher CAPEX with respect to growing corn, that can be significantly reduced by retrofitting and revamping existing plants.

Notably, both CO$_2$ and water can be recovered from the atmosphere with the benefit of reducing the concentration of two greenhouse gases, and this will make the synthesis of ethanol ubiquitous and not linked to a specification of soil and climate.

As neither route (bio or synthetic) prevails in a net manner and both have pros and cons (Table 2), considering that DAC-CO$_2$ cuts out debates on the origin of CO$_2$ and that synthetic ethanol is commonly labeled as a renewable fuel of non-bio-origin (RF-NBO), both routes can be equally useful for producing a liquid fuel that may have widespread applications in the land transport sector, or it could be used as raw material in the fuel
industry for producing longer-chain hydrocarbons \cite{25}, possibly reaching the state of fuel usable in the aviation sector.

| Category                          | Bio-Ethanol | Synthetic |
|----------------------------------|-------------|-----------|
| Soil fertility                   | High        | Low \textsuperscript{1} |
| Soil C-use                       | High \textsuperscript{2} | NO       |
| Land extension                   | Low         | High      |
| CAPEX                            | High        | Low       |
| OPEX                             | High        | Low       |
| Energy consumption               | Neutral \textsuperscript{3} | Neutral \textsuperscript{3} |
| GHG emission                     | Agrochemicals \textsuperscript{4} | Neutral      |
| Water consumption                | High        | Low       |
| Geographical location            | Climate dependence | Any place |
| Production cost                  | Neutral     | Neutral   |
| Waste production, recovery and utilization | Waste biomass utilization | Heat recovery |

\textsuperscript{1} Marginal areas can be used; \textsuperscript{2} Agricultural practices cause soil C pauperization; \textsuperscript{3} In the future, both practices may have levelled energy requirements; \textsuperscript{4} The production/use of agrochemical cause large emission of GHGs.

5. Conclusions

A revolution is in front of us, based on an industrial and energetic transition that will progressively reduce the use of fossil C most likely to 30\% of the current use by 2040–2050 (in my view, fossil C-based fuels will still be used till the end of this century, even if at a much lower rate), and technologies closer to nature will be implemented, improving the quality of our lives and preserving our planet. Table 3 lists the potential substitutes for fossil C and its actual production/or TRL.

| Fuel               | World Volume Consumed/y or TRL | Expected Consumption by 2030          | Ref. |
|--------------------|---------------------------------|---------------------------------------|------|
| Bio-ethanol        | 132 BL in 2020                  | 137 BL                                | \cite{26} |
| Bio-diesel         | 48 BL in 2020                   | 50 BL                                 | \cite{27} |
| Bio-jet fuel       | 15 ML in 2020                   | 500 ML                                | \cite{28} |
| Bio-Gas            | 31 Mtoe in 2018                 | 41 Mtoe; 78 Mtoe                      | \cite{29} |
| Bio-methane        | 1 Mtoe in 2018                  | 46 Mtoe; 114 Mtoe                     | \cite{29} |
| e-H\textsubscript{2} | 320 kt                         | 73–158 Mt                             | \cite{30} |
| e-CH\textsubscript{4} | 7–9                            | Depends on e-H\textsubscript{2} cost | \cite{31} |
| e-CH\textsubscript{3}OH | \textit{600 t}              | Depends on e-H\textsubscript{2} cost | \cite{31} |
| S-fuels            | TRL 2–3                         | Research is needed                    | \cite{19} |

(BL = billion liters; ML = million liters; Mtoe: million ton oil equivalent; Mt = million tons; TRL: Technological readiness level, 1–9 scale).

As biomass cannot satisfy the energy needs of our society (this has been the case since the start of the Industrial Revolution), developing man-made technologies that can complement the bio-production of chemicals and energy products is of fundamental importance. The use of atmospheric C in both cases will greatly reduce the environmental impact of anthropic activities and mitigate climate change.
A key point that the regulations have to make clear is that subventions and benefits cannot be given twice to the same or to different recipients for the same action: either the carbon capture is subsidized or captured CO₂ is converted into marketable products.

**Funding:** This research received no external funding.

**Institutional Review Board Statement:** Not applicable.

**Informed Consent Statement:** Not applicable.

**Conflicts of Interest:** The authors declare no conflict of interest.

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