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Biochar and activated carbons preparation from invasive algae *Sargassum spp.* for Chlordecone availability reduction in contaminated soils

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Keywords

Sargassum; Activated carbon; Biochar; Chlordecone; Pollution sequestration; Food safety

Abstract

This work aims to valorize an invasive brown macroalga (*Sargassum spp.* consisting of two species *Sargassum fluitans* and *Sargassum natans*) by producing biochars (BCs) and activated carbons (ACs). Its abundant and frequent occurrence along the Caribbean coastlines, Florida, Gulf of Mexico during the last past nine years, have triggered human health concerns and have negatively impacted local economy, ecology and the environment. In this paper, BCs and ACs were developed to assess the reduction of chlordecone (CLD) environmental availability in artificial and tropical contaminated soils. Such innovative approach was proposed to limit CLD bio-availability to fauna and outdoor reared-animals. The BCs were prepared by pyrolysis at 700 °C while ACs samples by chemical or physical activation. Textural characterization, has evidenced that bi-modal structures with micro-and mesopores, various surface and high pore volumes were successfully obtained. Finally, the environmental availability tests resulted in various ability of BCs or ACs to significantly sequester CLD on artificial contaminated soils and on a natural nitisol. In particular, the BCs prepared with a 3 h pyrolysis time, exhibited the highest porosities properties and was the best candidate to efficiently sequester CLD in soil samples.

1. Introduction

In Guadeloupe Islands (French Antilles) and other Caribbean islands, washing ashore of tons of pelagic *Sargassum spp.* (i.e. *Sargassum natans* and *Sargassum fluitans*, a complex of two co-occurring species of floating brown macroalga), have been regularly occurring since 2011 [1]. Such events are more than likely due to both global warming and anthropic activities. For instance, the islands of Guadeloupe faced in 2019 one of its worst event that had never been described in the past. Algae dramatically accumulated on the beaches windward coastlines (Fig. 1). Hydrogen sulfide and ammonia gases are produced from the *Sargassum* decomposition, causing health effects

and having indirect impacts on tourism, fisheries and recreational activities [2,3].

Recently, there has been a burgeoning trend in the Caribbean to transform such *Sargassum spp.* into added value matters. Herein, we propose to use this local and highly available resource to produce bio-chars (BCs) and activated carbons (ACs), thus contributing to solve the environmental problem of soils contamination by chlordecone (CLD) [4–6]. In fact, CLD was used during more than two decades (from 1972 to 1993) in the French West Indies, Africa and in other countries of central and South America, to treat banana plantations against the weevil *Cosmopolites sordidus* [7]. Whilst its toxicity

has been recognized since 1975, CLD was identified as a persistent organic pollutant only in May 2009 and as such was included in the Stockholm Convention on Persistent Organic Pollutants. The molecule has a high molecular weight (490.64 g mol⁻¹), a very low solubility in water (2.7 mg L⁻¹ at 25 °C), a high affinity for organic compounds (log K_{ow} = 4.5) and is highly stable due to its caged chlorinated structure [8,9]. As a result, CLD can easily bind to soil particles and residues have been found in sediment layers as well as in various living organisms (i.e. human beings, animal and plants) from the land and coastal areas [10].

Unfortunately, the presence of CLD in water, soils and sediments of banana crops production zones, should last at least several decades up to seven centuries (for the most polluted Andosols) [11]. Therefore, in this work, the development of BCs and ACs derived from *Sargassum* spp. has been investigated to propose an eco-friendly opportunity of "biomass waste valorization" and an innovative

strategy for limiting CLD bio-availability to fauna and outdoor reared-animals.

We have produced and characterized BCs and ACs, (with different textural and physico-chemical properties) to assess their potential for sequestering CLD in artificial and natural soils. While previous works have focused on using BCs to limit pollutants transfer [12–14], to improve soil water retention [15], for microbial activities [16] and to limit the volatility of nitrogen compounds needed for plant growth [17, 18]; few have examined the strategy of using BCs for preventing pollutant transfer to livestock. Recent papers [19–21] have reported on the great potential of ACs and BCs to sequester CLD in Antillean soil. In our previous work, the amendment of Nitisol by lignocellulosic ACs, such as Coconut shell or Oak wood which resulted in significant reductions of CLD-availability [21]. However, to our knowledge, no sequestration data are currently available concerning BCs or ACs derived from *Sargassum* spp.

2. Material and Methods

2.1. Preparation of activated carbons from raw material

French West Indies) in July 2017. The samples were initially dried at room temperature and then in an oven at 105 °C for 48 h to remove the moisture content. They were then grounded and finally sieved into several fractions [22]. The fraction containing particle size ranging from 0.4 to 1 mm was selected as precursor for producing the ACs and BCs. In this work, BCs were obtained from the pyrolysis of the *Sargassum* spp. precursor (i). The ACs were obtained either by physical activation of BCs (ii), or by chemical activation (iii) [22].

(i) *Sargassum* spp. precursor was pyrolysed in a nacelle which was placed in a tubular furnace (Carbolite Gero® CTF 12/75/700) under a nitrogen flow of 80 mL min⁻¹ at 700 °C for 1 h or 3 h, with a heating rate of 10 °C.min⁻¹. The pyrolysis temperature chosen is that for which the yield η (%) remains constant [23].

Where

w_i weight of initial dry precursor (g).
 w_f mass of carbon after pyrolysis (g).

$$\eta(\%) = 100 \times (w_i - (w_i - w_f)) / w_i$$

After the pyrolysis, the as-prepared BCs were washed under a fume hood with HCl (5 M) to remove impurities. Subsequent washing was carried out with deionized water (10 μ S cm⁻¹) in a beaker with stirring, until pH of 6–7 was reached [24,25]. Samples were finally purified by Soxhlet extraction during 24 h with deionized water, giving rise to BCs labeled BioSarg 1 h and BioSarg 3 h, depending on the pyrolysis time.

(i) For physical activation, approximately 3 g of BioSarg 1 h was heated at 10 °C min⁻¹ to the final temperature of 600 °C with carbon dioxide (CO₂), a CO₂/H₂O mixture and steam (H₂O), using nitrogen as carrier gas at a flow rate of 80 mL.min⁻¹, during 19 h, 20 h or 14 h respectively. Different time span and a temperature of 600 °C were chosen to reach a burn-off of approximately 50% and avoid the eventual collapse of some pore walls in the core of the ACs [26].

Samples were labeled SargCO₂, SargCO₂/H₂O and SargH₂O. The burn-off (%) refers to the percentage of mass loss, which indicates efficient activation [27,28]. This is calculated according to the following equation:

$$(1) \quad \text{<<burn - off >>\%} = 1 - W_{\text{final}}/W_{\text{initial}} * 100 \quad (2)$$

with

W_{initial} weight of initial dry BCs (g).
 W_{final} mass of carbon after activation (g).

(i) Chemical activation was carried out with approximately 9 g of *Sargassum* spp. precursor. For this purpose, the latter was impregnated in phosphoric acid (H_3PO_4 , 85 wt%) for 24 h to ensure optimal wettability, resulting in an impregnation ratio of 0.5:1 (g H_3PO_4 /g precursor) [22]. Thereafter, the samples were dried in an oven at 110 °C during 4 h, followed by a pyrolysis step using the same conditions as those described in BCs preparation. After cooling, until ambient temperature, the ACs thus obtained were washed with distilled water until stabilization of the pH, and then dried overnight using a drying oven at 110 °C.

In our study, chemically activated samples are referred to as SargP0.5.

2.2. Characterization of the ACs and BCs

Thermogravimetric analysis was carried out using a Labsys Evo-Setaram instrument from 35° to 900 °C at a rate of 10 °C.min⁻¹, under an inert nitrogen atmosphere flow of 80 mL min⁻¹. The morphological features of samples were analyzed by scanning electron microscopy (SEM, Hitachi S-2500, accelerating voltage of 20 kV).

Textural properties of ACs and BCs were determined using a Micro-meritics ASAP 2020 apparatus. Prior to analysis, the samples were degassed for 24 h at 300 °C. To estimate the surface area (S_{BET}), the Brunauer, Emmett and Teller (BET) model [29] was utilized and the Dubinin–Radushkevich (D-R) equation was used to determine micro-porous volume [30]. The mesoporous volume was estimated by applying the Barrett–Joiner–Halenda (BJH) method on the desorption branch [31,32].

Raman spectroscopy analysis was determined using a HR 800 Horiba multi-channel spectrometer.

FT-IR spectra were collected using a Perkin Elmer Spectrum One FT-IR Infrared Spectrometer, equipped with an ATR sampling accessory (diamond/ZnSe composite crystal). The spectra were recorded in a range of 4000–550 cm⁻¹.

X-ray photoelectron spectroscopy was investigated using a Kratos

2.3. Assessment of CLD sequestration potential of *Sargassum* biochars and ACs

In this study CLD sequestration efficiency of each *Sargassum* BCs or ACs was assessed through environmental availability assays, which allow to determine the CLD part which may be extracted by fauna or biota as described in [21]. These assays were first applied to artificial soils in order to select the most efficient *Sargassum* BCs or ACs. In a second step, the most efficient *sargassum* BCs was added to an Antillean CLD contaminated nitisol.

2.3.1. Environmental availability assays

Environmental availability assays were performed on each artificial and natural soil sample. The protocol was adapted from the XP ISO/TS 16 751 Part B to fit a 2 g soil-sample instead of 5 g as previously described [21]. Briefly, 600 ± 25 mg of Tenax® (60–80 mesh, Sigma-Aldrich, Saint-Louis) and 28 mL of ultrapure water were added to soil samples. After a 20 h agitation period Tenax® was recovered and then extracted 3 times with acetone (LV-GC, Biosolve, Lyon) using an ultrasonic bath. All extracts were combined and concentrated prior GC-MS analyze (Agilent 7890) to quantify CLD content.

2.3.2. Artificial soils preparation

Artificial soils were prepared accordingly to the OECD guideline 207 [33] as described in [21]. All artificial soils were spiked to reach a concentration of 5 µg of CLD (Kepone, Supelco, Sigma Aldrich, Saint-Louis) per g of DM. Then BCs or ACs were added (2% dry mass basis of whole soil) and hand-mixed thoroughly. All artificial soils were aged during 80 days prior the following experimental parts [21].

2.3.3. Nitisol amendment

Nitisol sample preparation and amendment were performed as described in [21]. Briefly, Nitisol contaminated soil was sampled in Guadeloupe [34] and dried at room temperature until mass stabilization. Then, 2% of the most efficient condensed materials (dry mass basis) was added to 2 g subsamples of this soil (n = 3). Six non-amended subsamples were used as control. Finally, ultrapure water was added to reach 18% of mass of the wet soil. All soils were aged during 80 days at room temperature (20 ± 2 °C) prior the following experimental parts.

2.3.4. Statistical analysis

In order to assess the impact of sargassum BCs and ACs on CLD environmental availability, an

ANOVA was performed. The ANOVA procedure and the Tukey–Kramer post-hoc test of R version 3.2.3 (R Foundation for Statistical Computing, Vienna, Austria) were used. Differences were considered significant at $P < 0.05$.

3. Results and Discussion

3.1. Characterization of the ACs derived from *Sargassum fluitans*

3.1.1. Thermogravimetric behavior

To anticipate AC and BC samples preparation procedure, thermal behavior of the sargassum precursor was studied by a thermogravimetric (TG) technique. Fig. 2(a) shows TG (weight loss, %) and DTG (derivative of TG) curves, while Fig. 2(b) shows the corresponding DSC (differential scanning calorimetry) curve with both endo- and exothermic peaks. Analysis of these thermograms revealed three different stages of the alga degradation: (i) stage I - from initial temperature to 150 °C, (ii) stage II - between 200 and 400 °C and (iii) stage III - from 450 °C to 900 °C. During the first stage, an endothermic peak was observed (at around 135 °C), which could correspond to the desorption of both cellular water and external water bound by surface tension. This represents a total mass loss of 4.3 wt% (calculated from the TG plot). In the second stage, the main pyrolytic process occurred and majority of weight loss was observed. The DSC curve shows an endothermic peak at 212 °C followed by an exothermic peak at 270 °C, corresponding to a mass loss of 8.6 wt% and 27.7 wt%, respectively. Previous work has shown that those phases could be attributed to the removal of alginic acid and fucoidan, two major constituents of algal biomass [35,36]. The following mass loss of 39.1 wt%, at around 328 °C, can be assigned to the decomposition of other carbohydrates [37]. Third stage starting from 450 °C with an endothermic effect, was probably caused by the loss of volatile metal and by carbonate decomposition [38,39]. Such multi-compounds decomposition of *Sargassum* spp., at temperature below 700 °C, suggests a complex structure. It should be noted that after 700 °C, no significant weight loss was observed, thereby suggesting thermal stability of the sample was reached. Therefore, a pyrolysis temperature of 700 °C can be chosen for the BCs preparation.

3.1.2. Scanning electron microscopy (SEM)

Surface morphology of the *Sargassum* samples was analyzed by scanning electron microscopy (SEM) as shown in Fig. 3. A relatively dense

sample surface with rod-like tissues could be observed, which essentially contains the proteins and cell wall of the algae (Fig. 3a). Unlike lignocellulosic precursors, it seems that the brown alga does not have macropores intrinsic to its matrix, which is probably due to its limited fibrillar microstructure [40]. Upon pyrolysis, the sample underwent microstructural modification, as indicated by the appearance of a rough surface in which macropores were clearly identifiable (Fig. 3b, c). Such thermal treatment has successfully led to release volatile compounds contained in the seaweed, giving rise to a porous structure. However, at such magnification, it is clear that SEM analysis did not provide evidences of the presence of smaller micropores that could participate to CLD sequestration. This feature will be discussed in the next sections using appropriate characterization techniques.

3.1.3. BET analysis

Adsorption/desorption of N₂ at 77 K was measured for the BCs and ACs *Sargassum* spp. and the corresponding isotherms are displayed in Fig. 4. Hysteresis loops were observed in all samples (except for BioSarg 1 h), indicating type IV isotherms according to the classification established by De Boer et al. [41]. The well-defined hysteresis loops (especially observed for BioSarg 3 h and SargP0.5), which taken in conjunction with the initial uptake of volume adsorbed at low partial pressure ($P/P_0 < 0.2$); indicated that the pore size distribution for these samples was bimodal with both micro- and mesopores present. On the other hand, BioSarg 1 h exhibited a type I isotherm, characteristic of exclusively microporous materials. Such trends were confirmed by analyzing the textural parameters (specific surface area, pore volumes and pore width) depicted in Table 1. BioSarg 1 h exhibited only micropores combined with the lowest total volume ($V_T = 0.019 \text{ cm}^3$), while BioSarg 3 h and SargP0.5 contained both micropores and meso-pores combined with highest total volumes ($V_T = 0.99$ and $0.76 \text{ cm}^3 \text{ g}^{-1}$). One can note that, comparison of BioSarg 3 h and Biosarg 1 h clearly shows the influence of the pyrolysis time as more porosity was developed in Biosarg 3 h. In addition, BioSarg 3 h exhibited one of the largest pore size ($DP = 4.54 \text{ nm}$) whilst BioSarg 1 h the lowest one ($DP = 1.56 \text{ nm}$). It is

expected that such features of BioSarg 3 h could contribute to enhance CLD bioavailability [21]. Finally, in term of surface areas, the descending order was as follows: SargCO₂ > SargP0.5 > BioSarg 3 h > SargH₂O > Sarg H₂O/CO₂ > BioSarg 1 h. The physically activated SargCO₂ has the largest surface area (SBET = 969 m².g⁻¹) with a large fraction of small mesopores compared to its microporous volume.

3.1.4. **Part 2**

ACs and BCs microstructures were characterized by Resonance Raman spectroscopy and the spectra are shown in Fig. 5. The Raman “G” peak referring to the bond-stretching vibrations of sp² carbon sites is generally found around 1580 cm⁻¹ in crystalline graphite [42]. The “D” peak at around 1355 cm⁻¹ is a signature of disorder and/or defects in diamond carbon and amorphous carbon due to sp³ breathing vibrations [43,44].

The D-mode is attributed to lattice defects and finite crystal size occurring inside the graphene atomic layer which induce a breaking of the 2D translational symmetry [45]. ID/IG represents the ratio R of the integrated intensity of the D mode (ID) over that of the G mode (IG) deduced from the Raman analysis. R is commonly used to characterize disorder in graphene and has usually the same trend as the ratio of the area under the D-peak over that under the G-peak [46]. BCs have the highest ID /IG ratio = 1.04 and 1.02 (Table 2). This means that the D band is related to formation of defects (disorder) and a high proportion of sp³ carbon and results from breaks in the sp² hybridized (out of plane) carbon bonds [47], according to the fact that BCs are only submitted to a one step pyrolysis [21].

3.1.5. FTIR spectroscopy

To identify chemical surface groups of the prepared samples, FT-IR spectroscopy was carried out. Fig. 6 shows the resulting spectra of the ACs (SargP0.5, SargCO₂, SargH₂O and Sarg CO₂/H₂O) along with that of BCs (BioSarg 1 h and BioSarg 3 h). It is clearly seen that there are similarities and differences in chemical makeup by comparing the BCs and ACs samples.

In ACs spectra, features bands were identified: (i) at 2981 and 2930 cm⁻¹ which are associated with asymmetric and symmetric CH₂ stretching vibrations, respectively; (ii) between 1400 and 1600 cm⁻¹ which are ascribed to the skeletal C=C vibrations in aromatic rings bands; (iii) at around 1400 cm⁻¹ which can be partially described by OH deformation vibration; (iv) in the region of 1350–

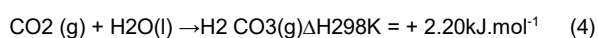
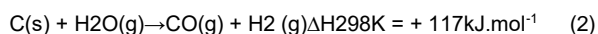
1200 cm⁻¹ which can be attributed to C–O–C vibrations; (v) at 871 cm⁻¹ can be attributed to C^{*}CH₂ bending out of-plane [48,49].

In BCs spectra, characteristic bands were identified at 1200 and 1530 cm⁻¹ which can be related to the hydroxyl and C–O–C groups stretching vibration in pyranose ring skeletal or stretching in aromatic ring and C=C groups (sp²-hybridized carbon) respectively [50].

We have to bear in mind that ACs prepared by physical activation were produced from BCs (activation of BioSarg 1 h with steam or CO₂). The significant absorption band in the region 1350–1200 cm⁻¹, suggests that the activation with steam or CO₂ favored the development of C–O groups on the surface of the ACs. This agrees well with observation reported in literature [51,52]. Overall, FT-IR results showed that the differences in surface chemistry of the ACs are relatively small and could lead to similar effect in the reduction of CLD bioavailability.

3.1.6. X-ray photoelectron spectroscopy (XPS)

The produced ACs and BCs were further analyzed via XPS spectroscopy to identify chemical elements and functions present on their surfaces. As shown in Table 2, the main elements detected are O, Na, Cl, Ca, S, Mg, Si and C, which correspond to typical marine residues [35]. Although the as-prepared samples were thoroughly washed in deionized water, persistence amounts of salt (NaCl) remain in some of the sample matrices (i.e. BioSarg 3 h, SargCO₂, SargH₂O/CO₂ and SargH₂O). Noteworthy, the salt content was higher in sample SargCO₂ than for those activated with steam (SargH₂O and SargH₂O/CO₂). This suggests that the activation with water vapor preferentially removes the NaCl during the process. This is supported by the fact that steam activation requires less energy than CO₂ activation as shown in Eqs. (2) and (3) SargH₂O/CO₂ sample confirms this trend due to the competition between the CO₂ and H₂O molecules as shown in Eq. (4).



As seen in Table 2, the ratio O1s/C1s increases by 1.5 fold when comparing SargH₂O against SargCO₂, indicating that during the activation

process, additional oxygenated functional groups were formed

However, the activation with H₂O/CO₂ mixture, shows an opposite effect, as the ratio O1s/C1s decreased by a factor close to 0.6. This observation is in good agreement with the work published by Mangun et al. [55], where the oxidation steps during physical activation were described as follows:



The deconvolution (not shown here) of the C1s spectrum consists of five components with binding energies corresponding to (i) the graphite type and amorphous carbon (C-C, C-H) at 284.6 eV, (ii) carboxyl and ester groups (O-C) at 285.7 eV, (iii) carbonyl group (O=C) at 287.2 eV, (iv) acidic group (O = C-OH) at 289.7 eV, (v) and a peak at around 292.1 eV belonging to the π - π^* transition band (HOMO-LUMO) in the aromatic carbon [56,57]. The O1s oxygen peak spectrum was also deconvoluted into two peaks at 531.4 eV and 532.9 eV. The peak at 531.4 eV can be ascribed to the oxygen attached to the carbonyl group (O = C) and the peak at 532.9 eV can be attributed to the oxygen in the epoxy and/ or hydroxyl (C-O-C, C-O - H) [58].

According to the results presented in Table 2, BioSarg 1 h and Bio-Sarg 3 h showed a higher content of disordered carbons C-C, C-H (54.39% and 74.34%, respectively) than the ACs materials. This correlates well with the RAMAN analysis, which showed that ACs matrices were the most structurally organized (with lowest % of disordered carbon ranging from 47.92% to 49.24%). In addition, ACs sample made by physical activation reveals a slightly higher percentage of carbonyl groups (C=O, O-C=O, HO-C=O). (Table 3).

3.2. Assessment of CLD sequestration potential of sargassum biochars and ACs

The first set of assays involved 7 OECD soils amended by one of the Sargassum ACs or BCs. The results are presented in Fig. 7. One of the significant figures is that $84.2 \pm 2.1\%$ of the whole CLD content was retrieved from Artificial soils (n = 6) without amendment indicating that the main part of soil CLD was easily available. No significant reduction of CLD availability was observed when the artificial soils were amended by the different Sargassum ACs or by BioSarg 1 h. The obtained values are ranging from 69% to 96% of environmental availability. Interestingly, a much better response ($p < 0.001$) was obtained when the

artificial soil was amended by the BioSarg 3 h (17% of CLD environmental availability compared to 84.2% without amendment). This reduction corresponds to 4/5 reduction of CLD environmental availability compared to the non-amended artificial soil.

In the second set of assays, a naturally contaminated Nitisol sample amended by BioSarg 3 h was studied. Significant reduced level of environmental availability was observed similarly to as in artificial soils, revealing the great sequestration potential of this BCs (Fig. 7). Thus, BioSarg 3 h presented more than 80% reduction of CLD environmental availability in Nitisol (19% environmental availability), confirming the effectiveness of this amendment to sequester CLD even in a contaminated soil.

3.2.1. Impact of Sargassum ACs or BCs characteristics on CLD sequestration

Interestingly, the present study reveals that one BCs (BioSarg 3 h), was able to efficiently sequester CLD, either in the OECD artificial soils or in the CLD contaminated Nitisol. In contrast to previous results obtained from lignocellulosic precursors, where the BCs did not present a strong sequestration potential [21], the BioSarg 3 h used in this study appears to be a valuable candidate for this purpose. Indeed, BCs is prepared in an eco-friendly way with a unique pyrolysis step that may induce, even if the sample must be effectively washed, a reduced cost of preparation at industrial scale when compared to physical or chemical activation.

The explanation of the sequestration capabilities is related on the fact that BioSarg 3 h exhibits numerous interesting characteristics when compared to others samples. Indeed, textural characteristics like specific BET surface area, total pore volume, and micro- and mesopore volume and mean pore diameter were the highest when compared to all other BCs and ACs. These physico-chemical characteristics are well known to play a major role for organic pesticide sequestration [21]. CLD availability reduction is far much higher than the values obtained for samples SargP0.5, SargCO₂ that exhibited as well a high surface area value. In addition, textural properties like micro- and mesopore volumes appear also to play a role in this reduction. Indeed SargCO₂ has the highest surface area of 968 m².g⁻¹, but was not able to retain CLD in this study, that may be due to the very low amount of micropores that may entraps CLD. Indeed, the BioSarg 3 h displayed a high proportion of microporosity and mesoporosity compared to the other matrices. A similar

observation was also previously noticed in another set of BCs and ACs [21,59]. On the other hand SargP0.5 that contains as well micropores in slightly lower quantity than BioSarg 3 h is not able to retain significantly CLD. Even with a very low surface area, and low micropore volume, SargH2O gives a CLD availability reduction of 32%, showing that textural characteristic is not the only factor affecting CLD sequestration by carbon material. Chemical groups at carbon samples, may as well play a role in CLD availability reduction in soil. Indeed, surface chemistry is known to play a major role in chlorinated pesticides adsorption in water, for example [60,61] that is not observed in the present work, where CLD is removed from soil instead of water. Although several surface characterizations were carried out in this work, no

conclusion could be drawn at this stage concerning the relationship between surface chemistry and CLD-bioavailability.

Overall, these results suggest that a significant CLD environmental availability reduction can be expected only when the condensed material is highly porous but it depends as well on other factors. Therefore, influence of carbon chemistry, that includes surface groups polarity, acido-basic character, which are dependent on soil characteristics such as soil pH, moisture, etc., needs further investigations.

4. Conclusion

This study revealed that pyrolysis of Sargassum spp. can result in condensed materials with high level of porosities. In particular, BioSarg

3 h (Sargassum precursor pyrolysed during 3 h) displayed both the highest total volume in pore and one of the best BET surface. Thus, this biochar appeared to be the best candidate to sequester efficiently CLD in soil samples, suggesting that textural properties of the materials played a major role. This really promising result indicates that a

wider use of this biochar, with a lower preparation cost and easy preparation process, could contribute to strongly limit the CLD transfer to the food chain. Additional works (i.e. influence of pH soils, effect of CLD loading in soils, contact time, etc.) are now in progress to provide better understanding of the obtained results. Overall, these results can be considered as a major first step toward further bioavailability assays aiming to study CLD transfer from soils to both plants and food producing animals.

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Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Table 1. Textural parameters of Sargassum spp. BCs and ACs calculated from N₂ sorption analysis.

Sample	Burn-off (%)	BET Surface area g ⁻¹	V _{Micropores} (D-R) (cm ³ g ⁻¹)	V _{Mesopores} (BJH) g ⁻¹	Total volume in pore g ⁻¹	Average pore width D _p (nm)
BioSarg 1 h	72	54	0.021	—	0.019 (NLDFT ^a)	1.56
BioSarg 3 h	70	872	0.44	0.55	0.99	4.54
SargP0.5	51	877	0.30	0.46	0.76	3.46
Sarg H ₂ O/ CO ₂	47	80	0.10	0.36	0.46	2.29
Sarg H ₂ O	46	109	0.12	0.09	0.13	4.79
Sarg CO ₂	46	969	0.041	0.34	0.38	1.57

^a NLDFT: Non-local density functional theory.

Table 2. Calculated ID/IG ratio, graphitic and disordered carbon proportions in ACs and BCs samples.

Sample	BioSarg 1 h	BioSarg 3 h	SargP0.5	SargCO ₂	SargH ₂ O	SargH ₂ O/CO ₂
I _D /I _G	1.02	1.04	0.94	0.96	0.97	0.92
% graphitic carbon	49.50	49.02	51.55	51.02	50.76	52.08
% disordered carbon	50.50	50.98	48.45	48.98	49.24	47.92

Table 3. Atomic concentration (%) of the chemical elements and functional groups present Sargassum spp. BCs and ACs surfaces (obtained from XPS spectra).

Sample	Atomic Concentration (%)															
	Na 1 s	O 1 s	Ca 2p	C 1 s	Cl 2p	Mg 2p	S 2p	Si 2p	O 1 s/C 1 s	C-C; C-H	C-O	C ⁺ O	O-C ⁺ O	HO-C ⁺ O	π-π ⁺	π-π ⁻
BioSarg 1 h	16.93	32.98	6.69	30.41	9.13	—	3.87	—	1.08	54.39	29.26	16.35	—	—	—	—
BioSarg 3 h	—	46.29	—	28.71	—	15.08	—	9.93	1.61	74.34	19.13	6.52	—	—	—	—
SargP0.5	—	41.41	7.89	32.34	—	8.25	8.08	2.03	1.28	—	—	—	—	—	—	—
SargCO ₂	25.37	19.53	7.11	23.01	21.01	3.49	—	—	0.85	48.19	25.71	—	26.11	—	—	—
SargH ₂ O	11.83	22.78	9.31	41.27	9.44	3.89	1.46	—	0.55	46.23	10.66	26.19	—	—	12.33	4.56
SargH ₂ O/CO ₂	10.7	23.72	—	49.26	8.52	6.71	1.10	—	0.48	48.6	24.27	14.1	—	5.63	7.4	—



Figure 1. Sargassum spp. accumulation on Saint-Felix beach, Guadeloupe Island(French Antilles)

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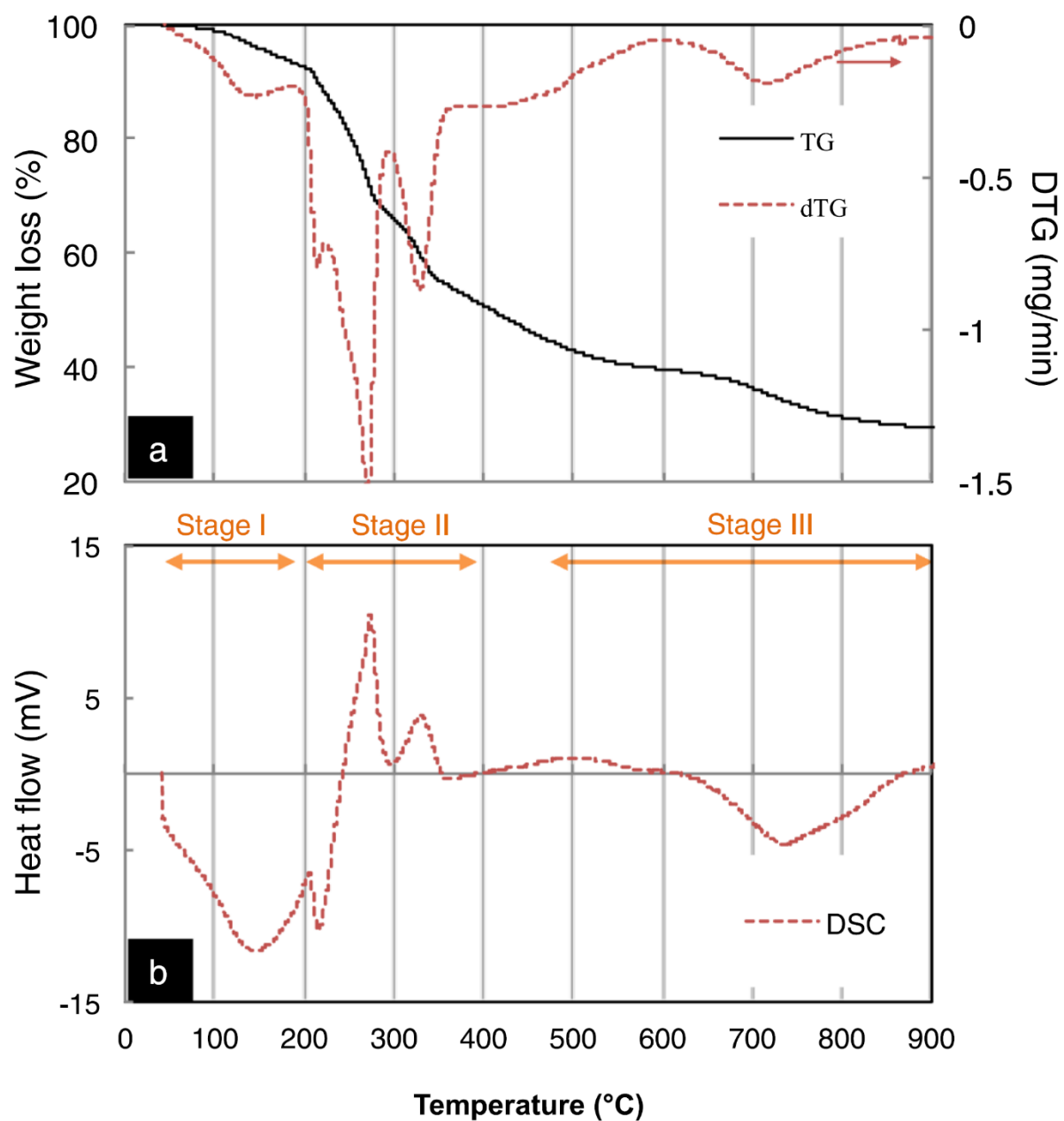


Figure 2. Thermogravimetric plots of Sargassum spp. precursor (a) TGA and DTG, (b) DSC.

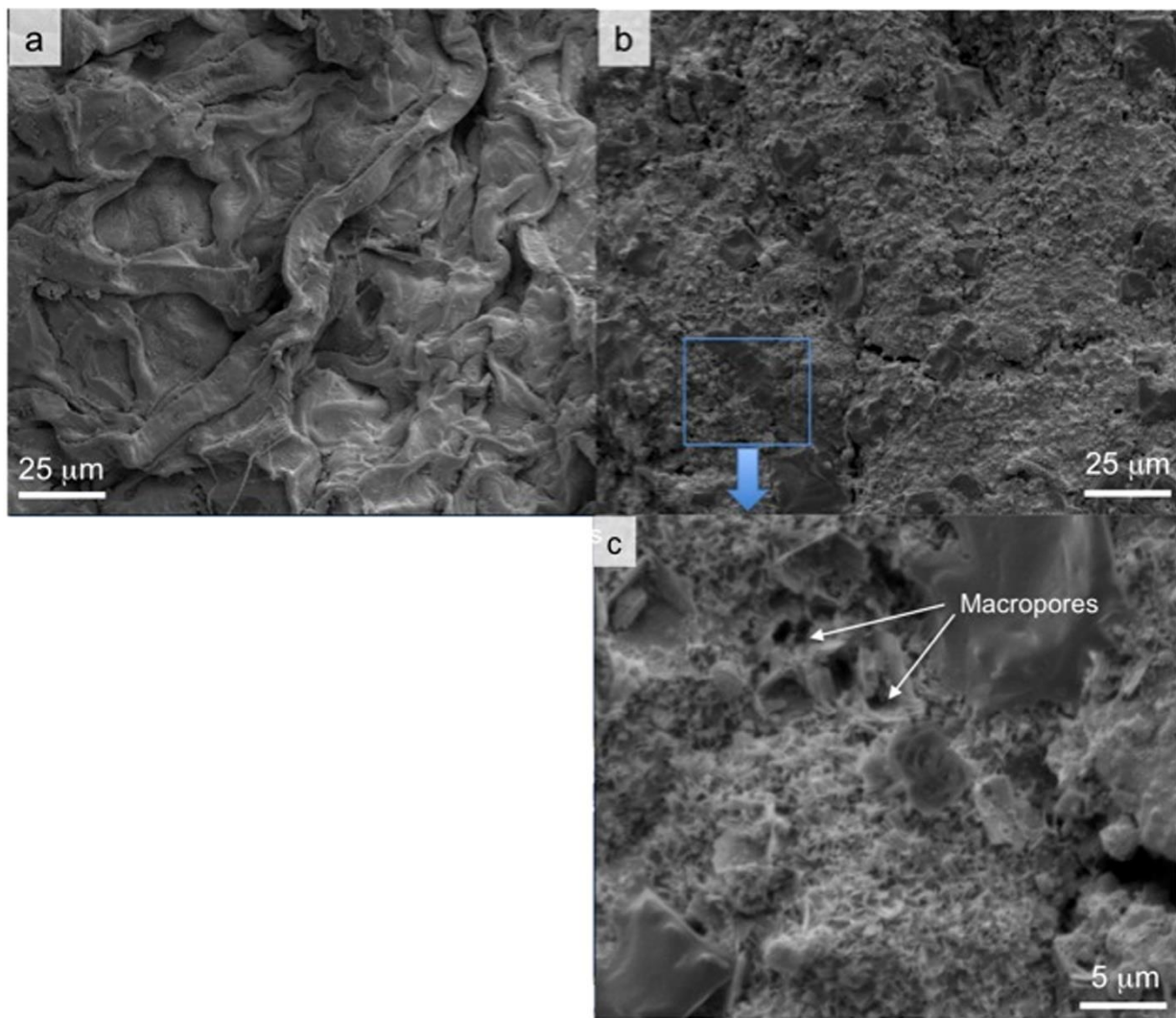


Figure 3. SEM images of *Sargassum* spp. before (a), after (b, c) pyrolysis at 700°C.

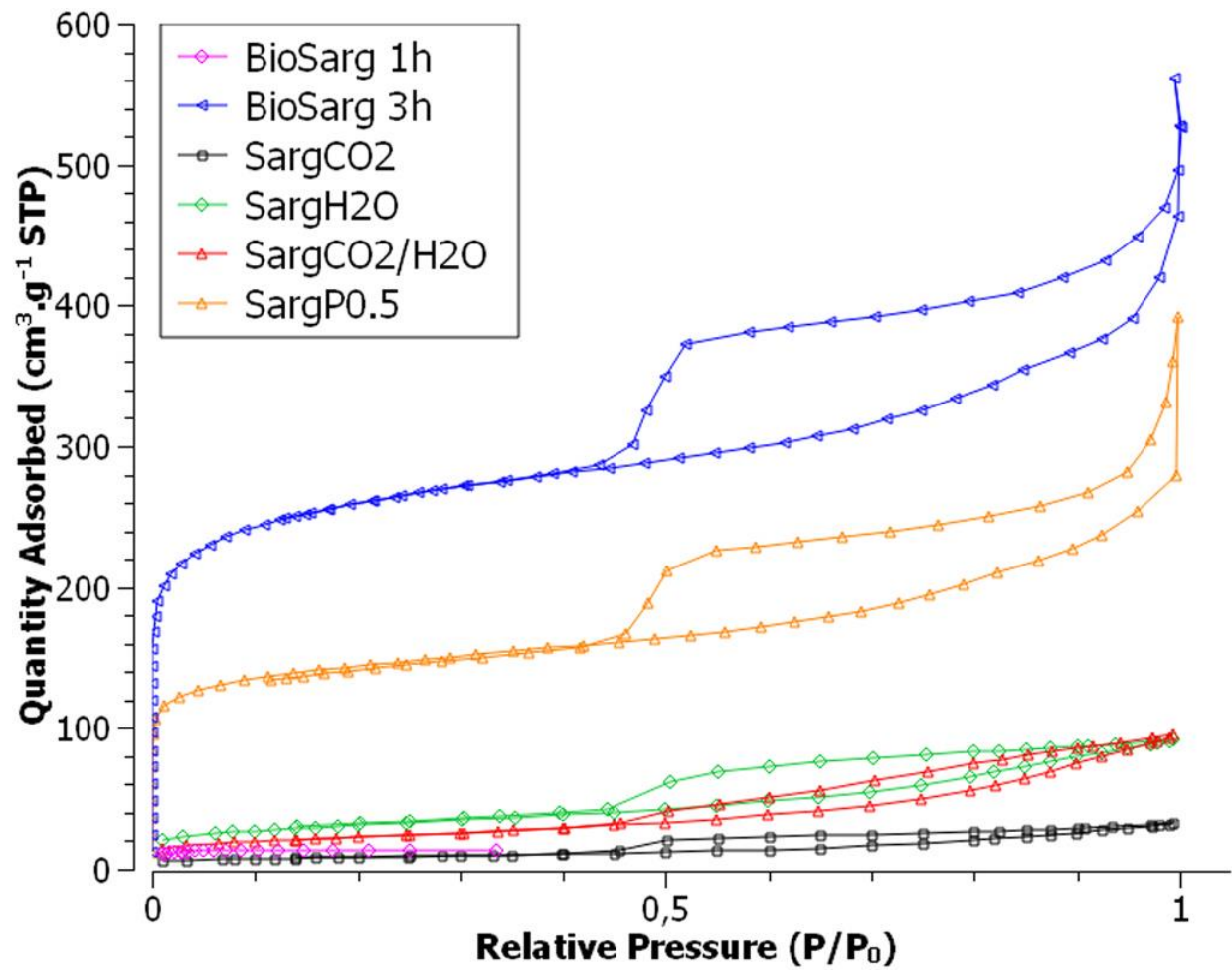


Figure 4. N₂ adsorption-desorption isotherms of Sargassum spp. BCs (BioSarg 1h, BioSarg 3h) and ACs (SargCO₂, SargH₂O, SargCO₂/H₂O and SargP0.5).

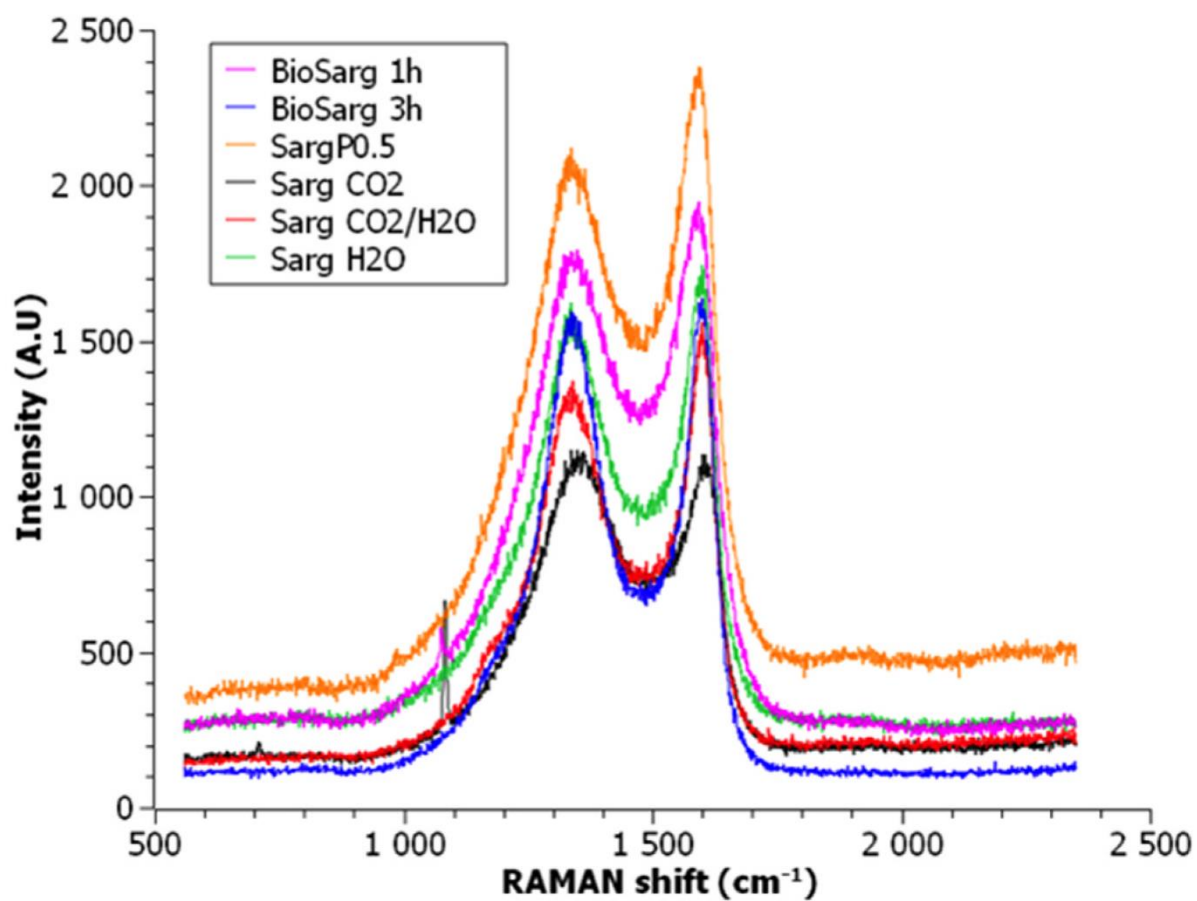


Figure 5. RAMAN spectra of *Sargassum spp.* BCs and ACs.

Indication: Numbers correspond to the PCB congeners. Framed congeners are dioxin-like PCBs. Bold numbers were congeners transferred at a high level ranking from 38 to 78% and from 30 to 80% respectively for milk and eggs.

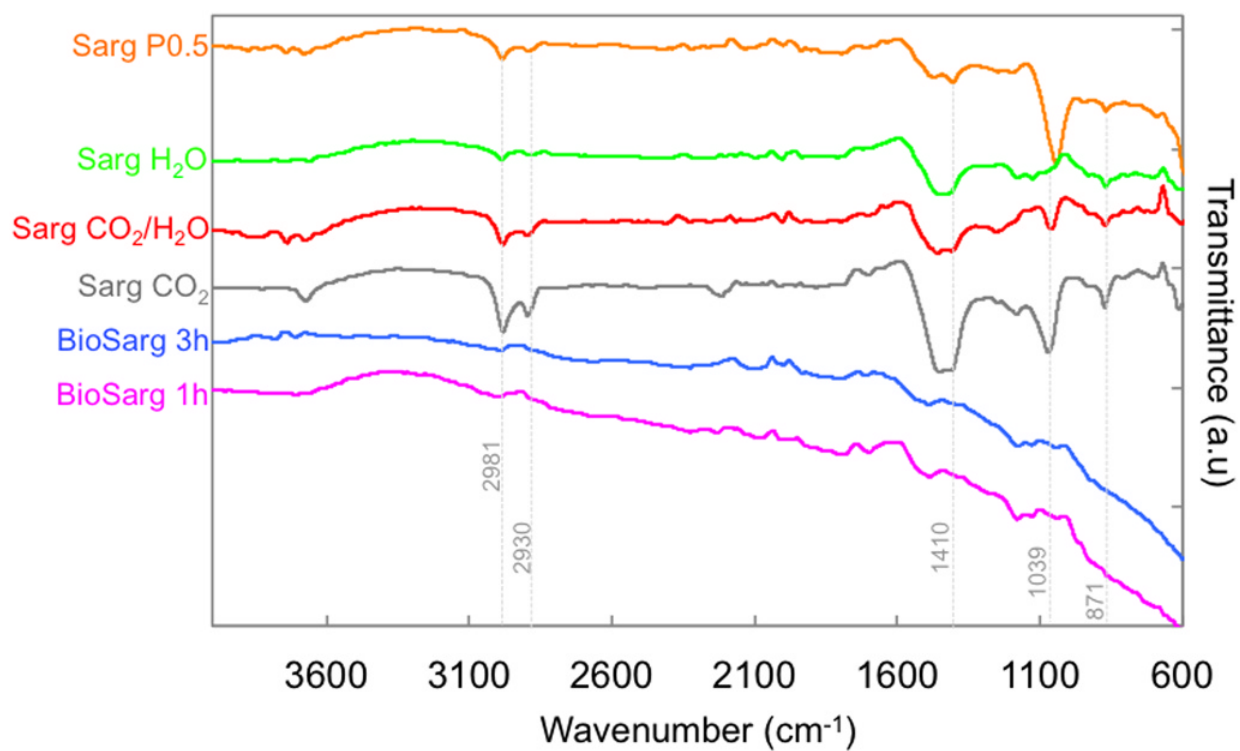


Figure 6. FT-IR spectra of *Sargassum* spp. based samples: biochars BCs (BioSarg1h, BioSarg 3h) and activated carbons ACs (SargCO₂, SargH₂O, SargCO₂/ H₂O and SargP0.5).

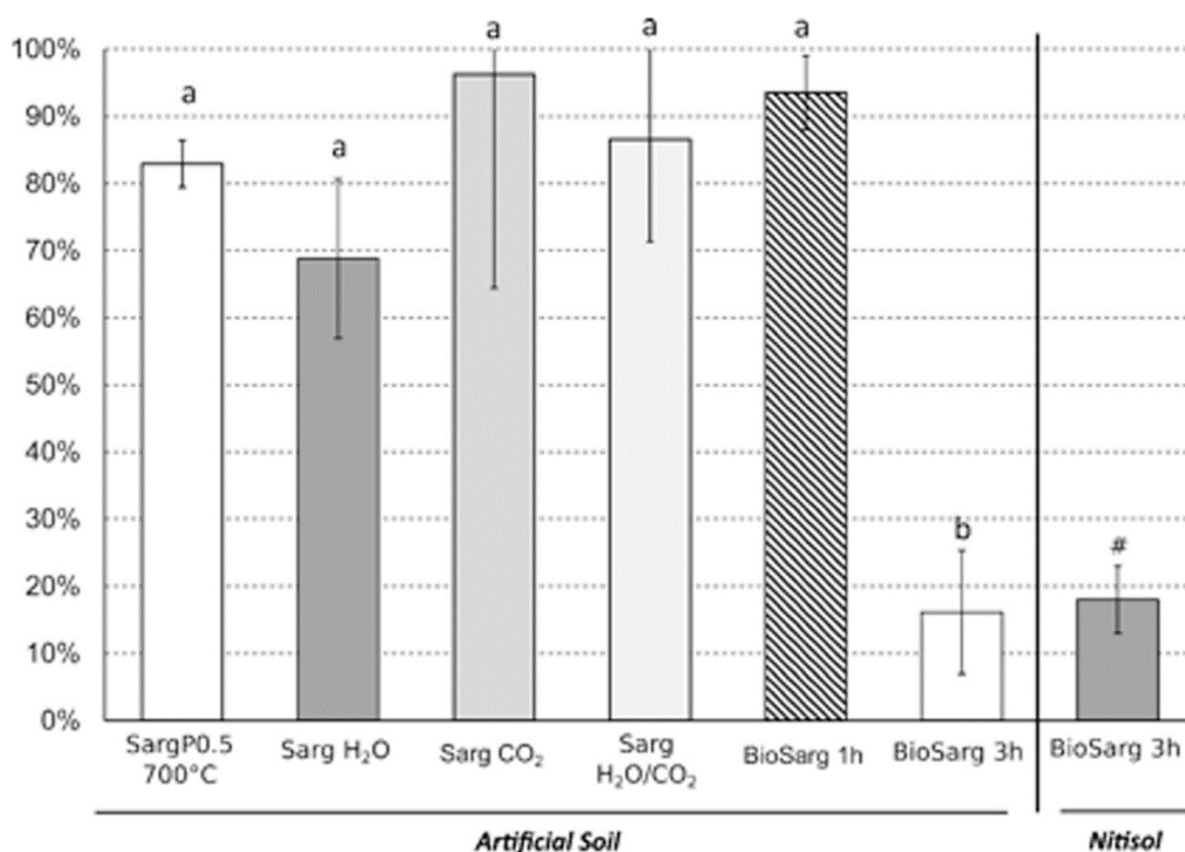


Figure 5. CLD availability in OECD artificial soil and Nitisol. CLD availability is expressed in % of CLD availability of corresponding soil without amendment (n = 6).

Values correspond to the mean \pm SD (n = 3). Mean values with different superscript letters (a, b) are statistically different ($P < 0.05$). #: Amended Nitisol presented significantly reduced levels ($p < 0.001$) compare to Nitisol without amendment. Statistical analysis was performed using the one-way ANOVA procedure of R software and Tukey post-hoc test.

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