

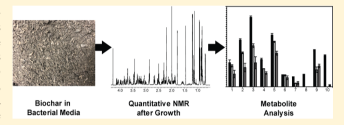
Effect of Biochar on Microbial Growth: A Metabolomics and Bacteriological Investigation in E. coli

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Supporting Information

ABSTRACT: Biochar has been proposed as a soil amendment in agricultural applications due to its advantageous adsorptive properties, high porosity, and low cost. These properties allow biochar to retain soil nutrients, yet the effects of biochar on bacterial growth remain poorly understood. To examine how biochar influences microbial metabolism, Escherichia coli was grown in a complex, well-defined media and treated with either biochar or activated carbon. The concentration of metabolites in the media were then



quantified at several time points using NMR spectroscopy. Several metabolites were immediately adsorbed by the char, including L-asparagine, L-glutamine, and L-arginine. However, we find that biochar quantitatively adsorbs less of these metabolic precursors when compared to activated carbon. Electron microscopy reveals differences in surface morphology after cell culture, suggesting that Escherichia coli can form biofilms on the surfaces of the biochar. An examination of significant compounds in the tricarboxylic acid cycle and glycolysis reveals that treatment with biochar is less disruptive than activated carbon throughout metabolism. While both biochar and activated carbon slowed growth compared to untreated media, Escherichia coli in biochartreated media grew more efficiently, as indicated by a longer logarithmic growth phase and a higher final cell density. This work suggests that biochar can serve as a beneficial soil amendment while minimizing the impact on bacterial viability. In addition, the experiments identify a mechanism for biochar's effectiveness in soil conditioning and reveal how biochar can alter specific bacterial metabolic pathways.

■ INTRODUCTION

Biochar (BC) is often produced from renewable organic waste that is converted to a carbonaceous material through pyrolysis, and it is comparatively cheaper than activated carbon (AC). 1-3 BC, a carbon source, is often used in agriculture as a soil additive due to its adsorptive properties, porous surface, and low cost. BC improves soil fertility and structure, sequesters carbon (mitigating climate change), and enriches beneficial soil microorganisms. 4-9 BC's adsorptive properties vary with feedstock and pyrolysis conditions, which can be tailored for optimal removal of specific chemicals such as pesticides, phosphates, organic dyes, heavy metals, and pharmaceuticals from wastewater and soil. 3,9-14

Previous studies have focused on bulk properties of soil after treatment with BC rather than focusing on the behavior of individual metabolites. These properties—pH, adsorption, nitrogen content, etc.—can change how organisms grow in BC-treated soil. For example, addition of BC to soil can alter metal mobility, either positively or negatively, depending on its influence on soil pH and dissolved organic matter.3,11,13 BC adsorptive properties also improve soil physical and chemical properties by increasing pH, aeration, and water holding capacity. 1,3,15,16 Moreover, BC has been shown to restore nutrients derived from feedstock by modulating cation

exchange capacity (CEC), increasing specific surface area, and influencing hydrophobicity. 10,11,17

Physical and chemical properties of BC increase net nitrogen mineralization, increase nitrification, and enhance denitrification during microbial-mediated transformation of nutrients in soil. 11,18 Microbial activity itself is influenced by BC treatment. This has been linked to nutrient and pH changes, as well as nitrogen transformations. 10,15 BC-treated soil can also influence microbial composition by enhancing plant growth and mineralization of the soil's organic carbon. 11,15 While it is clear that BC can dramatically influence organisms in the soil by altering soil properties, much less is known about how BC influences microbial metabolism. Presumably, biochar's metabolic influence favors organisms that prefer basic growing conditions or disfavors organisms that prefer acidic growing conditions, as the populations of organisms change with the addition of BC into the soil. 11,15 BC is also effective in controlling foliar and soil-borne pathogens as well as showing a neutral or negative effect on plant pathogens. 4,5,11,16

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Understanding the metabolomics of soil microorganisms promises to enhance our knowledge of BC's influence in the microbial community. Metabolomics is the study of small molecules that make up a network of metabolic pathways, and it provides key insights into gene-environment interactions. 19,20 Mass spectrometry (MS) and nuclear magnetic resonance (NMR) techniques have been used extensively to study metabolomics. In the metabolomics field, MS is commonly used because of its high sensitivity, detection of many metabolites in a single measurement, and reliable metabolite identification through tandem mass analysis; however, MS is destructive to samples and the identities for a major fraction of detected metabolites are frequently not known.²¹ NMR spectroscopy is a useful tool to quantify the extracellular metabolites since it is nondestructive, highly reproducible, and provides a snapshot of what is occurring in the sample. 21-27 The extracellular metabolites in the media were monitored to determine the uptake of nutrients such as glucose, adenine, and asparagine, as well as the accumulation of fermentation products and other metabolites secreted by the bacteria. 22-24,26,27 This approach has the advantage of directly monitoring nutrients in the environment. NMR spectroscopy monitors and quantifies the more abundant compounds present in biological fluids and cell extracts, and it has the advantage that it can detect metabolites that are difficult to ionize by MS. 21,28-30

Additionally, a 1D proton NMR spectrum is often sufficient for resolving the components of a complex mixture with little overlap of the constituent compounds.³¹ The detected signals can also be directly analyzed by statistical tools to identify different subgroups (i.e., healthy vs diseased, untreated media vs treated media, etc.), eliminating the need to identify the chemical components of the sample.³² Alternatively, other statistical tools, like principal component analysis (PCA),³³ can be used to identify chemical differences when individual compounds are being investigated. Thus, NMR-based metabolomics is a mature and useful approach for understanding the behavior of bacteria in the presence of BC.

In this study, we treated RPMI, a complex, well-defined bacterial growth media, with BC and AC and compared how *E. coli* grew in treated media versus untreated media. While primarily useful as a model organism, *E. coli* survives readily in soil³⁴ and is associated with fertilization of crops.³⁵ We hypothesized that BC and AC would have similar effects on bacterial growth compared to untreated RPMI media. In addition to comparing the different metabolomic profiles, statistical analysis was used to highlight differences and similarities between the media types, and the overall effect on bacterial growth rates was measured. BC and AC surfaces were characterized to interpret the difference in the cell growth rates. Our results provide insight on how BC can support the development of microbiota, and they reveal a mechanism by which BC can influence microbial populations in the soil.

■ MATERIALS AND METHODS

Treatment of Media and Bacterial Preparation. *E. coli* (BL21(DE3) with an ampicillin-resistance gene) was grown in RPMI 1640 medium with L-glutamine (Fisher Scientific, Pittsburgh, U.S.A.) for the overnight stock culture and bacterial growth. Biochar Supreme, LLC (Everson, WA) generously provided Douglas fir, ultra-rinsed Black Owl biochar (BC). To prepare this char, auger-fed, Douglas fir wood chips were placed into an air-fed updraft gasifier at 900–1000 °C for 1–

10 s. 36,37 The particles were collected and washed with water several times to remove fine particulates and impurities before drying at room temperature. 36,37 The biochar was sieved to a particulate size of 75–150 μ m. This char has been characterized extensively in previous studies, $^{36-38}$ and our observations of point of zero charge (PZC), scanning electron microscopy (SEM), and surface porosity were consistent with these prior investigations. This confirms that the biochar used was consistent between separate batches of starting material.

Cultures were inoculated from a fresh plate of Luria—Bertani (LB) media, plus 100 μ g/mL ampicillin. The overnight stock culture was shaken for 12 h at 200 rpm at 37 °C. The final optical density (O.D.) (Nanodrop One/One^c, ThermoFisher Scientific, Pittsburgh, U.S.A.) of the overnight stock culture at 600 nm was 0.855. The O.D.₆₀₀ was measured to assess the rate of bacterial growth for the overnight stock culture, BC- or AC-treated media, and untreated RPMI media. To 25 mL of RPMI media, 5% (by weight, 1.25 g) BC or 5% (1.25 g) Norit Activated Carbon (AC) was added. The overnight stock culture was then divided equally into the three different media (RPMI, BC, and AC media) to begin bacterial growth. In this set of experiments, bacteria grew in the presence of BC, AC, or untreated RPMI media throughout the time course (24 h) of each experiment.

An additional set of experiments was performed where BC or AC was removed from media before inoculation (Method S1). For these experiments, 10% (by weight, 3.0 g) BC or 10% (by weight, 3.0 g) AC was added to 30 mL of RPMI media. The weight-by-mass percentage was doubled (from 5 to 10%) to allow more of the media to adsorb to the particles before filtering BC or AC from RPMI media. Doubling the weight-bymass percentage did not alter the growth rate of E. coli, as both growth rates for treated media were similar between the two experiments. The media was placed in a floor shaker (Innova 43 Incubator Shaker Series, New Brunswick Scientific, Edison, NJ) at 200 rpm at 25.0 °C for 24 h. This allowed ample time for BC and AC to adsorb metabolites from the media in the absence of bacterial growth. During this time, no significant growth was observed based on O.D. measurements. After 24 h of shaking, the media was filtered through a 25 mm poly(ether sulfone) (PES), 0.2 μ m sterile syringe filter. The pH for each media was measured and adjusted to have a starting pH of 7.5. The overnight stock culture was divided equally into the three filtered media to begin the bacterial growth, and experiments were performed as those described above.

Bacterial Growth. To begin the bacterial growth, the overnight stock culture was diluted into the media. From the 50 mL overnight stock culture, 15 mL portions were transferred into separate flasks: one containing 1.25 g of BC in 25 mL of RPMI media, one containing 1.25 g of AC in 25 mL of RPMI media, and a control sample of 25 mL of RPMI with no treatment. The flasks were swirled to mix BC and AC into the media. For the filtered experiments, 15 mL of the 50 mL overnight stock culture was transferred into the separate flasks similarly described for the unfiltered experiments. In addition, 2 mL of media was transferred to a sterile microcentrifuge tube to represent the sample before treatment. The media was returned to the shaker to continue incubation. The average starting O.D.₆₀₀ value for each flask was 0.285.

An aliquot of each sample was taken every 30 min for the first 3 h and every 3 h for the next 9 h. A final aliquot was taken at 24 h. For each time point, 2 mL of sample was transferred to sterile microcentrifuge tubes. All samples were placed on ice to

slow the bacterial growth, allowing BC and AC to sediment. After 5 min on ice, most of BC and AC had sedimented to the bottom of the tube so that 300 μ L of the sample could be carefully transferred to a second sterile microcentrifuge tube. The sample was then diluted with 700 μ L of filtered RPMI media to allow for O.D.₆₀₀ measurements in the 0.1–1.0 range. This was necessary to prevent BC and AC from interfering with measurements of optical density. The sample was carefully mixed using the pipet tip and then transferred to a 1 cm cuvette. The O.D.₆₀₀ was measured for each time point to monitor the bacterial growth.

NMR Sample Preparation. The remaining aliquot after O.D. measurement for each sample was centrifuged at 21000 g for 12 min in preparation for NMR spectroscopy. After centrifugation, the samples were sterile filtered and 400 μ L of the supernatant was transferred to another clean, sterile microcentrifuge tube. A 200 μ L portion of 200 mM phosphate buffer at pH of 7.0 with 1.000 mM trimethylsilylpropanoic acid (TMSP) in 50% D₂O was added to the supernatant as the internal NMR reference and lock solvent. For NMR analysis, 550 μ L of the sample was transferred to a clean 5 mm NMR tube (Wilmad Lab Glass, U.S.A.).

NMR Spectroscopy. The ¹H NMR spectra were obtained at 600 MHz and at a temperature of 298 K on a 600 MHz Bruker Avance III cryoprobe-equipped NMR spectrometer. A 1D-NOESY (noesypr1d) pulse sequence was used to obtain the spectra. The pulse sequence used presaturation on the water signal during the 4 s relaxation delay (d1) and a 50 ms mixing time (d8) to suppress the water signal at 4.75 ppm (o1p). A total of 64 free induction decays (FIDs) were collected with a 20 ppm spectral width. The TMSP reference peak was used as an internal standard.

The TMSP peak for each spectrum was calibrated to 0 ppm, and then peaks were manually phased and baseline corrected. Compounds in the processed spectra were identified and quantified using AMIX-Viewer v3.9.14 software (Bruker Biospin GmbH). Peaks were identified by comparison to a library of pure, 3.000 mM standard compounds. In addition to serving as a chemical shift reference, the TMSP peak was also used as an analytical standard. The library was created inhouse, and it contained a total of 56 common metabolites. These metabolites were not targeted to a specific pathway but instead were chosen to focus on glycolysis, the tricarboxylic acid cycle, and soluble amino acids. The library contained chemical shift data and peak integrations for each compound, although some peaks in crowded spectral regions were excluded from analysis. The output from AMIX was a listing of concentrations for each of the 56 compounds at a given time point. Concentrations versus time were collected to form a large data matrix for AC-treated, BC-treated, or untreated RPMI media, and this matrix was then used for statistical analysis, described below.

BET and SEM Analysis. The Brunauer–Emmett–Teller (BET) analysis was used to measure the adsorption of nitrogen gas on the surface area of BC and AC. BET surface areas, pore volumes, and averaged pore diameters of BC and AC were examined on a Micromeritics TriStar II Plus 3030 surface area analyzer using a nitrogen adsorption isotherm at 77 K.

Scanning electron microscopy (SEM) was used to determine BC and AC morphology. Sample preparation for SEM was prepared as described above; however, reagent amounts and volumes were scaled down for microscopy. 150 mg of BC or AC was added to 3 mL of RPMI media in a sterile 15 mL

conical tube. This set of samples was placed in the shaker for 24 h with no bacteria to characterize BC and AC morphologies in RPMI media alone. The 10 mL overnight stock culture had a final O.D.₆₀₀ of 0.732. A 500 mg portion of BC or AC was added to 10 mL of RPMI media, and then the overnight culture was equally divided into the three media types. The starting O.D.₆₀₀ value for each media was 0.224. The O.D.₆₀₀ after 24 h of growing was 0.815 for RPMI media, 0.668 for BCtreated media, and 0.375 for AC-treated media. Each sample was vacuum filtered using a Buchner funnel. The samples were each placed into a sterile glass vial and dried in the oven for 3 days at 67 °C. To analyze the samples on SEM, carbon tape was placed onto a specimen holder, and the tape was gently coated with an even layer of BC or AC. Excess particles were removed using compressed air. Once all the samples were mounted onto the specimen holder, the holder was placed under a vacuum and coated with 20 nm of Pt in a sputter coater. The surface morphologies of BC and AC were then examined by SEM using a JEOL JSM-6500F FE-SEM at 5 kV.

Statistical Analysis. Statistical analysis was conducted using MetaboAnalyst.³⁹ The average metabolite concentrations of three replicates for each sample was determined for each incubation time and media type. MetaboAnalyst normalized the data with Pareto scaling, which is mean centered and divided by the square root of the standard deviation of each variable. Pareto scaling was used to reduce the influence of the intense peaks and emphasize the weaker peaks to interpret the biological relevance.³³ Once the data were normalized, multivariate analysis (PCA) and cluster analysis (heatmaps) were used to perform data analysis.

For PCA analysis, the data set was divided into principal components (PCs) to identify the statistical differences between the classes. The first component (PC1) was the linear combination of the original predictor variables that captured the maximum variance in the data, whereas the other components (second, third, fourth, etc.) were the linear combination that captured the remaining variance in the data set and were orthogonal to the first component. 40,41 A biplot was used to identify metabolites that were significantly different between RPMI, BC, and AC samples. 42 The metabolites with the largest variance were easily identified in the biplot due to a larger vector magnitude. Significant metabolites (p < 0.05) were determined using a two-sample ttest with unequal variances. The metabolites identified as statistically significant were colored yellow in the biplot; insignificant metabolites were colored gray.

A heatmap of normalized, relative concentrations was used to visualize the relationship between two sets of data by using a color scale. The heatmap employed a hierarchical clustering analysis of the metabolites that was based on Pearson's correlation and Ward's linkage. Pearson's correlation determined the linear correlation between the variables, and Ward's linkage algorithm was based on the Euclidean distance that minimized the sum of the squares of any two clusters. The heatmap used a color scale to represent the variance in concentration for each metabolite in the sample.

RESULTS AND DISCUSSION

Initial Adsorption of Metabolic Compounds. The adsorptive properties of BC and AC make them ideal as soil additives in agriculture because BC and AC enhance the soil's physical and chemical properties by increasing the water holding capacity, retaining nutrients, enhancing CEC, increas-

ing surface area, and influencing hydrophobicity. ^{1,3,10,15} Variations in BC and AC feedstock and pyrolysis conditions produce materials with varied adsorptive properties. This allows chars to be tailored to enhance specific soil conditions. Varying the adsorptive properties of BC is an effective strategy for removing specific chemicals in soil and water. Woody chars typically have a higher CEC, allowing them to be used as a carbon sequestration agent as well as a soil amendment. ^{46,47} Woody BCs are typically used to remove metals, pesticides, and pharmaceuticals from soil and water. ^{46,48–50}

Both organic and inorganic BC components influence adsorption. S1 Chars that contain a higher percentage of ash and low carbon content are not as energy efficient and are less useful for adsorbing compounds. A7 BC and AC containing a higher carbon content, usually found in woody and plant materials, demonstrates an increased adsorption of water, metals, and metabolites relative to chars containing a lower carbon content. High pyrolysis temperatures effect the sorption of BC by increasing surface area, microporosity, and hydrophobicity, whereas the low pyrolysis temperatures remove inorganic and polar organic contaminants by oxygencontaining functional groups, electrostatic attraction, and precipitation. In this study, we used ultra-rinsed black owl biochar, a commercially available soil amendment manufactured from Douglas fir waste wood.

Both BC and AC adsorbed many metabolites when introduced into RPMI media (Figure 1) including amino acids and D-glucose. Figure 1 represents a few of the amino acids that were adsorbed by BC and AC that are not part of the

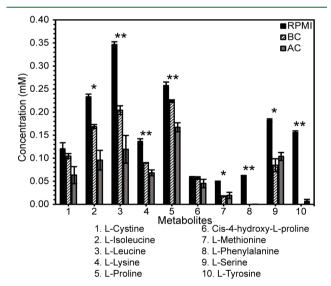


Figure 1. Initial metabolite concentration in media immediately after treatment with biochar or activated carbon. The initial concentrations of (1) L-Cystine, (2) L-Isoleucine, (3) L-Leucine, (4) L-Lysine, (5) L-Proline, (6) Cis-4-hydroxy-L-proline, (7) L-Methionine, (8) L-Phenylalanine, (9) L-Serine, and (10) L-Tyrosine in RPMI (control, solid black), BC (diagonally striped, black and white), and AC (dark gray) media 5 min after initial mixing. Error bars are the standard error of the mean of three independently prepared samples. One-way ANOVA with Tukey's HSD test was used to identify the significant differences between RPMI and either BC- or AC-treated media. A single asterisk (*) indicates that the RPMI concentration is significantly different from both AC and BC media at a *p*-value <0.05. A double asterisk (**) indicates a *p*-value <0.005. The differences shown between concentrations in AC- and BC-treated media were not statistically significant for initial adsorption.

tricarboxylic acid (TCA) cycle. One-way ANOVA was used to identify significant amino acids that were adsorbed after initial treatment with BC or AC. Five amino acids (L-leucine, L-lysine, L-proline, L-phenylalanine, and L-tyrosine) were strongly adsorbed by AC and BC (p < 0.005), suggesting a significant interaction with these compounds. L-isoleucine, L-methionine, and L-serine were less significant (p < 0.05) but still strongly adsorbed. Other amino acids, including L-lysine, and cis-4-hydroxy-L-proline, showed a decrease in concentration but were not statistically significant. Overall, the addition of BC and AC into the RPMI media introduced an environmental factor that altered the bacterial growth rate (Figure 2).

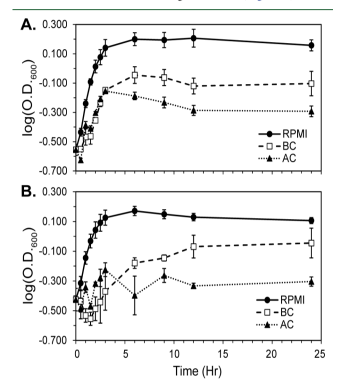


Figure 2. Growth curves of *E. coli* in RPMI media. Growth curves of *E. coli* in (A) unfiltered and (B) filtered RPMI media (black dots with solid black line), BC-treated RPMI media (open black squares with a dashed black line), and AC-treated RPMI media (black triangles with a dotted black line). In the filtered experiments, BC and AC are removed before growth occurs. Error bars are the standard error of the mean of three identically prepared samples.

E. coli Growth in RPMI vs Treated Media. E. coli was grown in RPMI media for the overnight stock culture up to 12 h and used as the starting culture with an initial O.D.600 of 0.285. RPMI media was treated with BC and AC to observe how their presence affected the bacterial growth rate. In unfiltered experiments, BC and AC remained in RPMI media during bacterial growth. In filtered experiments, BC and AC were filtered from RPMI media before bacterial growth. From the O.D.₆₀₀, it was observed that *E. coli* grew more readily in RPMI media with more abundant metabolites than treated RPMI media, likely indicating that the adsorption of nutrients by BC and AC influenced the bacterial growth rate (Figure 2). For the first 3 h, BC- and AC-treated media had a similar growth rate, whereas after 3 h, E. coli adapted to BC-treated media and reached a stationary phase for unfiltered AC-treated media (Figure 2A). The stationary phase for both RPMI and unfiltered BC-treated media was reached around 6 h. The

filtered BC- and AC-treated media showed qualitatively similar results as unfiltered media, although the initial O.D.₆₀₀ values were slightly noisier when media was filtered. In the filtered experiments, AC reached stationary phase faster than BC (Figure 2B), but the trend in final O.D.₆₀₀ values was identical to the unfiltered experiments. The slower growth rate seen in BC and AC was from the adsorption of metabolites that were needed for bacterial growth. For example, the starting concentration of D-glucose decreased when BC and AC were introduced into RPMI media. Furthermore, L-asparagine and L-glutamine also had a significant decrease in concentration once BC and AC were introduced into RPMI media.

Bacterial Interaction with BC and AC Particles. After confirming the effects of AC and BC on bacterial growth, we investigated the structural interaction of bacterial cells with BC and AC surfaces using scanning electron microscopy (SEM). SEM was used to measure the surface morphology of BC and AC without RPMI media, with RPMI media, and with *E. coli* in RPMI media. The surface of BC was observed to be flat and smooth while AC was porous and coarse (Figure 3 and Figure

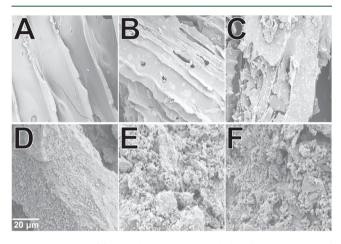


Figure 3. SEM of bacterial interactions with biochar and activated carbon. (A) Douglas fir, ultra-rinsed Black Owl biochar (BC), (B) BC in RPMI media, (C) BC in RPMI media with *E. coli*, (D) Norit activated carbon (AC), (E) AC in RPMI media, and (F) AC in RPMI media with *E. coli*. The surface morphologies were studied at 5 kV. The 20 μ m scale in (D) was used for all the other figures as well.

S1), agreeing with previous comparisons of AC and BC. 36,37 SEM images of BC showed a morphology change as RPMI and E. coli were introduced to the char, with the smoother microscopic texture of BC making it easier to observe the attachment of material to the surface (Figure 3A and Figure S1). When BC was introduced to RPMI media and E. coli cultures, there were small patches of adsorbed metabolites and bacterial cells visible on the surface of the char (Figure 3B,C). After treatment with bacteria, the biochar appeared to have numerous bacterial colonies (Figure 3C), and several of these colonies contained clusters of 3-4 1 µm sized cells. E. coli biofilms can take up to several days to form, 52 but Sugimoto et al. observed similarly sized clusters during the early formation of E. coli biofilms. 53' This suggests that biofilms may form on the flat surfaces present in biochar. The formation of biofilms is due to bacterial cells producing an extracellular polymeric matrix comprising of polysaccharides, proteins, lipids, and nucleic acids. 54 Each of these components would likely enhance attachment to charred biomass, which is known to contain not only hydrophobic interacting surfaces⁵⁵ but also

hydrogen bond donors/acceptors⁵⁵ and ionizable groups.⁵⁶ Importantly, most of these chemical features differ significantly in AC, which may make it less amenable to bacterial attachment.⁵⁴ The physical basis of surface attachment is influenced by many factors, including electrostatics and metal ion content,⁵⁷ and further studies are needed to understand why bacteria attach to some surfaces and not others.

The coarse surface of AC made it difficult to observe any adsorbed metabolites or bacteria (Figure 3, D-F). AC was observed to have a larger surface area than BC (Figure 3), and overall adsorption of metabolites was found to be higher (Figure 1). The absence of bacterial clusters, however, suggests that biofilms will form much more slowly on the surface of AC, if at all. The adsorption of metabolites on BC and AC introduced an environmental stress to the media, where bacteria had to compete for nutrients (Figure 2). It is possible that this stress factor significantly altered the bacterial growth, making biofilm formation more likely on BC surfaces than AC surfaces. As noted above, the surface charge of BC and AC strongly influences the adsorption of metabolites and bacteria. 54,57 Measurement of the point of zero charge (PZC) revealed that BC had a higher PZC than AC (pH $9.793 \pm 0.005 \text{ vs pH } 8.920 \pm 0.003, \text{ Figure S2 and Method}$ S2). The higher PZC for BC likely enhances the capacity to adsorb charged species by favoring electrostatic interactions with metabolites and cell surface components at neutral pH, where our studies were performed.⁵

The pore size of \overline{BC} and \overline{AC} can potentially modulate adhesion of bacteria to the material. Previous studies have suggested a relationship between pore size and bacterial adhesion. An overly large or small pore size will diminish adhesion due to pore curvature being too small for the bacteria to adhere (large pore size) or due to physical exclusion from the pores (small pore size). Brunauer, Emmett, and Teller (BET) analysis was used to estimate the surface area and average pore size of BC and AC. This analysis does not quantify surface curvature but rather measures the surface areas of solids by physical adsorption of N_2 gas molecules.

BET analysis revealed that BC had a smaller average pore size than AC (15.0 \pm 0.1 Å vs 16.1 \pm 0.6 Å, respectively; Table 1). Moreover, AC has a higher surface area than the BC used

Table 1. BET Particle Size Analysis^a

	surface area (SA) (m^2/g)	total pore vol (cm³/g)	mean pore size (Å)
black owl biochar	510 ± 9	0.21 ± 0.01	15.0 ± 0.1
activated carbon	704 ± 16	0.42 ± 0.07	16.1 ± 0.6

^aStandard error of the mean of three independently prepared samples.

in this study. These observations are consistent with the fact that AC is able to adsorb more small compounds overall, suggesting that surface area and adsorption capacity play the primary role in modulating bacterial growth following treatment with BC or AC. Surface area and pore size may also play a secondary role in controlling bacterial attachment, even though the BET pore sizes are much smaller than even a single bacterial cell. This secondary role follows because the BET pore size will influence which cell surface components of the *E. coli* outer membrane are able to interact with the BC or AC surface. Additionally, it is known that selective adsorption of nutrients themselves are able to assist in bacterial

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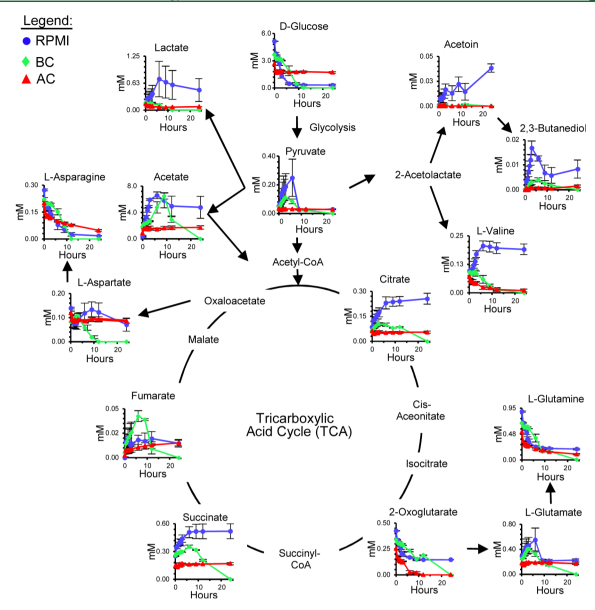


Figure 4. Tricarboxylic acid (TCA) cycle of *E. coli*. The extracellular metabolite concentrations of *E. coli* in RPMI (blue), BC (green), and AC (red) media arranged with respect to common metabolic pathways. Error bars are the standard error of the mean for three identically prepared samples.

accumulation to a surface.⁵⁸ Therefore, the surface characteristics of BC and AC on a molecular scale can potentially have a larger-scale influence on bacterial colonization. While it is difficult to correlate BET measurements with surface roughness,⁵⁹ the BET surface area approximately reflects the smoothness of the surface as visualized by SEM (Figure 3 A–C). Thus, it is possible that pore size in BC is more conducive to binding *E. coli* cell surface components. This would in turn make the smooth, flat surfaces of BC more easily able to accommodate bacterial attachment and colonization. Additionally, the BC used in this study is prepared by pyrolysis of wet biomass. This treatment is thought to increase surface area as pockets of steam and other volatiles rupture during pyrolysis. ^{36–38,46,50} It is possible that this wet pyrolysis treatment also plays a role in bacterial attachment.

Metabolite Excretion and Adsorption During Bacterial Growth. To investigate the effect BC and AC have on glucose metabolism, we examined the concentration depend-

ence of several key compounds in glycolysis and the TCA cycle. The TCA cycle is a good indicator of bacterial growth in the presence of external stressors, because it represents the major pathways for energy flux in the cell. Deglucose is one of the main carbon sources in RPMI media that was depleted and was shown to have an inverse-relationship with bacterial growth (Figure 2). The process of glycolysis rapidly converts Deglucose to pyruvate in *E. coli*, and both compounds can be quantified using NMR (Figure 4). Deglucose was completely depleted in RPMI and BC-treated media (an indication of bacterial growth), whereas the concentration of Deglucose in AC-treated media was only slightly reduced (minimal bacterial growth). When Deglucose is depleted as part of the TCA cycle, the concentration of byproducts like acetate and pyruvate increases.

Examination of the TCA cycle revealed several similarities and differences between cells treated with AC or BC (Figure 4). For example, acetoin, lactate, and L-valine showed a

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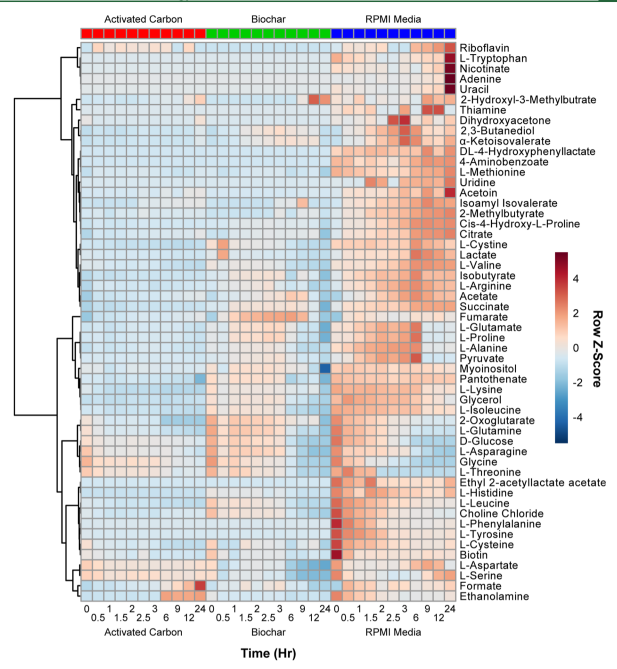


Figure 5. Heatmap of extracellular metabolites' relative concentrations versus time in unfiltered media. The row Z-score for each feature was used to color code the map: absent to low concentration metabolites were colored blue and high concentration metabolites were colored red. Ward's linkage algorithm ranked the metabolites in a clustering tree. The red boxes were labeled as AC-treated media as time increases from 0 to 24 h going left to right. The green boxes were BC-treated media, and the blue boxes were RPMI media. Data are the mean values of metabolite concentrations of three replicates after being normalized using Pareto scaling.

decrease in concentration for BC- and AC-treated media, whereas RPMI media had a higher concentration than treated media for these compounds. The citrate concentration in BC- and AC-treated media was fairly constant during the experiment, whereas in RPMI media, the concentration of citrate increased over the same time frame. L-glutamine, D-glucose, and L-asparagine decreased slowly in concentration for BC-treated media and RPMI media, whereas in AC-treated media the concentrations for these metabolites decreased more quickly. In the TCA cycle, 2-oxoglutarate for BC-treated media had a similar behavior compared to RPMI media. Fumarate, L-aspartate, L-glutamate, and acetate for BC-treated media

behaved differently compared to those for AC-treated and RPMI media. The patches of adsorbed metabolites on the surface of BC and AC seen in the SEM images may explain why we observed a difference in concentrations for a few metabolites in RPMI media (Figures 3 and 4).

The concentrations of the extracellular metabolites were also visualized using a clustered heatmap (Figure 5, Figure S3, and Figure S5C). A heatmap graphically represents the change in metabolite concentrations versus time with a color scale being used to represent the row Z-score. This heatmap covers the complete list of 56 metabolites used in this analysis, including the glycolysis-related compounds already discussed.

In general, AC-treated media had very low concentrations for most of the metabolites in comparison to BC-treated and RPMI media. AC adsorbed more metabolites than BC, as evidenced from the first time point (0 h) before the cultures were inoculated. Interestingly, adsorption profiles for AC and BC differ significantly, with compounds like L-isoleucine, choline chloride, L-leucine, and 2-oxoglutarate adsorbing more strongly to AC. Over time, however, the metabolic profiles for BC- and AC-treated media were seen to be similar when compared to untreated RPMI media. For example, glycine and L-asparagine in BC- and AC-treated media had very similar trends in decreasing concentration compared to untreated RPMI media.

Several compounds, including acetate, L-glutamate, L-proline, L-alanine, and pyruvate, peaked in concentration in RPMI media at 6 h. After this, they rapidly decreased as key metabolites, including D-glucose (a carbon source) and L-glutamine (a nitrogen source), were utilized by *E. coli* cells. Given that cells replicate more rapidly in RPMI media, this peak may reflect the point at which the initial nutrients are depleted, and cells must adapt to a more competitive environment. In AC- and BC-treated media, the overall nutrient concentrations are already reduced, and this effect becomes less pronounced. It is weakly visible in BC-treated media, but such an effect is not observed in AC-treated samples.

The heatmap clusters into two general classes of compounds: those that increase in concentration in RPMI media and those that decrease over time. Smaller clusters reflect differences in behavior of those compounds in AC- and BC-treated media. Across all three types of media, the concentrations were generally observed to correlate, although several exceptions to this were observed. The increasing concentration of fumarate in BC-treated media, already discussed above, is one example of this. The increase of formate and ethanolamine in AC-treated media at longer time points is another. For these two compounds, the concentration remains approximately constant in BC-treated media, and in RPMI, the concentration is observed to decrease. It is not clear why these compounds behave differently in media treated with AC. Importantly, the compounds do not appear to form clusters based on simple chemical properties such as charge and/or hydrophobicity: this suggests that the concentration dependence originates from changes in bacterial behavior and not selective surface interactions. To investigate chemical trends among metabolites, compounds in the heatmap (Figure 5) were rearranged based on their chemical properties. Amino acids, vitamins, sugars, constituent bases, and nucleosides were grouped in an alternative organization of the heatmap (Figure S3). No significant trends were observed within each group, suggesting that the concentration dependence arises due to metabolic changes and not the adsorptive properties of AC or BC, which should be similar for similar compounds. This hypothesis is further supported by experiments with filtered media, as described below.

Analysis of the initial adsorption of metabolites revealed that both BC and AC depleted some nutrients needed for bacterial growth. Overall, BC adsorbed a lower quantity of metabolites than AC. The filtered and unfiltered media had similar characteristics in metabolite adsorptions. PCA analysis had very similar results for unfiltered and filtered media, whereas BC-treated media had similar characteristics to AC-treated and RPMI media (Figure 6 and Figure S5). The heatmaps overall

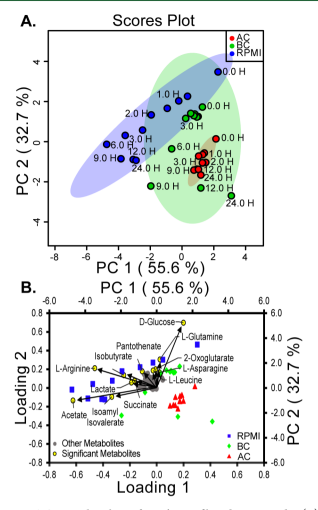


Figure 6. Statistical analysis of *E. coli* in unfiltered RPMI media. (A) Principal component analysis (PCA) scores plot on *E. coli* growth in unfiltered media used to identify statistical differences between each media type. The shaded oval represents the 95% confidence limit for each media type (RPMI, blue circles; BC-treated media, green circles; AC-treated media, red circles). Time points for each hour are labeled, but points at half hour intervals are not labeled for clarity. (B) Metabolites in the loading plot (left and lower axes) overlaid on the PCA biplot (right and upper axes) for unfiltered media. Yellow dots represent the significant metabolites (p < 0.05), i.e., those making the largest contribution to each principal component (RPMI, blue squares; BC-treated media, green diamonds; AC-treated media, red triangles).

had similar metabolic profiles between filtered and unfiltered media (Figure 5 and Figure S5). For unfiltered samples, the adsorbed metabolites can potentially be returned to the media, leading to persistent differences through the growth cycle. However, filtration removes AC or BC, and those samples permanently lose the initial concentration of adsorbed metabolites.

Since filtered and unfiltered samples both have similar profiles, the desorption and resorption of metabolites to the surface during the experiment do not appear to alter the bacteria growth significantly. Rather, it is the initial adsorption of metabolites to the char's surface that seems to most dramatically influence the bacterial growth rate. This is significant, as it suggests that supplementing BC with several key nutrients may be an effective strategy for offsetting any adverse effects in soil bacterial growth rates.

Statistical Analysis of Metabolites Present After Growth. The metabolic data were further analyzed using statistical analysis provided by the MetaboAnalyst suite. PCA was used to determine statistically meaningful differences between three media types (score plot shown in Figure 6A and Figure S5A). Removal of BC or AC by filtration before inoculation made essentially no difference in the principal components (Figure 6A). Both filtered and unfiltered growth showed a clear differentiation between RPMI and AC-treated media as determined by the ellipses representing 95% confidence intervals (Figure 6A and Figure S5A). However, the behavior of BC-treated media showed similarities to both AC-treated and RPMI media, demonstrated by its overlap on the scatter plot.

The first and second components for unfiltered and filtered media accounted for the largest variation between the media. When combined, these two components account for 88.3% of the variance in the data, with the first two components (PC1 and PC2) accounting for 55.6% and 32.7% of the variance, respectively. This observation that BC-treated media retains characteristics of AC-treated media as well as untreated RPMI media is consistent with qualitative comparisons previously noted above. Since AC adsorbs more metabolites than BC, AC can be viewed as a more aggressive amendment, whereas BC retains some of the adsorptive capabilities of AC while promoting conditions favorable for bacterial growth.

An examination of the scatter plot (and biplot) reveals that the early time points of BC-treated media cluster near RPMI media, whereas later time points cluster with AC-treated media. After approximately 6 h, the data points corresponding to BC-treated media diverge from the cluster of early time points on the scatter plot and trend toward the cluster of AC-treated media. It is this multimodal behavior that gives rise to a substantially larger confidence interval for BC-treated media (Figure 6A and Figure S5A). On the basis of the loadings (Figure 6B and Figure S5B), D-glucose, L-arginine, and acetate contribute significantly to the second principal component, and the depletion of these compounds leads to the transition from one cluster to the other on the biplot.

The component loadings identify which metabolite concentration differences contribute to PC1 and PC2 (Figure 6B and Figure S5B).³⁹ The metabolites with the largest variance in the metabolic profile were identified and labeled as significant metabolites (yellow dots, Figure 6B and Figure S5B). The magnitude of the vectors for the significant metabolites quantifies the change in the concentrations over time for each principal component. Therefore, the longer vectors represent a larger relative change in metabolite concentrations and the shorter vectors represent a smaller change in metabolite concentrations. These largest contributors are the metabolites that deviate in behavior from the insignificant metabolites, which are clustered near the origin of the loading plot (Figure 6B, gray dots). The size of each metabolite's loading is based on its trend during the growth. For example, D-glucose decreased in concentration over time while acetate increased in concentration, resulting in these metabolites being on opposite ends of loading 1 and loading 2 (Figure S4). The largest contributors to the first loading were acetate, L-arginine, isoamyl isovalerate, and D-glucose, whereas the second loading was dominated by acetate, D-glucose and L-arginine. These data indicate that, following treatment with BC or AC, significant differences are observed in the extracellular metabolite concentrations of these compounds. BC and AC alter the

media in complex ways leading to significant differences in several extracellular metabolite concentrations, and these changes cannot be explained by the adsorption of simply one or two compounds. Moreover, some metabolite concentrations remain unaffected or show little discriminatory power between samples (Figure 6B, gray circles). This suggests that treatment with AC and BC leaves many compounds unaffected during bacterial growth.

To summarize, based on our measurements, there were statistically significant differences between RPMI and ACtreated media, whereas BC-treated media shared characteristics with both RPMI and AC-treated media. This suggests that the favorable properties of BC, namely, its ability to retain nutrients, may not adversely affect the bacterial life cycle in the way that other adsorbents, such as AC, might. This is especially true at early time points, where extracellular metabolite concentrations for BC-treated cells were observed to behave similarly to cells growing in untreated media. Together, these results have implications for biochar when used as a soil amendment. BC's adsorption properties enhance soil fertility, add needed nutrients like carbon and nitrogen to the soil, and enhance soil microorganisms.⁴⁻⁷ Moreover, BC allows crops to grow larger and faster because BC enhances the soil's physical and chemical properties by increasing pH, aeration, water holding capacity, and adding nutrients; BC also adsorbs metabolites that alter bacterial growth. 3,11,13

This study begins to explain these effects at the level of the bacterial metabolome. Here, the behavior of metabolites demonstrates that BC can effectively adsorb certain compounds, similar to AC, while leaving others available for utilization by soil microbiota. It is this property that allows BC to exhibit the favorable properties of RPMI media during the early time points when bacteria are doubling rapidly. At the same time, BC is known to adsorb hydrophobic compounds and retain nitrogenous fertilizers, 11-13 making conditions favorable for plant growth. Combined, these properties create conditions that favor growth of both crops and soil microbiota. While BC can adsorb nutrients needed by microorganisms, as demonstrated by the initial depletion of D-glucose, oxoglutarate, and several other compounds, this effect appears to be minor compared to AC, where aggressive adsorption of these compounds leads to poor bacterial growth. Interestingly, the selective adsorption of metabolites could lead to conditions that favor one particular organism over another. While we have not explored the influence of BC on different populations of microbes, this work strongly suggests that BC can alter metabolite concentrations to favor one particular population of microbe. For example, the addition of BC or AC mainly had a positive effect on the abundance of mycorrhizal fungi, but in some cases, BC or AC limited the nutrients and the mycorrhizal fungi was hindered.⁶² It was also observed that the relationship between mycorrhizal fungi and BC sequesters carbon (mitigating climate change) to restore the ecosystem.⁶²

In summary, as a soil amendment, biochar's metabolic effects on soil microbiota have been largely uninvestigated. Here, we explored how biochar can alter the metabolomic profile of *E. coli*. There were significant differences between RPMI and AC-treated media, whereas BC-treated media shared characteristics with both RPMI and AC-treated media. This suggests a mechanism whereby BC can adsorb nutrients without adversely affecting soil microbiota. While cell culture does not represent soil conditions, the experiments capture the essential properties of adsorption by BC and AC, evidenced by

the concentration of metabolites. Efforts are underway to understand how BC influences the populations of soil biota.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.est.8b05024.

A detailed description of the methods used for treatment of media, SEM figures of the different biochar morphologies, the statistical analysis of bacterial growth in filtered media, and tables for the metabolite concentrations in the RPMI, BC-treated, and AC-treated media (PDF)

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Notes

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ABBREVIATIONS

AC, activated carbon; BC, biochar; NMR, nuclear magnetic resonance; TCA, tricarboxylic acid; PCA, principal component analysis; CEC, cation exchange capacity; LB, Luria—Bertani; PES, poly(ether sulfone); TMSP, trimethylsilypropanoic acid; FIDs, free induction decays; BET, Brunauer—Emmett—Teller; SEM, scanning electron microscopy

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