Reply to Reviewers’ Comments (bg-2017-509) by Weng et al.

Dear Associate Editor and Reviewers:

We have carefully prepared our response and revised our manuscript accordingly. Some major revisions made in the main text can be found in: P2 L1-15; P13 L6-22; P14 L1-5; P15 L1-18; P16 L5-14; P17 L3-5 (rephrased). Some minor revisions made in the main text are in: P6 L9; P8 L20-21; P9 L8-9; P9 L16; P12 L6; P14 L6-22; P15 L22; P16 L16. Please check the revised manuscript for details in the attachment. Two figures are added into the supplementary information.

Our rebuttal letter is lengthy, and we appreciate the patience of Associate Editor, Reviewer#1 and Reviewer#2. Abbreviation used here are: organic carbon (OC), short-range-order (SRO), soil organic matter (SOM), soil organic carbon (SOC)

Response to Reviewer#1:
We use an old (3500 years) pyrogenic OC-mineral consortium to demo the in-situ OC-mineral interplay in a relatively simple mineralogy context. In-situ evidence reveals abundant mineral nano particles, in dense thin layers or nano-aggregates/clusters, instead of crystalline, micron or clay-size minerals on or near OC surfaces, the key working minerals for C stabilization are essentially SRO minerals and/or poorly crystalline submicron size clay minerals. Supported by FTIR spectroscopic results, the studied OC is not merely in criss-cross co-localization with reactive SRO minerals. There is a significant degree of binding between OC and minerals. We also have other sets of unpublished 3-D tomography data for samples from various mineralogy background, and the minerals found associated with OC surface are coincidentally in line with nano particles/poorly crystalline minerals. ‘Mineral physical protection for OC stabilization may be more important than previously thought’ is not a solid conclusion, the statement is deleted.

Technology Progress for In-Situ Evidence:
This high resolution 3-D tomography tool we introduce has unprecedented resolution, is promising for new insight on in-situ organo-mineral association, the interior 3-D structure of microaggregates, and the fate of mineral nano particles including heavy metals in natural environment. Though some 3-D tomography research have been done on soil microstructure and porosity using X-ray micro-CT, the best achieved resolution only reaches tens of microns (Quin et al., 2014 (with biochar amendment, 70 microns); Kravchenko et al., 2015 (13 microns)), which exceeds the size of clay minerals (<2 microns), and not fit for capturing the microstructure of clay, and the submicron
assemblage of SRO minerals such as Fe oxyhydroxides. High resolution X-ray scanning and 3-D tomography is highly demanding in terms of technology, multidisciplinarity and big-data analysis (Lafond et al., 2015). Except for the adoption of high resolution X-ray objective, fidelity of 3D tomography relies on the accurate alignment of the 2D projections in correct three-dimensional positions. However, non-negligible mechanical imperfection of the rotational stages at nanometer level, or the thermal effects may significantly degrade the spatial resolution of reconstructed tomography. We’ve developed a markerless image auto-alignment algorithm for fast projection matching (Faproma, Wang et al., 2017) to surpass these difficulties, and accomplished accurate reconstruction of 3-D tomography down to nanometers.

**Carbon Stabilization:**

Soil organic C can be: (1) physically stabilized, or protected from decomposition, through microaggregation; or (2) intimate association with silt and clay particles in organo-mineral complexes; and (3) can be biochemically stabilized through the formation of recalcitrant SOC compounds (Six et al., 2002). Three main mechanisms of Soil organic matter stabilization are: (1) chemical stabilization, result of chemical or physiochemical binding between SOC and soil minerals (especially clay and silt in current opinions), aka organo-mineral complexation; (2) physical protection, which is predominantly at the microaggregate level, built on top of chemical organo-mineral complexation, and (3) biochemical stabilization, which is often experimentally defined, or equated to the nonhydrolyzable fraction (Stevenson, 1994; Christensen, 1996; Six et al., 2002). Experimentally to quantify C of different stabilized levels, a size/density fractionation approach with external force is routinely used for bulk soil, and three or more density fractions are divided: (a) free particulate OM; (b) occluded particulate OM; and (c) colloidal or clay associated OM, in the fine and dense fraction (Golchin et al., 1994, Sollins et al., 2009).

The stabilization of OC by the soil matrix is considered a function of the chemical nature of specific soil mineral fraction, the presence of multivalent cations, the presence of mineral surfaces capable of adsorbing organic materials, and the micro architecture of the soil matrix (Edwards and Bremner, 1967). The degree and amount of protection attributed by each mechanism depends on the chemical and physical properties of mineral matrix, the morphology, chemical nature and structure of organic matter (Baldock and Skjemstad, 2000). However, each mineral’s unique capacity to stabilize organic C is rarely recognized in bulk sample analysis, or within size/density fraction. Thus, quantifying the protective capacity of a soil requires a careful consideration of all mechanisms of protection and the implications of experimental procedures.
It is found that accumulation and subsequent loss of organic C were largely driven by changes in the millennial scale cycling of mineral-stabilized carbon (Torn et al., 1997, *Nature*). The protection of SOC by silt and clay particles in soils is well established for a positive relationship between old/stabilized organic C and the content of silt and/or clay in fine and dense fraction (Sorensen, 1972; Feller and Beare, 1997; Hassink, 1997; Trumbore, 1997). In addition to the clay content, clay type (2:1 vs 1:1 clay vs allophanic/short-range-order (SRO) minerals) has strong influence on the stabilization of organic C (Torn et al., 1997). Yet little is known about how spatial and temporal variation in soil mineralogy (especially type, phase, reactivity) controls C long term stabilization in terms of quantity and turnover (Oades, 1994, Torn et al., 1997). Quantification of interactive relationships is imperative, measurement and information using 2-D and 3-D quantitative tool will have great benefit in understanding soil C and aggregate dynamics (Six et al., 2004).

Very limited information is available on the in-situ distribution of clay-size or reactive minerals in soil, and subsequent organo-mineral micro assemblage in soils. For example, a substantial parts of mineral surfaces are considered likely not covered by organic matter (Ransom et al. 1998; Arnarson and Keil 2001; Mayer and Xing 2001; Kahle et al. 2002). Recently, nano scale two-dimensional mapping using Nano SIMS generated direct evidence, revealing that less than 19 percent of the clay-sized mineral surfaces co-localize with OC (Vogel et al., 2014, *Nature Communication*). Views of carbon sequestration in soils and the widely used carbon saturation estimates have been revolutionized to recognize likely only a limited proportion of the clay-sized surfaces contribute to OM sequestration in real soil environment. However, this research could underestimate the actual contact surface area between OC and mineral due to the limitation of sample treatment and two-dimensional approach. Exactly which reactive mineral in the clay-sized fraction interact with OC remains unknown.

The cornerstone research done by Torn et al. (1997) revealed a positive relationship between non-crystalline minerals and organic carbon in soils through the climate gradient. In couple with extraction using ammonium oxalate and/or sodium dithionite-citrate-bicarbonate (DCB), SRO minerals, which may only be a small portion in the fine and dense fraction, are found of great importance for C stabilization, and further discovered preferentially associated with aromatic/lignin-like C (Kaiser et al., 2002; Eusterhues et al., 2005; Kleber et al., 2005; Mikutta et al., 2005, 2006, 2007; Rasmussen et al., 2005; Schneider et al., 2010; Cusack et al., 2012). The SRO minerals are known of high reactivity due to small size, large surface area and charges, their encounter with reactive OC surface may develop strong organo-mineral complexation (Eusterhues et al., 2008, 2011). Yet, the actual key mineral player (micron size clay vs nano scale SRO minerals) for interaction with OC, and their spatial and temporal variation is not
known in real soil environment (Vogel et al., 2014). On the other hand, a three-dimensional functional view of carbon turnover dynamics at the microscale has gained ground (Kleber et al., 2018), which consider a multitude of largely independent microreactors within the soil. For example, anaerobic microsites are discovered of great importance for C stabilization (Keiluweit et al., 2017). New perspectives call for observational evidence on OC-mineral micro assemblage in details at nano scale.

Using high resolution 3-D tomography and synchrotron-based spectroscopic approach, we unveil:

i) There is a high heterogeneity within OC-mineral consortium, including many nano/submicron size mineral particles, porosity and microsites.

ii) Most of the particulate OC surface is coated by minerals.

iii) The minerals on the OC-mineral interplay surface are essentially nano size SRO minerals, and or submicron poorly crystalline clay minerals, which are highly reactive and may have large surface for C absorption.

iv) Very few micron size/clay-size, euhedral/crystalline minerals are observed.

v) The 3500-year-old pyrogenic OC is highly aromatic (1596 and 1386 cm⁻¹), when at the same time has highly reactive carboxyl functional group (1706 cm⁻¹).

vi) Broad FTIR bands indicate a significant degree of organo-mineral association between OC and SRO minerals, which are not just criss-cross co-localization.

vii) The OC surface is often coated by dense and thin layers of mineral nano particles, pointing to likely mineral absorption.

viii) Within regions microns from particulate OC surface, abundant mineral nano particles develop nano-aggregates/clusters, pointing to likely OC-mineral co-precipitation.

Our results suggest:

i) The coated particulate OC surfaces by SRO mineral nano particles are stabilized.

ii) Minerals physical protection contributes to the long term persistence of OC (3500-year-old) in the sub-tropical environment.

iii) Substantial mineral surfaces are likely available for C stabilization.

iv) Large amount of nano-aggregates/clusters likely interact/co-precipitate with relatively free OC, which is difficult to prove directly.

v) Current electron microscopy/spectroscopy/fractionation approach focusing on clay-sized mineral may lead to underestimation of C stabilization attributed by mineral physical protection.
We propose:

i) High resolution 3-D tomography is a powerful, suitable tool for nailing down the correct key working minerals for C stabilization.

ii) Two-dimensional TEM may be applicable in revealing nano particles, amorphous/poorly crystalline minerals, on certain focal plane, but not for a whole view.

iii) XRD is powerful for detecting crystalline minerals but very poor in resolving SRO minerals, cautions should be taken in sample preparation and mineralogy pattern interpretation.

iv) More research should be done to prove: i) whether it is a general phenomenon that the minerals interact with OC are essentially nano particle SRO minerals and or poorly crystalline submicron size clay minerals, ii) whether the major mineral for C stabilization is nano size SRO minerals instead of clay-sized minerals.

v) Perspective on SOM dynamics may have to take account the role of SRO minerals into modelling parameters, instead of drawing conclusion mainly on clay type and content (Kleber et al., 2010).

Carbon Models and Carbon Saturation:

Most current models of SOM dynamics try to describe soil heterogeneity by defining several pools, typically three to five, experimentally applicable with supposed different intrinsic decay rates. Meanwhile, first order kinetics are assumed for the decomposition of different conceptual pools (Parton, et al., 1994; Paustian, 1994; McGill, 1996), which infers equilibrium C stocks are linearly proportional to C inputs (Paustian et al., 1997). These models predict that soil C stocks can theoretically be increased without limit, provided that C inputs increase without limit, leading to no assumptions of soil C saturation. Such models have been largely successful in representing SOM dynamics under some studied conditions and management practices, and native soils (e.g. Parton et al., 1987, 1994; Paustian et al., 1992; Powlson et al., 1996). Notably, the representation of the model pools (or quality spectrum) is primarily conceptual in nature (Christensen, 1996; Elliott et al., 1996; Six et al., 2002). The individual pools are generally only loosely associated with measurable quantities obtained with existing analytical methods. The notion of carbon saturation is developed on the assumption that mineral physical protection could be limited by soil physiochemical characteristics such as silt and/or clay content, microaggregate, and surface area (Kemper and Koch, 1966; Hassink, 1997). Six et al. (2002) proposed a conceptual model which use the physicochemical characteristics inherent to soils to define the maximum protective capacity of different SOC pools, which may limit the increases in C sequestration with
increasing organic residue inputs. This conceptual model includes four measurable pools: (1) a biochemically-protected C pool; (2) a silt and clay-protected C pool (less than 53 micron organomineral complexes); (3) a microaggregate-protected C pool (53 to 250 micron aggregates); and (4) an unprotected C pool. Each pool is supposed to have its own dynamics and stabilizing mechanism, which is in turn hypothesized to determine a level at which soil C becomes saturated. The silt and clay protected C pool, which is protected by association with the mineral particles, is defined as hydrolysable fraction and the biochemically-protected C pool is experimentally defined as nonhydrolyzable fraction. Biochemical stabilization is understood as the stabilization of SOM due to its own chemical composition (e.g. recalcitrant compounds such as lignin and polyphenols) and through chemical complexing processes (e.g. condensation reactions) in soil. The unprotected C pool is considered labile, an important nutrient source, which is very sensitive to management practices. Interestingly, the hierarchy for carbon stabilization based on protection capacity/level, from low to high has been ranked as: silt + clay protected pool, microaggregate protected pool, biochemically protected, and non-protected pool on the top to cap the maximum carbon saturation.

Six et al. (2002) pointed that there are major gaps in knowledge and proposed a research priority in terms of soil C saturation on mechanistic level, if it exists, especially for unprotected and biochemically protection pools.

As an outsider, we share a few points:

i) The heterogeneity of microaggregates is little understood and minimal quantitative data is available.

ii) The unprotected pool actually involves some mechanism separation and dispersion, which is not truly unprotected.

iii) Six et al. (2002) shared our thought that non-hydrolyzable fraction is not only biochemically stabilized but is also partially stabilized by association with clay and silt particles, and the silt and clay protected pool could also be partially stabilized by incorporation in microaggregates.

iv) The aged pyrogenic OC (3500-year-old) in our study is not only biochemically stable, but also partially stabilized by association with submicron minerals.

Reviewer #1 suggested some interesting research on biochemically stabilized organic compounds free of minerals, which appears a large expansion to very solid organic geochemistry. Many models for SOM dynamics indeed incorporate a huge ‘inert’ but relatively uncharacterized pool, which may include biochemically stabilized organic compounds free of minerals, or not. Perspective on carbon stabilization may have to expand from a time scale of centennials and millennials to an even longer geological
time scale in current C saturation models. Derenne et al. (1991) carried out research on biochemically stabilized organic compounds free of minerals, especially on kerogen/fossil-like organic compounds, in the perspective of kerogen/fossil biochemical transformation, condensation and formation. It takes millions of years for the formation and transformation of kerogen/fossil-like organic compounds. We suggest research on biochemically protected pool also expand into black carbon/pyrogenic organic carbon, which are biochemically stable due to their high aromaticity, have similar chemistry and structure with kerogen and coal, and can persist over millennia under natural exposure. Long term abiotic/biotic degradation, aerobic/anaerobic alteration can render development of reactive surface functional groups (Liang et al., 2006), and lead to consequent mineral physical protection and C stabilization over millennia, even in the topical environment (Liang et al., 2008).

Response to Reviewer#2:
We thank Reviewer #2 for recognizing our work in methodology development and contribution to the understanding of C stabilization. In-situ evidence reveals abundant mineral nano particles, in dense thin layers or nano-aggregates/clusters, instead of crystalline micron/clay-size mineral on or near OC surfaces. The key working minerals for C stabilization are essentially nano size SRO minerals and/or poorly crystalline submicron size clay minerals. Inspired by reviewer’s suggestions, efforts have been taken to identify micro assemblage features which indicate adsorption or co-precipitation in our natural OC-mineral consortium using TXM images. In the all-depth 2-D X-ray absorption-contrast image (Fig. S2), sheet-like mineral coating is observed on OC surface, which is a dense and thin layer, pointing to likely a high level of physical protection. We propose such texture originate from absorption. Another distinct texture is recognized as OC-mineral nano-aggregates/clusters, with either minerals in the core and OC around (Fig. S2), or vice versa (Fig. S3). This type of texture indicates possible OC-mineral co-precipitation at microsites. Many clusters of various shapes are observed (Fig. S3). However, to fully explore this long-standing question on the structural development of OC-Fe mineral assemblages attributed by absorption vs co-precipitation, we suggest future research with carefully designed samples (such as distinct lab-made OC-Fe consortiums) and enough 3-D tomography observations are needed for reliable evidence and answers.
We’ve added Figures S2 and S3 into the supplementary information, please check for details, thank you.
Mineral physical protection and carbon stabilization in-situ evidence revealed by nano scale 3-D tomography

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Abstract

An approach for nano scale 3-D tomography of organic carbon (OC) and associated mineral nano particles was developed to illustrate their spatial distribution and boundary interplay, using synchrotron-based transmission X-ray microscopy (TXM). The proposed 3-D tomography technique was first applied to in-situ observation of a lab-made consortium of black carbon (BC) and nano mineral (TiO₂, 15 nm), and its performance was evaluated under dual-scan absorption contrast and phase contrast modes. Then this novel tool was successfully applied to a natural OC-mineral consortium from high mountain soil at a spatial resolution down to 60 nm, showing the fine structure and boundary of OC, distribution of abundant minerals at nano size, and in-situ 3-D organo-mineral association. The stabilization of aged natural OC was found mainly attributed to
the physical protection of **nano size** Fe-containing minerals (Fe oxyhydroxides including ferrihydrite, goethite, and lepidocrocite), and the strong organo-mineral complexation. In-situ evidence reveals abundant mineral nano particles, in dense thin layers or nano-aggregates/clusters, instead of crystalline clay-size minerals on or near OC surfaces. The key working minerals for C stabilization are essentially reactive short-range-order (SRO) mineral nano particles and/or poorly crystalline submicron size clay minerals. The sorption of OC to SRO mineral (or vice versa) through organo-mineral multiple complex bonds could occupy and consume their respective reactive surface sites, leading to enhanced stabilization. The ubiquity of abundance of mineral nano particles on OC surface, and their heterogeneity in natural environment could have been seriously underestimated by traditional study approach. Our in-situ description of organo-mineral interplay at nano scale provides direct evidence to substantiate the importance of mineral physical protection for OC long term stabilization. **Mineral physical protection for OC stabilization may be more important than previous understanding.** This high resolution 3-D tomography tool is promising for new insight on the interior 3-D structure of micro-aggregates, in-situ organo-mineral interplay, and the fate of mineral nano particles including heavy metals in natural environment.

**Introduction**
Mineral association with organic carbon (OC) may be an important stabilization mechanism for carbon long-term sequestration, yet little is known about their in-situ interplay and extent of association on aggregation level either chemically or physically (Baldock and Skjemstad, 2000; Cusack et al., 2012; Mikutta et al., 2006; Torn et al., 1997; Vogel et al., 2014). Traditional fractionation methods based on size and external force for dissecting the association strength between OC and minerals in soils are limited to bulk sample. High resolution information and in-situ knowledge is required for interpretation of fractionation results and modeling (Kaiser et al., 2002; Kleber et al., 2007; Sollins et al., 2009). Nano scale two-dimensional isotopic mapping discovered that only a limited amount of the clay-sized surfaces contributed to OC sequestration (Vogel et al., 2014). Understanding OC interplay with minerals in the fine fraction warrants study in a three dimensional way (Kinyangi et al., 2006; Lehmann et al., 2007; Lehmann et al., 2008; Solomon et al., 2012). Detailed in-situ association information between OC and minerals may lead to breakthrough on mineral physical protection mechanism for OC long term stabilization. To overcome the limitations of commonly used electron microscopic methods (such as only on the surface layer, and undesirable artifacts due to pretreatments), non-destructive high-resolution X-ray 3-D tomographic technique will be used for exploring the fine structure of OC and boundary interplay with mineral nano particles.

High resolution Synchrotron-based TXM has been demonstrated as a powerful tool for understanding the internal 3-D structure of particles down to nano meter scale, due to its large penetration depth and superior spatial resolution (Kuo et al., 2011; Wang
et al., 2015). This technique was successfully applied to reveal the discrete three-dimensional micro-aggregation structure of clay (kaolinite) in natural aqueous environment, and generated remarkable tomography that revealed precise inter-particle structure (Zbik et al., 2008). Clay particles with diameter below 500 nm were clearly visible and their pseudohexagonal symmetry was recognized in details in a three dimensional way.

The synchrotron-based TXM at the beamline BL01B1 of Taiwan Light Source (TLS), which has been used in this study, provides two-dimensional imaging and three-dimensional tomography at a spatial resolution of 30/60-nm with tunable energy (8-11keV). It provides unprecedented opportunity for studying OC boundary interplays with mineral particles at nano meter scale. Two image acquisition modes, absorption contrast and phase contrast, can be used alternatively for recognizing OC and nano minerals. Conventionally, X-ray images are taken in the absorption contrast mode, and the resulting image contrast only depends on the difference of X-ray attenuation coefficient between materials. This mode is especially useful for materials consisted of high atomic number compositions. However, in organic materials, the difference of X-ray attenuation coefficients between specimen and air is too small to distinguish each other. For this reason, the structure of organic materials is often hard to be recognized due to low contrast in absorption contrast images. Alternatively, phase contrast technique transfers optical path length differences (optical phase) inside specimens into intensity contrast, can be used for imaging low atomic number materials, which are poor to absorb X-rays. It provides a unique opportunity to observe fine structures of organic specimens such as
OC. Little study has been done on OC and mineral nano particles using high-resolution 3-D X-ray tomography, though non-synchrotron-based 3-D X-ray microscopy was used to observe occluded carbon in phytolith structure and kerogen in micrometer scale (Alexandre et al., 2015; Bousige et al., 2016). We aim to develop a new dual-scan method using phase contrast and absorption contrast modes of the TXM alternatively for the observation of OC and mineral consortiums inside lab-made and natural samples in nanometer scale. Lab-made OC in forms of black carbon (BC) will be examined in the artificial consortium with added nano mineral (TiO₂) particles using synchrotron-based TXM for the first time.

Black C/biochar has received increasing research interest globally due to its importance in global carbon cycling, soil fertility improvement and environmental pollutant remediation (Bond et al., 2013; Jeffery et al., 2015; Kuhlbusch, 1998; Lehmann et al., 2007; Liang et al., 2006; Liang et al., 2008; Schmidt, 2004). On top of method development for 3-D tomography at nanometer scale, this study provides in-situ evidences on the minerals physical protection on natural OC, and to explore the C stabilization mechanism in natural soil.

**Methodology**

**Sample preparation and background**

Black C was made in lab using leguminous plant (*Sesbania roxburghii*) of 80 days' harvest, which was first oven-dried (65 °C) and charred inside a muffle furnace at 300 °C
in loosely sealed stainless containers (Chen et al., 2014b). This consortium of low
temperature BC and mineral nano particles was constructed by dry deposition of
commercial TiO$_2$ (15 nm) on lab-made BC (3 mm chunk), and then embedded in Gatan
G-1 epoxy. The block were cross sectioned to a thickness of 100 to 200 µm using a
microtome (Leica Reichert Ultracut E ultra-microtome) and subsequently hand-polished
to a thickness of 30 to 50 µm. Each section was transferred onto Kapton tape and
mounted on a stainless steel sample holder for TXM observation. Before TXM analysis,
gold nano particles (50-150 nm or 400-500 nm in diameter) were deployed on the section
surface for image registration before 3-D tomography reconstruction.

Thin section of natural OC and mineral consortium (NH) was prepared using
micron to millimeter size particulate sample from high mountain soil. Particulate organic
matter of mm-size with minerals embedded inside was taken from the lower dark layer at
depth of 72-93 cm in a Typic Humicryepts soil profile, located in Mt. Nanhua, Nantou
County, Taiwan (24°03'00", 121°17'02"). On top of this dark layer, iron stain was observed
within the depth of 63-72 cm in the profile. The soil has developed on top of sandstone
and slate, with some features of inceptisol and spodosol. The sampling elevation is 3092
m, the annual temperature is 7.57 °C, and the yearly rainfall is 2203.1 mm. The major
vegetation is arrow bamboo (*Yushania nittakayamensis*), with sporadic Hemlock (*Tsuga
chinensis*), fir (*Abies kawakamii*), and spruce (*Picea morrisonicola*).

The sequestration environment represents weak leaching and inactive chemical
weathering conditions. The age of soil organic C has been estimated to 3500 years B.P.
Working conditions of TXM

A superconducting wavelength shifter source provides a photon flux of $4 \times 10^{11}$ photons s$^{-1}$ (0.1% bw)$^{-1}$ in the energy range of 5-20 KeV at the BL01B1 beamline. A double crystal monochromator exploiting a pair of Ge (111) crystals selects X-rays within the energy range of 8-11 KeV. The specimen is imaged using a Fresnel zone plate, which is used as an objective lens for an image magnification of 44× by the first order diffraction mode. Conjugated with a 20× downstream optical magnification, the TXM provides a total magnification of 880× with a field of view of 15×15 μm$^2$. By acquiring a series of 2D images with the sample rotated 1° stepwise, 3-D tomography datasets is later reconstructed based on 151 sequential image frames that are captured with azimuth angle rotating from -75° to +75°.

Image acquisition for 3-D tomography

Under the most frequently used absorption contrast mode, 2-D images are recorded based on the projection of the different X-ray absorption coefficient integration along the optical pathway through samples on a detector. The absorption mode is useful for materials of high absorption coefficient, such as minerals or high atomic number materials, but it is poor for the observation of low atomic number materials, such as organic or polymer materials. In order to recognize the OC structure more accurately, 2-
D/3-D images for the same sample region are recorded using absorption contrast and phase contrast modes, respectively. In the phase contrast mode, the gold-made phase ring positioned at the back-focal plane of the zone plate is used to retard or advance the phase of the non-diffractive light by $\pi/2$, generating (Zernike's) phase contrast images recording at the detector. The light diffracted by specimen is interfered with the retarded non-diffractive light, generating phase contrast image. The intensity difference in a phase contrast image shows the combination of optical phase difference and absorption difference through specimens. This ability is especially important for the observation of OC which has a low X-ray absorption coefficient.

### 3-D reconstruction and analysis

Three dimensional tomography reconstruction is performed using homemade software, which is coded based on iterative image registration (Faproma) (Wang et al. 2017) and filtered back projection (FBP) reconstruction algorithms. Firstly, a serial of single TXM image captured from -75º to +75º at rotational increments of 1º is loaded to do image registration automatically using Faproma algorithm. Finally, the reconstruction is processed using FBP algorithm. The reconstructed dataset is exported in cross-sections, and later used for 3-D visualization using Amira. The intensity contrast of reconstructed datasets is inversed for better visualization. Compositions with higher absorption coefficients are shown in higher intensity and those with low absorption.
coefficients are shown in lower intensity. The exported cross-section of 3-D tomography (reconstructed datasets) shows the real distribution details and boundary interplay of OC and mineral particles. The final 3-D tomographic structures for visualization and illustration are generated using Amira 3-D software for image post-process and computation (Fig. S1).

Elemental mapping by SEM-EDS

For correlated spatial distribution of selected elements (C, O, Fe, Al etc) in natural OC particles from the high mountain soil, a low-vacuum scanning electron microscope (JEOL W-LVSEM, JSM-6360LV) equipped with an energy dispersive X-ray spectrometer (Oxford EDS) and a cathodoluminescence (CL) image detector (Gatan mini-CL) was used for elemental mapping, at an accelerating voltage of 15 KeV.

X-ray Diffraction for Mineralogy

To analyze the forms of minerals associated with natural OC, particulate OC (with minerals on the surface and embedded inside) was grounded and injected into capillary tubes (Special Glass 10, Hampton Research, CA) for synchrotron high resolution X-ray diffraction analysis at 09A beamline at Taiwan Photon Source (TPS), which is equipped with a set of high-resolution monochromator (HRM). The wavelength is 0.8266 Å at the energy of 15 KeV. The XRD spectra were recorded under room temperature for 240s accumulation time and specific X-ray diffraction peaks and patterns were assigned ICDD using PDF-2/4 program.
Carbon functionality and interfacial mineral forms using SR-FTIR

For FTIR analysis, mineral-bearing OC (NH) particles were grounded, dried (60 °C overnight), and mixed with potassium bromide (KBr) at a ratio of 1:100, and molded into disks using a hydraulic press. During the pressing process, a vacuum pump was used for evacuating air and water. The samples were measured using Infrared Microspectroscopy (IMS) at the BL14A1 beamline of the National Synchrotron Radiation Research Center (NSRRC), Taiwan. The FTIR spectra were collected up to 1024 scans in the mid-infrared range of 4000-400 cm⁻¹ with a spectral resolution of 4 cm⁻¹, using a FTIR spectrometer (Nicolet 6700, Thermo Fisher Scientific, Madison, WI, USA) with a self-equipped light source. The automatic atmospheric suppression function in OMNIC (OMNIC 9.2, 2012; Thermo Fisher Scientific Inc., Waltham, MA, USA) for bulk sample analysis was activated for data analysis, to eliminate the rovibration absorptions of CO₂ and water vapor in ambient air.

Results and Discussions

Distinguish the fine structure of BC and boundary interplay with mineral nano particles

High resolution 2-D X-ray photographs were captured for the identical regions in lab-made BC and nano mineral consortium using dual-scan absorption contrast and phase contrast modes (Fig. 1, a, e). The cross-section views exported from the
reconstructed 3-D datasets reveal subtle details of BC and mineral nano particles, and clearly outline the fine boundary of BC and the distribution of TiO$_2$ nano particles (Fig. 1). The shape, size, and distribution of mineral nano particles can be identified accurately using absorption contrast mode due to their high X-ray absorptivity (Fig. 1, b, c and d). In comparison, the BC structure and contour of its boundary can be revealed much more clearly using phase contrast mode (Fig. 1, f, g and h). However, the bright halo artifacts in phase-contrast image enhance the intensity of margin texture for nano minerals, and may lead to overestimation of their volume (Fig. 1, e, f, g and h). Use of dual-scan mode allows cross-checking of details and validation.

Cross-section views of the reconstructed 3-D tomography share consistent and comparable features of BC and nano minerals in multi-angles (Fig. 2). According to the display of different slicing planes (XY, XZ, YZ), it can be recognized that TiO$_2$ nano particles deposit inside BC only sporadically contact with BC boundary (Fig. 2, b, e, c, and f) due to the treatment of dry deposition. The nano scale gap between BC and nano minerals has been clearly observed in absorption and phase-contrast images (Fig. 2, b, e, c, and f). It is feasible to calculate the interplay surface and mineral volume quantitatively by examining each cross-section views in a selected region. Our approach is a success in thorough exploration of OC and minerals 3-D distribution, and verification of their real in-situ spatial correlation under nano scale resolution.

3-D tomography for illustrating in-situ distribution of BC and mineral nano particles
3-D tomography for visualization has been computed and generated to illustrate the spatial correlation between BC and minerals based on post-process of reconstructed 3-D datasets. Unprecedented details of 3-D in-situ distribution of BC and mineral nano particles are revealed in computed 3-D tomography (Fig. 3; Fig. SMOV1, 2). Results from absorption mode and phase contrast mode are consistent and comparable. The fine boundary feature of BC is contoured to a more completeness in the phase contrast mode. The OC was rendered by transparent mode and high absorptivity materials (such as minerals and gold particles) were rendered by solid mode with various colors. All renderings are combined to visualize their interaction. The illustration of 3-D computed tomography allows randomly tilted and set angles for image and animated video exports, thus any region of interest inside a specimen may be explored thoroughly.

The lab-made consortium was successfully tested by this dual-scan methodology using both absorption contrast and phase-contrast acquisition modes (Figs. 1, 2, 3). Low temperature BC, which is more similar to natural OC (especially recalcitrant OC) than that made at high temperature, was especially made to test its applicability under absorption contrast mode. Results show that the fine structure and boundary of low temperature BC can be clearly observed under absorption contrast mode. Thus for environmental OC samples, the use of absorption contrast mode is probably sufficient for capturing organo-mineral features.

Different from field samples, the minerals observed within the lab-made consortium often distribute in clusters and are only sparsely in association with BC surface. The preservation of plant-like structures in BC could play a role for carbon stabilization in
natural environment, as their porosity and reactive surface provide large areas and sites for mineral coating, which may contribute to their long residence and physical endurance (Eusterhues et al., 2008; Rasmussen et al., 2005; Rawal et al., 2016).

Interplay of OC and minerals and C stabilization in high mountain soil

Nano scale 3-D tomography revealed a high heterogeneity within natural OC-mineral consortium, and most of the particulate OC surface is coated by minerals, provides new insight for the mineral physical protection mechanism of OC in soil. Natural OC exhibited strong organo-mineral association on its surface at nano scale in high mountain soil (Fig. 4; Fig. SMOV3). Abundant short-range order minerals in forms of subhedral particle or anhedral nano-aggregate have direct association with the boundary of OC, and develop coating on the tracheid surface (Fig. 4 b and c) (Mikutta et al., 2006). Sheet-like mineral coating is observed on OC surface, which are dense and thin layers, likely originated from absorption (Fig. 4 b, Fig. S2). Another distinct texture is recognized as nano-aggregates/clusters of various shapes, possibly formed by OC-mineral co-precipitation. Mineral aggregation by poorly crystalline nano particles renders natural sub-micron porosity, which may contribute to elevated sorption capacity in soil (Rawal et al., 2016). The densely-packed mineral texture suggested significant physical protection on OC surface (Kaiser and Guggenberger, 2007). The sorbed minerals not only can form physical protection, but also could shield OC from chemical weathering (Mikutta et al., 2006). The key working minerals for OC-mineral interplay are essentially SRO mineral nano particles, and/or poorly crystalline submicron
size clay minerals, instead of crystalline, micron/clay-size minerals. Another distinct texture is recognized as nano-aggregates/clusters of various shapes, indicating possible OC-mineral co-precipitation (Fig. 4b, Fig. S3). Mineral aggregation by poorly crystalline nano particles renders natural sub-micron porosity, which may contribute to elevated sorption capacity in soil (Rawal et al., 2016).

The nature of associated minerals was confirmed to be mainly SRO Fe oxyhydroxides, specifically ferrihydrite (ICDD 01-073-8408), goethite (ICDD 01-073-6522), lepidocrocite (ICDD 00-044-1415), and quartz (ICDD 00-033-1161) (Fig. 5; Table S1), analyzed using high resolution synchrotron-based X-ray diffraction. Quartz may be at most a minor component on OC surface, considering their chemistry and particle size. Yet siliceous mineral surfaces may become coated with a veneer of hydrous Al- and Fe- oxides, which could confer net positive charge and promote their reactivity in tropical environments (Chen et al., 2014a; Sposito, 1989).

Considering their high surface area and reactivity, the abundant nano scale Fe oxyhydroxides could play a significant role for OC long-term stabilization through chemical bonding and physical shielding (Eusterhues et al., 2005; Kaiser et al., 2002; Kiem and Kogel-Knabner, 2002; Mikutta et al., 2006), as well as cation sorption in soil, and contribute to the longevity of OC in high mountain soil. According to elemental mapping results, aluminosilicates may also be present, however, their portion and crystallization level should be low due to their minimal signal in the XRD spectra (Figs. 5, 6). The FTIR analyses reveal the chemistry of organo-mineral association (Fig. 7; Table S2). The aged OC is highly aromatic when at the same time highly reactive, likely
originated from pyrogenic C, as broad bands are observed centered at 1596 cm\(^{-1}\) and 1706 cm\(^{-1}\) for aromatic C=C stretching and carboxylic \(\nu\) C=O, (Özçimen and Ersoy-Meriçboyu, 2010; Sharma et al., 2004). Both aromatic and carboxyl C functional groups normally have high affinity with Fe (III) (Mikutta et al., 2007; Zhao et al., 2016). Broad bands indicate a significant degree of association between OC and minerals (Chen et al., 2016; Gu et al., 1994; Kaiser and Guggenberger, 2007). The sorption of OC to Fe oxyhydroxides through organo-mineral multiple complex bonds such as ‘ligand exchange’ could occupy their reactive surface sites on OC and Fe-oxyhydroxides, tune down their activity and enhance their respective stabilization (Chorover and Amistadi, 2001; Cornell and Schwertmann, 2006; Hall et al., 2016; Kaiser and Guggenberger, 2007; Mikutta et al., 2007). The discovery of short-range-order SRO mineral ferrihydrite in air-dried OC particles and later ground-samples indirectly validates its stabilization due to organo-mineral interplay. As a SRO/metastable mineral, ferrihydrite is hard to estimate accurately in dry soil samples due to its transient nature and the limitation of traditional extraction and spectroscopic methods (Cornell and Schwertmann, 2006). The specific mineral phase in direct contact with OC on surface at nano scale warrants future study (Fig. SMOV3). In-situ mineral mapping of different SRO minerals/Fe oxyhydroxide on OC surface will provide mechanistic understanding on OC stabilization. Mineral physical protection on OC may represent the end stage of carbon stabilization, especially in a weak leaching and weathering environment.

Our in-situ description of organo-mineral interplay at nano scale provides direct evidence on the importance of mineral physical protection for OC long term stabilization.
High amounts of ferrihydrite and other Fe oxyhydroxides were also found associated with lignin-like OC in soil under an aquic moisture regime (Eusterhues et al., 2011). The abundance of mineral nano particles, and their high heterogeneity and short-range-order nature could be common in humid environment, however, they could have been seriously underestimated by traditional analysis methods, such as electron microscopy, X-ray diffraction and fractionation approaches, which focus on clay-size minerals (Mikutta et al., 2005). Mineral physical protection for OC stabilization may be more important than previous understanding. More research is proposed to explore: i) whether it is a general phenomenon that the minerals interact with OC are essentially SRO mineral nano particle and or poorly crystalline submicron size clay minerals, ii) whether the major mineral for C stabilization is nano size SRO minerals instead of clay-size minerals in soils. Perspective on C stabilization and saturation should be revolutionized to take the role of SRO minerals into account in modelling soil C dynamics, in addition to parameters such as clay type and content.

In summary, a high resolution 3-D tomography tool is required for exploring the in-situ interplay of OC and mineral nano particles in natural environment. Nano scale 3-D tomography provides direct evidence and new insight for the mineral physical protection mechanism of OC in soil. This high resolution 3-D tomography approach is a promising technique for probing the multi interfacial features between OC and minerals in lab and field samples, and may provide new perspective on the fate of nano particles including heavy metals in natural environments.
**Figure Captions**

**Figure 1.** The 2-D X-ray images for BC and mineral nano particle consortium. Two images are taken for the same region using absorption contrast mode (a) and phase contrast mode (e), respectively. Cross-section views of the reconstructed 3-D tomography under each mode at different depths relative to the position of gold nano particle along Z-axis as a reference. (b) and (f) are sections extracted at the position of the gold particle. (c) and (g) are sections extracted at 800 nm above the gold particle. (d) and (h) are sections extracted at 800 nm below the gold particle. The scale bar is 5 μm.

**Figure 2.** Three-directional orthogonal sections of lab-made BC and mineral nano particle consortium. The upper row sections are extracted from absorption contrast tomography (a, b, c), and the lower row sections are extracted from phase contrast tomography (d, e, f), specifically (a) and (d) are for XY plane, (b) and (e) are for YZ plane, and (c) and (f) are for XZ plane. The scale bar is 5 μm.

**Figure 3.** 3-D tomography illustration of lab-made BC and mineral nano particle consortium observed at -45° (a, d), 0° (b, e), and +45° (c, f) azimuthal viewing angles under absorption contrast (a, b, c) and phase contrast mode (d, e, f). The scale bar is 5 μm.

**Figure 4.** Three-directional orthogonal sections of high mountain mineral-bearing OC from absorption contrast tomography (a for XY plane, b for XZ plane, and c for YZ plane). The scale bar is 5 μm. Minerals mainly present two types of textures, subhedral particles
and anhedral nano-aggregates. The lower row images highlight the free surface of specimen (red line in d), the boundary of OC (green dotted-line in e), and the subhedral mineral particles (pink arrow in e and f).

**Figure 5.** The X-ray diffraction pattern of minerals within OC particles from high mountain soil. Highly reactive Fe oxyhydroxides are identified and denoted with lines of different colors: ferrihydrite (ICDD 01-073-8408, orange), goethite (ICDD 01-073-6522, blue), and lepidocrocite (ICDD 00-044-1415, green). Q stands for Quartz (ICDD 00-033-1161). Details are included in Table S1.

**Figure 6.** Elementary mapping by SEM-EDS for mineral-bearing OC from high mountain soil. Left: SEM backscattering image (The bright spots inside are gold nano particles for coating). Right: Elemental mapping of C, O, Fe and Al. Scale bars are 20 μm.

**Figure 7.** The FTIR spectra for the chemistry of organo-mineral association. The aged OC is highly aromatic (1596 and 1386 cm⁻¹), and highly reactive with obvious carboxyl functional group (1706 cm⁻¹). The broad bands point to possible significant degree of association between OC and minerals. Some minor bands near 1274, 1062, 1024, and 989 cm⁻¹ indicate the lignin-derived nature of OC. Those bands near 476, 534, 798, 910 and 1025 cm⁻¹ have similar characteristics of soil inorganic/mineral matrix. More details are included in Table S2.

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**References**


Figure 1.

Figure 2.
Figure 3.
Figure 4.
Figure 5.

Figure 6.
Figure 7.