

1 **Characterization of Iron Speciation in Urban and Rural Single Particles using**
2 **XANES Spectroscopy and Micro X-ray Fluorescence Measurements:**
3 **Investigating the Relationship between Speciation and Fractional Iron**
4 **Solubility**

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Abstract

Soluble iron in fine atmospheric particles has been identified as a public health concern by participating in reactions that generate reactive oxygen species (ROS). The mineralogy and oxidation state (speciation) of iron have been shown to influence fractional iron solubility (soluble iron/total iron). In this study, iron speciation was determined in single particles at urban and rural sites in Georgia USA using synchrotron-based techniques, such as X-ray Absorption Near-Edge Structure (XANES) spectroscopy and microscopic X-ray fluorescence. Soluble and total iron content (soluble + insoluble iron) of these samples was measured using spectrophotometry and synchrotron-based techniques, respectively. These bulk measurements were combined with synchrotron-based measurements to investigate the relationship between iron speciation and fractional iron solubility in ambient aerosols. XANES measurements indicate that iron in the single particles was present as a mixture of Fe(II) and Fe(III), with Fe(II) content generally between 5 and 35% (mean: ~25%). XANES and elemental analyses (e.g. elemental molar ratios of single particles based on microscopic X-ray fluorescence measurements) indicate that a majority (74%) of iron-containing particles are best characterized as Al-substituted Fe-oxides, with a Fe/Al molar ratio of 4.9. The next most abundant group of particles (12%) was Fe-aluminosilicates, with Si/Al molar ratio of 1.4. No correlation was found between fractional iron solubility (soluble iron/total iron) and the abundance of Al-substituted Fe-oxides and Fe-aluminosilicates present in single particles at any of the sites during different seasons, suggesting solubility largely depended on factors other than differences in major iron phases.

1. Introduction

Iron is an important component in atmospheric aerosols due to its potential impacts on human health (Smith and Aust, 1997). Adverse health effects associated with aerosols, such as cell and DNA damage, can stem from toxic levels of reactive oxygen species (ROS; e.g. hydrogen peroxide, hydroxyl radical, superoxide anion and organic peroxides, etc.) that form as a consequence of redox cycling of trace metals (Kelly, 2003, Vidrio et. al., 2008). In comparison to other trace metals, iron has been reported as a significant source of ROS via metal-mediated pathways (Shafer et. al., 2010, Smith and Aust, 1997, Zhang et. al., 2008). The role of metals in adverse health impacts associated with aerosols depends largely on the fraction of total metal content that is readily soluble (Costa and Dreher, 1997, See et. al., 2007, Valavanidas et. al., 2008), thus, primary factors and mechanisms that alter iron aerosol solubility must be understood to further link aerosol iron to adverse health effects.

A growing body of knowledge has emerged on various control factors that impact iron aerosol solubility. Soluble iron in aerosols varies between 0 to 80% of total iron, showing no general trend with total iron concentration (Mahowald et. al., 2005). While several chemical mechanisms and physical particle properties have been shown to influence iron solubility, there is still significant uncertainty on the primary factors that control fractional iron solubility (Baker and Croot, 2010). Modeling, laboratory and field studies have suggested that iron particles in dust may undergo atmospheric transformations (e.g. acid-processing) that may enhance fractional iron solubility (Meskhidze et. al., 2005, Shi et. al., 2009). On the other

hand, Baker and Jickells (2006) observed no relationship between atmospheric acid processing and iron solubility in coarse and fine crustal particles in a marine environment, but instead observed a relationship between increasing fractional iron solubility and decreasing dust mass concentration, which was likely accompanied by decreasing particle size. In this particular study, the observed increase in fractional iron solubility was attributed to the large surface area available for iron dissolution that is characteristic of small particles (e.g. large surface area to volume ratio). However, Shi et. al. (2010) later demonstrated that differences in particle size alone cannot explain the increase in fractional iron solubility observed in Baker and Jickells (2006), suggesting that other processes (e.g acid-processing or mixing with other anthropogenic particles) may play a more dominant role or work synergistically with particle size to promote fractional iron solubility. A few recent laboratory studies have observed a strong relationship between iron speciation and fractional iron solubility in crustal and industrial fly ash particles (Journet et. al., 2008, Cwiertny et. al., 2008, Schroth et. al., 2009). Cwiertny et. al. (2008) showed that Fe(II)-containing solid phase minerals may contribute to a significant portion of soluble iron in crustal sources. In addition, Schroth et. al. (2009) showed that soluble iron content from industrial combustion sources, comprised mainly of iron sulfates, was significantly greater (~80% of total iron) than the soluble content of crustal particles (~0.04-3% of total iron), which were mainly comprised of iron oxides and silicates. These results are also consistent with aerosol data from a field study in Korea that showed enhanced fractional iron solubility in anthropogenic combustion sources rather than crustal sources (Chuang et. al., 2005). However,

there was insufficient data to determine whether unique speciation or acid-processing of iron in the combustion particles caused enhanced solubility. In addition, other studies have shown a positive relationship between organically-complexed iron (iron oxalate complexes) and fractional iron solubility (Paris et. al., 2011). Furthermore, Furukawa et. al. (2011) demonstrated that a majority of oxalate in ambient samples from Japan exists as metal complexes. Although the relationship between iron speciation and fractional iron solubility has been established in source particles (e.g. crustal and industrial), it is not clear in ambient particles. A comprehensive knowledge of iron speciation in relation to solubility in urban and rural aerosols would help to further understand its association with public health.

Relatively few analytical tools are available to provide detailed characterization of iron, which are described in detail by Majestic et. al. (2007). Typically, studies of aerosols rely upon chemical extractions or spectroscopic techniques that provide oxidation and/or mineralogy information on bulk properties of iron in a sample. Spectrophotometry (e.g. ferrozine) and high performance liquid chromatography (HPLC) have been used to quantify Fe(II) and Fe(III) in bulk aerosol samples, but yield little information on mineralogy (Johansen et. al., 2000, Zhuang et. al., 1992). Mossbauer spectroscopy has been successfully used to directly characterize the oxidation state and mineralogy in aerosol samples (Hoffmann et. al., 1996); however, collection of aerosol over a several month period is required to obtain sufficient mass for analysis (~1 g). Recent innovations in synchrotron-based X-ray absorption spectroscopy, specifically X-ray Absorption

Near Edge Structure (XANES) and Extended X-ray Absorption Fine Structure (EXAFS) spectroscopies, have made it possible to explore both the oxidation state and mineralogy of iron. These methods require minimal sample preparation and are capable of single particle analysis. XANES and EXAFS have been widely used to probe iron speciation in soil samples (Marcus et. al., 2008, Prietzel et. al., 2007). Werner et. al. (2007) recently extended EXAFS to atmospheric aerosols to identify oxidation state and mineralogy of chromium in urban California. A few studies have demonstrated the feasibility and benefits of synchrotron-based X-ray spectroscopic techniques using a low energy X-ray beam for the analysis of iron in aerosol samples, but primarily focused on oxidation state characterization (Majestic et. al., 2007, Takahama et. al., 2008). In this study, particles collected on Teflon filters from urban and rural sites were investigated using synchrotron-based methods, XANES and microscopic X-ray fluorescence, to characterize the oxidation state, elemental association and mineralogy of single iron-containing particles. Soluble iron was quantified using the ferrozine method (Stookey, 1970) to investigate the link between speciation and solubility properties.

2. Methods

2.1 Sample Collection and Storage

Iron particles collected on Teflon filters (Whatman, Piscataway, New Jersey: 47mm-diameter, 2µm pore size) were analyzed using XANES and microscopic X-ray fluorescence. Twenty-four hour integrated PM_{2.5} filters were collected during different seasons at three urban sites and one rural site (Table1) for the ongoing Assessment of Spatial Aerosol and Composition in Atlanta (ASACA) air quality study (Butler et. al., 2003) and used in this analysis. Briefly, ambient air at a nominal flow rate of 16.7 L/min was pulled through a cyclone (URG, Chapel Hill, North Carolina USA), selecting for particles with an aerodynamic diameter less than 2.5 µm (PM_{2.5}), then directed through a series of two annular glass denunders (URG, Chapel Hill, North Carolina USA), removing acidic and alkaline gases. The particles were then collected onto the Teflon filter (Whatman, Piscataway, New Jersey: 47mm-diameter, 2µm pore size). Samples were stored in sealed polyethylene bags in a dark freezer (~-20°C) immediately after collection and were analyzed within 1 to 11 months. Because iron is non-volatile, sampling artifacts are likely associated with changes in iron oxidation state during sample storage. Majestic et. al. (2006) studied this specific artifact in aerosol samples and observed minimal Fe(II) loss on samples stored in a dark freezer for periods up to 6 months. In addition, Takahama et. al. (2008) found no evidence for significant Fe(II) loss in samples stored in freezing temperatures over extended periods of time (> 1 year). Although Fe(II) loss due to chemical conversion is possible on these samples, it is not expected to be significant based on the sample storage time and conditions employed in this study.

Before XANES and solubility analysis, filter samples were cut with ceramic scissors into half portions with each portion used for either synchrotron-based analyses or fractional iron solubility measurements.

2.2. Single Particle Analysis: Synchrotron-based X-ray Spectroscopy

Synchrotron-based X-ray spectroscopy is based on the principle that every element has characteristic absorption edges that correspond to the binding energy of electrons in individual quantized shells (e.g. K, L₂, and L₃). In this technique, incident X-rays of sufficient energy bombard atoms, ejecting the electrons from an electron shell. Subsequently, an outer shell electron may relax into the vacated position, emitting a characteristic fluorescence signal. K-edge XANES spectroscopy, used in this study, specifically explores the absorption edge associated with the innermost, K-shell electron. The ejected electrons of the innermost K-shell interact with neighboring atoms. These interactions are influenced by the type, oxidation state and structural arrangement of atoms in a particle and are reflected in XANES spectra (Ingall, 2011). Thus, XANES spectra provide information on both oxidation state and the mineralogical structure associated with the element of interest.

A total of 221 iron-containing particles deposited on the Teflon filters were analyzed on the 2-ID-D beamline at the Advanced Photon Source at Argonne National Laboratory in Argonne, Illinois, USA. The 2-ID-D beamline uses an energy dispersive Si-drift detector (Vortex EM, with a 50 mm² sensitive area, and a 12.5 µm Be window; SII NanoTechnology, Northridge CA, USA) to measure X-ray fluorescence of the sample. All measurements were conducted under a helium atmosphere in order to minimize absorption and fluorescence artifacts caused by

low-Z elements in air. A randomly selected area of each filter sample ($\sim 0.5 \text{ cm}^2$) was placed over a slot of an aluminum sample mount for direct spectroscopic analysis of the iron particles on the filter. The sample was initially analyzed in microscopic X-ray fluorescence mode to identify regions on the filter with detectable iron concentrations (e.g. iron-containing particles). In this mode, a monochromatic X-ray beam with a diameter of ~ 400 nanometers was scanned over a filter area (typically $\sim 40 \times 40 \text{ }\mu\text{m}$) at a step size of $0.4 \text{ }\mu\text{m}$ and 0.4 s dwell to produce an elemental distribution map of the filter. These maps were produced by setting the X-ray energy to 7200 eV , which allowed for the collection of K-edge X-ray fluorescence data on elements with masses from aluminum to iron (Al, Si, P, S, Cl, K, Ca, Ti, V, Cr, Mn, and Fe). The fluorescence data was converted into concentration data ($\mu\text{g}/\text{cm}^2$) for each element using a calibration with National Bureau of Standards (NBS) reference material. Minimal interference was caused by PTFE and Zeflour filters as seen in figure 1 (low background signal). In addition to locating iron-containing particles, calibrated data from these maps was used to characterize the association of other elements with iron. An energy scan (e.g. XANES analysis) was subsequently collected for iron-containing particles identified in microscopy mode (typically 30 iron-containing particles/filter). The X-ray energy scale was calibrated to the iron K-edge (7112.0 eV) using an iron metal foil before XANES measurements were performed. The incident X-ray energy was varied from 7090 to 7180 eV in 0.5 eV increments using a monochromator for a $0.5\text{-}3.0 \text{ s}$ dwell to produce an energy scan near the iron K-edge of a given iron-containing particle.

2.3 XANES spectra analysis using ATHENA software (2.1.1)

ATHENA software (version 2.1.1) was used to process the raw energy spectra. Individual energy scans were smoothed using a three-point algorithm for 10 iterations. The energy scans were subsequently normalized using the edge step normalization option to avoid mathematical discrepancies caused by directly dividing the fluorescence signal of incident X-ray beam by the signal in the upstream ionization chamber. The pre-edge centroid of the XANES spectra was the primary spectral feature used to determine oxidation state. The pre-edge centroid position was only determined from high intensity spectra (>5000 intensity counts: 103 spectra) to avoid any interferences caused by the low signal to noise ratio in low intensity spectra. The pre-edge feature was normalized by subtracting the pre-edge absorption from the background absorption, calculated by interpolating a cubic spline equation through the absorption 1 eV before and after the pre-edge feature. A Gaussian equation was fit to the normalized pre-edge feature using peak fitting program in Igor software (version 6.1) to determine the pre-edge centroid position. Figure S1 provides a detailed demonstration on how the pre-edge centroid feature was extracted in this study. In addition, XANES energy scans of a wide range of Fe(II) and Fe(III) minerals (augite, pyrite, iron (II) oxalate, iron (II) sulfate, goethite, hematite, iron (III) sulfate, iron (III) sulfide) were collected at the 2-ID-D beamline under similar sampling conditions as ambient sample analysis during February 2010. Table S1 provides a detailed description (classification and origin) of each iron mineral standards. Powder from each iron mineral standard was mounted on aluminum sample stick using double-sided tape for analysis. Similar pre-edge centroid analysis was applied to XANES standard data of Fe minerals to compare to

ambient sample data. Oxidation state was determined by the relationship between oxidation state and pre-edge centroid position. In this study, a linear equation was interpolated through Fe(II) (augite, pyrite, iron (II) sulfate, iron (II) oxalate) and Fe(III) (goethite, hematite, iron (III) oxalate and iron (III) sulfate) mineral data with the mean pre-edge centroid position of Fe(II) and Fe(III) minerals representing 0% Fe(III) and 100% Fe(III), respectively. The pre-edge centroid position determined from single particles was converted to % Fe(II) content using this interpolation, using equation 1. Several studies have used a similar approach to convert pre-edge centroid position of K-edge XANES spectra of octahedral-coordinated Fe minerals into % oxidation state (Bajt et. al. , 1994, Wilke et. al., 2001).

$$\%Fe(II) = 100 - \frac{\text{Centroid Position} - 7112.9}{0.0143}$$

Equation 1. Linear interpolation used to determine oxidation state.

2.4. Fractional Iron Solubility Analysis

Soluble iron on the filter samples was measured using the ferrozine technique by Stookey (1970), based on the absorption of light by the Fe(II)-ferrozine complex at 562nm to quantify Fe(II) in solution. A DTMini-2 equipped with a dual deuterium and tungsten halogen bulb (Ocean Optics: Dunedin, Florida, USA) provided light in the UV/VIS range (200-800nm), and a USB2000 spectrophotometer (Ocean Optics: Dunedin, Florida, USA) was used for light absorption measurements. A flow-through 100cm Liquid Waveguide Capillary Cell (LWCC) (World Precision Instruments: Sarasota, Florida, USA) provided a long liquid absorption path length to enhance measurement sensitivity. The

spectrophotometer was calibrated using five ammonium Fe(II) sulfate standards ranging from 0 to 20 Fe(II) ppb liquid concentration (typical $r^2=0.9999$) before and after soluble analysis. The de-ionized water sample leach used in this study showed minimal interference with the ambient sample (< 1% measurement interference).

Sample preparation and analysis used in this study are similar to the protocol described by Majestic et. al. (2006) for the analysis of soluble iron on 24-hr integrated filter samples. In our study, one half of the filter sample was placed in an acid-cleaned 30 ml amber Nalgene bottle and was subsequently diluted by 15 to 20 ml of de-ionized water (>18.0 M Ω). PM_{2.5} was extracted into solution via 30 minutes of ultra-sonication. A 10 ml aliquot of the extracted sample was filtered through a 0.45 μ m PTFE filter (Fisher Scientific: Pittsburgh, Pennsylvania, USA) to remove insoluble particles (>0.45 μ m diameter) from the solution. Ferrozine (5.1 mM) was added to the sample aliquot (100 μ l ferrozine/10 ml sample) and pulled through the LWCC after 10 minutes of incubation time. Light absorption was immediately measured at 562nm (max light absorption of Fe(II)-Ferrozine complex) and 700nm (background measurement) to yield a 10-minute operationally-defined soluble Fe(II) measurement. Hydroxylamine (HA) was subsequently added to the remaining filtrate (100 μ l HA/10 ml sample) to reduce soluble Fe(III) to Fe(II). After 10 minutes of incubation time, the light absorption measurements were repeated following the same procedure as the Fe(II) measurements, yielding the total soluble iron (Fe(II) + Fe(III)) content of the filtrate. Fe(III) concentration was determined by subtraction of the Fe(II) soluble concentration from the total soluble iron concentration. The method limit of detection is estimated to be 0.11 ng m⁻³

(Majestic et. al., 2006), which is well below soluble iron concentrations typically observed in urban aerosols.

Bulk total iron concentration was determined by micro X-ray fluorescence measurements. In this approach, total iron measured on a given elemental map (typically 40 X 40 μm) was multiplied by the total sample area divided by the elemental map area to determine the total iron concentration of the sample. When 2 or more elemental maps were collected for a given sample, the average of the total iron concentration from all elemental maps was used to determine the concentration of the sample using micro X-ray fluorescence measurements. Uncertainty was determined by the variability of total iron measured on different elemental maps for the same filter sample. In this study, there was moderate uncertainty (20-55% standard deviation) associated with this measurement approach. These uncertainties were predominately associated with an uneven distribution of iron-containing particles on the filter. Fractional iron solubility was determined by normalizing soluble iron to total iron concentrations (e.g. soluble iron/total iron). The uncertainty in fractional iron solubility was estimated for each filter sample by propagating the error associated with soluble iron (1-3%) and total iron (20-55%) measurements.

3. Results and Discussion

3.1 Identification of Iron-Containing Particles

Using microscopic X-ray fluorescence, several areas were scanned on each urban and rural filter (1 - 4 maps per filter) to map out the spatial distribution and concentration of elements from aluminum to iron (Al, Si, P, S, Cl, K, Ca, Ti, V, Cr, Mn, and Fe), referred to as elemental maps. Iron-containing particles were identified in this analysis and were subsequently analyzed using XANES spectroscopy. In addition, the elemental maps provided data on the elements that were associated with iron in each particle. The combination of XANES spectra and microscopic X-ray fluorescence were used to characterize mineralogy. Figure 1 shows an example of iron, aluminum, and silicon elemental maps from the South Dekalb winter (11/11/08) filter. The fourth map presented in Figure 1 shows the combined signal of all 3 elemental maps, indicating that both aluminum and silicon were associated with certain iron-containing particles in this sample.

3.2 Iron Oxidation State and Mineralogy

The pre-edge centroid position is the primary XANES spectra feature used to determine oxidation state and coordination chemistry of a given iron particle. It has been widely used to study iron in common minerals in soils (Prietz et. al., 2007, Wilke et. al., 2001) and continental shelf particles in the ocean (Lam and Bishop, 2008). The energy of the pre-edge centroid position shifts anywhere from 1.4 to 3 eV for a change of one valence electron (e.g. Fe(II) to Fe(III)).

The pre-edge centroid position was calculated for 103 particles (e.g. particles with high intensity spectra, >5000 raw counts) from the filter samples. **Figure S2**

shows the distribution of pre-edge centroid positions for single iron-containing particles from our samples. The pre-edge centroid position varied by 2.05 eV among filter samples, ranging from 7112.75 to 7114.8 eV, with an average of 7114.0 \pm 0.3, indicating significant oxidation state variability among our samples. The majority of the pre-edge centroid data for the urban and rural Fe particles falls between the centroid positions observed for Fe(II) and Fe(III) minerals (Figure S2). Figure 2 shows corresponding percent Fe(II) to total Fe of single particles, based on pre-edge centroid position, on each of the 8 filters identified by sampling site and season. The Fe(II) fraction in single particles from both urban and rural sites was generally between 5 and 35%, with the majority of the particles consisting of roughly 25% Fe(II). The rural site (Fort Yargo) during the winter had a much higher Fe(II) content than the other filters, having a mean Fe(II) content of 53%. In addition, a few particles (e.g. 6 out of 103) had much lower pre-edge centroid positions (7112.75 - 7113.15eV) compared to the average of the entire dataset (7114.0 eV), indicating iron in these particles was 100% Fe(II). The Fe(II) content of single particles in this study is greater than those observed by Takahama et. al. (2008), who showed a majority of marine and urban Fe aerosols exists as mixed-oxidation state agglomerations and surface-reduced particles, containing less than 10% Fe(II). The results presented in our study compared to those of Takahama et. al. (2008) suggests that large differences in iron redox state may characterize iron collected in different regions and seasons.

XANES spectra were similar for most of the Fe particles analyzed, regardless of season or site. Figure 3 shows a XANES spectra observed for a typical oxidized

and reduced particle (100% Fe(II) content, determined by pre-edge centroid position as seen in figure 3b, solid red line) observed in this study. Although the reduced particle shows a strong decreased shift in pre-edge and K-edge peak position, the shape of the spectra is similar to that of oxidized particles. Figure 4 shows the XANES spectra of a typical Fe particle (Sample 1, dashed blue spectra) observed in this study compared to the XANES spectra of several Fe(II) and Fe(III) compounds. The XANES spectra, for the most part, closely follows the spectra of iron oxides (e.g. goethite and hematite, blue solid spectra) and lacks a resemblance to other classes of Fe minerals, such as silicates (augite and biotite), sulfides (pyrite), organics (Fe(II) and Fe(III) oxalate), and sulfates (Fe(III) sulfate), suggesting that the majority of Fe in urban aerosols is iron oxides. Further separation into specific Fe oxides was difficult, since differences in spectral features amongst this mineral class are very subtle. Most of the XANES spectra of the reduced particles follow the spectra of iron oxides with a shift in edge position; however, a few (2 out of 13) spectra of “more reduced” (pre-edge centroid position <7113.6eV) particles show a strong resemblance to silicates (e.g. biotite) shown in Figure 4. The presence of Fe(II) (based on pre-edge centroid position) in iron-containing particles that appear to be iron oxides may suggest the presence of surface reduced species.

3.3 Elemental Composition of Iron-Containing Particles: Insight on Mineralogy

In addition to the XANES spectra, the elemental composition determined from microscopic X-ray fluorescence measurements of each iron-containing particle

was investigated to further understand iron mineralogy. The concentration of each element from aluminum to iron was converted into molar units (mol/cm²) and compared to the iron molar concentration of each particle. Collectively, the iron data showed no strong correlation with any element ($r^2 < 0.20$) for Fe mol vs. X mol, where X represents elements from Al to Mn. Although a portion of single particle elemental data had low elemental molar concentrations, two clear trends emerged when Fe (mol) was plotted against Al (mol) as in Figure 5.

The trends indicate that iron-containing particles could be divided into two groups. The first group, comprising the majority of particles (163 out of 221, 74%) (Figure 5: outlined by the blue area), were low in silicon (Si molar concentration < 0.1), yet contained a relatively consistent fraction of aluminum, in a 4.9:1 Fe:Al molar ratio ($r^2 = 0.81$, $p < 0.05$, e.g. within 0.05 statistical significance level). The aluminum content of these particles greatly exceeded trace aluminum levels that would be expected in pure iron oxide minerals, which ideally contain only Fe, O, and OH. Iron is commonly substituted by cations of similar size and charge, like aluminum, in iron oxide matrices, and is often observed in crustal particles (Cornell and Schwertmann, 2003). For example, aluminum-substitution observed in goethite can vary from 0-33% on a molar basis. This data coupled with the XANES spectra suggest that these particles are likely Al-substituted Fe-oxides.

The second group of particles (26 out of 221, 12%), shown in Figure 5, is characterized by lower iron concentrations and enhanced levels of silicon (Si molar concentration > 0.1) and aluminum (Al molar concentration > 0.1) relative to the first group of particles, referred to here as Fe-aluminosilicates. The silicon content

of these particles strongly correlates with aluminum, with a 1.4 Si/Al molar ratio, ($r^2=0.72$, $p<0.05$), which compares well to Si/Al molar ratios of common aluminosilicate minerals (typically 1 to 4) (Deer et. al., 1978). Though the XANES spectra of a few of these particles (2 out of 26 particles, Sample 2 spectra in Figure 4) resembled spectra of common iron-containing aluminosilicates, the majority of the spectra for these particles (24 out of 26 particles, Sample 1 spectra in Figure 4) are best matched by the common iron oxides. This result indicates that a majority of these particles contain a significant amount of iron in the form of oxides, which are oxidation products of Fe-aluminosilicates (Deer et. al., 1978). The Si/Al molar ratio coupled with XANES spectra, which indicates iron oxide, suggests these particles are processed Fe-aluminosilicates. The remaining 14% of iron-containing particles did not correspond to trends observed in either Al-substituted Fe-oxides or processed Fe-aluminosilicates, thus, their mineralogy was undetermined.

3.4. Mineralogy of Iron-containing Particles at Different Sites

Figure 5 also shows the distribution of Al-substituted Fe-oxides and processed Fe-aluminosilicates particles at urban and rural sites. Al-substituted Fe-oxides and processed Fe-aluminosilicates are observed at both urban and rural sites. For the urban sites, South DeKalb and Fort McPherson, show a mixture of both types of particles, regardless of season, while Fire Station 8 particles were exclusively associated with Al-substituted Fe-oxides for both winter and summer (Figure 5, with the exception of 1 point). In addition, Fort Yargo contained both types of particles during the winter, but was exclusively associated with processed Fe-aluminosilicates in the summer. The predominance of Al-substituted Fe-oxide

and processed Fe-aluminosilicate particles from our samples is consistent with a previous study showing iron aerosols collected in an urban area in Germany over a 5-month period are comprised of 78% iron oxides and 22% Fe silicates (Hoffmann et. al., 1996). Though the elemental Fe, Al, and Si molar concentration for a portion of single iron-containing particles was low (as shown in figure 5), thus, mineralogy was difficult to discern, our results clearly demonstrate that variations in iron mineralogy do exist in urban and rural PM_{2.5}. However, analysis of more samples (e.g. > 2 samples per site) is necessary to establish reasonable spatial and seasonal trends in bulk iron mineralogy.

3.5 Investigating Factors that Control Fractional Iron Solubility

Soluble and total iron content was determined for all the urban and rural filters for different seasons using the ferrozine and micro X-ray fluorescence measurements. Results from these analyses are presented in Table 2. A significant amount of variability was observed in soluble iron on filter samples collected in urban areas, with the concentrations ranging from 3.4 to 47.9 ng/m³, while soluble iron content on samples from rural areas were comparatively low ranging from 4.3 to 5.8 ng/m³. These concentrations are typical of fractional iron solubility in urban and rural aerosol in the Midwestern US (Majestic et. al., 2007). A wide range of total iron concentrations was also observed on the samples, ranging from 15 and 1734 ng/m³. Although the total iron calculated had moderate uncertainty (Table 2), the majority (e.g. 7 out of 8) of the iron from our samples was within the range (mean \pm standard deviation) of typical iron concentrations observed in urban and rural sites in the Southeastern US (Table S1). However, the iron concentration (1734 ng/m³)

observed at the urban site Fire Station 8 during the winter was much higher than typical concentrations observed in Atlanta, GA and urban Southeastern US sites. Although this concentration was observed at Fire Station 8, which is characterized as an urban Atlanta site with poor air quality (e.g. PM mass concentration generally 30% greater than other Atlanta sampling sites (Trail, 2010)), the total concentration was probably a direct result of an uneven distribution of iron on the filter. Iron on this particular elemental map may have been concentrated with respect to the remaining sample area, leading to an overestimation of total iron collected on the filter. This result reflects the moderate uncertainties associated with calculating absolute concentrations of aerosol components using synchrotron-based technology. With the exception of the Fire Station 8 winter sample, the majority of data are within the acceptable range of typical iron aerosol concentrations in the Southeastern US. However, for further statistical analysis, the Fire Station 8 sample was omitted.

To investigate solubility in relation to other variables (e.g. iron speciation), soluble iron concentration was normalized to total iron content determined by micro X-ray fluorescence measurements to yield fractional iron solubility (e.g. soluble iron/total iron content). Fractional iron solubility was between 2 and 38% (mean: 15.8 +/- 11.8 %) at individual urban and rural sites during different seasons (individual site data in Table 2). Though moderate uncertainty was associated with these solubility estimates (23-55% error, predominately due to total iron measurements), it did not appear to have a significant impact on the trends observed in solubility (see Table 2 and figure 6). Overall, the fractional solubility

levels observed in this study compare reasonably well to those found in common iron oxide (<1%) and silicate (3-6%) minerals (Journet et. al., 2008, Schroth et. al., 2009), suggesting our mineralogy data correspond well to expected fractional solubility levels.

Fractional iron solubility was compared to a number of variables to assess their influence. No clear relationship was found between fractional iron solubility and total iron content ($r^2=0.004$, $p>0.05$). These results are consistent with several studies, which have reported fractional iron solubility as an inconsistent fraction of total iron, ranging anywhere between 0 to 80% of total iron (Baker and Croot, 2010, Mahowald et. al., 2005). In addition, speciation (oxidation state and mineralogy) was also compared to fractional iron solubility. Figure 6 shows a moderate relationship between fractional iron solubility and median single particle Fe(II) data ($r^2=0.56$, Table 2); however, this trend is not statistically robust ($p>0.05$). In addition, given the limitations of this analysis (large variations observed in single particle Fe(II) content and small sample size), it is difficult to determine whether or not this moderate trend actually exists. However, our data suggests that Fe(II) content may not significantly impact fractional iron solubility. The absence of trend with speciation is more profound when fractional iron solubility is compared to variations in mineralogy observed at different sites. The most significant difference in the mineralogy of particles in this study was observed between Fire Station 8 site (summer and winter) and the Fort Yargo site during the summer, where particles were exclusively associated with Al-substituted Fe-oxides and processed Fe-aluminosilicates, respectively. No significant difference in fractional solubility was

observed at these two sites, suggesting iron speciation is not the only factor influencing solubility.

Although no clear relationship between fractional iron solubility and abundance of major iron phases observed in this study, several laboratory experiments on iron-containing crustal and oil fly-ash particles provide evidence supporting this relationship (Journet et. al., 2008, Schroth et. al., 2009). One reason explaining the lack of trend in this study is the two dominant mineral phases observed (e.g. Al-substituted Fe-oxides and processed Fe-aluminosilicates) have similar fractional iron solubility levels (iron oxides (< 1 %) and iron silicates (3 - 6%) (Journet et. al., 2008, Schroth et. al., 2009)). Thus, various mixtures of these two phases are not expected to yield large variations in solubility, which is the case in our data where fractional iron solubility is low and only slightly variable among different sites. A stronger association between mineralogy and solubility, however, is expected in areas where highly soluble iron minerals (e.g. iron sulfate) are the dominant source of iron in aerosols. For instance, iron oxides and silicates in crustal particles (< 6% fractional solubility) are significantly less soluble than iron sulfates in oil fly ash (~80% fractional solubility) (Journet et. al., 2008, Schroth et. al., 2009). No iron sulfates, which are observed in anthropogenic combustion sources, were observed in this study. However, Liu et. al. (2005) showed that various anthropogenic combustion sources comprise a small, but measureable component (< 10%) of Atlanta PM_{2.5}. Given the limited single particle analysis used in this study (e.g. limited sample size), the small fraction of iron-containing particles from anthropogenic combustion sources may not have been detected. This small fraction

of iron sulfates may have contributed to the somewhat enhanced solubility levels observed in our data at several sites (> 6%) compared to that of pure iron oxide (< 1%) or silicate (3-6%) minerals. In addition, low levels of iron sulfates may also explain the minor differences in solubility levels between different sites and seasons.

Another factor possibly affecting the relationship between mineralogy and solubility in this study is that ambient aerosol may have undergone a variety of atmospheric processes altering its chemical and physical properties. Oakes et. al. (2010) showed evidence for enhancements in Fe(II) aerosol solubility in acidic sulfate plumes in Georgia. Secondary sulfate is a significant portion of particle mass (~50%) in the Southeastern US (Liu et. al., 2005), which can form sulfuric acid in the absence of sufficient neutralizing cations. In this study, acid-processing mechanisms may play a more central role than speciation or work synergistically with speciation to influence iron solubility. This would be consistent with Shi et. al. (2011) who demonstrated that fractional iron solubility, in dust aerosols is highly sensitive to particle acidity (e.g. acid-processing mechanisms) and less dependent on other factors (e.g. particle source/mineralogy and size). In addition to acid-processing mechanisms, particle size can play an important role in fractional iron solubility. In this study, the size range of particles was limited (between 0.4 – 2.5 μm in aerodynamic diameter, lower size resolution based on the spot size of X-ray beam), and minor differences in particle size were difficult to detect. Thus, the relationship between particle size and fractional iron solubility could not be evaluated. More detailed studies involving particle pH, size, iron speciation, and fractional iron

solubility are necessary to better understand factors that influence iron solubility in ambient aerosols.

While the results of this study clearly demonstrate the value of single particle synchrotron-based analysis in determining aerosol speciation, there are uncertainties associated with this approach. One uncertainty is representing bulk sample properties with single particle measurements, which comprise a limited portion of the sample. Though the results in this study showed unique variations in single particle iron speciation, these properties may not be representative of the entire sample, making it difficult to compare to other bulk properties (e.g. fractional iron solubility). The results of this study show the need for both single particle and bulk iron speciation in order to fully understand its impact on fractional iron solubility in aerosols.

3.6 Atmospheric Implications: Insight on Human Health Toxicity

Fine aerosols that contain iron have been shown to generate toxic levels of ROS (Shafer et. al., 2010, Zhang et. al., 2008). Recent experiments have related toxicity to iron oxidation state in nanoparticles (diameters smaller than 100nm). Reduced iron in nanoparticles, either present as water-soluble or crystalline Fe(0) or Fe(II), has been shown to be more efficient than Fe(III) in ROS generation (Auffan et. al., 2008, Keenan et. al., 2009). For example, Auffan et. al. (2008) showed oxidation of Fe(0) and Fe(II) oxides (e.g. magnetite) immediately produce ROS, while Fe(III) oxides (e.g. maghemite) produced little to no ROS within one hour. Although particles in this study are larger (approximately 0.4-2.5 μm) and presumably less reactive than nanoparticles due to less surface area per mass, the

same mechanisms are likely involved in the formation of ROS via iron-mediated pathways (e.g. Fenton reactions). Thus, Fe(II) is a plausible precursor for immediate production of ROS in PM_{2.5}. The ambient particles we investigated contained various amounts of Fe(II) and Fe(III), with the Fe(II) fraction accounting for approximately ~5 to 35% of total iron. These results indicate that a significant portion of iron-containing particles is in a redox state that can produce ROS immediately. Although Fe(II) is not always soluble in ambient aerosols (a factor strongly associated with ROS formation), particle-bound Fe(II) may interact with specific species (e.g. acidic aerosol) during atmospheric transit, promoting its solubility.

4. Conclusions

We present a novel approach for exploring the speciation of iron in single atmospheric fine particles collected over urban and rural regions during different seasons using synchrotron-based XANES spectroscopy and microscopic X-ray fluorescence techniques. The majority of the particles contained mixtures of oxidized (Fe(III)) and reduced (Fe(II)) iron, with an average of 25% of the iron present as Fe(II). Particulate iron from urban and rural sites in Georgia was observed primarily in two phases, Al-substituted Fe-oxides and processed Fe-aluminosilicates. Though the composition of these aerosols was substantially different than pure minerals, it was consistent with modifications that occur during oxidation processes. Based on the techniques used in this study, variations in the abundance of Al-substituted Fe-oxides and processed Fe-aluminosilicates did not coincide with the fractional iron solubility. Fractional iron solubility may be

controlled by iron minerals from minor sources, for example, anthropogenic combustion sources of iron sulfates that were not detected by XANES as a component of overall mineralogy. In addition, other physical or chemical properties (e.g. particle acidity or size) may act in conjunction with mineralogy to influence solubility. These other properties may control the toxicity of iron-containing particles more than bulk mineralogy.

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Figure Captions

Figure 1. Elemental maps (30 X 30 μm) of iron (red), aluminum (green), and silicon (blue) from South Dekalb 11/11/08 filter sample are presented. The fourth map is a colocation map, where the iron map is superimposed on aluminum and silicon maps. The white particles on the colocation plot indicate that iron, aluminum and silicon are concentrated in this area. The yellow circles on the colocation plot indicate 3 iron-containing particles that are enriched in aluminum and silicon.

Figure 2. Percentage of Fe(II) to total Fe (Fe(II) + Fe(III)) observed in single particles on filter samples are plotted for individual sites separated by season. 50th percentile (black vertical line), 25th and 75th percentiles (upper and lower box), 10th and 90th percentiles (upper and lower whiskers) of each dataset are represented in this graph. The right axis represents the number of data points represented for each sampling site/season.

Figure 3. A) XANES spectra of a representative oxidized particle (blue line) and reduced Fe particle (red line) B) Example of the normalized pre-edge centroid position of a representative oxidized particle (blue line) and reduced particle (red line)

Figure 4. Comparison of XANES sample spectra (dashed line) to the spectra of common Fe mineral standards (solid line). Two sample XANES spectra are plotted. Sample 1 (dashed blue line) represents a typical Fe urban oxidized particle corresponding to the majority of the data. Sample 2 (dashed green line) represents a reduced Fe particle observed in a few particles. Fe standards that resemble the ambient XANES sample spectra are plotted (iron oxide group (solid blue line) and iron-containing aluminosilicates (solid green line)). The vertical lines on the graph represent the energy where key XANES spectral features were identified.

Figure 5. Scatter plot of iron and aluminum molar concentration in iron-containing particles identified on urban and rural filters. The color scale denotes the silicon content in the particles. The blue outline represents single particles that are Al-substituted Fe-oxides. Particles from urban are represented by open colored circles. Particles from Fort Yargo summer and Fire Station 8 sites are represented by colored diamonds and black Xs, respectively.

Figure 6. Fractional total (soluble + insoluble) Fe(II) (Fe(II)/Total Fe) measured by XANES (red bars) and fractional iron solubility (blue hatched bars) content in each filter sample. Error bars represent the $\pm 1\sigma$.

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792 **Table 1.** ASACA filter information

<i>ASACA sample site</i>	<i>Site Type</i>	<i>Local Sources</i>	<i>Winter Date</i>	<i>Summer Date</i>
Fire Station 8 Atlanta, GA (urban)**	Industrial	Rail yard Diesel Traffic	1/28/10	6/3/09
Fort McPherson Atlanta, GA (urban)*	Commercial/ Residential	Major Interstate Mobile	2/10/09	6/8/09
South Dekalb Atlanta, GA (urban)*	Commercial/ Residential	Major Interstate Mobile	11/11/08	9/9/09
Fort Yargo Winder, GA(rural)**	State Park	Forested Region	11/29/09	7/4/09

793 *Analyzed with XANES in October 2009

794 **Analyzed with XANES in February 2010

Table 2. Solubility Results for Urban and Rural Filters

<i>Site</i>	<i>Season</i>	<i>Soluble Fe(II)^a</i> (ng/m ³)	<i>Soluble Fe^{a,b}</i> (ng/m ³)	<i>Total Fe^c</i> (ng/m ³)	<i>N_{maps}^d</i>	<i>Total Fe(II)^e/</i> <i>Total Fe</i>	<i>N_{part}^g</i>	<i>Soluble Fe^b/</i> <i>Total Fe^h</i>
<i>Rural Sites</i>								
Fort Yargo	Summer	2.0 ± 0.08	4.3 ± 0.17	78.0 ± 43.1	2	0.47±0.09	7	0.055±0.03
	Winter	4.9 ± 0.2	5.8 ± 0.23	15.4 ± 3.6	3	0.83±0.28	8	0.38±0.09
<i>Urban Sites</i>								
Fire Station 8	Summer	4.6 ± 0.06	13.6 ± 0.18	52.6	1	0.45±0.16	6	0.26±0.10
	Winter	30.0 ± 1.0	49.9 ± 1.64	1734 ^e	1	0.50±0.12	23	0.028±0.01
Fort McPherson	Summer	4.0 ± 0.04	12.9 ± 0.14	88.1 ± 32.2	2	0.34±0.17	19	0.15±0.05
	Winter	6.9 ± 0.09	12.2 ± 0.16	96.5	1	0.49±0.10	11	0.13±0.05
South Dekalb	Summer	5.8 ± 0.1	10.8 ± 0.22	174.0 ± 86.3	2	0.40±0.23	14	0.062±0.05
	Winter	3.0 ± 0.06	3.4 ± 0.07	174.0 ± 39.5	3	0.39±0.09	30	0.020±0.004

a) Measured by ferrozine technique

b) Soluble Fe = Soluble Fe(II) + Soluble Fe(III)

c) Measured by XANES spectroscopy

d) Number of XANES elemental maps used to determine total Fe concentration

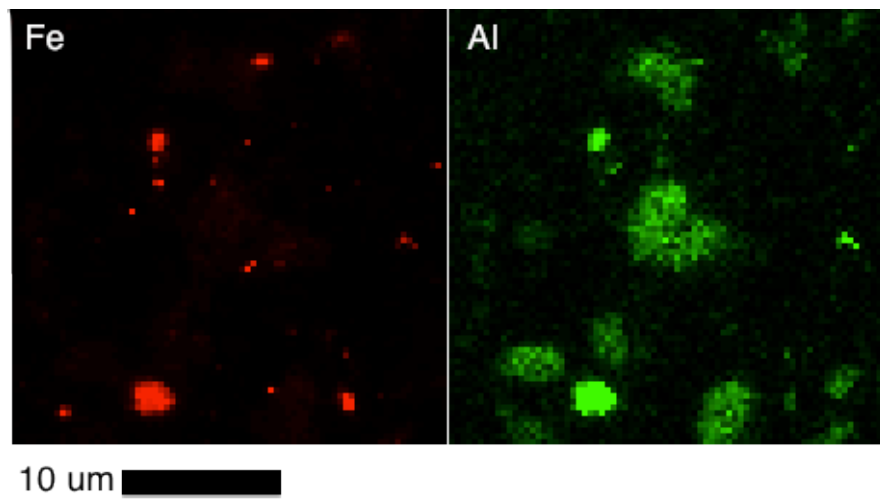
e) Measured by XANES spectroscopy, mean value of single particle oxidation state calculated for each filter

f) Outlier: Concentration is outside the standard deviation of typical total PM2.5 Fe concentration measured on filters collected in urban and rural areas in Southeastern US (seen in Table S1)

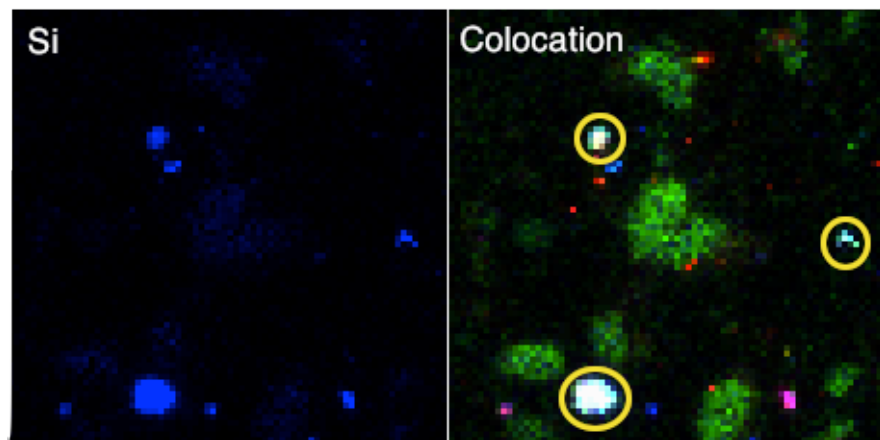
g) Number of single particles analyzed using XANES

h) Uncertainty propagated by the error in soluble and total iron measurements

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808



809 **Figure 1.**
810

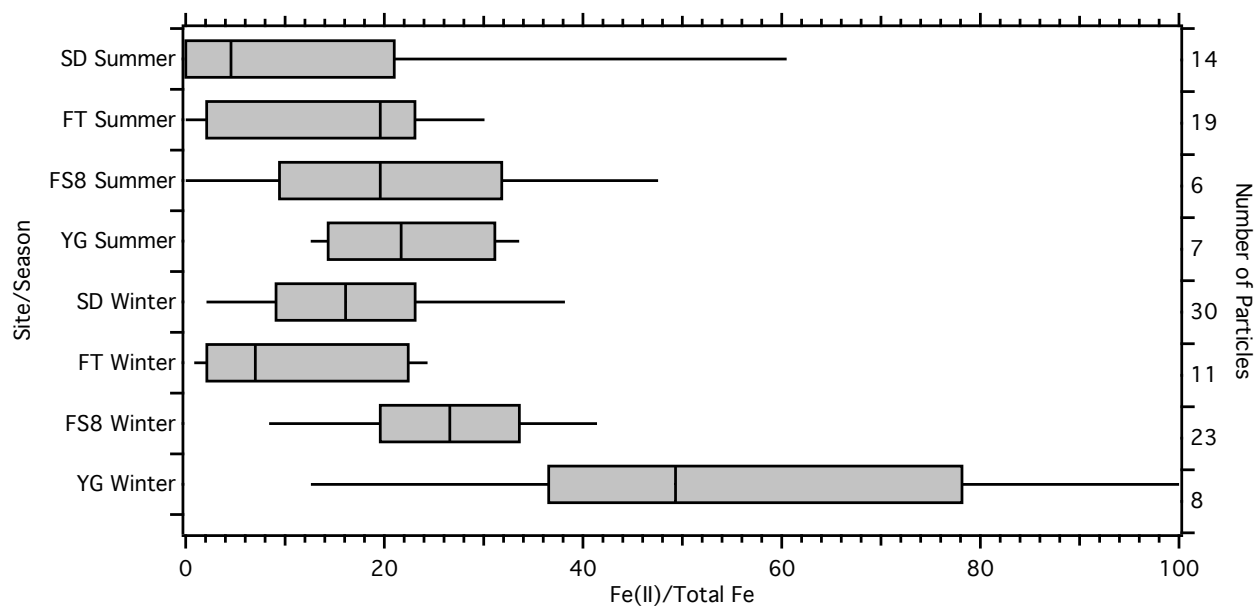


Figure 2.

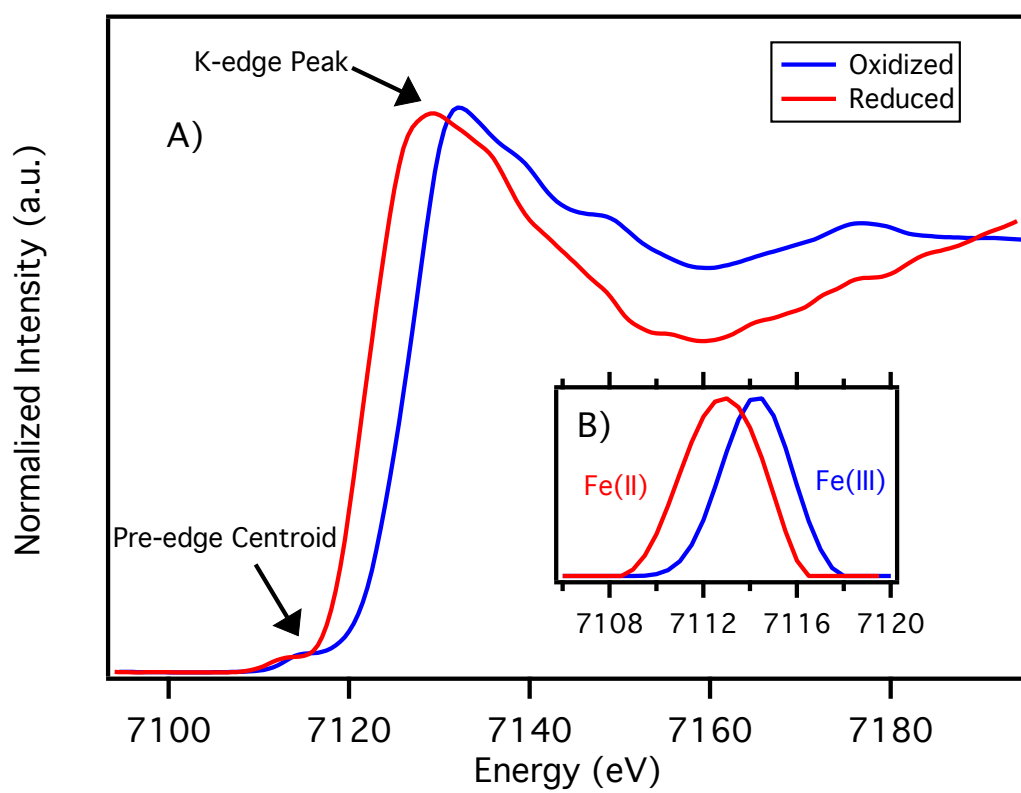


Figure 3.

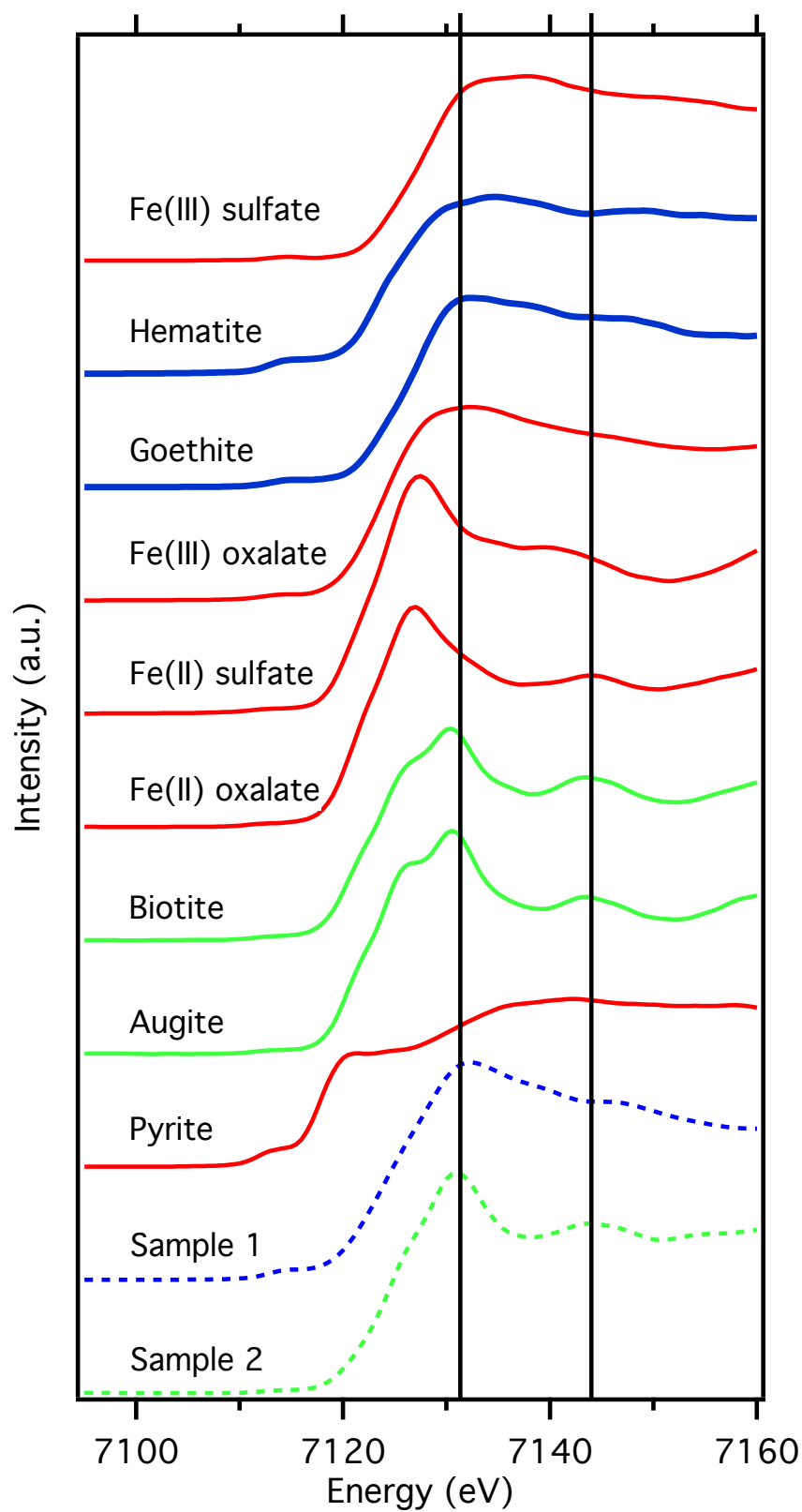


Figure 4.

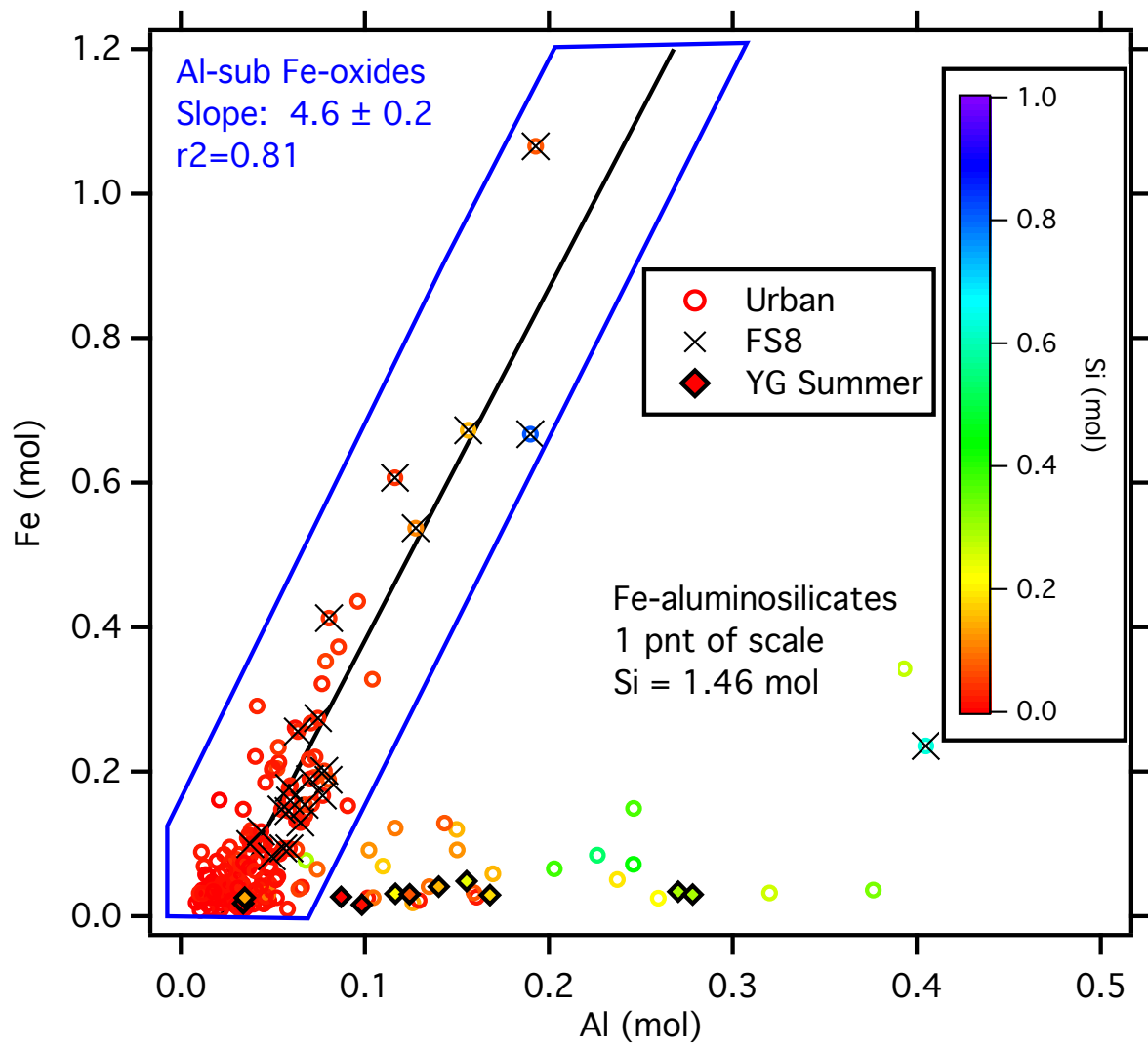


Figure 5.

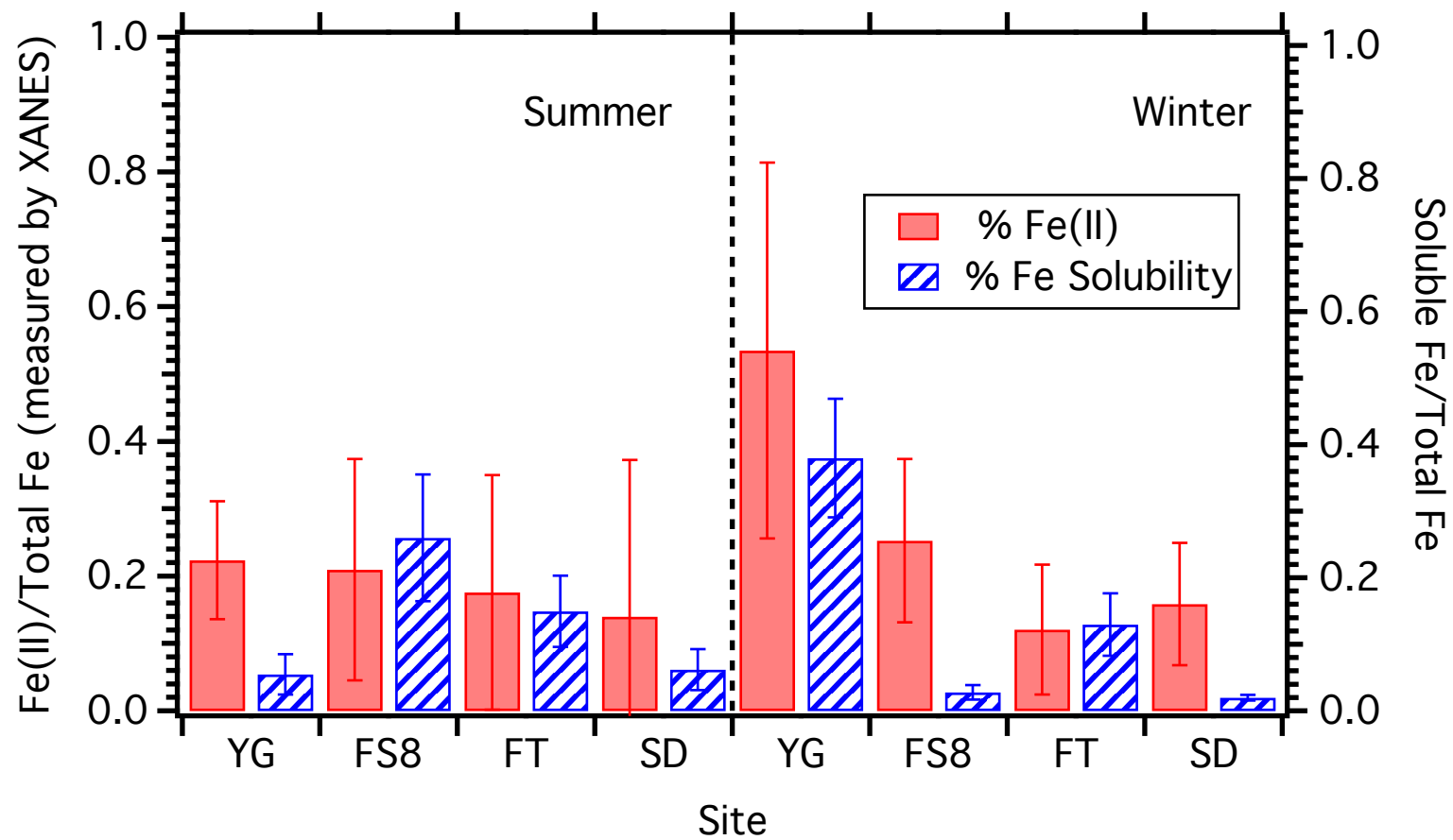


Figure 6.