



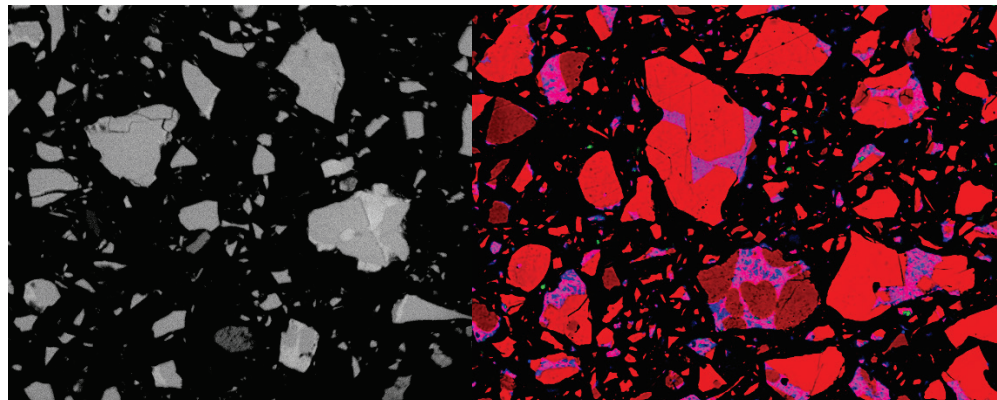
**US Army Corps
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Constituent Materials Characterization for Virtual Concrete Microstructure Generation

Jameson D. Shannon, Cody M. Strack, and Robert D. Moser

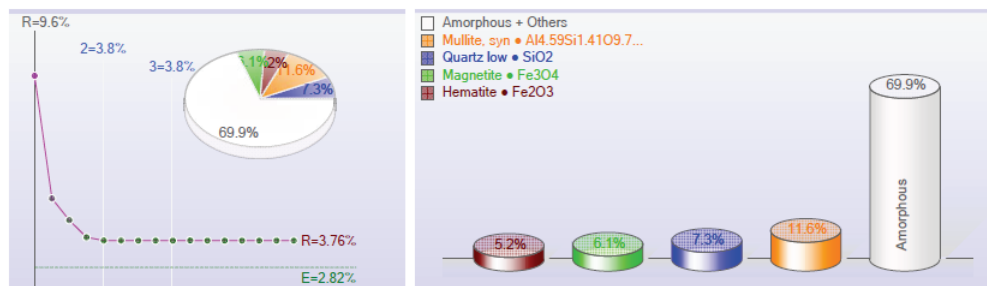
May 2019



Phase ID (4)	Chemical Formula	NA	NR	NP	Wt% (esd)	RIR
Quartz low	SiO ₂	---	12	9	7.3 (7.4)	3.02
Mullite, syn	Al _{4.59} Si _{1.41} O _{9.7}	---	29	10	11.6 (11.7)	0.83
Hematite	Fe ₂ O ₃	2	11	9	5.2 (5.3)	3.16
Magnetite	Fe ₃ O ₄	---	8	8	6.1 (6.2)	5.13
Amorphous + Others	SiO ₂				69.9 (123.5)	2.50

XRF(Wt%): Fe2O3=11.5%, SiO2=80.2%, Al2O3=8.5%

Refinement Halted (R/E=1.33), Round=3, Iter=6, P=37, R=3.76% (E=2.82%, EPS=0.5)



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Constituent Materials Characterization for Virtual Concrete Microstructure Generation

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Final report

Approved for public release; distribution is unlimited.

Abstract

This effort was undertaken as a part of the Materials Modeling for Force Protection (MMFP) Military Engineering 6.2 applied research program. One of the main philosophies of MMFP is the creation of a materials-by-design process for predictive properties and material selection capabilities. This report details the capability improvements to facilitate materials-by-design technologies and the collection and characterization of material data to compile and calibrate an integrated materials-by-design model. The Virtual Cement and Concrete Testing Laboratory (VCCTL) was evaluated for suitability in hydration model generation, and various programs were evaluated for initial virtualization of physical elements. To provide model inputs for particle size analysis, X-ray diffraction (XRD), X-ray fluorescence (XRF), scanning electron microscopy (SEM), and energy-dispersive spectroscopy (EDS) testing were conducted on a wide variety of constituent materials, and characterization data were processed included herein.

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Preface

This study was conducted for the U.S. Army Engineer Research and Development Center, Geotechnical and Structures Laboratory (GSL). The technical monitor was Dr. Jameson D. Shannon.

The work was performed by the Concrete and Materials Branch (GMC) of the Engineering Systems and Materials Division (GM), U.S. Army Engineer Research and Development Center, Geotechnical and Structures Laboratory (ERDC-GSL). At the time of publication, Mr. Christopher M. Moore was Chief, CEERD-GMC; Mr. Jeffrey G. Averett was Acting Chief, CEERD-GM; and Ms. Pamela G. Kinnebrew, CEERD-GTZ, was the Technical Director for Military Engineering. The Deputy Director of ERDC-GSL was Mr. Charles W. Ertle II, and the Director was Mr. Bartley P. Durst.

COL Ivan P. Beckman was the Commander of ERDC, and Dr. David W. Pittman was the Director.

Unit Conversion Factors

Multiply	By	To Obtain
cubic feet	0.02831685	cubic meters
degrees Fahrenheit	$(F-32)/1.8$	degrees Celsius
feet	0.3048	meters
inches	0.0254	meters
square feet	0.09290304	square meters
square inches	6.4516 E-04	square meters

1 Introduction and Background

1.1 Introduction

The Army has identified materials-by-design as a key area of research interest in recent years. Materials-by-design describes the complete process of the ability to transition between systems-level performance and analysis, and compositional material design seamlessly. This previously unattained computational capability would allow for virtual material design, fabrication, and optimization without the need for expensive and lengthy physical testing. The computational ability, once achieved, will allow for rapid analysis of system options based on available materials and for further optimization of material components. Using this system may also allow for new and advanced analysis of composites and additional performance increases of currently available products.

One of materials-by-design's key aspects is its integration of both top-down and bottom-up design considerations. However, the additional complications caused by this approach will require a large amount of build-up, and the original implementation will have to be studied and optimized to ensure the technique is viable to a wide range of materials. Overall, materials-by-design has great merit and if successful brings an immensely effective technique to Army engineering capabilities, but creation of a successful materials-by-design computational ability is ambitious and will require many years of research to fully implement.

1.2 Materials Modeling for Force Protection (MMFP) Background

This effort was undertaken as a part of the Materials Modeling for Force Protection (MMFP) Military Engineering 6.2 applied research program. Force protection solutions increasingly rely on the use of indigenous construction materials that are able to be obtained and produced in-theater. There is no current capability to design and develop high-performance construction materials in a virtual environment. Presently, available raw materials from areas of interest must be imported to U.S. laboratories, and iterative experimentation must be undertaken to create and optimize materials for protective design. To improve efficiency and resiliency, a materials-by-design approach is needed.

Materials-by-design determines material engineering properties needed to produce a specified structural response, and those material properties are then achieved through targeted formulation and synthesis processes. In the case of cementitious materials, the synthesis process is focused mainly on hydration of materials. Though multi-scale modeling approaches have been developed to predict weapons effects on structural elements, the capability to model the synthesis of the materials is missing. To date, material synthesis has been conducted solely by trial-and-error experimentation and characterization. Furthermore, advanced materials systems and structural elements are needed that provide improved protection with reduced logistics and cost.

A computational synthesis capability for advanced force protection materials does not currently exist. Though some material synthesis models are available, they do not account for the modern worldwide variety of constituent materials and the specialized curing conditions that are advantageous in advanced material formulations. Additionally, there is a current lack of multi-scale model coupling pathways to integrate synthesis, structure-property, and performance models. Because the needed model coupling does not exist, using a materials-by-design approach to develop protective components is not presently possible and represents a barrier to technical success. Finally, structural element optimization using indigenous materials cannot be performed virtually with current capabilities.

During the course of the MMFP research effort, a combination of computational model development and advanced experimentation using domestic and foreign raw materials will be performed to develop and validate an integrated materials-by-design capability. Synthesis models will be developed to predict material microstructures. Model coupling will be achieved by developing both analytical and computational approaches that predict material properties from the predicted microstructures, then predicted properties will feed performance models. Guided by simulations, new protective materials and structural elements will be fabricated, optimized, and validated experimentally.

Testing and validation of the materials-by-design capability will occur in two-ways: 1) structural element optimization based on selected sets of available raw materials (bottom-up), and 2) targeted material synthesis based on protective performance requirements (top-down). Integration with performance modeling efforts being conducted in other U.S. Army Engineer Research and Development Center (ERDC) programs will be

facilitated through semi-annual technical workshops, led by this work package, with broad participation from material developers, testers, modelers, and users across the ERDC direct and reimbursable programs as well as outside partners.

The remainder of this report discusses the first steps of this process being undertaken at the ERDC Geotechnical and Structures Laboratory. This phase of the effort focuses on improving materials characterization capabilities and increasing the amount of available data. These characterization data are then to be used as inputs in a cementitious material hydration model. Advancements in model coding and usage are necessary to achieve the desired results in a meaningful format that could be used to further feed higher scale models. Additional background on the hydration model and coding is given in Section 1.3. Characterization procedures and model usage, featuring inputs and outputs, are described in Chapters 3, 4, and 5.

1.3 CEMHYD3D and VCCTL background

CEMHYD3D is a package of programs designed by the National Institute of Science and Technology (NIST) to simulate cement hydration. The package functions by simulating a virtual cement in three dimensions and applying multiple hydration codes to form a virtual hydrated microstructure from which certain performance properties can be obtained. A two-dimensional image is first obtained experimentally and loaded into the program, from which a three-dimensional cube can be predicted. The virtual three-dimensional cube is formulated to reliably predict particle size distribution and volumes, surface areas, and autocorrelation functions of cementitious phases. Technical details of the interworkings of the CEMHYD3D package of programs are discussed in previous publications by Bentz (1995, 1997a, 1997b) and Bentz et al. (1996).

The Virtual Cement and Concrete Testing Laboratory (VCCTL) was created to improve the usability of the CEMHYD3D by providing a Graphical User Interface (GUI) and expanded control, input, and output options. VCCTL functions in the same way as the CEMHYD3D code it was built upon, beginning first with a 2-D segmented image and building a 3-D virtual mixture. Figures 1 and 2 illustrate examples of the imported 2-D image and virtualized 3-D microstructure. Chemical phases are identified on a pixel-by-pixel basis in the imported image and are transitioned into 3-D pixels (voxels) in the 3-D microstructure.

Figure 1. Imported 2-D image with “false color” (from Garboczi et al. 2004).

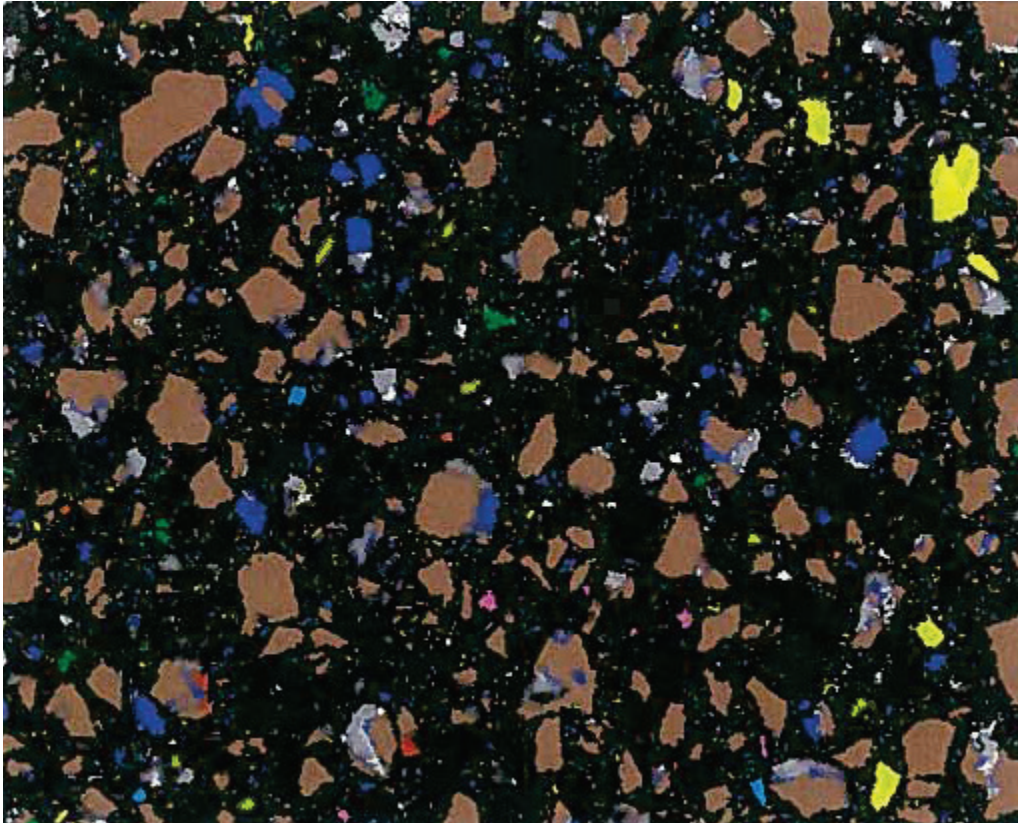
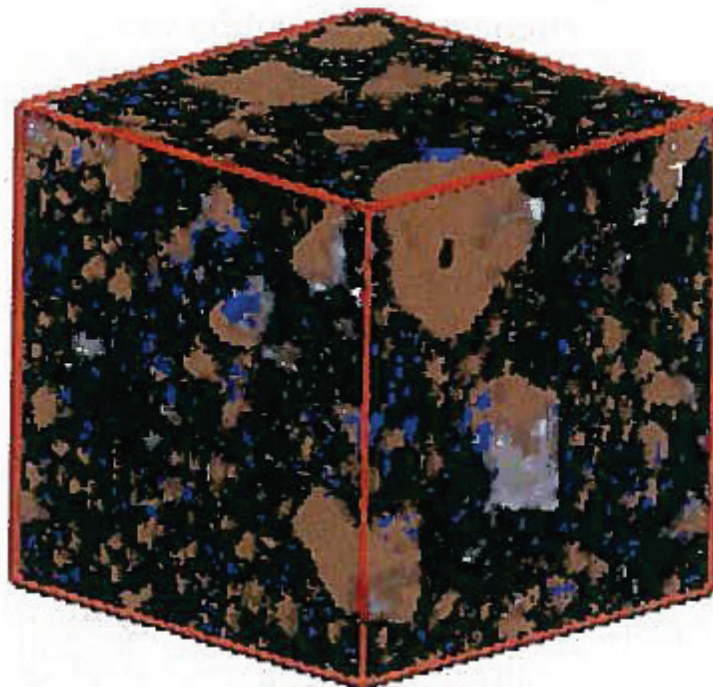


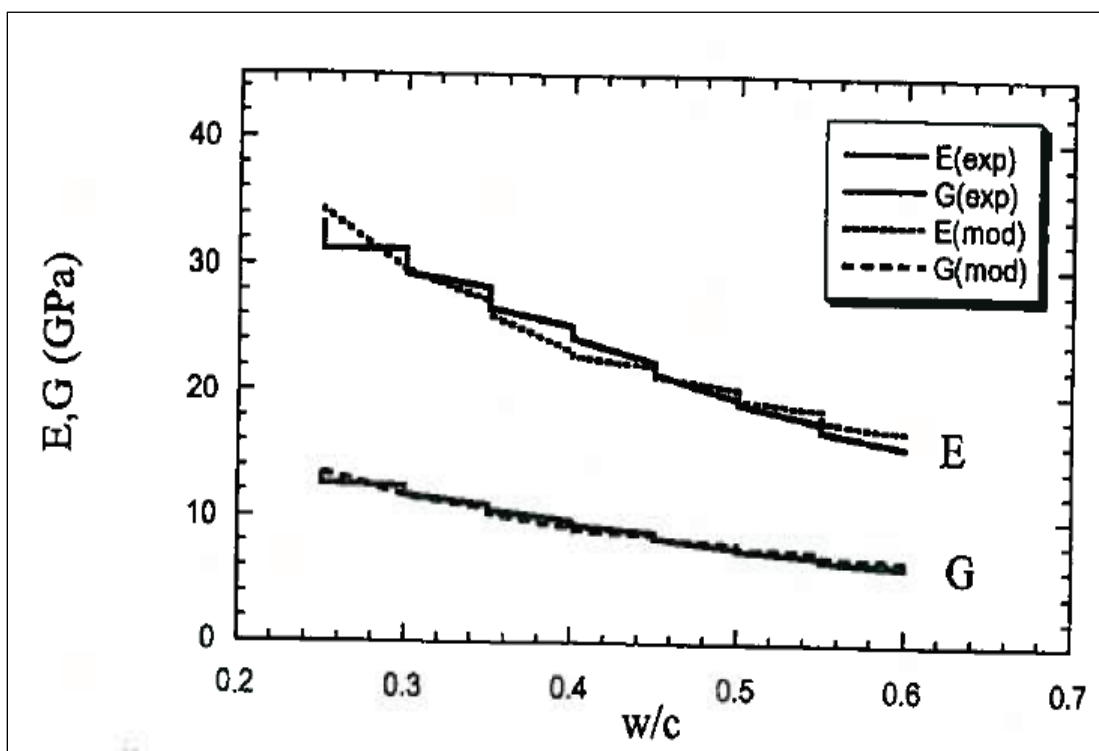
Figure 2. 3-D virtualized microstructure created from 2-D image (from Garboczi et al. 2004).



The VCCTL software package includes pre-installed cement, supplementary cementitious materials, and aggregate data to facilitate virtual testing. Materials unique to a certain project must be added to the software before modeling. Due to the large size differences between aggregates and cement paste particles, the two systems are handled separately. The cement paste portion is first calculated and stored, and an aggregate randomization program is used to determine fine and coarse particle locations in a matrix. The paste portion is then added to fill gaps between aggregate particles.

Running the hydration model “dissolves” the individual voxels, allowing them to react with water and produce a 3-D hydrated microstructure where each voxel becomes a phase of the cement paste. The hydrated image can then be used to calculate properties such as set point, temperature changes, chemical shrinkage, desiccation, and diffusion. Values for mechanical properties, such as elastic moduli, are then calculated by utilizing finite element techniques. A comparison of moduli results showing experimental values (exp) compared to model values (mod) is shown in Figure 3. Other concrete properties, such as compressive strength, may be calculated by using empirical or semi-empirical relations to the model’s calculated gel-space ratio or Young’s modulus.

Figure 3. Experimental vs model moduli results (from Garboczi et al. 2004).



2 Material Characterization

2.1 Characterization data required for model

In order to accurately produce results with the model, certain characterizations of the materials are necessary. For cement, a segmented indexed image file is required, most likely obtained by scanning electron microscopy (SEM) and adjusted using imaging software. Other necessary inputs include chemical property data, most likely obtained using x-ray diffraction (XRD) techniques, and a physical particle size distribution (PSD), which can be obtained by a variety of methods.

Other cementitious materials, such as silica fume, fly ash, slag, etc., require similar chemical composition and particle-size distribution inputs, but do not require their own segmented indexed image file. Aggregate input requirements include basic particle shape information and particle-size distribution files. However, shape requirements appear to be estimable without negative effects on the model as long as aggregates are roughly normally shaped for their type, e.g., no or a low content of flat or elongated particles.

Although the above requirements represent the minimum inputs required to use the model, additional information may be useful or desired. For example, characterization techniques such as x-ray fluorescence (XRF) can be used to further validate chemical compositions. Other characterization areas, such as particle-size distribution information that can be determined by multiple methods, may need to be optimized to provide the most accurate inputs. Depending on material type, various different methodologies may produce the most relevant particle size distributions, and as is the case with all virtual models, improving input accuracy increases the overall reliability.

2.2 Materials tested

At the time of publishing this report, 79 sets of material characterization data have been developed and processed into a digital material properties data storage location. An additional 59 sets of material characterization data have been collected and are awaiting processing into digital data storage. Table 1 lists materials selected for characterization and status of characterization data. Black checkmarks indicate data that have been

collected and processed. Green checkmarks indicate data that have been collected and awaiting processing. Material ID numbers were assigned at the time the material was received and are used to reference specific materials for the remainder of the report.

2.3 Advances in characterization capabilities

The majority of materials' characterization techniques were performed using existing equipment present in the Concrete and Materials Branch (CMB) of the Geotechnical and Structure Laboratory (GSL). XRD and XRF measurements were taken using a PANalytical X'Pert Pro diffractometer and PANalytical Axios spectrometer, respectively. SEM imaging was conducted using a FEI Nova NanoSEM 630 scanning electron microscope with energy dispersive x-ray (EDX) capability. EDX measurements were used to ensure the SEM image phases were identified appropriately.

2.3.1 Scanning electron microscopy and energy dispersive x-ray spectroscopy

Cementitious materials characterized with scanning electron microscopy (SEM) and energy-dispersive X-ray spectroscopy (EDX) measurements were observed in an unhydrated state in order to more accurately model pre-hydration microstructure. Samples were mixed with a quick setting epoxy blend and allowed to set overnight in 1-in.-diameter plastic molds. Epoxy-blended specimens were then cross-sectioned and polished up to a 1 micron suspension and sputter coated. SEM examination was focused on the microstructure of the paste. Imaging was performed using an FEI Nova NanoSEM 630 capable of high-resolution imaging on non-conductive materials. Imaging was performed in low-vacuum mode at pressures of 0.1-0.5 mbar and accelerating voltage of 15-30kV.

Table 1. Material selection and status of data collection.

Material Type	Description	ID	Property Data			Microstructure Images
			PSD	XRF	XRD	
Cement	Type V (Hanil Plant)	150146	✓	✓	✓	
Cement	Type I/II (Hanil Plant)	150147	✓	✓	✓	
Cement	Type III - National (Ragland Kiln)	160160	✓	✓	✓	
Cement	South Korea	170001	✓	✓	✓	
Cement	White cement - South Korea	170002	✓	✓	✓	
Cement	White cement - Lehigh	170009	✓	✓	✓	
Cement	Type I - National (Ragland Kiln)	170010	✓	✓	✓	
Cement	Type I/II MMC (St. Gen Kiln)	170052	✓	✓	✓	✓
Cement	Type H	170065	✓	✓	✓	✓
Cement	Type I/II MMC (St. Gen Kiln)	170096		✓	✓	
Cement	Type I/II MMC (St. Gen Kiln)	170098			✓	
Cement	PLC	170099			✓	
Slag	Lafarge Chicago	170080	✓			
Slag	South Korea	150150	✓	✓	✓	
Slag	Guam	---		✓	✓	
Fly Ash	Cook	80159		✓	✓	
Fly Ash	Meuth	100015		✓	✓	
Fly Ash	Gaston	100069		✓	✓	
Fly Ash	Nashville	120013		✓	✓	
Fly Ash	GHR	120083		✓	✓	
Fly Ash	China Lake	120104		✓	✓	
Fly Ash	Monroe	120222		✓	✓	
Fly Ash	Cumberland	130032		✓	✓	
Fly Ash	Cook	140022		✓	✓	
Fly Ash	NHS	140054		✓	✓	
Fly Ash	Pro Ash Ecotherm	150108		✓	✓	
Fly Ash	Pro Ash Ecotherm	150109		✓	✓	
Fly Ash	Activated Carbon Addition	150110			✓	
Fly Ash	Activated Carbon Addition with Re	150111			✓	
Fly Ash	Limestone Addition	150112			✓	
Fly Ash	Trona Addition	150113			✓	
Fly Ash	Guam			✓	✓	
Fly Ash	NNSS			✓	✓	
Fly Ash	South Korea (Handuk)	170003	✓	✓	✓	
Silica Fume	F10k (CMB)	130008	✓	✓	✓	
Silica Fume	Elkem 970D	130050	✓	✓	✓	
Silica Fume	Elkem ES 900W	140092	✓	✓	✓	
Silica Fume	Elkem Sidistar	140093	✓	✓	✓	
Silica Fume	Elkem 955U	140094	✓	✓	✓	
Silica Fume	Angelfort - White	150048	✓	✓	✓	
Silica Fume	Angelfort - Concrete	150049	✓	✓	✓	
Silica Fume	RED Hydraulic - Original Sample	150050	✓	✓	✓	
Silica Fume	RED Les Clavaux	150051	✓	✓	✓	
Silica Fume	RED Laudun	150052	✓	✓	✓	
Silica Fume	Hanil Plant	150148	✓	✓	✓	
Silica Fume	South Korea	170003	✓		✓	
Silica Fume	Ecotech85 (Guam)	Ecotech	✓	✓	✓	
Silica Fume	F10k (Guam)	F10	✓	✓	✓	
Silica Fume	Z3-95	170107	✓	✓	✓	
Silica Fume	D-1000		✓	✓	✓	
Silica Fume	Elkem 971U		✓	✓	✓	
Silica Flour	Sil-co-sil	---	✓		✓	
Metakaolin	Power Pozz	140015	✓	✓	✓	
Metakaolin	Hanil Plant	150149	✓	✓	✓	
Metakaolin	South Korea	170004	✓	✓	✓	
Sand	Silica Sand - F50	---	✓	✓		

All images were acquired using a backscattered electron detector to improve phase contrast. Imaging was performed at random sites and at various levels of magnification to capture the overall microstructure along with specific regions of interest. In conjunction with SEM imaging, chemical microanalysis using energy-dispersive X-ray spectroscopy (EDX) was performed using a Bruker Quantax AXS detector in order to characterize the chemical composition of phases of interest observed, particularly focusing on the composition of the cement paste.

2.3.2 X-ray diffraction analysis

X-ray diffraction (XRD) analysis was performed to identify the mineralogy of each component present in the concrete according to ASTM C1365 – Standard Test Method for Determination of the Proportion of Phases in Portland Cement and Portland-Cement Clinker Using X-ray Powder Diffraction Analysis (2018). The specimen was prepared into random powder-pack sample holders for XRD measurements. Diffraction patterns to be used for qualitative phase identification were obtained using a Panalytical X'Pert Pro materials research diffractometer equipped with a Co-K α X-ray source operated at 45kV and 40mA. Diffraction patterns were obtained over a period of 2 hr from 20° to 70° with a step size of 0.02°. Phase identification was performed using MDI Jade 2010 powder diffraction file (PDF) reference databases.

2.3.3 Particle size analysis

No capability for particle-size distribution analysis was available in the GSL at the onset of this research effort and, as this was expected to be a long-running effort, it was decided to invest in the capability. A Mastersizer 3000 laser diffraction particle size analyzer was obtained and installed on the premises. The laser diffraction method utilizes a controlled laser that interacts with particles with measurements being calculated based on the deflection of the laser off of the particle surfaces.

The unit is capable of determining particle sizes of materials as a solution in a liquid, as a dry powder, and in quantities less than 3 ml. These capabilities ensured that the GSL could produce accurate particle-size distribution data on a wide variety of materials. Figures 4-6 show the Mastersizer 3000 with dry powder analyzer (Figure 4), dispersed liquid analyzer (Figure 5), and small volume analyzer module (Figure 6).

Figure 4. Mastersizer 3000 with dry powder analyzer.



Figure 5. Mastersizer 3000 with dispersed liquid analyzer.



Figure 6. Small volume analyzer module.



3 Data Collection and VCCTL Model Inputs

3.1 Data collection archive

Collection of material characterization data on a large scale is a sizable portion of the MMFP research effort. Working with other researchers in the GSL, a priority list was created with materials of key interest. So far, 12 cements, 19 fly ashes, 4 metakaolins, 15 silica fumes, and 1 slag cement have been identified for characterization as well as aggregate materials such as standard sands, gravels, and silica flour. A large archive of material properties for the identified key materials is in the process of being created both for MMFP modeling and for future access needs.

Particle size distribution, XRF, and XRD experiments are being performed and analyzed, and SEM imaging is completed for the identified cements to be used in the model calculations. During the continued collection of the characterization data, virtual model experiments are also being conducted simultaneously so that, when the data collection process has been completed, a majority of the model validation will have already been conducted. Figures 7-10 illustrate some of the different types of data being collected and materials of interest.

Figure 7. Laser diffraction particle -size distribution of silica fume 1500149.

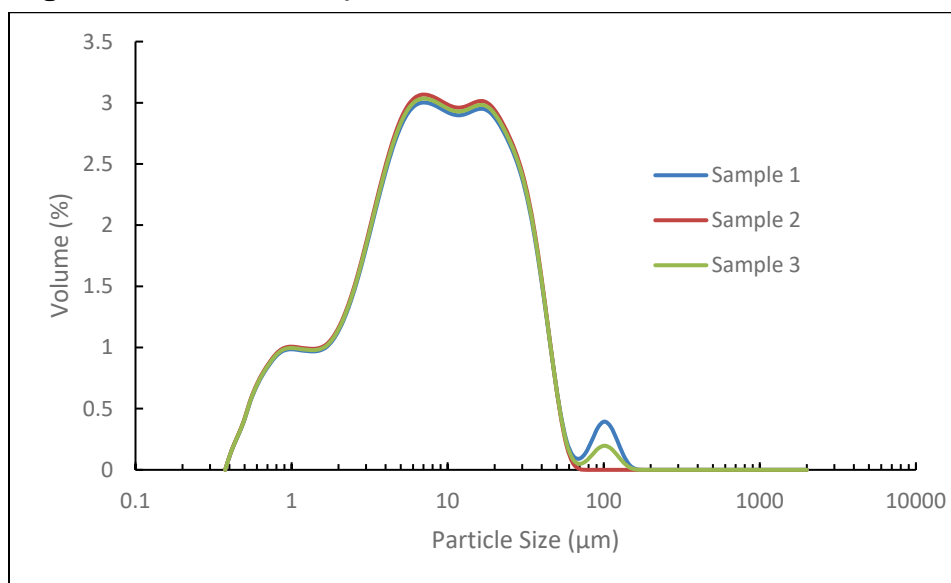


Figure 8. X-ray fluorescence (XRF) data of cement 170052.

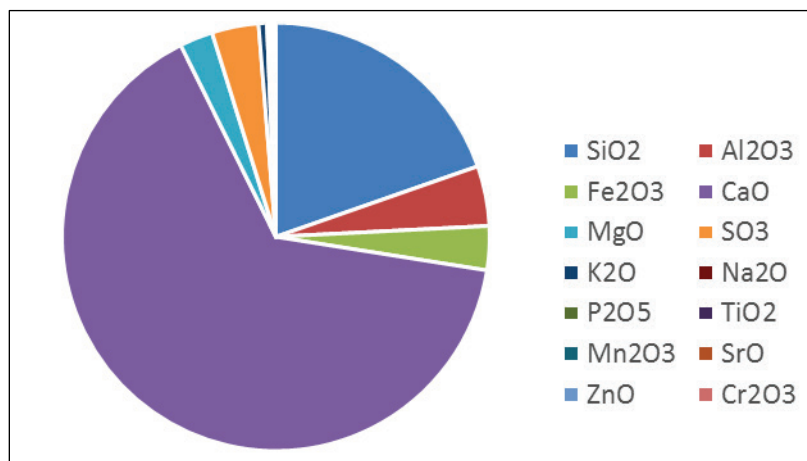


Figure 9. X-ray diffraction (XRD) data of fly ash 100015.

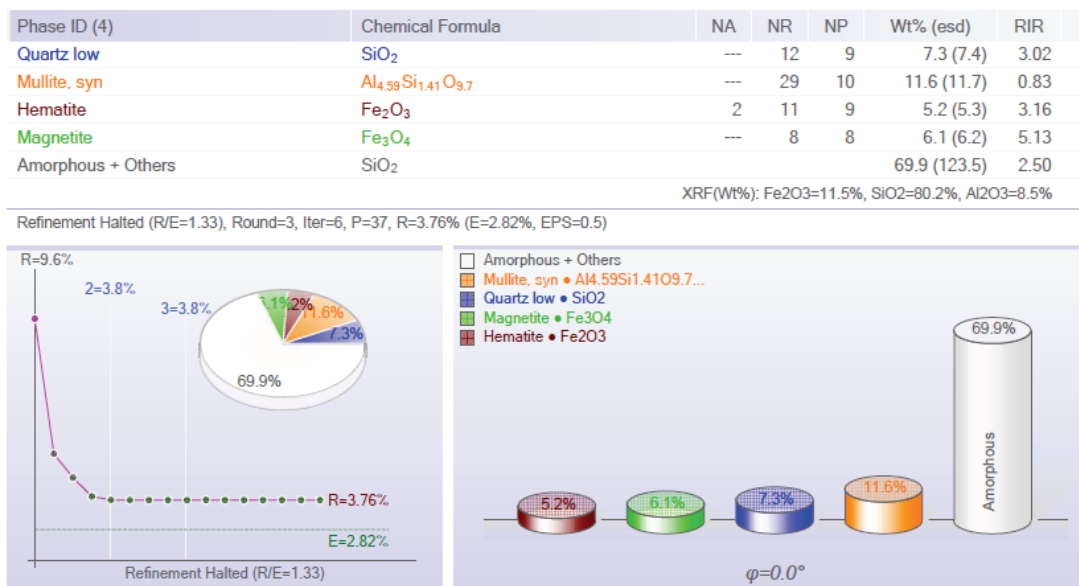
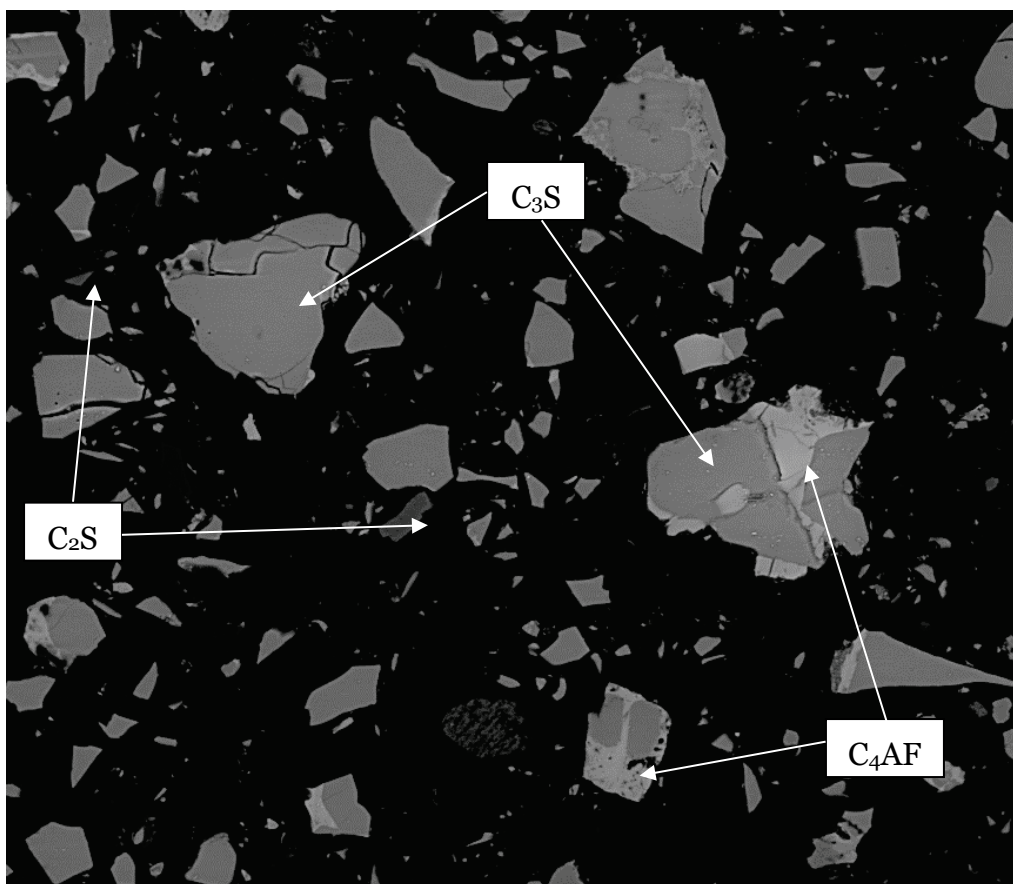


Figure 10. Scanning electron microscopy (SEM) image of type H cement (170065).



*MAG: 1000x; HV: 15kV; WD: 3.9mm; HFW: 285 mm.

3.2 Image processing and phase identification

Multiple images were taken of a single section with each image highlighting a different set of element parameters. An example set of images is shown in Figure 11. The six images represent the backscattered SEM image (a), Silicon highlighted (b), Sulfur highlighted (c), Iron highlighted (d), Aluminum highlighted (e), and Magnesium highlighted (f). The images can then be used to visually identify certain phases in the sample by taking into account which overlapping elements make up each particle

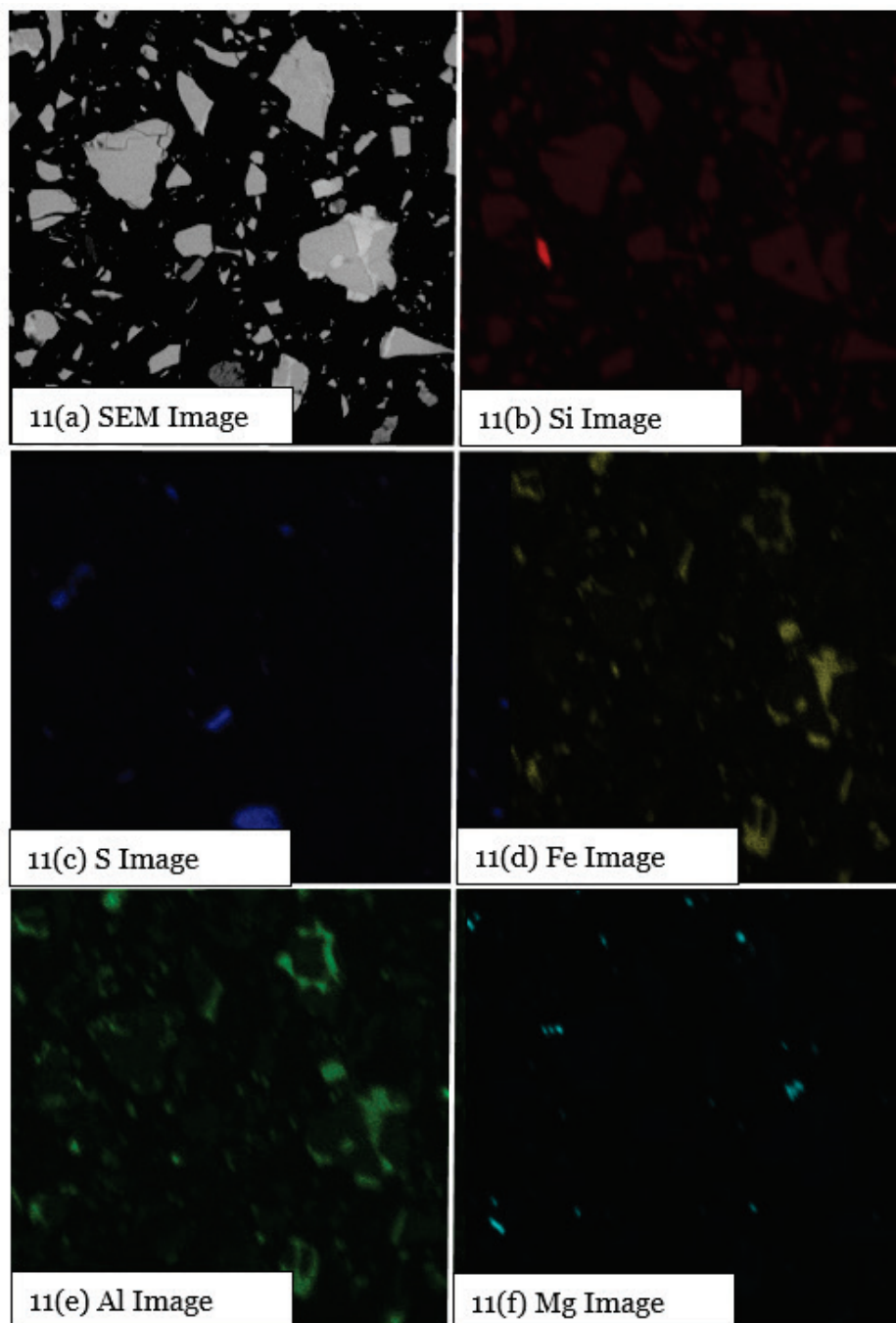
An investigation into digital image processing methodology was conducted using SEM and EDX images and a combination of the programs Imagej and Multispec. Using multiple color patterns and no color layers, it was possible to identify each of the phases present in the sample. For example, setting the base image to red and the aluminum image to blue showed the C_3A phase as light red, C_2S phase as dark red, and any Ferrite phases as purple. Using this methodology, phases can be processed, identified, and

labeled using software. An example of this methodology is shown in Figures 12 and 13.

A cement supplied by the Cement and Concrete Reference Laboratory (CCRL), given the designation 17705, was imaged by SEM. Separate images containing the complete gray-scale SEM image and element images of aluminum, magnesium, sulfur, potassium, and sodium were obtained. Images were assigned a color value of either red, blue, green, or none and placed on top of each other to form a single image file, shown in Figure 12a. Using knowledge of cement phases and color values, a label field was assigned to groups or areas of like pixels. This is illustrated in Figure 12b.

The software is then used to determine a calculated set of rules based on the user assigned values. Applying the same value field rules over the entire image produced a false color image of the sample in which each color aligns to a specific cement phase. The calculated false color image for the labeled combined image in Figure 12 is shown in Figure 13. The program also features the ability to calculate the percentages of each phase present in the image. Using the Multispec imaging process as a measure appears to allow for a greater level of certainty in detecting the phase portions of a gray-scale SEM image than visual interpretation.

Figure 11. Suite of SEM and EDX images using the same location and parameters as Figure 10.



*MAG: 1000x; HV: 15kV; WD: 3.9mm; HFW: 285 mm.

Figure 12. Combined Multispec image of multiple layers of cement 17705 without (a) and with (b) fields labeled.

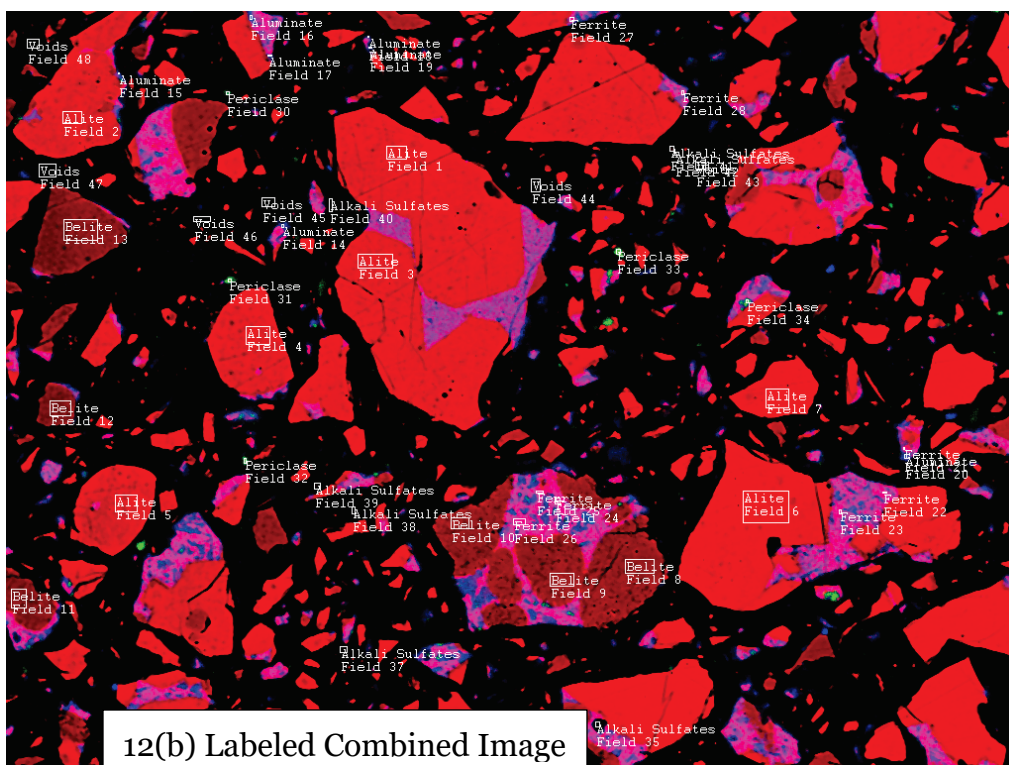
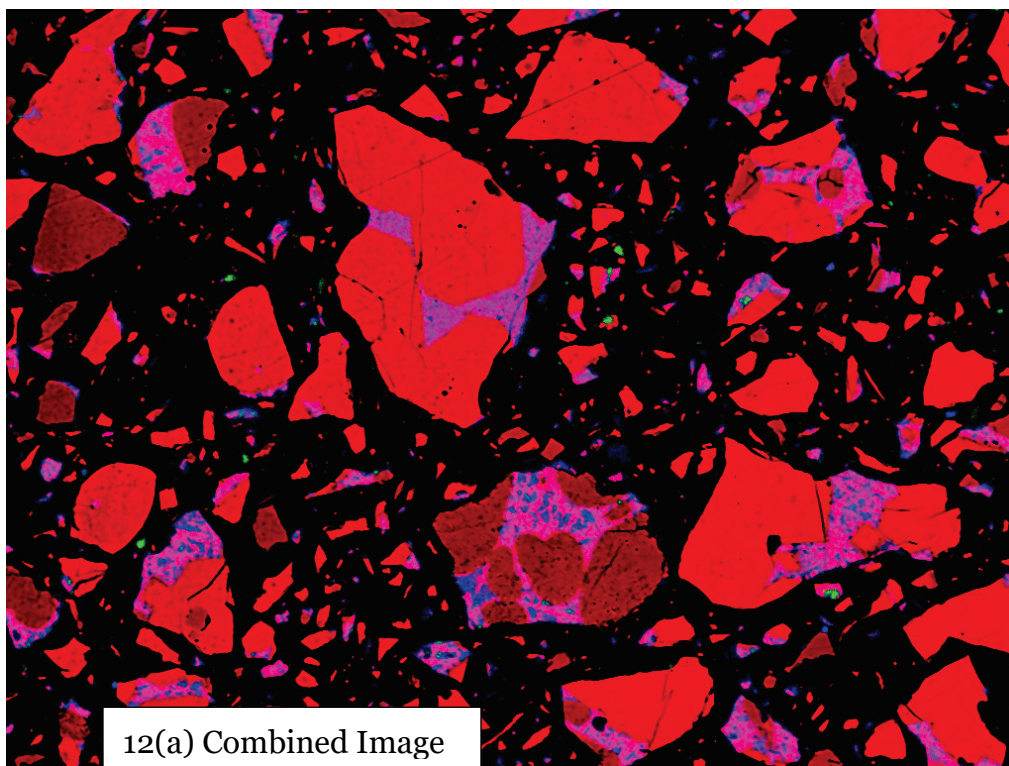
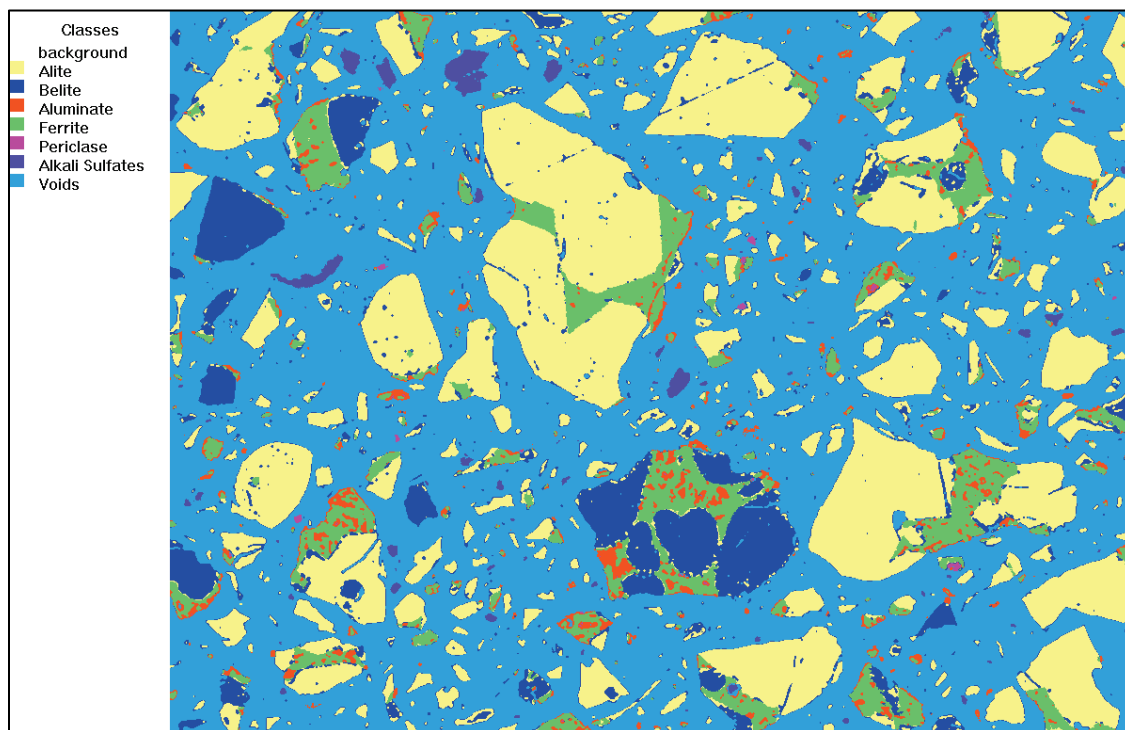


Figure 13. Calculated false color image with values based on image and labels from Figure 12.



3.3 VCCTL inputs

Input requirements into the program are described in Section 2.1. For the remainder of this report, a full example of input requirements and output data is discussed. Materials used in this virtual microstructure include a standard Type I/II cement with 20% replacement by fly ash, a 0.45 water-to-cementitious material (w/cm) ratio, 57 stone coarse aggregate, and standard C33 sand fine aggregate.

Selection of materials that are already present in the model database is typically straightforward and can be accomplished using the VCCTL GUI. The GUI allows for selection of a digitized image, particle size distribution file, and chemical composition. The selection can be features of the same material or comprised of multiple material data entries. For example, the user can choose all three files for a single cement or use the chemistry of one cement, the image of a second cement, and the particle size distribution of a third cement. This allows for some experimentation to investigate cement property relationships to performance.

In the case of supplementary cementitious materials, the user only need input particle size distributions and very basic chemistry data, and for

aggregates only, basic shape information and particle size is required. Due to the simpler inputs, a supplementary cementitious material or aggregate can be entered “on the fly” as the user is selecting the mix proportions. Cement data entries must be made via command line scripts, or uploading property files prior to using the VCCTL GUI.

After all materials are entered, the virtual mixing process begins. The user can specify the desired proportions of each material, the desired w/cm ratio, length of reaction time, and which outputs to be created. The processing time of the program is directly related to length of reaction time and outputs.

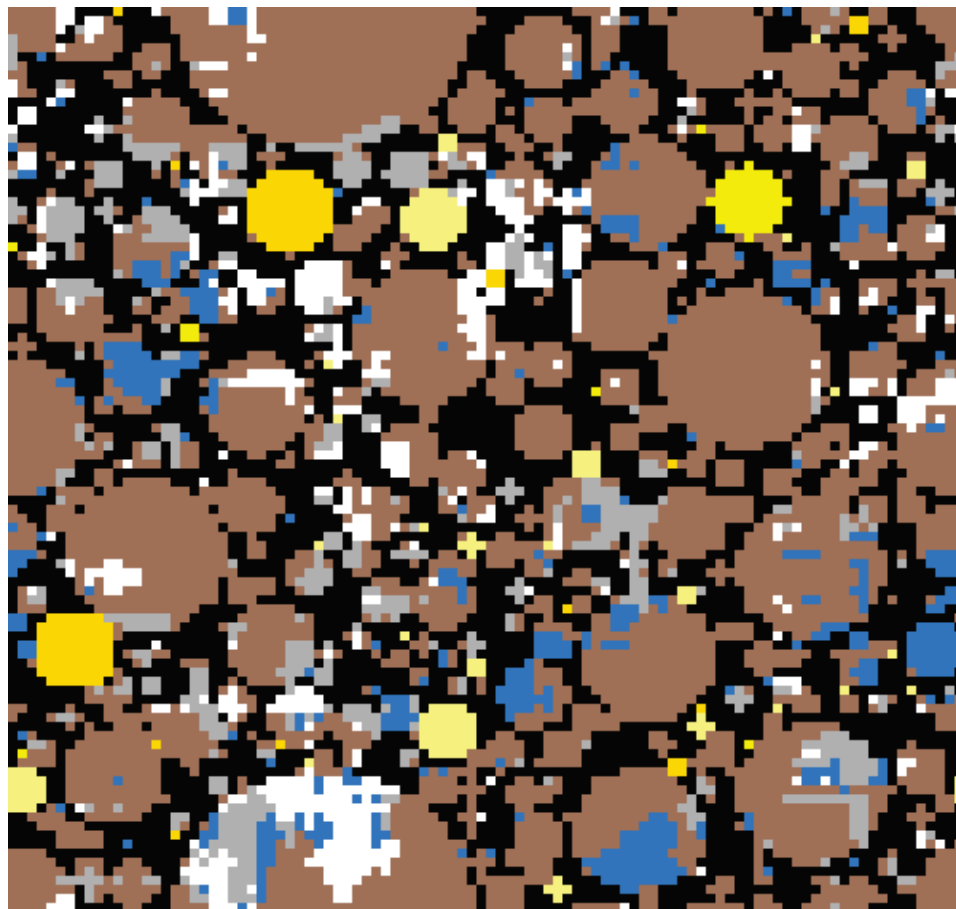
4 VCCTL Model Outputs

4.1 Property data outputs

Outputs for the program include microstructure image files and mechanical and transport property data files. The mixture is converted into a 100 by 100 by 100 voxel cube, and each voxel is given a color to represent the phase present in that voxel. Microstructure image files are provided as 2-D slices of the 3-D microstructure. One hundred images are created such that each represent one voxel depth of the cube. Figure 14 shows the first voxel depth of a virtually generated unhydrated microstructure.

The mixture design shown includes a standard CCRL type I/II cement, silica fume (ID #140092), and silica sand. The CCRL cement was already included in the program while the silica fume and silica sand were characterized at the ERDC and added to the program database. The mixture contained 38% cementitious material by mass (80% cement and 20% silica fume) at a 0.45 w/cm ratio. This mixture was given the designation 140-900W-O-F50. The VCCTL legend for relating voxel colors to microstructure phases is shown in Figure 15. The same mixture and virtual microstructure in Figure 14 was hydrated using the program and is shown in Figure 16.

Figure 14. Unhydrated microstructure image of a cement paste for mixture 140-900W-0-F50.

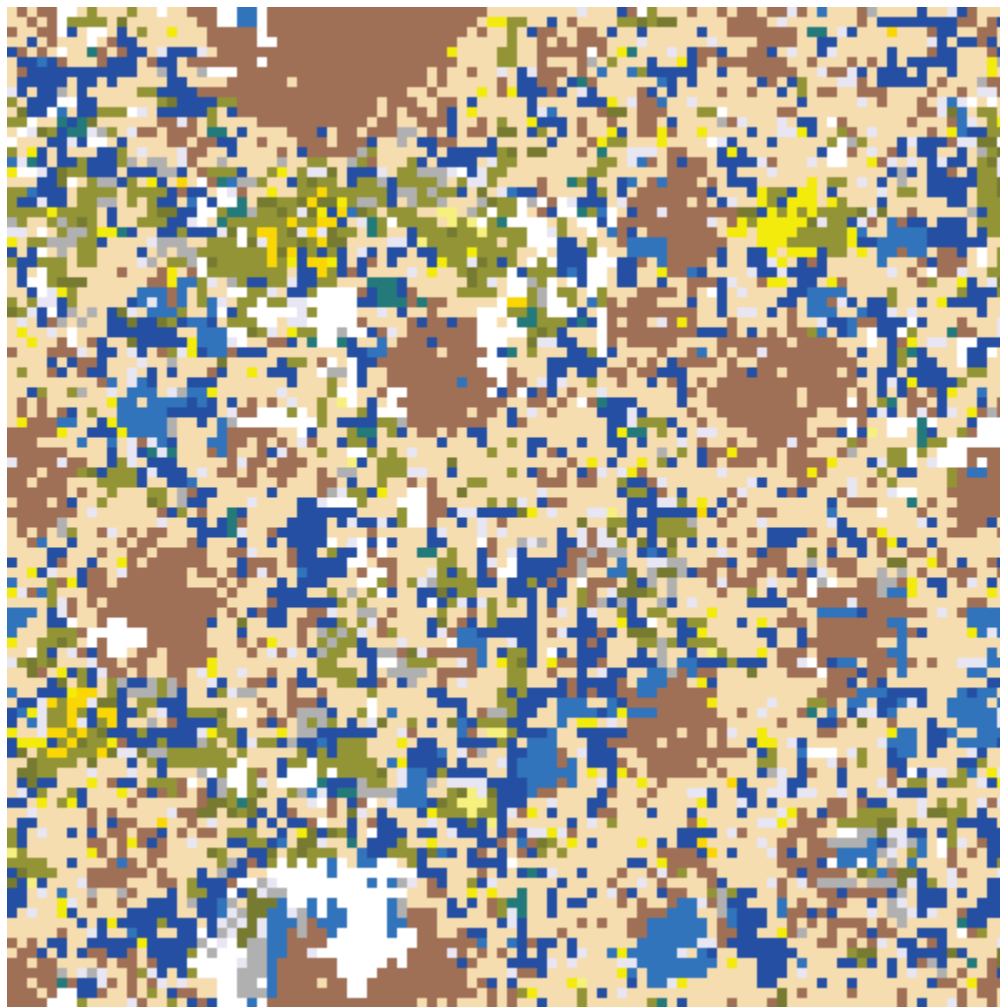


* Size is 100 x 100 micrometers

Figure 15. Legend for VCCTL microstructure image phases.

							
C_3S	C_2S	C_3A	C_4AF	K_2SO_4	Na_2SO_4	Gypsum	Hemihydrate
							
$CaCO_3$	Free Lime	Silica	Aggregate	CAS_2	Slag	ASG	Inert
							
Ettringite	AFm	Stratlingite	$CaCl_2$	Friedel Salt	C-S-H	Slag C-S-H	Pozz C-S-H
							
					FH_3	Brucite	MS
							
						Other Solid	Empty Porosity

Figure 16. Hydrated microstructure image of a cement paste for mixture 140-900W-O-F50.



* Size is 100 x 100 micrometers

As seen in Figure 17, the CSH portion forms during hydration and encompasses most of the available pore space from the unhydrated image (Figure 14). CSH growth can only occur at the edge of unhydrated, but potentially reactive, materials and must occur one voxel at a time. Essentially, the model investigates constituent material boundaries and performs a numerical check to determine if the materials react and, if a reaction occurs, what phase forms in the voxel along that boundary. If a reaction occurs, checks then continue using the reacted voxel material and its adjacent voxels until no new reactions are predicted.

The hydrated microstructure can also be used to calculate additional outputs such as degree of hydration (Figure 17), porosity (Figure 18), gel-space ratio (Figure 19), and heat release (Figure 20). Overall, 68 property

measurements can be obtained using the hydration microstruture image. The following four figures illustrate the aforementioned properties over the first 20 hr for the virtual microstructure 140-900W-o-F50.

Figure 17. Degree of hydration over the first 20 hr for mixture 140-900W-0-F50.

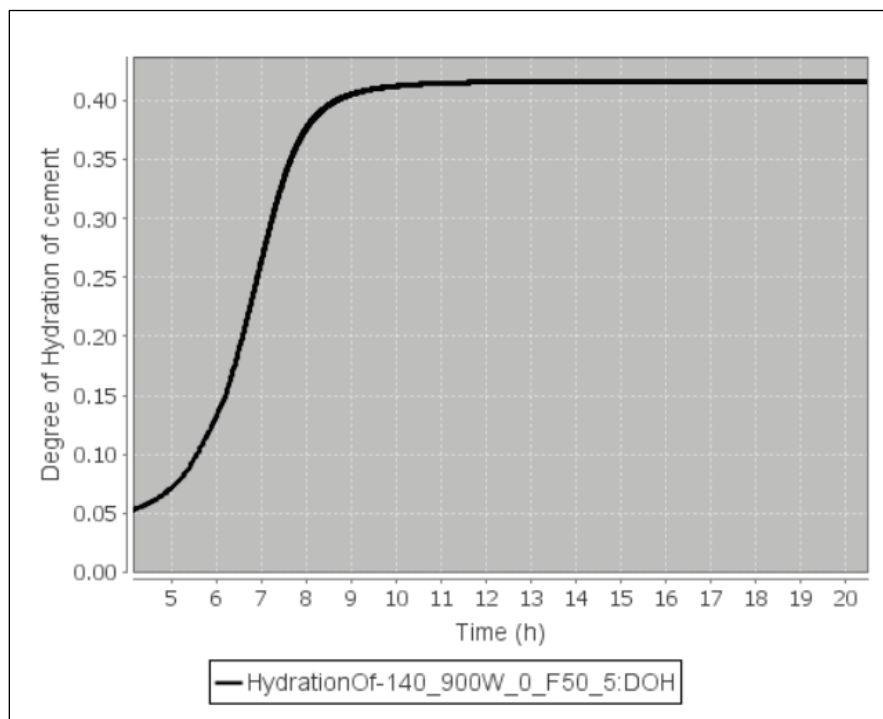


Figure 18. Porosity over the first 20 hr for mixture 140-900W-0-F50.

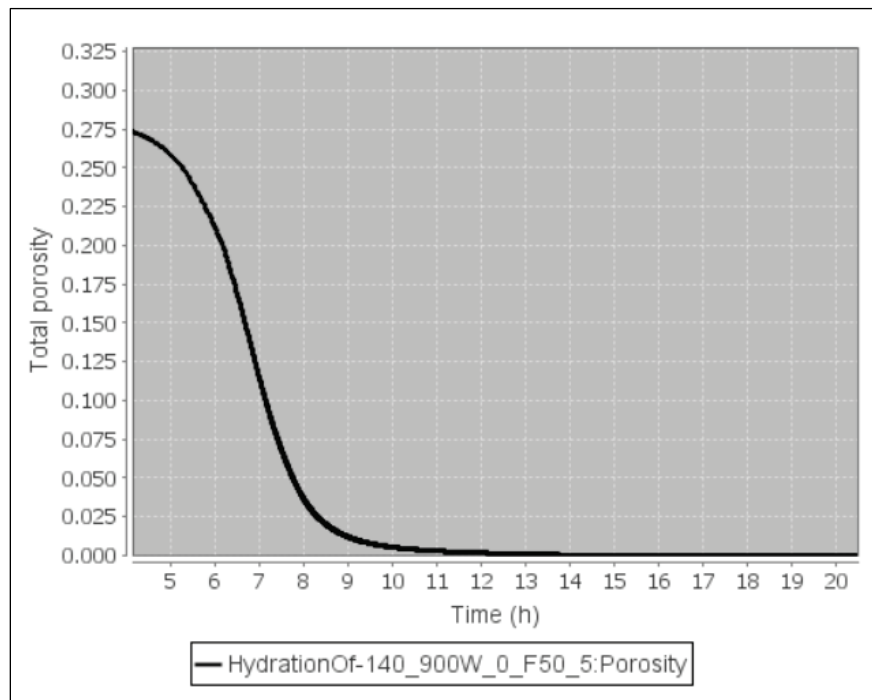


Figure 19. Gel-space ratio over the first 20 hr for mixture 140-900W-0-F50.

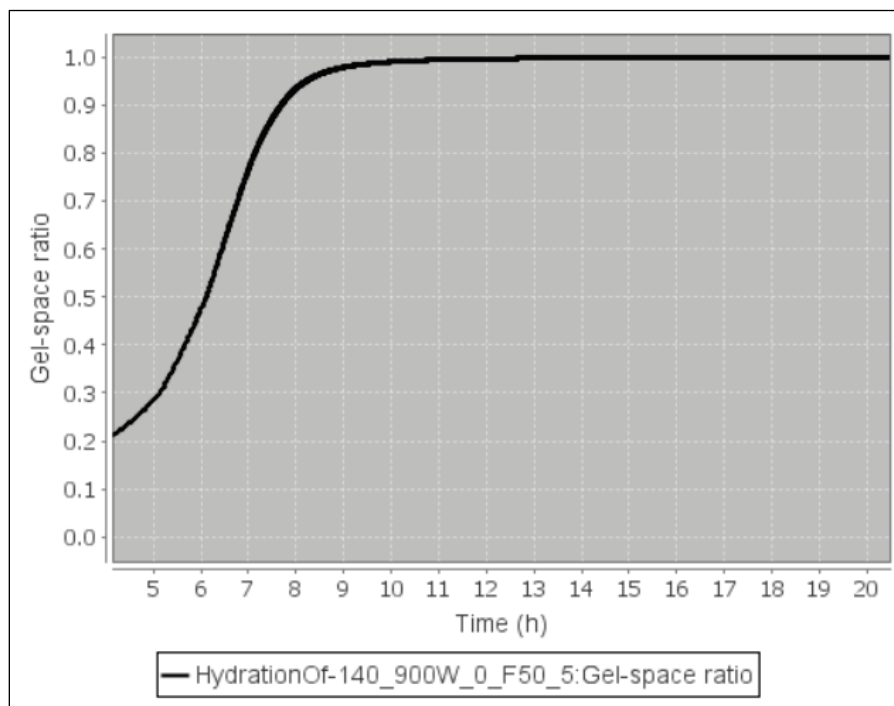
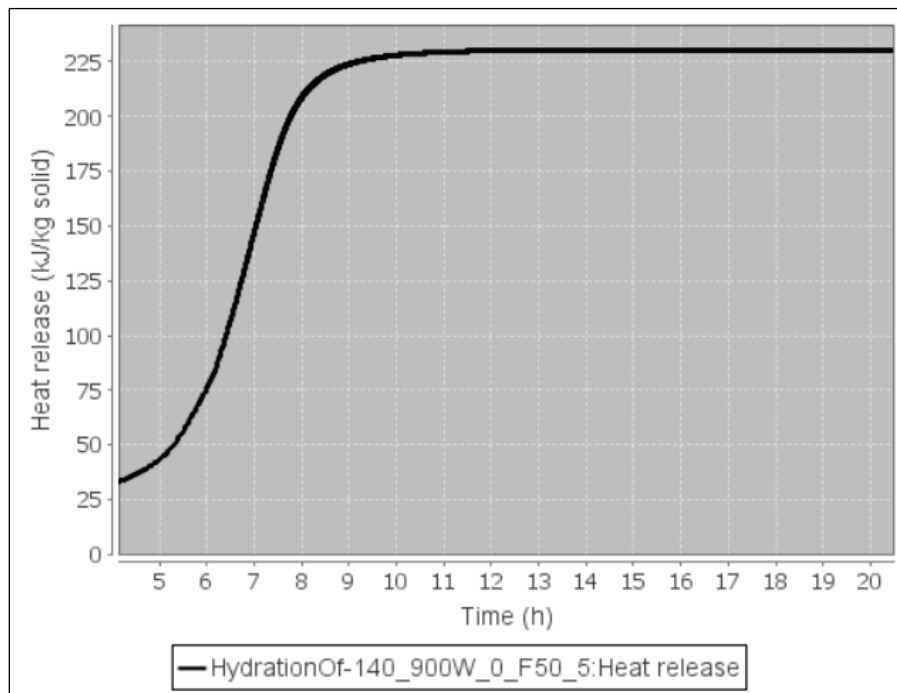


Figure 20. Heat release over the first 20 hr for mixture 140-900W-0-F50.



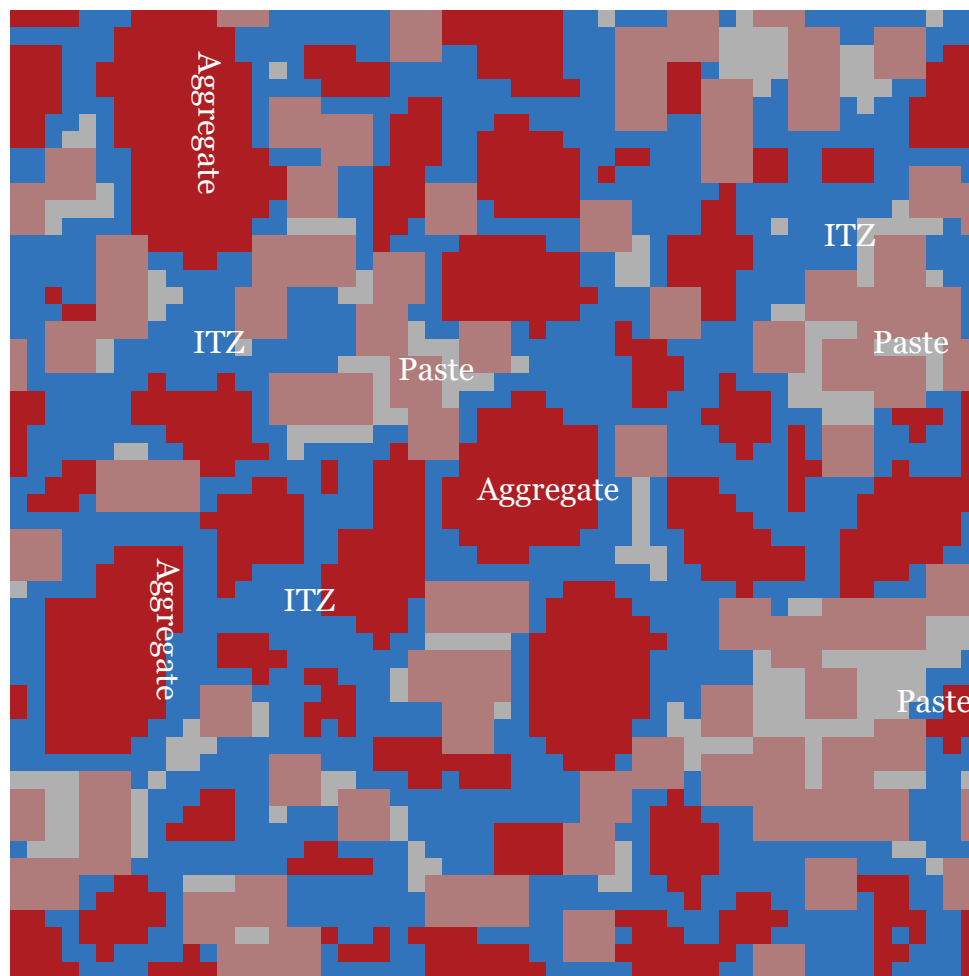
Moduli results are calculated and provided in tabular format. Moduli results from the sample microstructure are shown in Figure 21. Aggregate results are produced separately due to scaling issues. Aggregates are processed using a random number generator and predicted density relationships. The aggregate microstructure image is shown in Figure 22. The image is in the same style as the cement paste in that the image is a 2-D representation of 1-voxel depth slides of a 3-D cube. In this image, the large red particles represent coarse aggregates, the small dark red particles represent fine aggregate, and the gray represent a continuum, which is the cement paste.

Figure 21. VCCTL calculated moduli results for mixture 140-900W-0-F50.

CEMENT PASTE ELASTIC MODULI:

bulk_modulus 32.1 GPa
shear_modulus 16.9 GPa
Youngs_modulus 43.1 GPa
Poissons_ratio 0.27

Figure 22. Virtual microstructure of aggregate for mixture 140-900W-0-F50.



* Size is 2.28 mm x 2.28 mm; each pixel represents .04 mm

4.2 Data outputs for next level models

The overarching goals of the MMFP project include a materials-by-design focus in which it is possible to transition from micro-sized grains to full-scale concrete elements and vice versa with predictable performance properties being measurable throughout the entire process. To achieve this, the results of the VCCTL program must be compatible in higher length scale models. VCCTL features text-based image outputs for both the aggregate and cement paste microstructure files. These outputs contain a list of one million numbers, each representing a voxel in the 100 by 100 by 100 voxel cube. These numbers represent a certain phase of the mixture and are ordered so that the cube could be reconstructed using only the text number file. These files along with the calculated physical properties are thought to be key in transitioning to a higher-level model.

For example, a text hydrated microstructure file that begins with the numerals 1, 1, 0, 0, 4 represents consecutive pixels of C_3A , C_3A , porosity, porosity, C_4AF . Reconstructing the entire 1 million number file into a 3-D cube in another scalable program, such as a finite element or lattice discrete particle model, would allow for further research into effects on crack propagation and applied forces. Possible methods to achieve this are currently being studied in the next phase of the MMFP research.

5 Conclusions

Overall, the MMFP program has already increased capabilities in material characterization and the properties being collected may provide substantial benefit and time savings to other programs. VCCTL appears to have the ability to accurately predict hydrated microstructure from raw material properties and fills a much-needed role in the materials-by-design process. Moving on to higher scale models and correctly incorporating hydration model results will be key in progress moving forward.

The appendices provide processed characterization data for XRD, XRF, and particle-size experimentation results. The appendices represent the current data availability at the time of this report, but do not reflect the total amount of data to be collected over the course of the research effort. At the time of this report, the characterization data archive features the processed results for 29 particle size distributions, 27 XRF experiments, and 16 XRD experiments.

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References

- American Society of Testing and Materials (ASTM). 2018. Standard test method for determination of the proportion of phases in portland cement and portland-cement clinker using x-ray powder diffraction analysis. Designation C1365-18. West Conshohocken, PA: ASTM International.
- Bentz, D. P. 1995. A three-dimensional cement hydration and microstructure program. Hydration rate, heat of hydration, and chemical shrinkage. NISTIR 5756. Washington, DC: U.S. Department of Commerce.
- Bentz, D. P. 1997a. Three-dimensional computer simulation of portland cement hydration and microstructure development. *Journal of the American Ceramic Society* 80(1):3-21.
- Bentz, D. P. 1997b. Guide to using CEMHYD3D: A three dimensional cement hydration and microstructure development modelling package. Gaithersburg, MD: Building and Fire Research Laboratory.
- Bentz, D. P., J. R. Clifton, and K. A. Snyder. 1996. A prototype computer-integrated knowledge system: predicting service life of chloride-exposed steel-reinforced concrete. *Concrete International* 18(12):42-47.
- Garboczi, E. J., J. W. Bullard, and D. P. Bentz. 2004. Virtual testing of cement and concrete. *Concrete International* 39(12)33-37.

Appendix A: Particle-Size Distribution Results

Figure A1. PSD Results of Cement (150146).

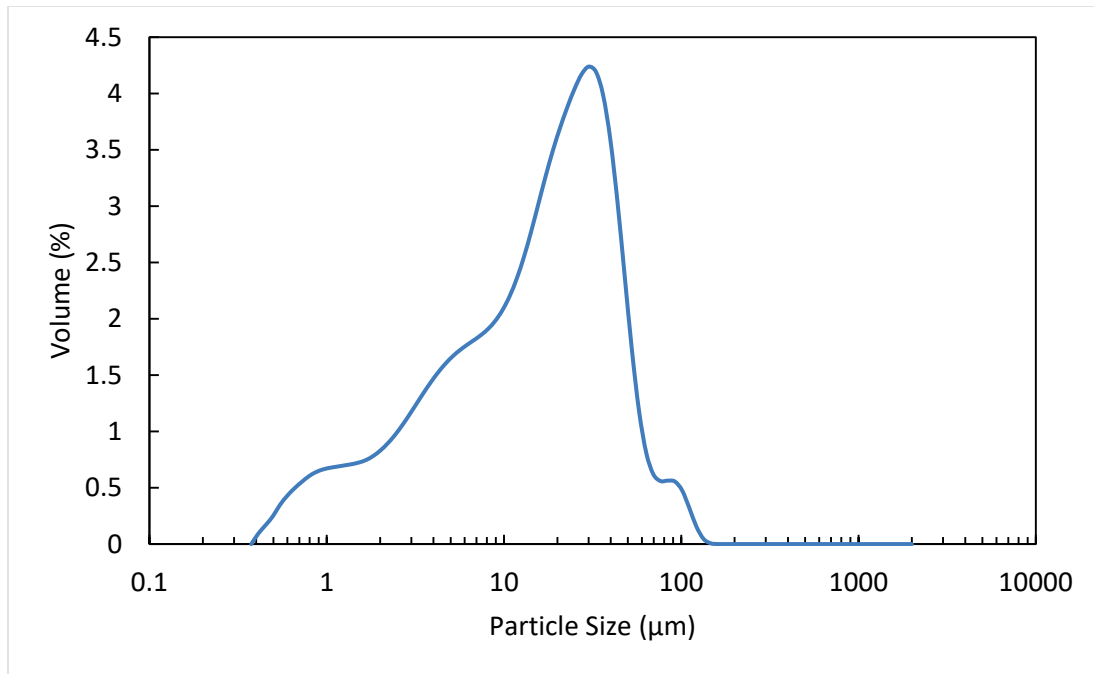


Figure A2. PSD Results of Cement (150147).

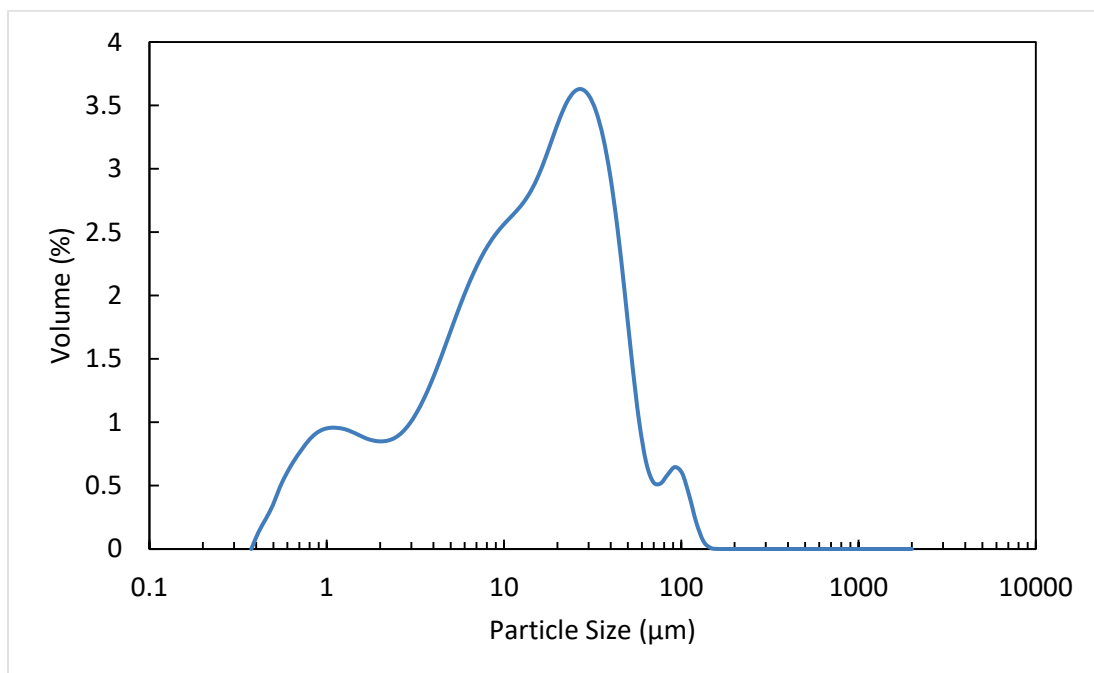


Figure A3. PSD Results of Cement (150148).

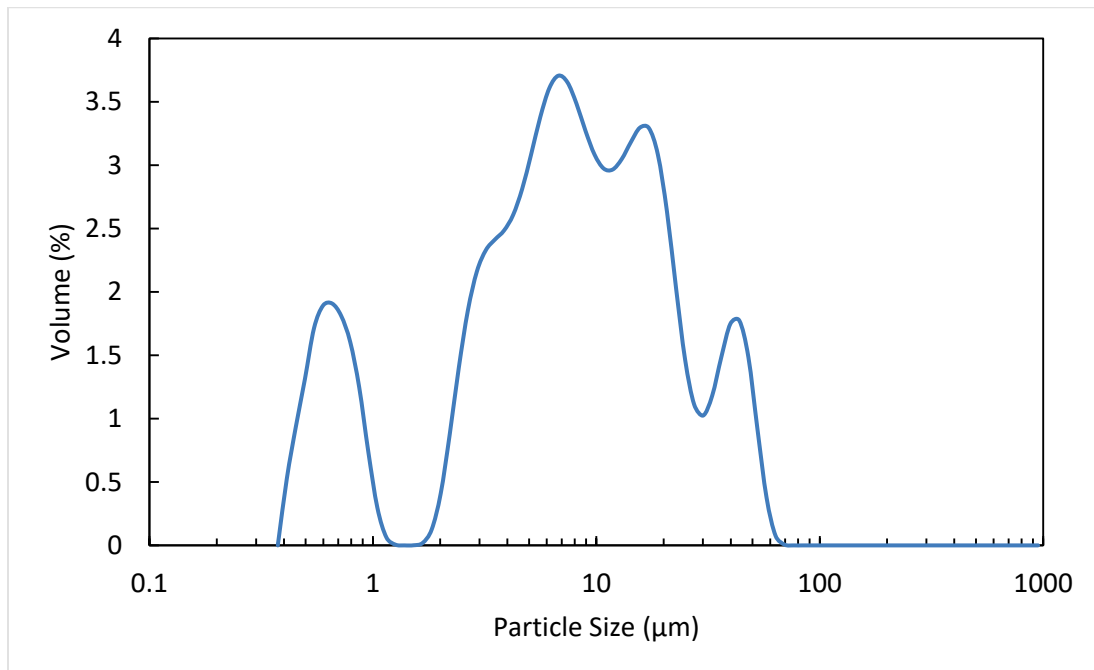


Figure A4. PSD Results of Cement (160160).

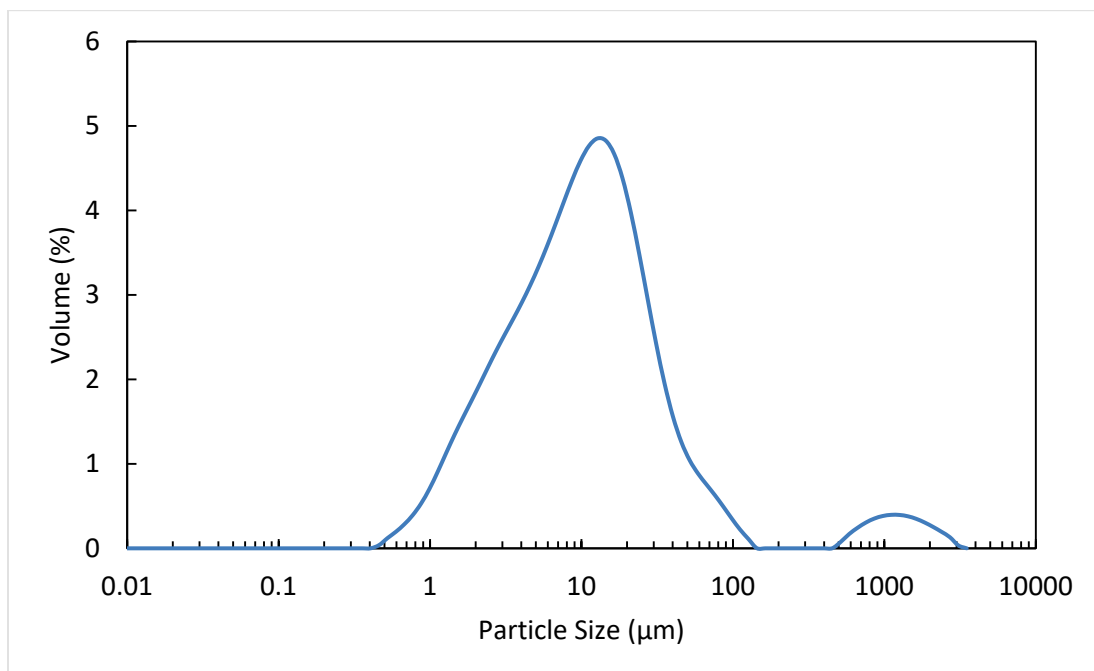


Figure A5. PSD Results of Cement (170001).

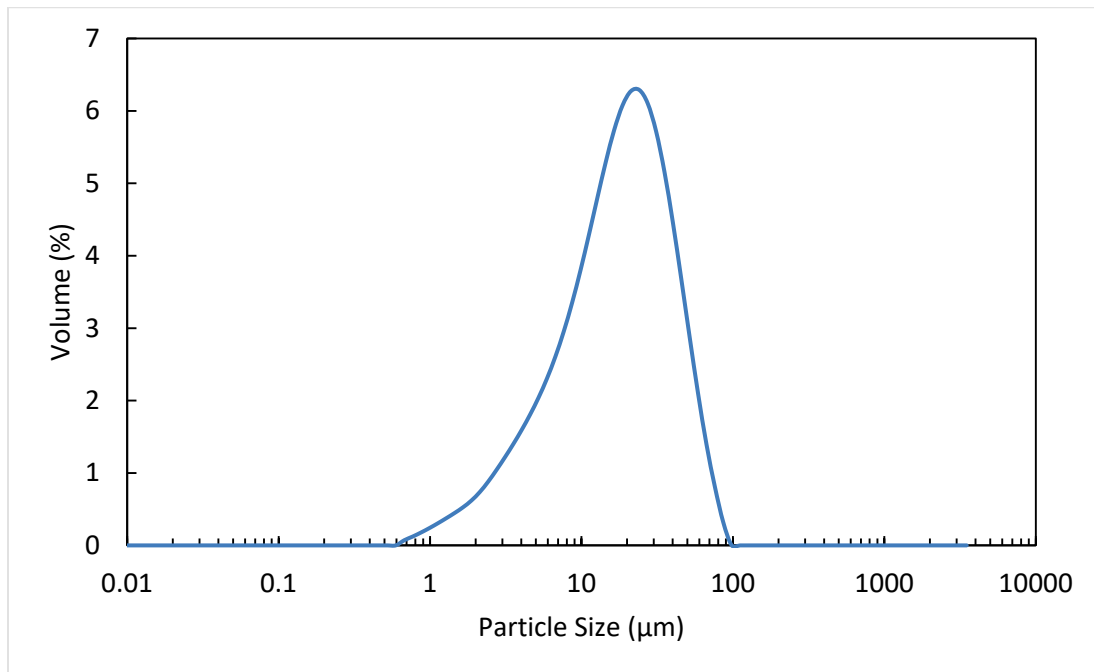


Figure A6. PSD Results of Cement (170002).

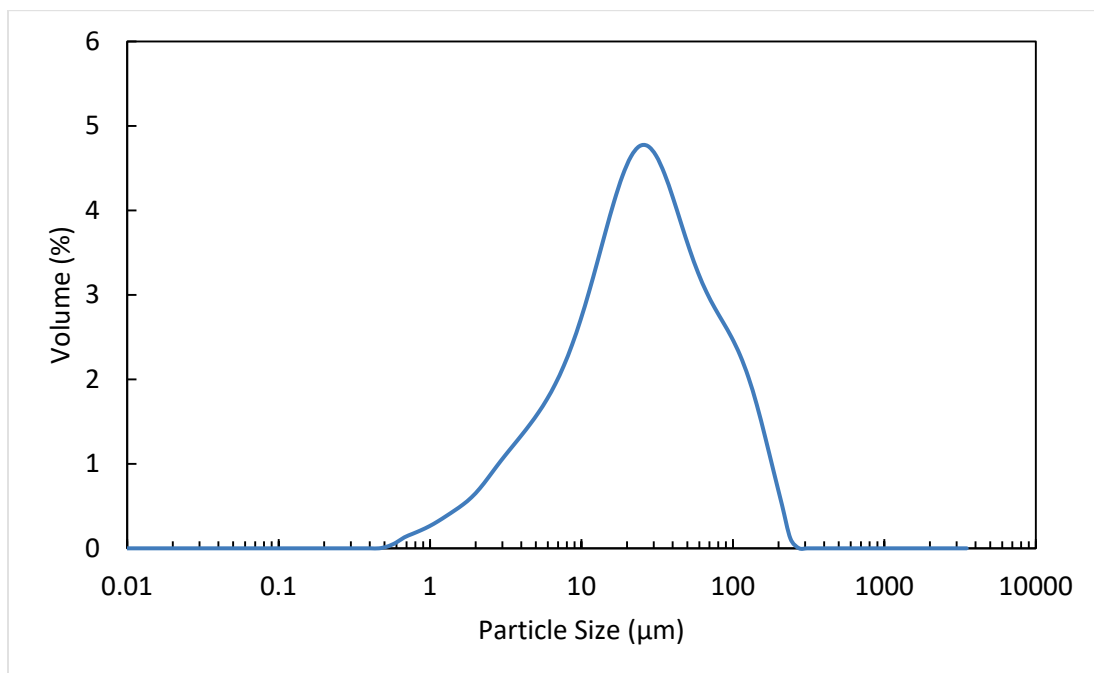


Figure A7. PSD Results of Cement (170009).

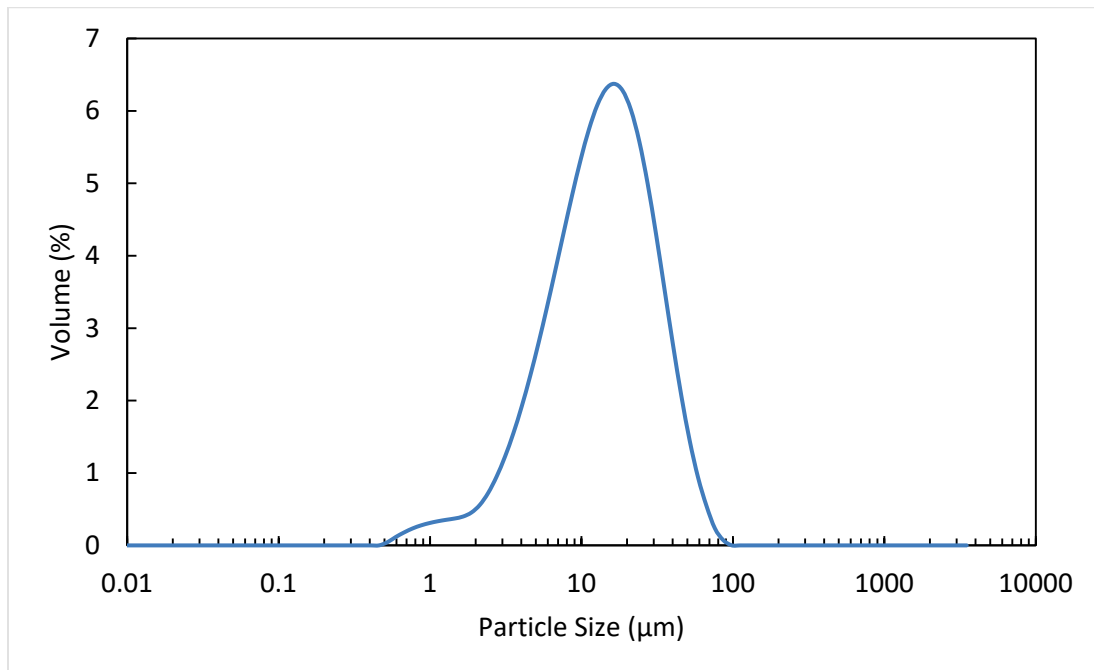


Figure A8. PSD Results of Cement (170010).

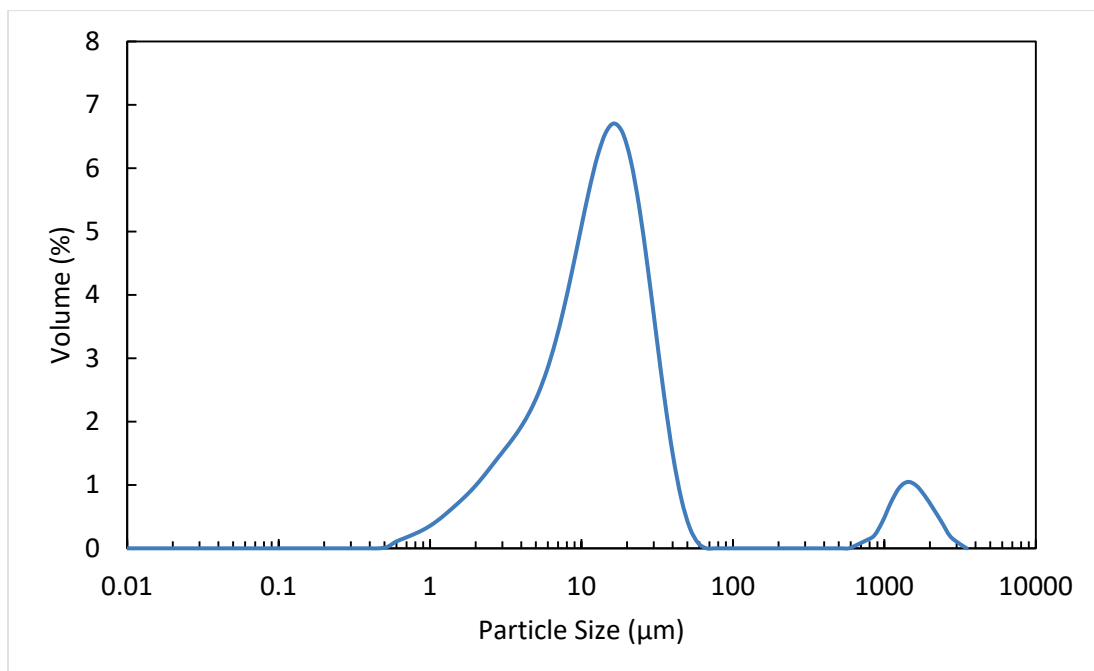


Figure A9. PSD Results of Cement (170052).

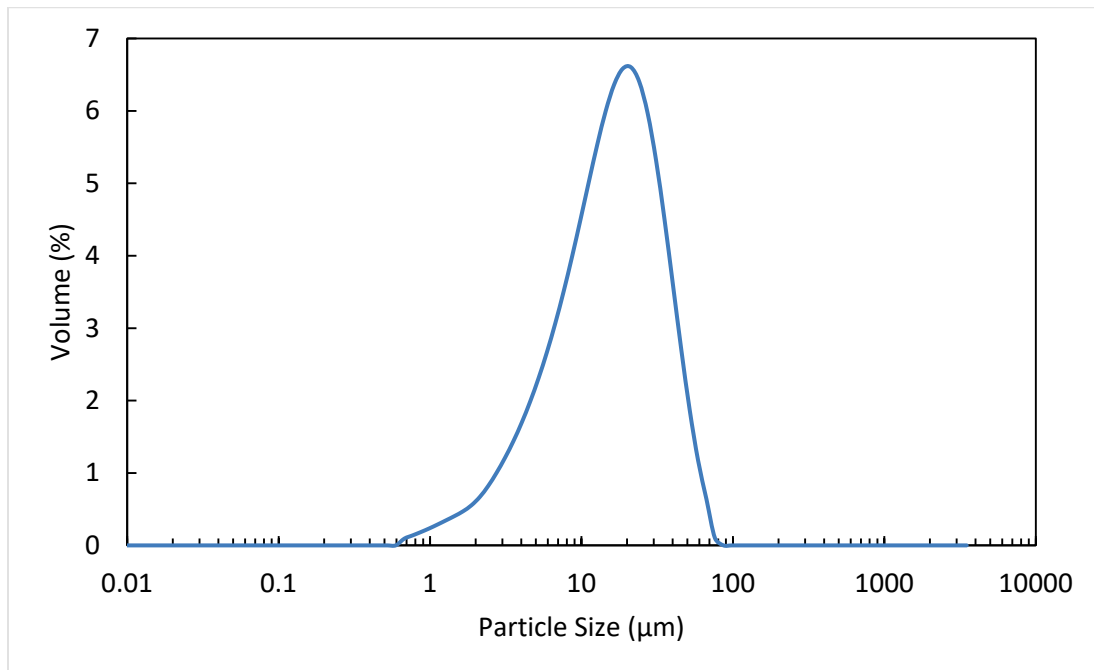


Figure A10. PSD Results of Cement (170065).

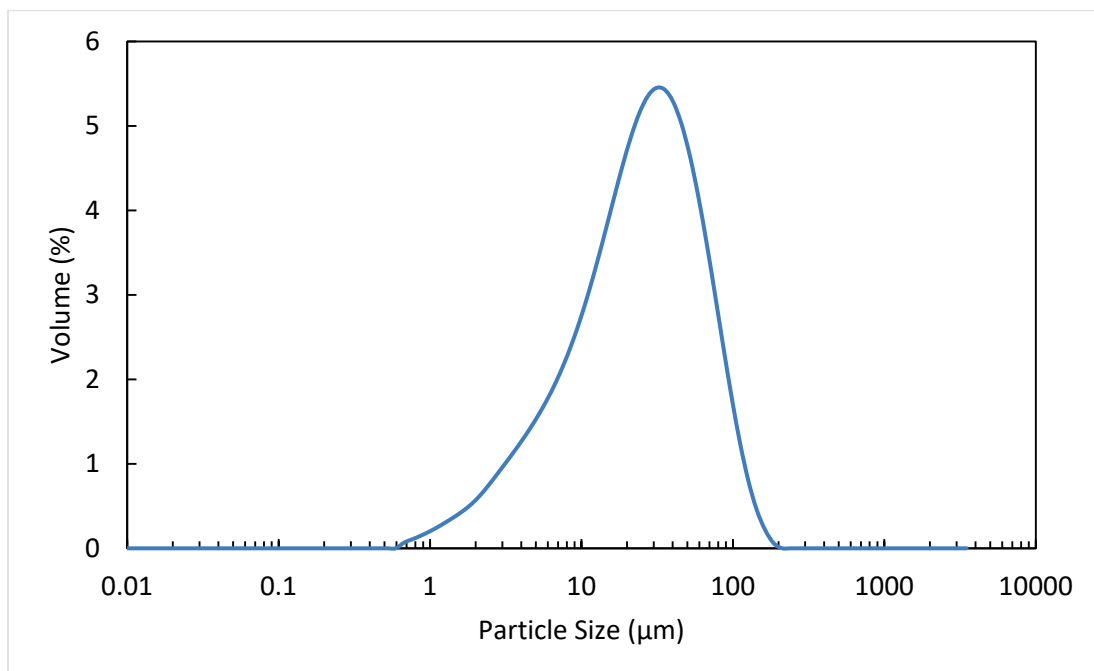


Figure A11. PSD Results of Metakaolin (140015).

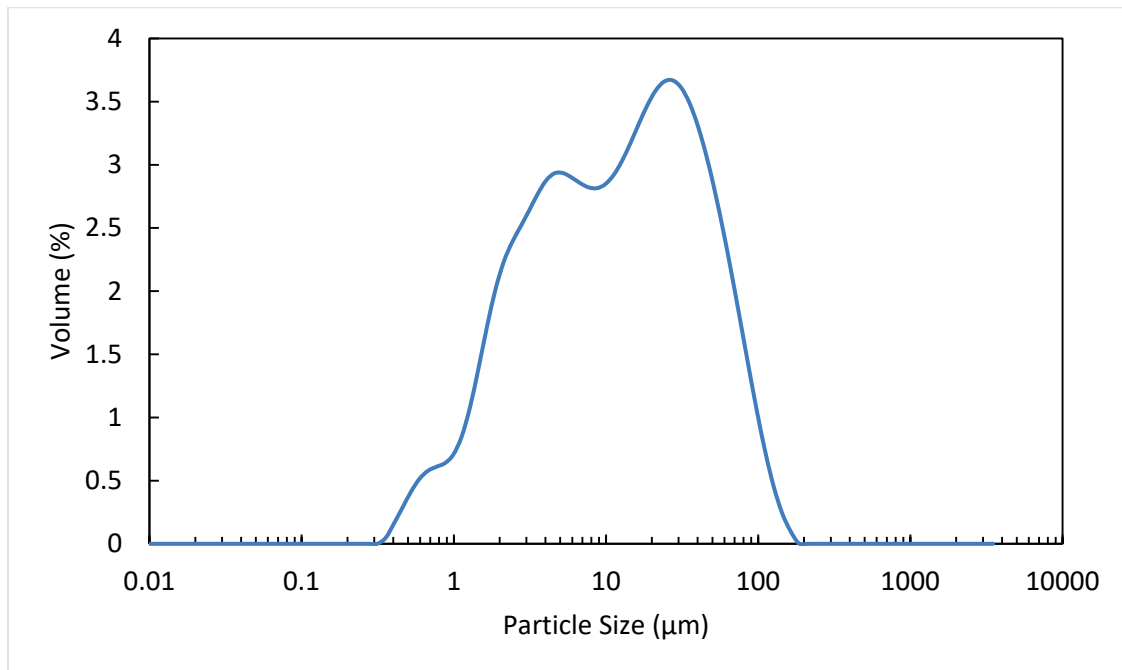


Figure A12. PSD Results of Metakaolin (150149).

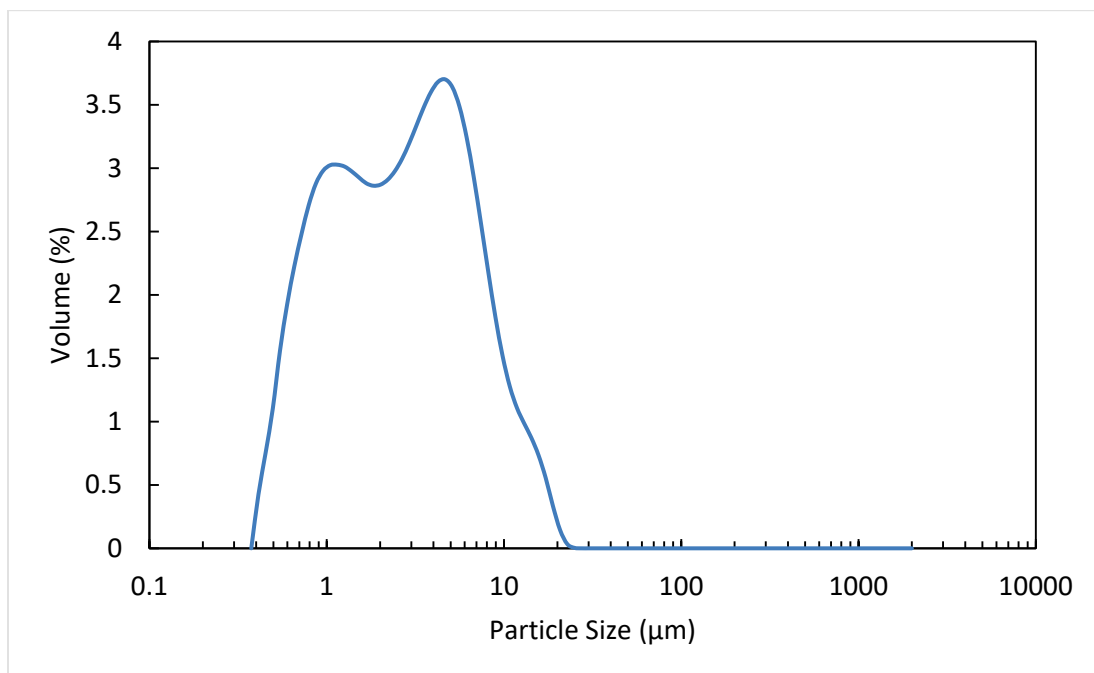


Figure A13. PSD Results of Metakaolin (170004).

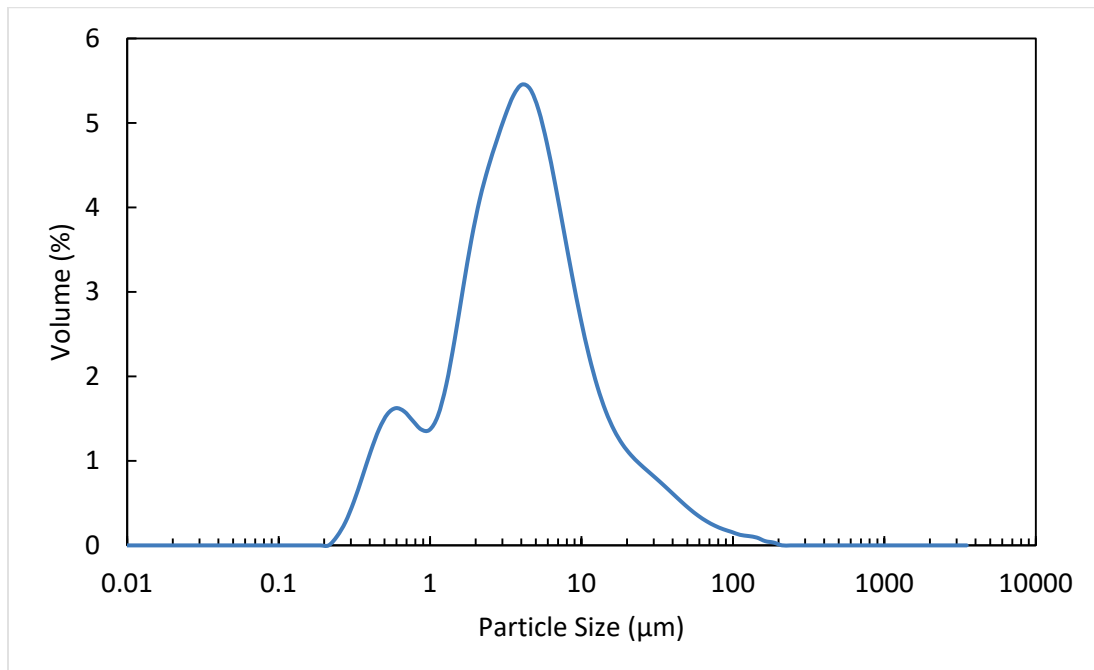


Figure A14. PSD Results of Silica Fume (130008).

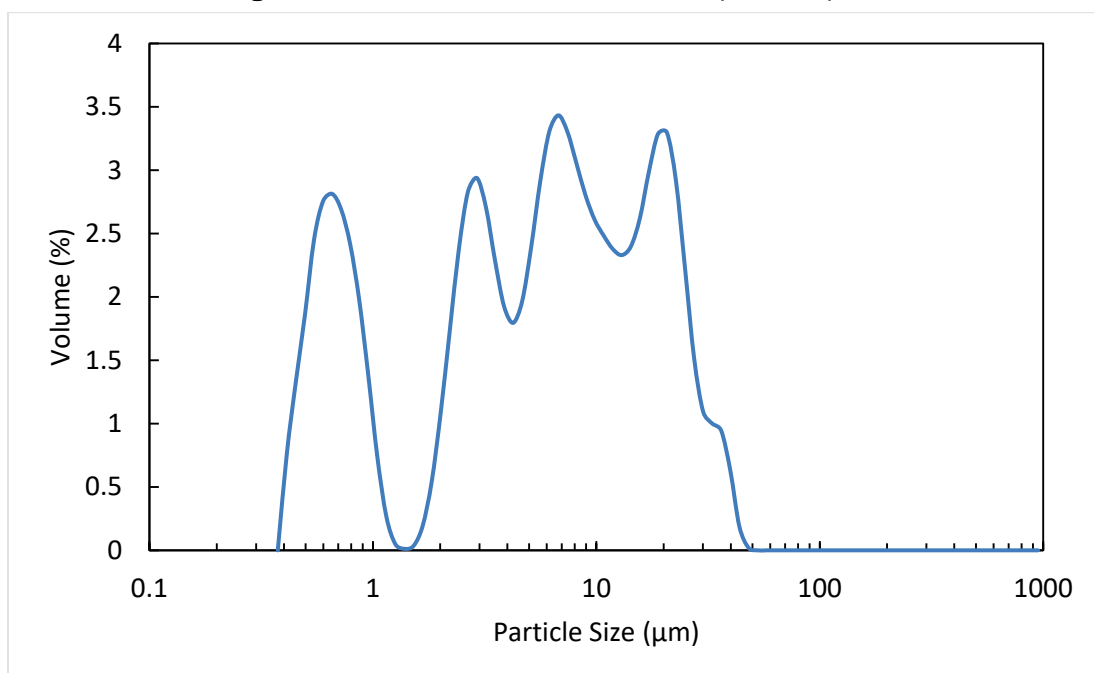


Figure A15. PSD Results of Silica Fume (140092).

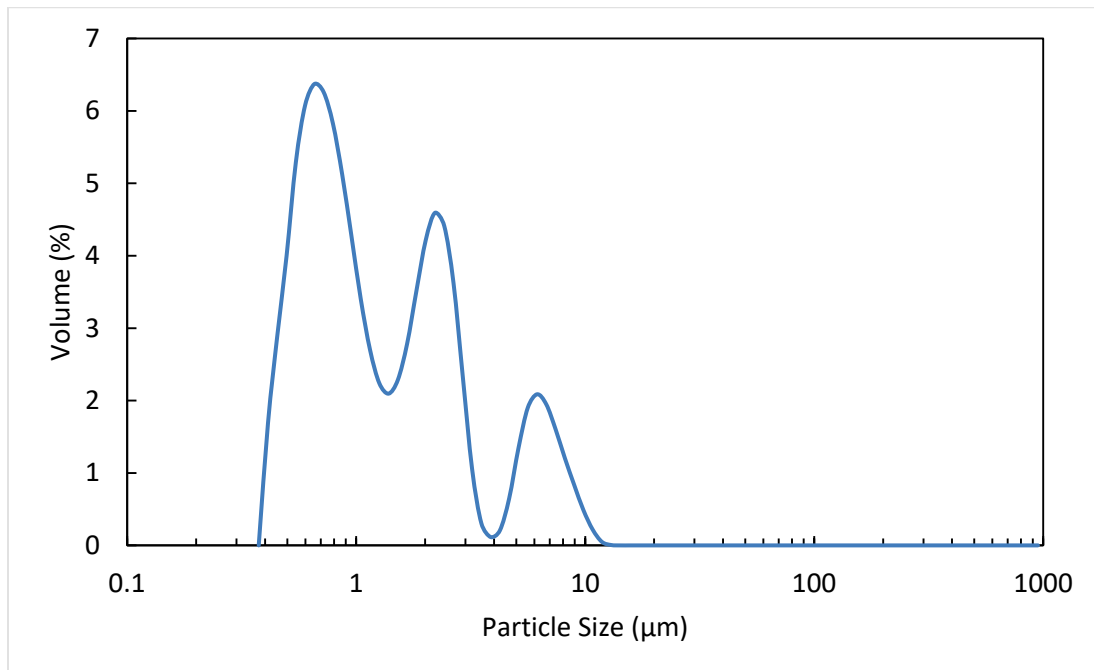


Figure A16. PSD Results of Silica Fume (130050).

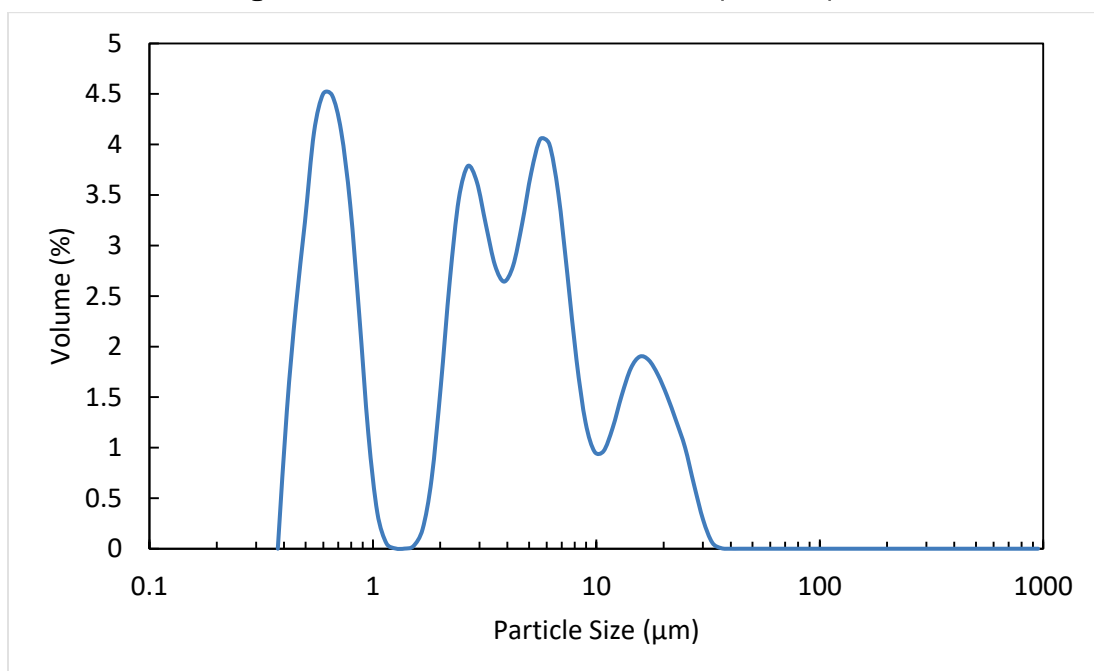


Figure A17. PSD Results of Silica Fume (140093).

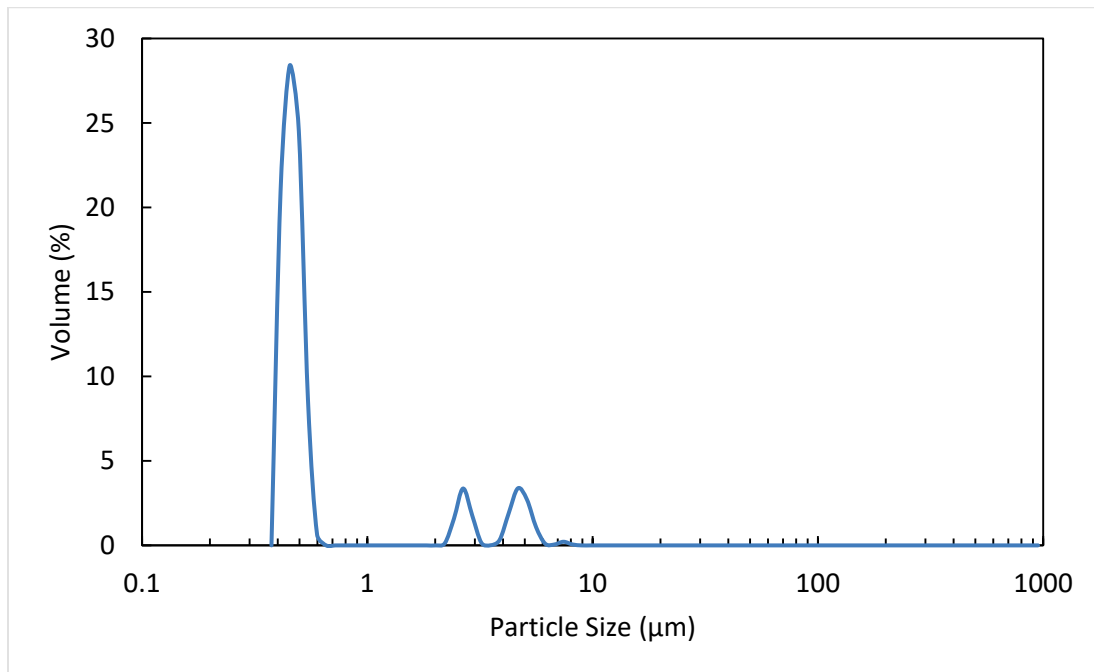


Figure A18. PSD Results of Silica Fume (140094).

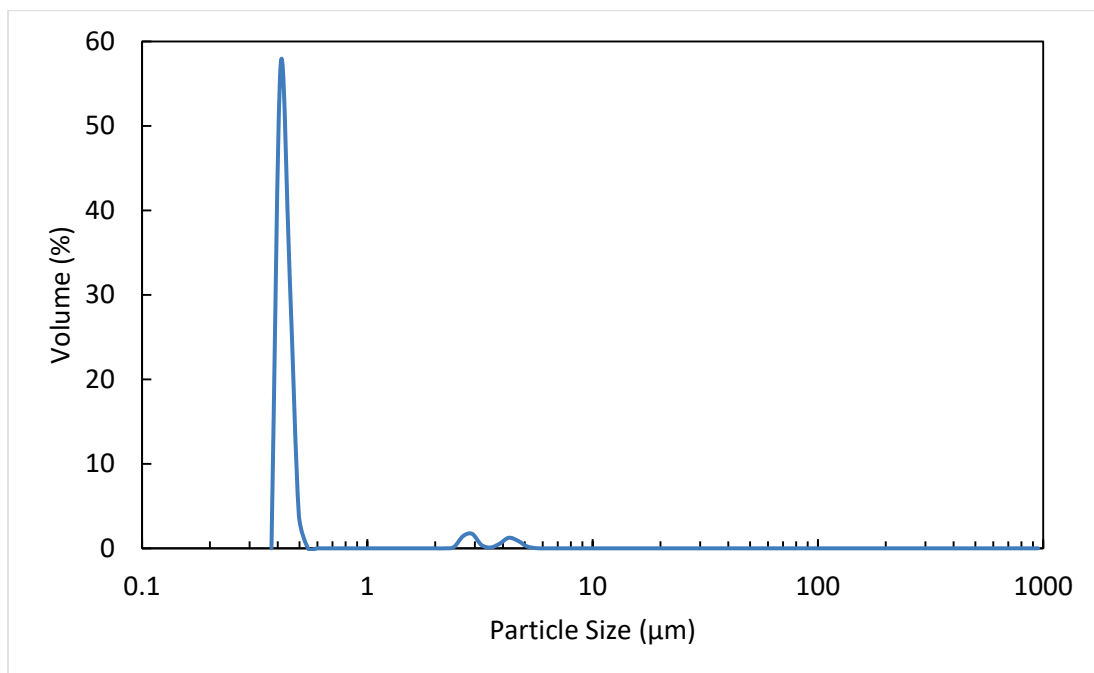


Figure A19. PSD Results of Silica Fume (150048).

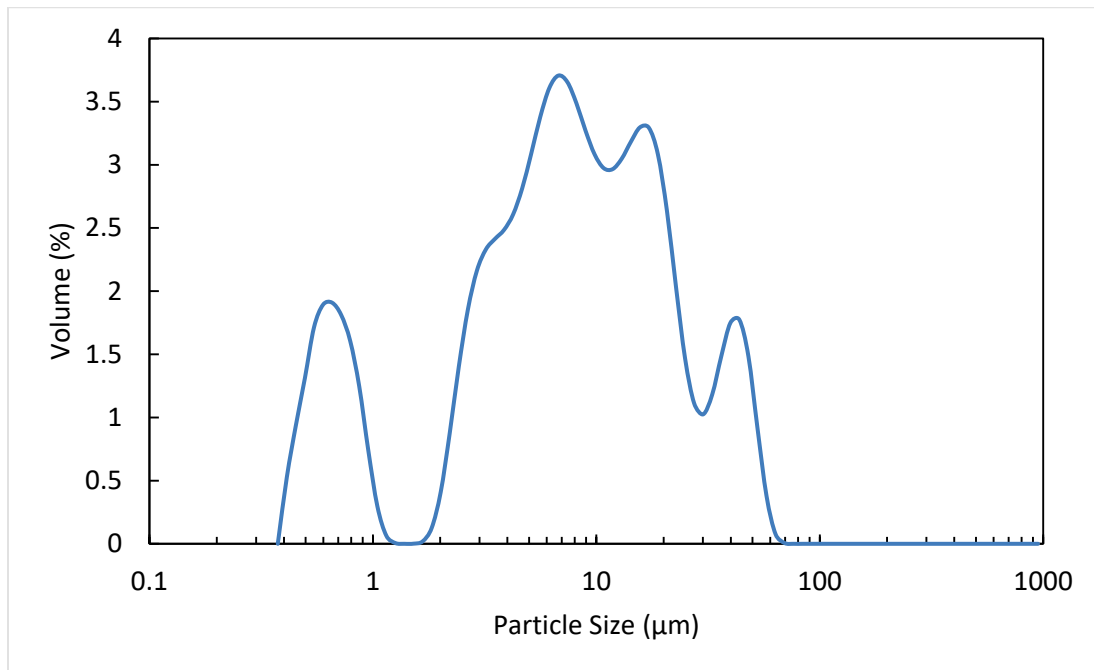


Figure A20. PSD Results of Silica Fume (150049).

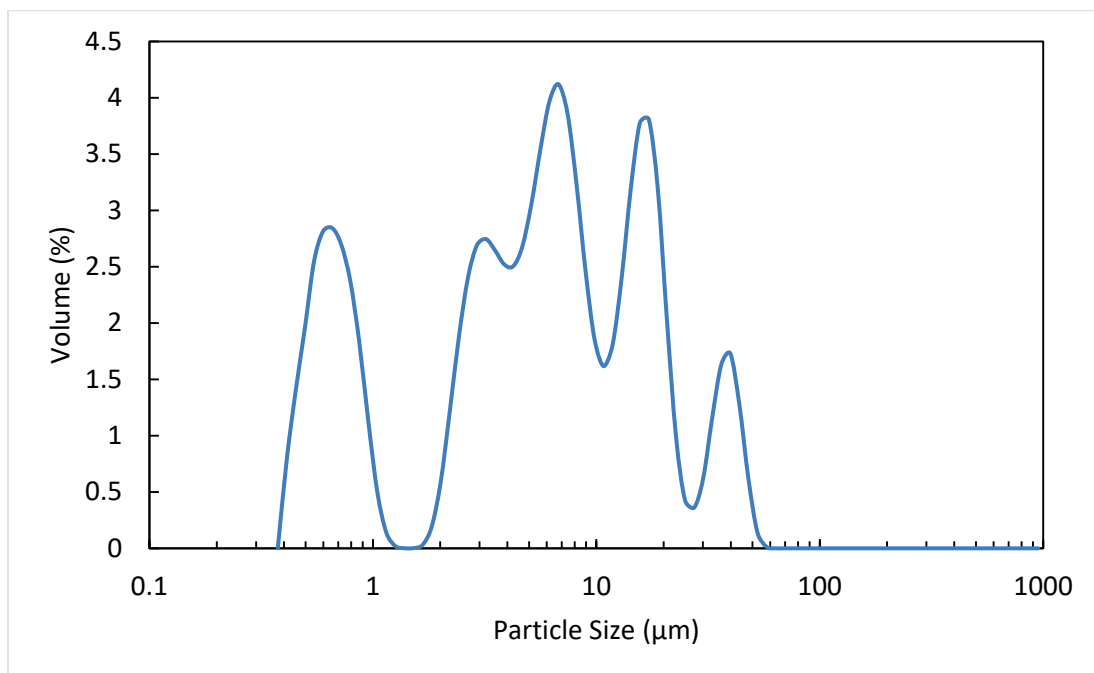


Figure A21. PSD Results of Silica Fume (150050).

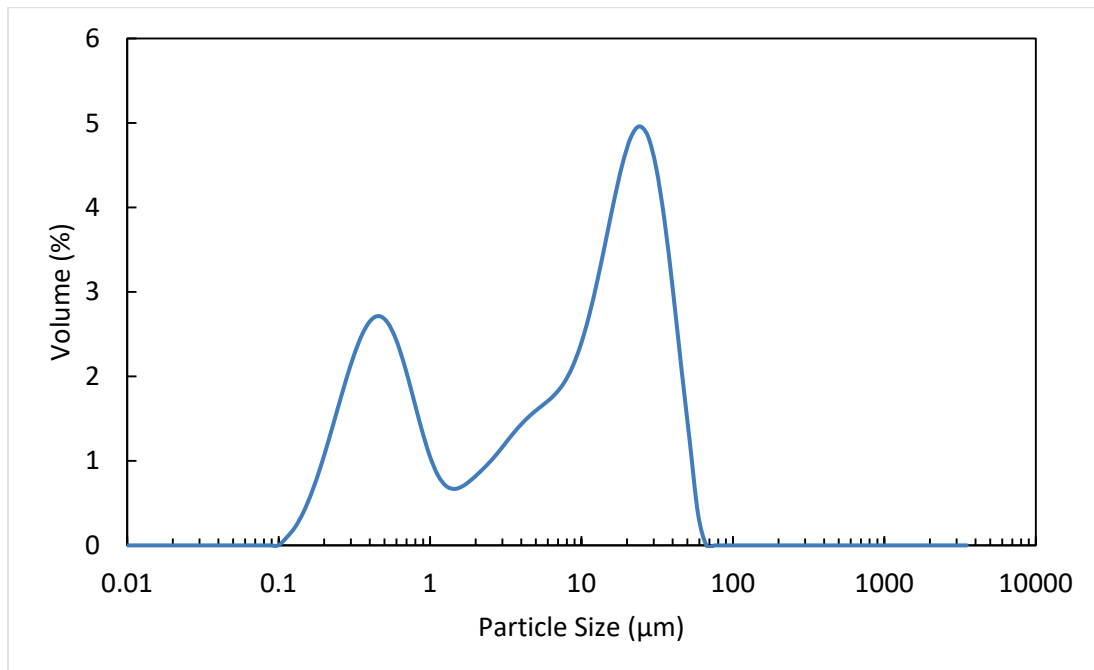


Figure A22. PSD Results of Silica Fume (150051).

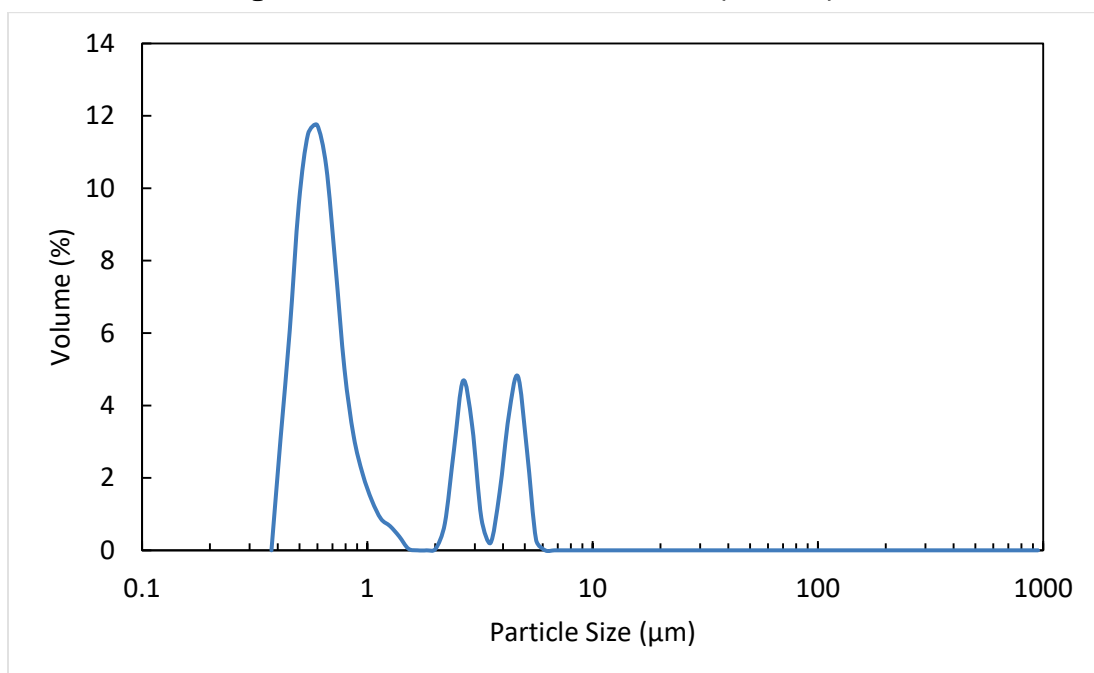


Figure A23. PSD Results of Silica Fume (150052).

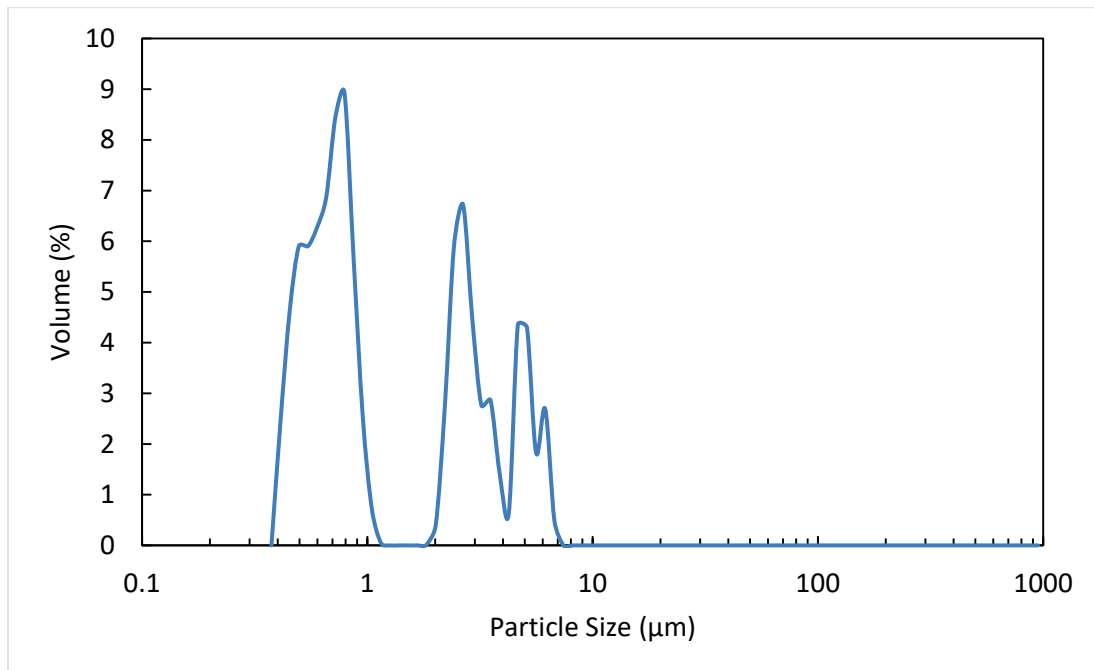


Figure A24. PSD Results of Silica Fume (170003).

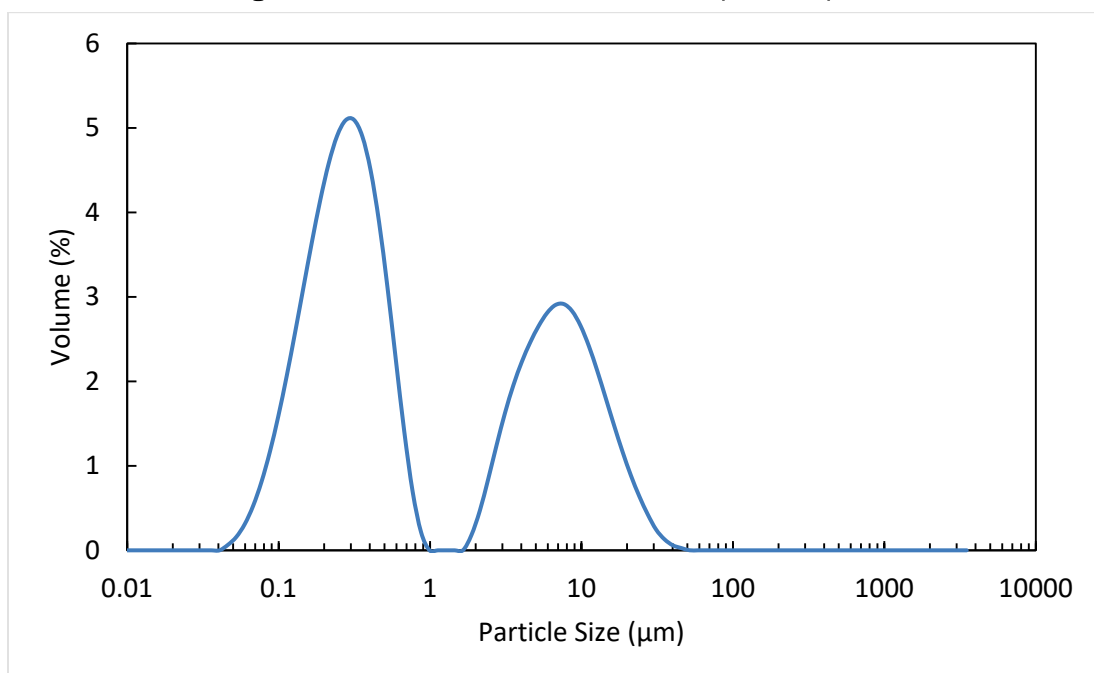


Figure A25. PSD Results of Silica Fume (Guam F10).

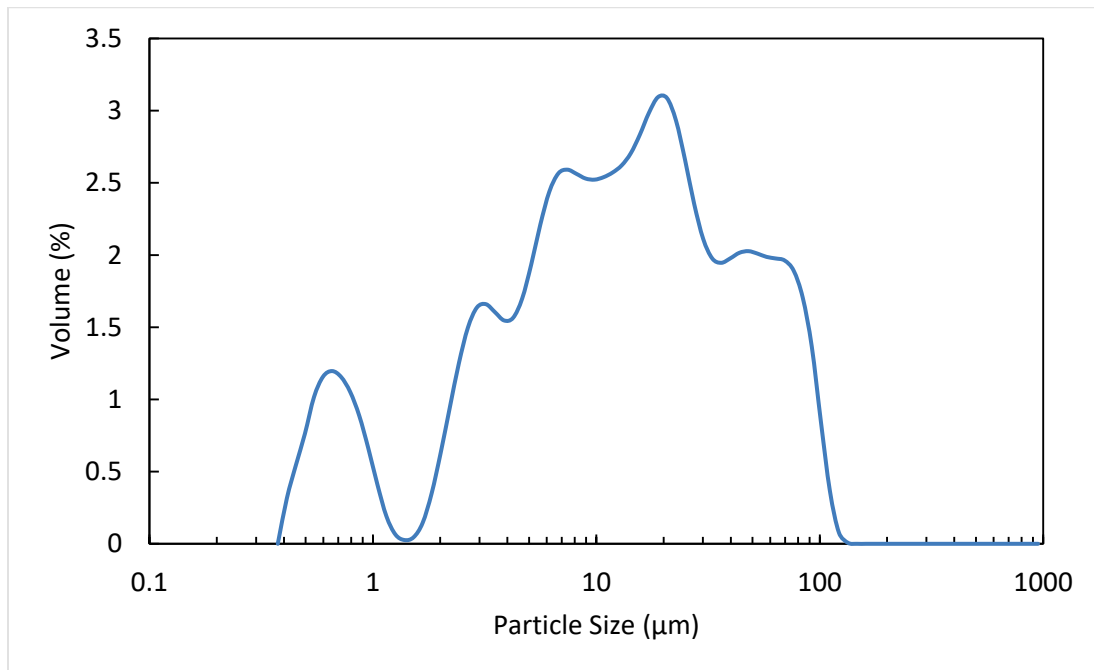


Figure A26. PSD Results of Silica Fume (Ecotech85).

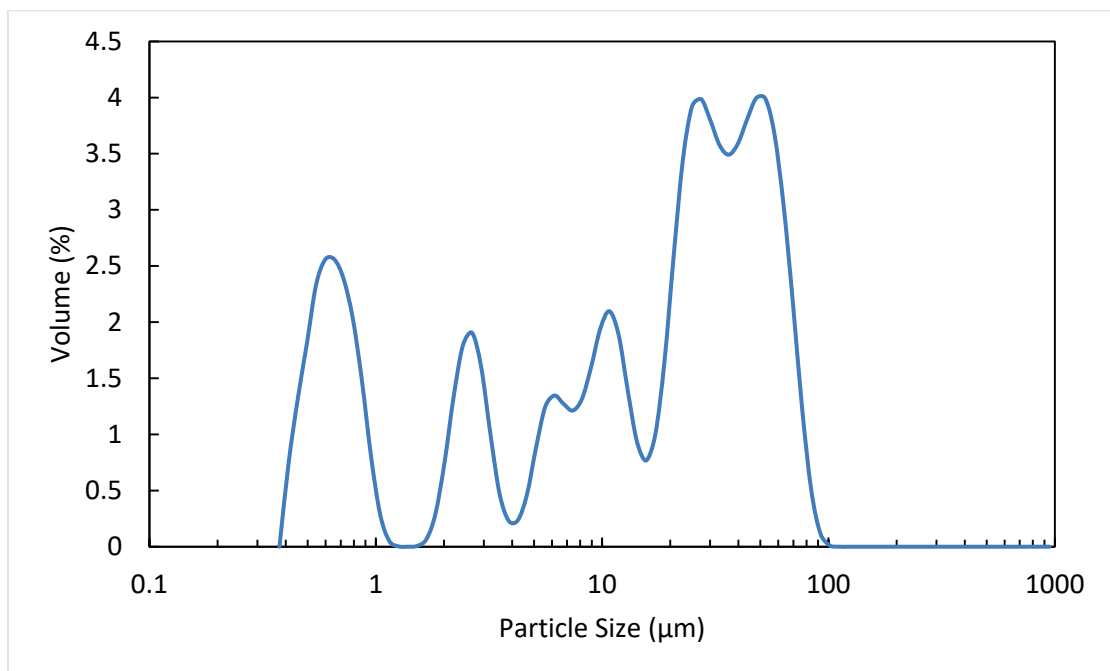


Figure A27. PSD Results of Silica Flour (Sil-co-sil).

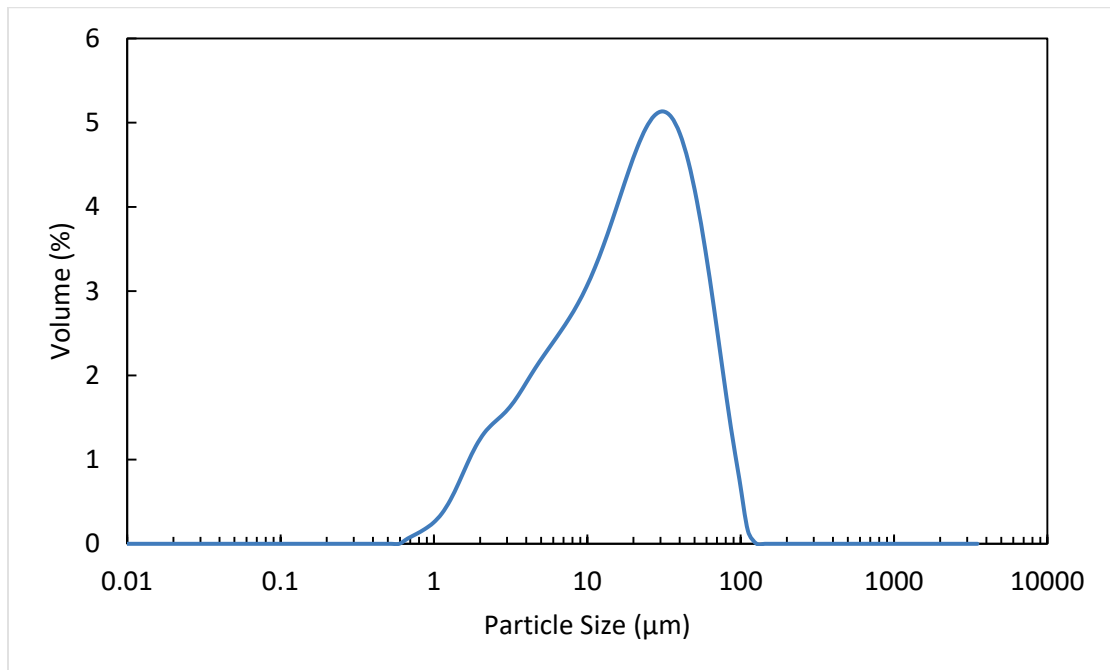


Figure A28. PSD Results of Silica Sand (F50).

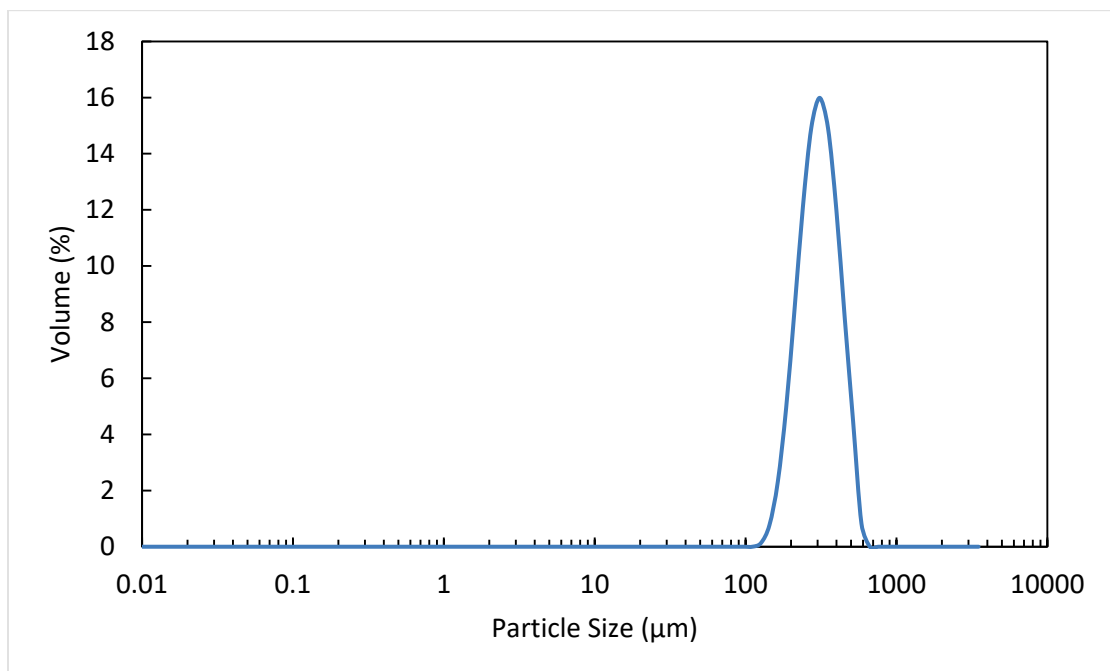
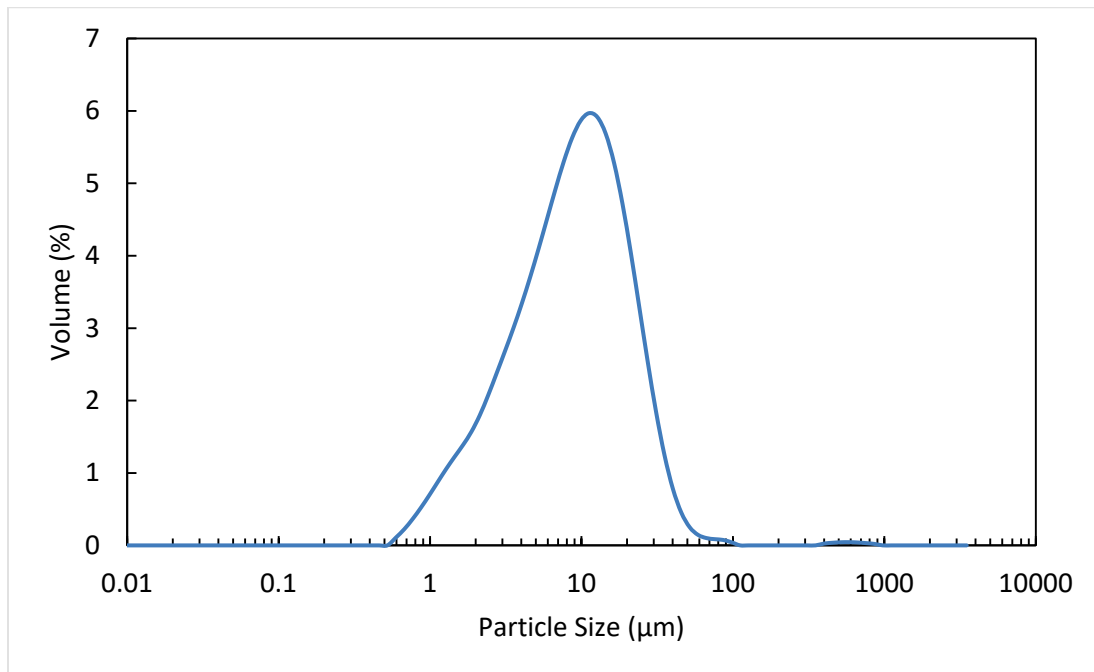


Figure A29. PSD Results of Slag (170080).



Appendix B: X-Ray Fluorescence Results

Figure B1. XRF Results of Cement (170052).

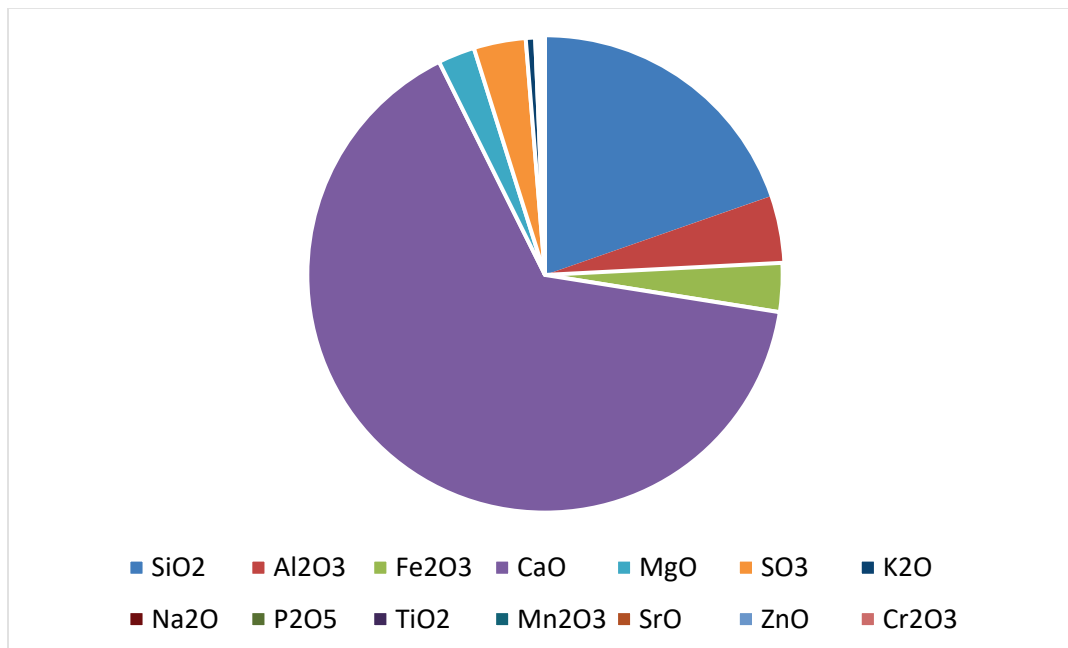


Figure B2. XRF Results of Cement (170065).

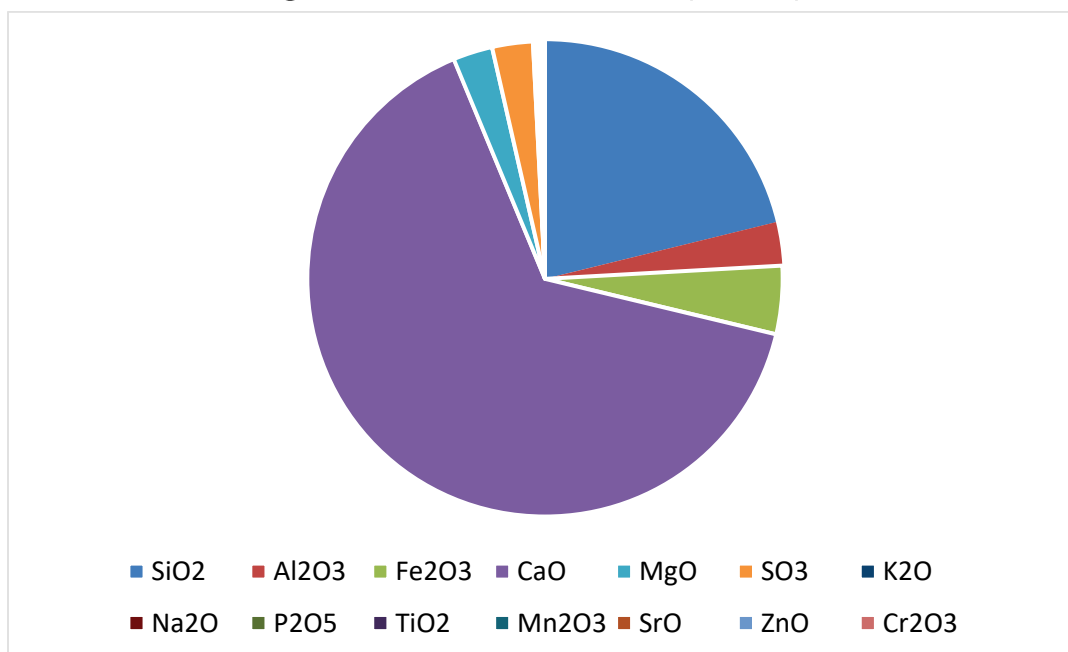


Figure B3. XRF Results of Fly Ash (080159).

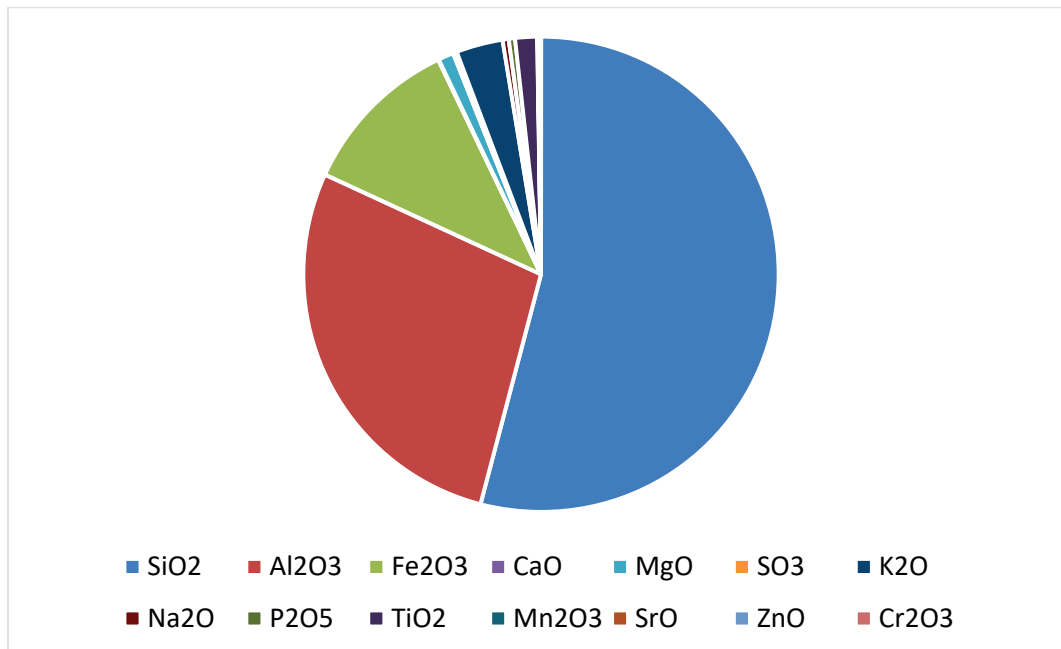


Figure B4. XRF Results of Fly Ash (100015).

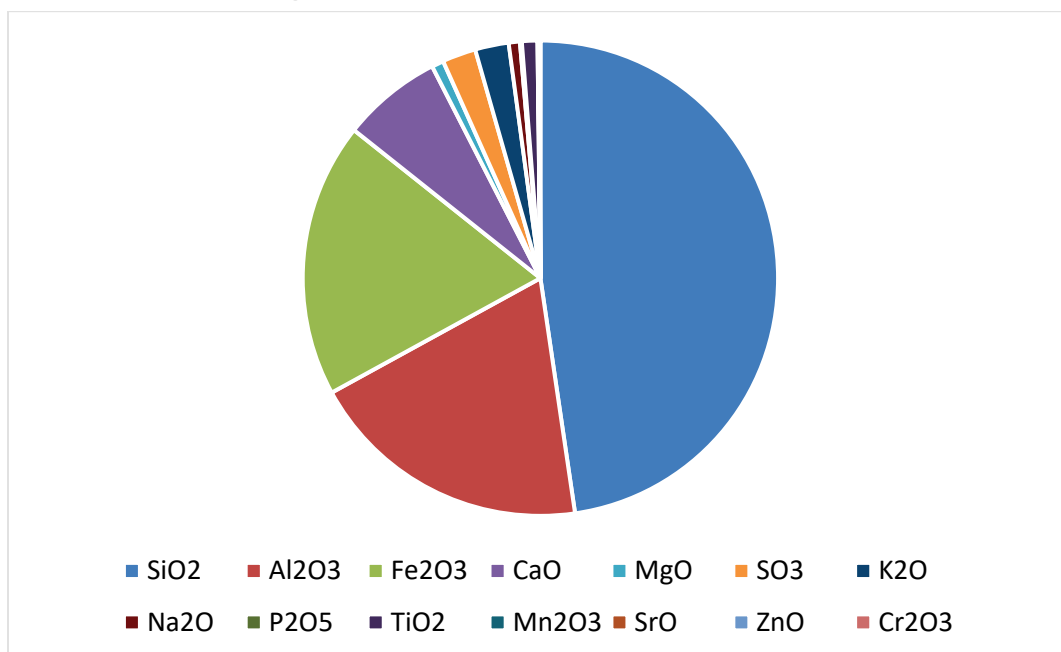


Figure B5. XRF Results of Fly Ash (100069).

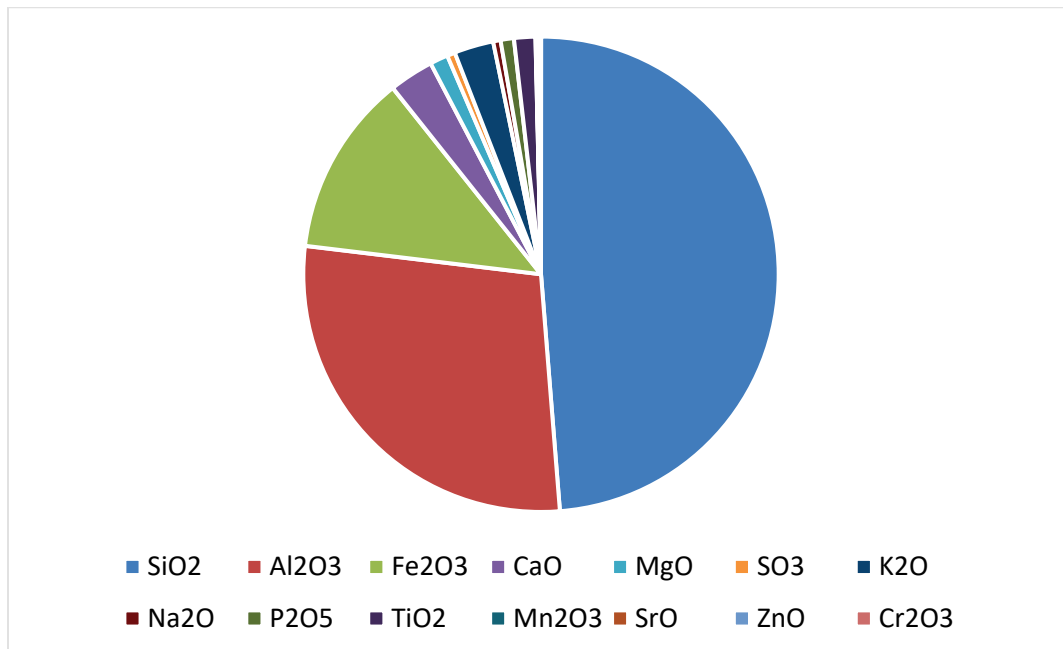


Figure B6. XRF Results of Fly Ash (120013).

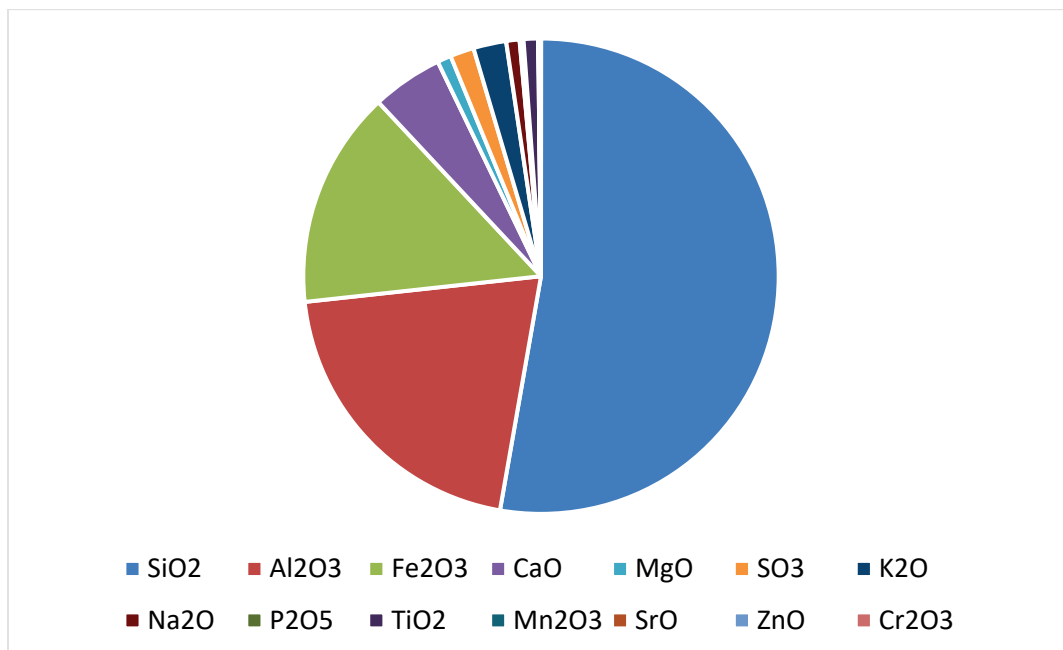


Figure B7. XRF Results of Fly Ash (120083).

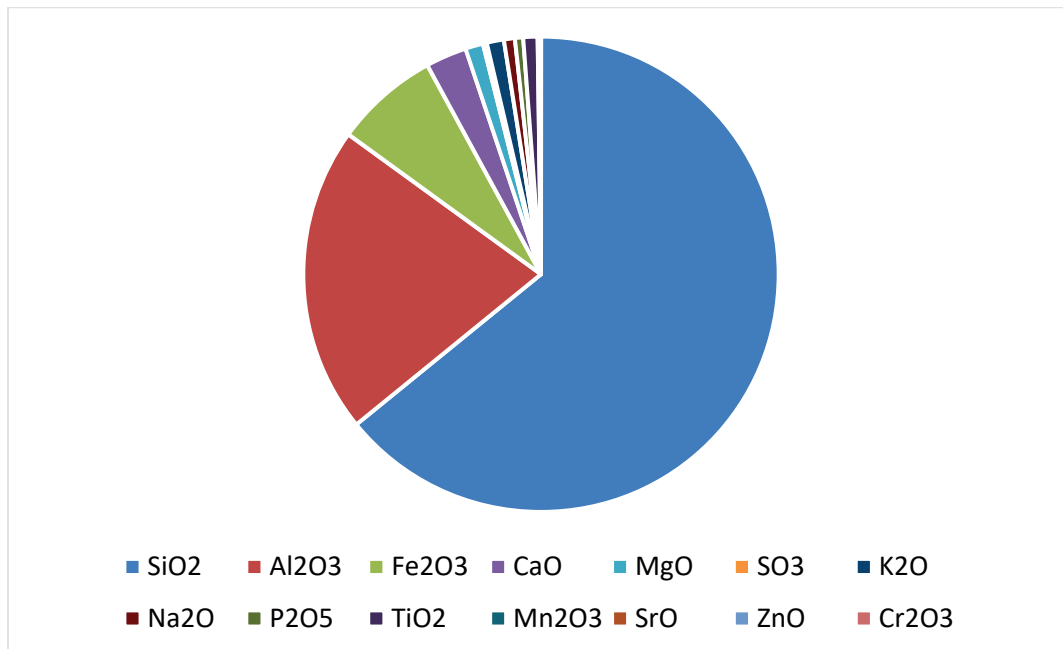


Figure B8. XRF Results of Fly Ash (120104).

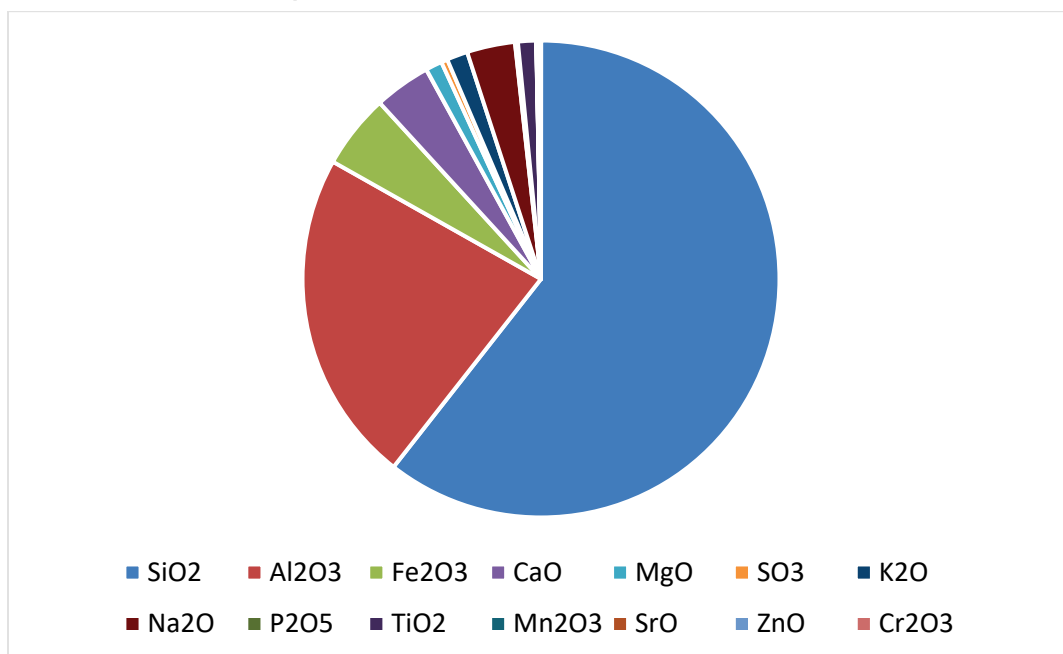


Figure B9. XRF Results of Fly Ash (120222).

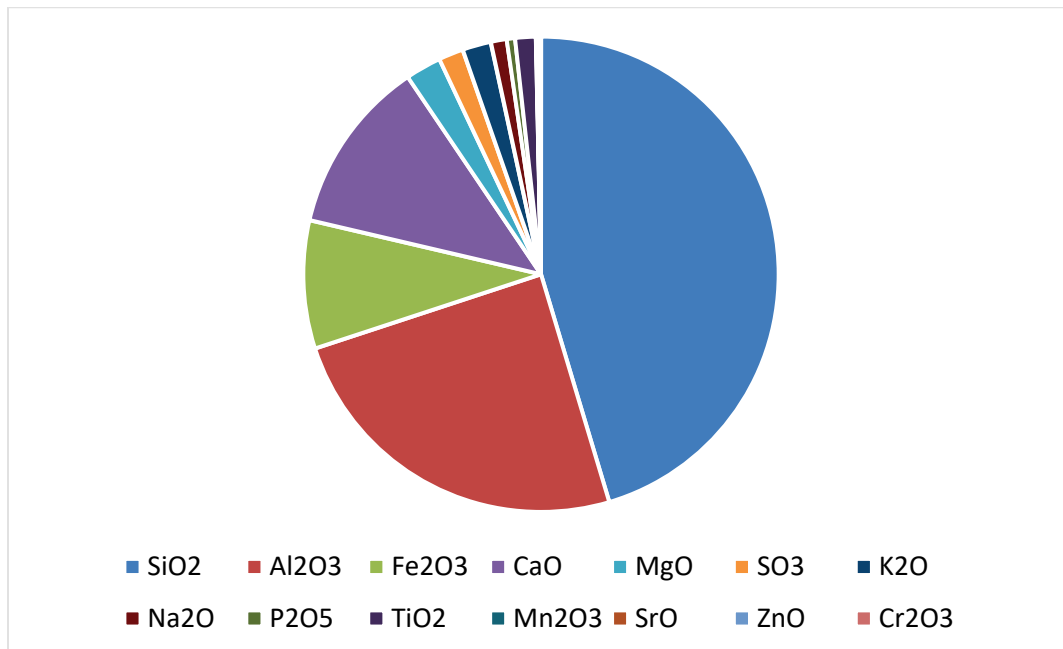


Figure B10. XRF Results of Fly Ash (130032).

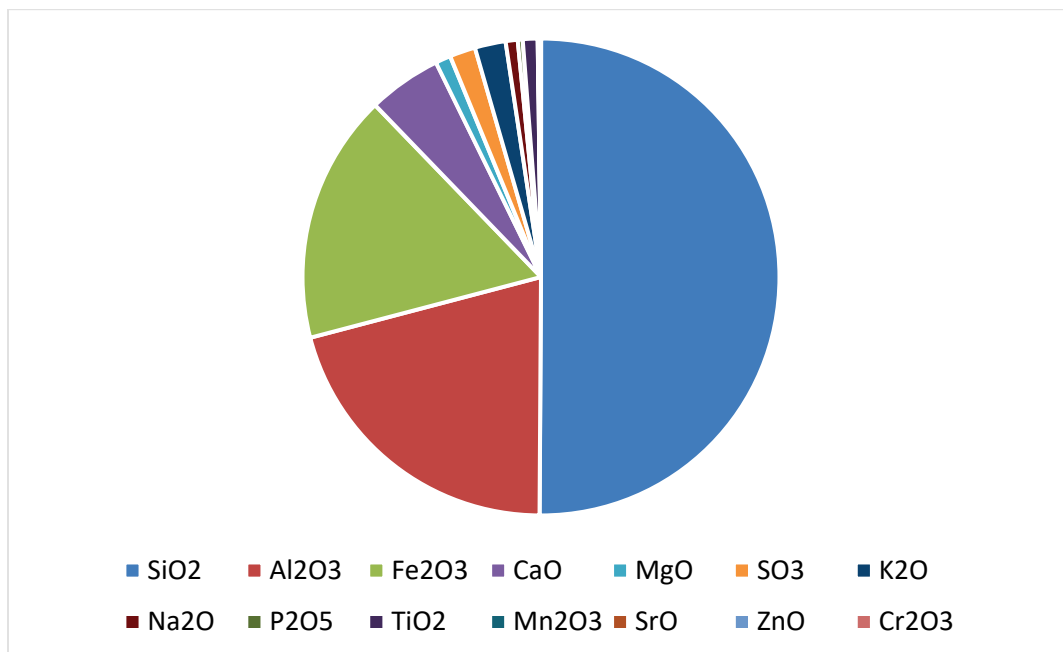


Figure B11. XRF Results of Fly Ash (140022).

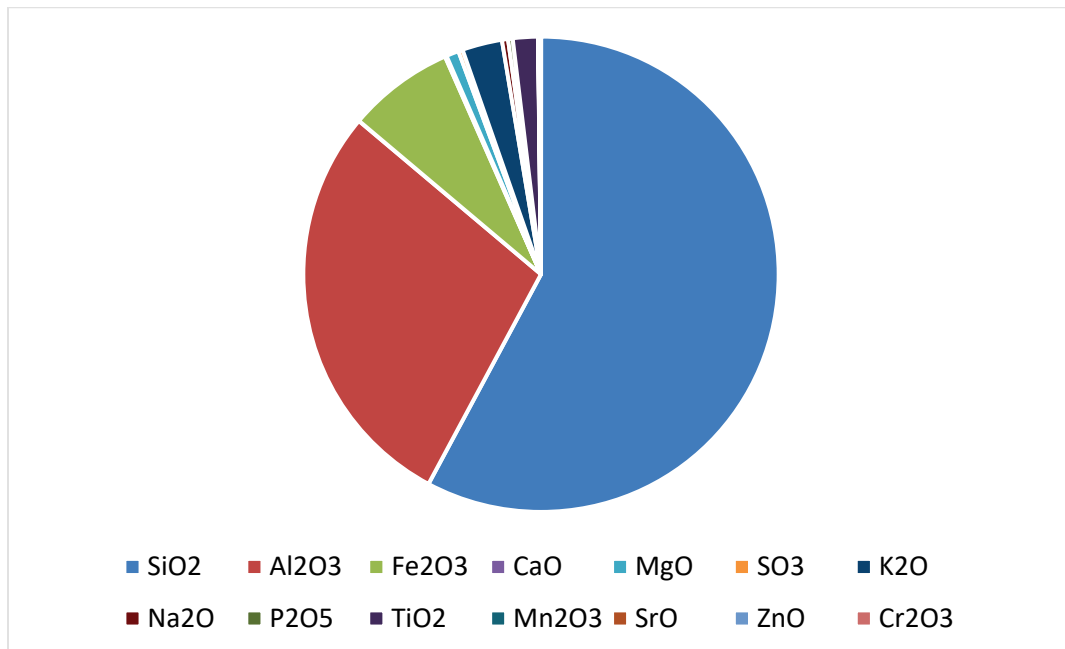


Figure B12. XRF Results of Fly Ash (140054).

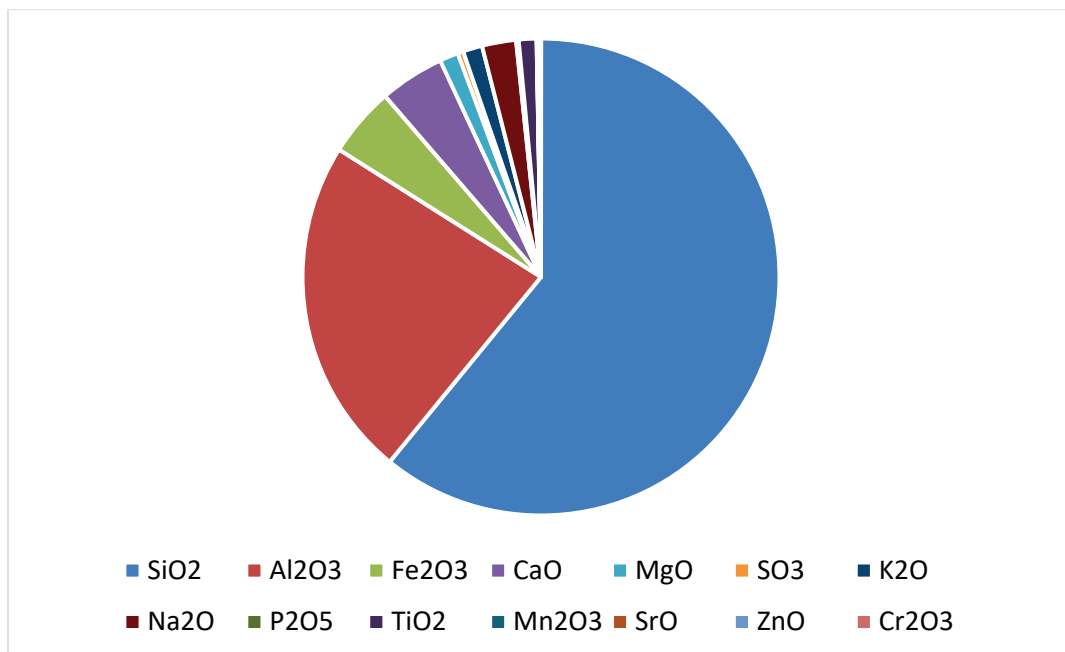


Figure B13. XRF Results of Fly Ash (150108).

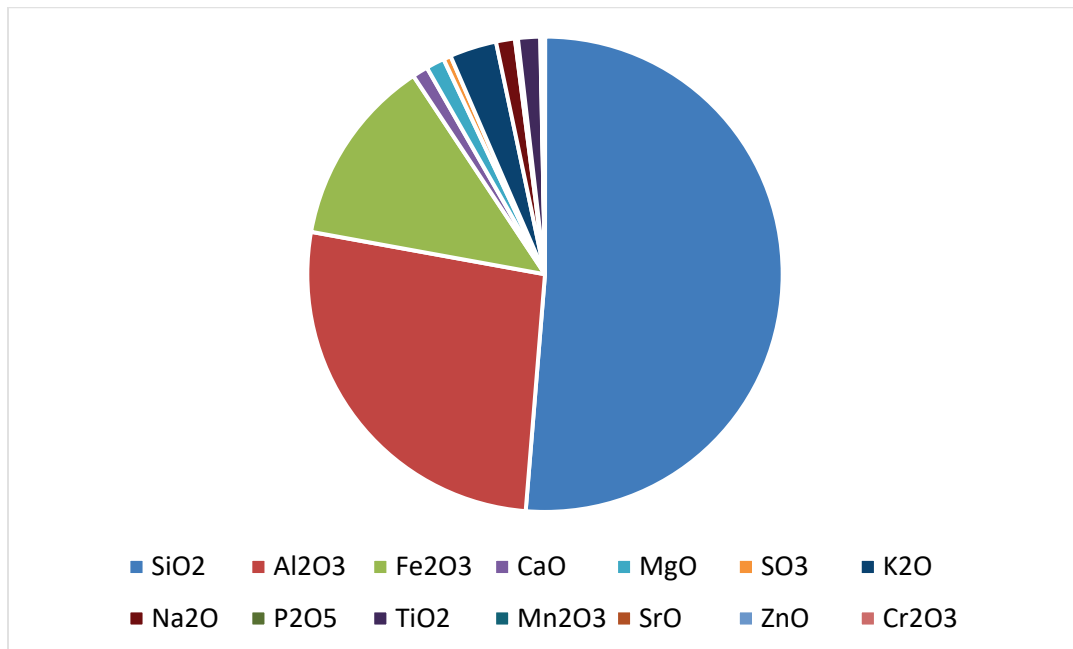


Figure B14. XRF Results of Fly Ash (150109).

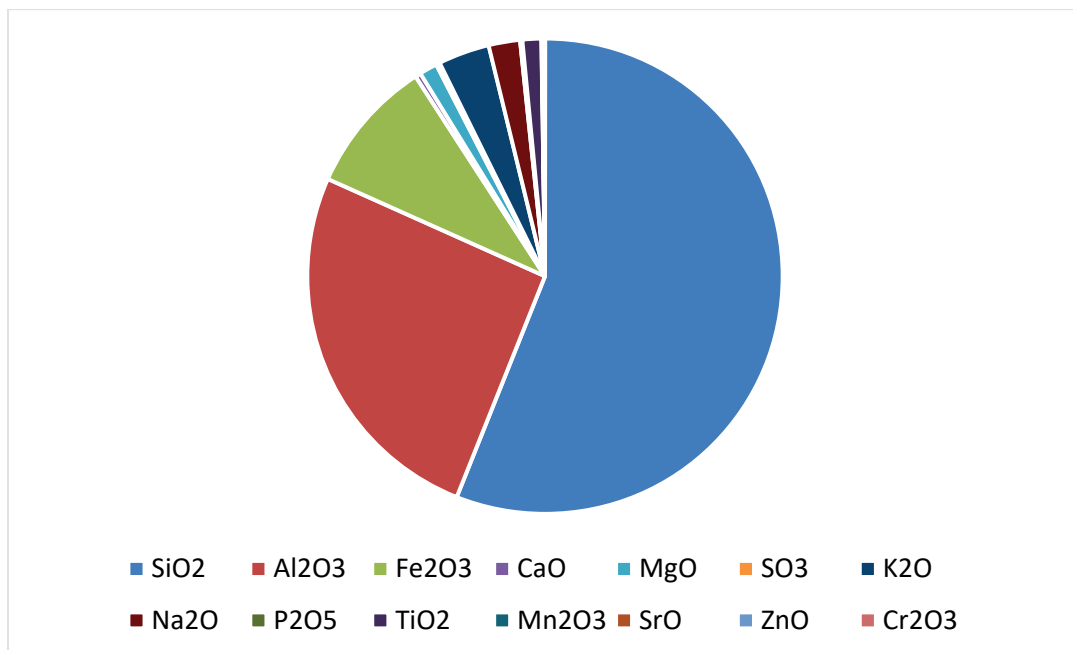


Figure B15. XRF Results of Silica Fume (Guam F10).

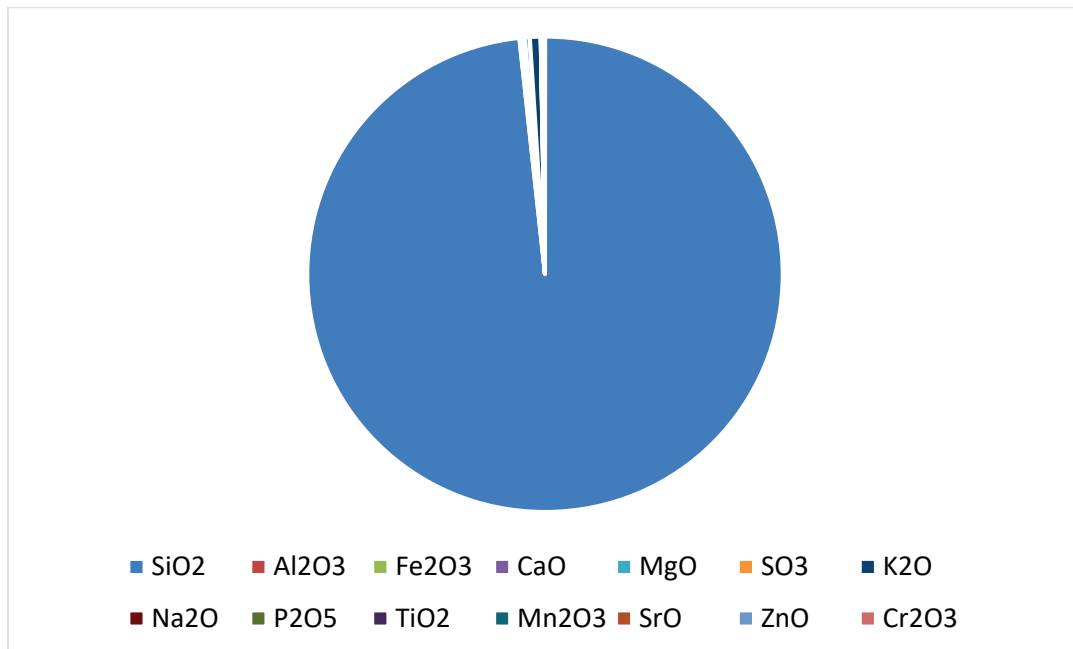


Figure B16. XRF Results of Silica Fume (130008).

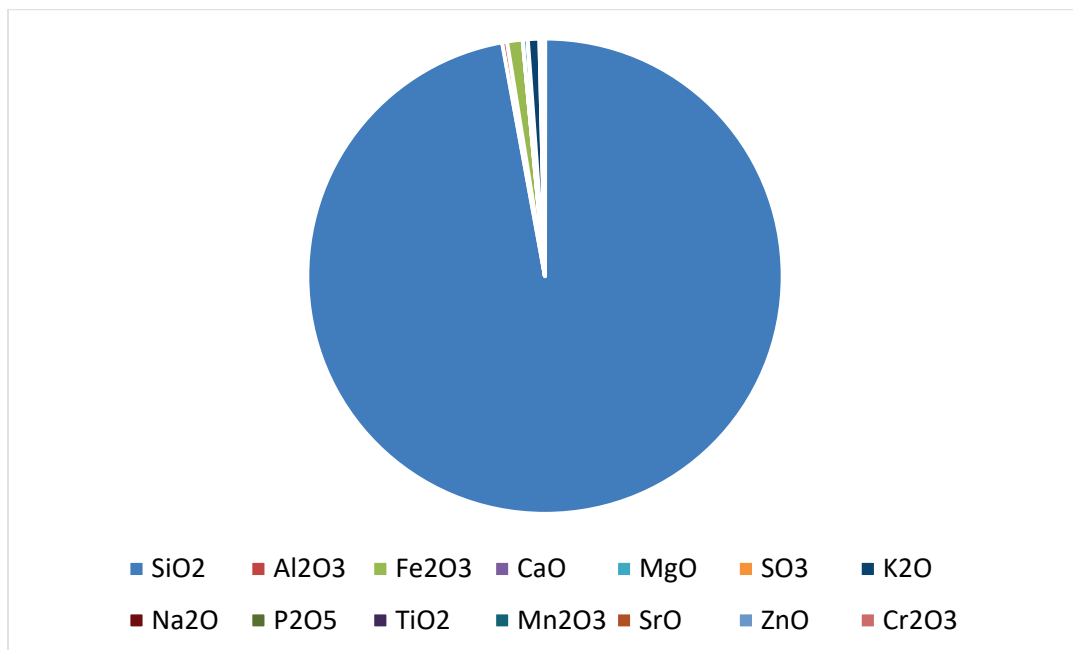


Figure B17. XRF Results of Silica Fume (130050).

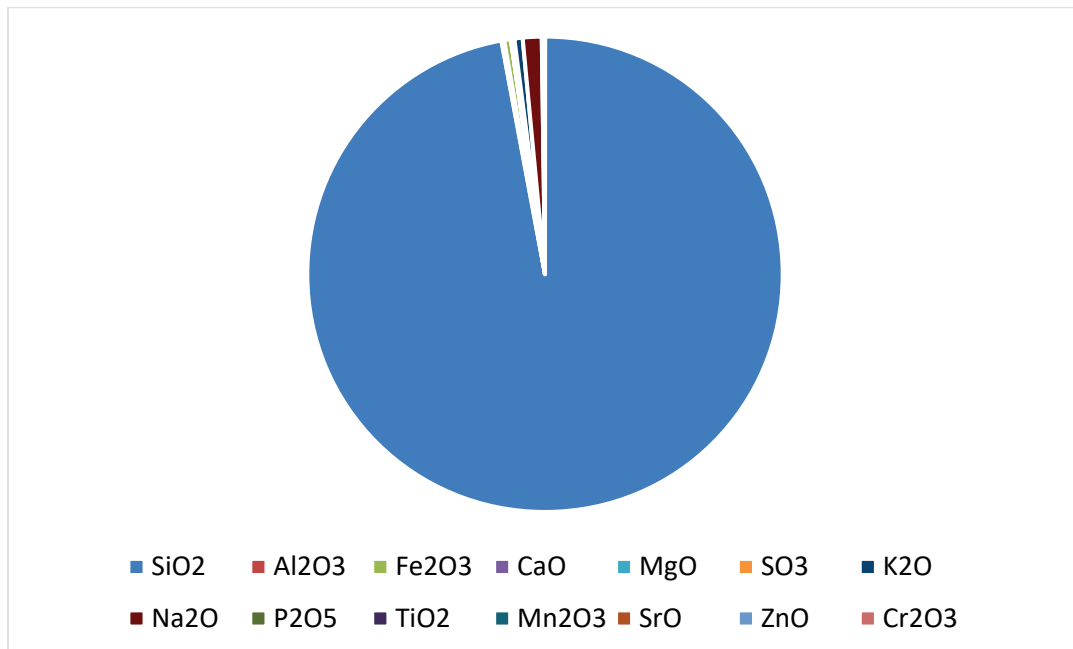


Figure B18. XRF Results of Silica Fume (140092).

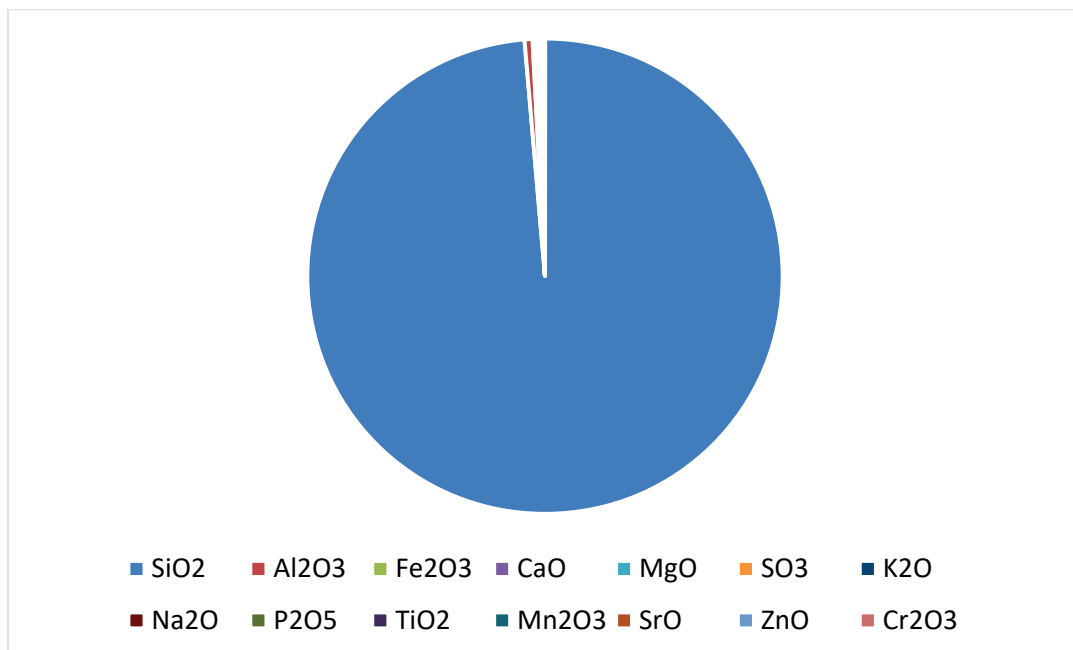


Figure B19. XRF Results of Silica Fume (140093).

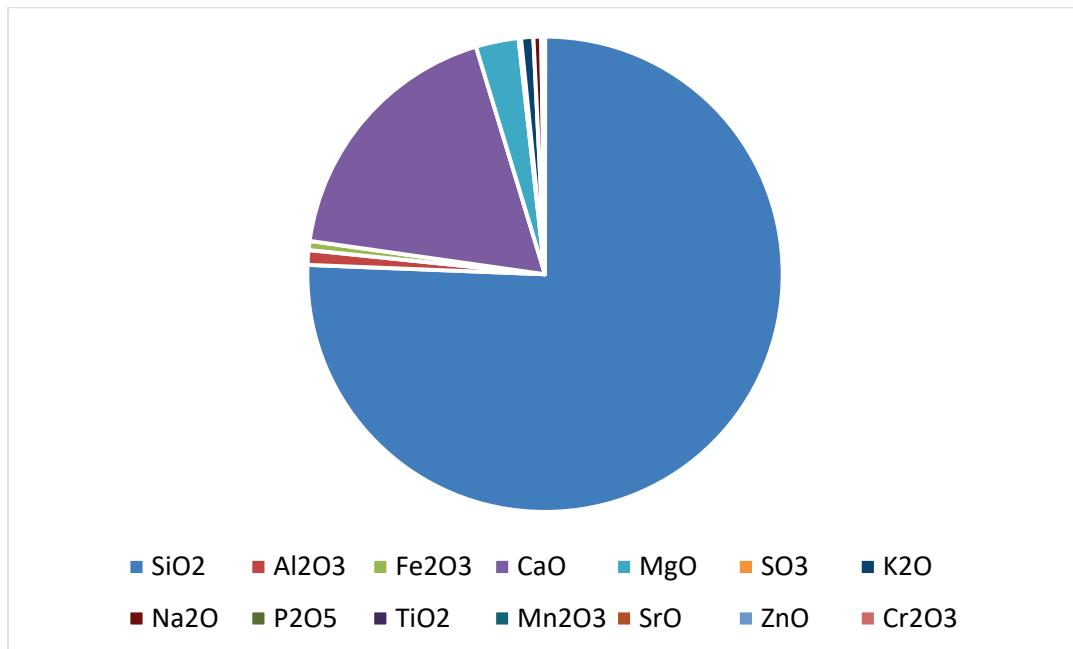


Figure B20. XRF Results of Silica Fume (140094).

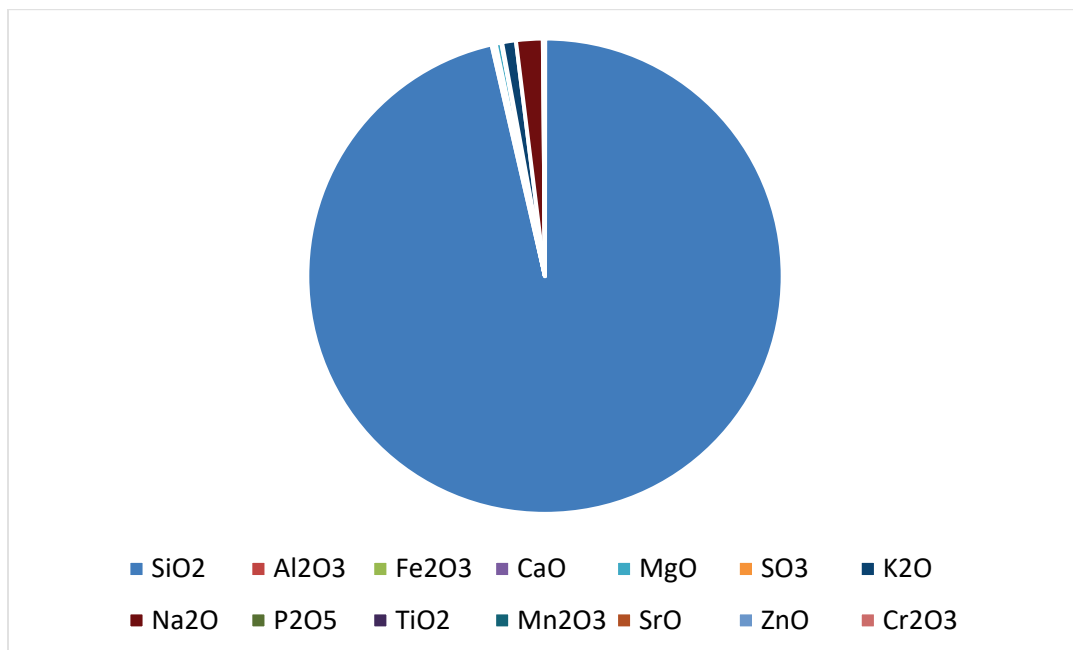


Figure B21. XRF Results of Silica Fume (150148).

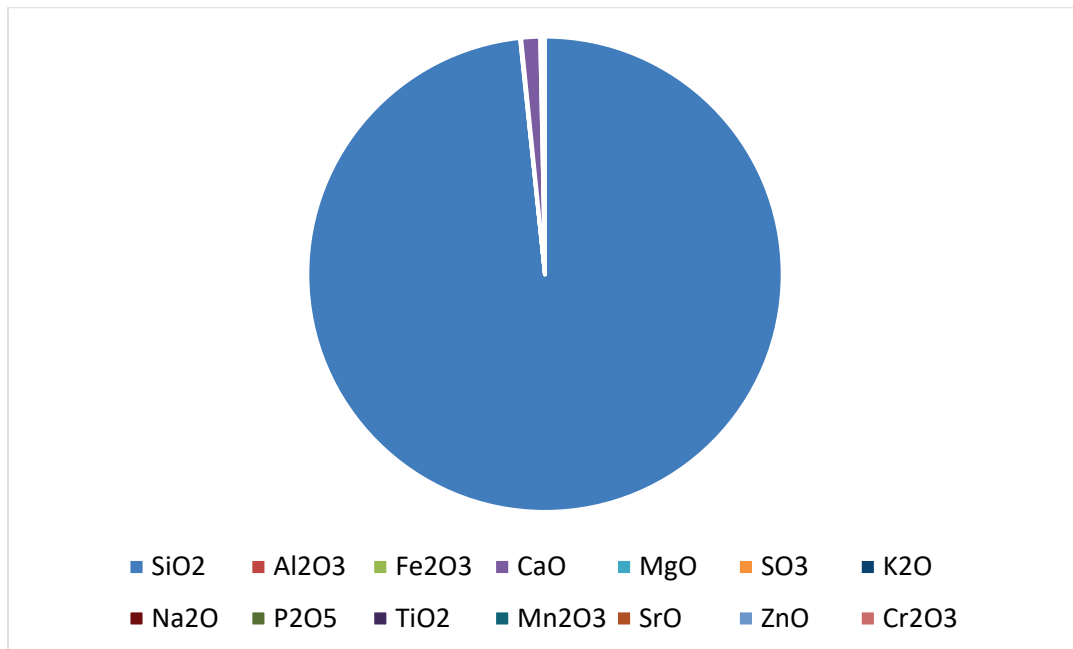


Figure B22. XRF Results of Silica Fume (150049).

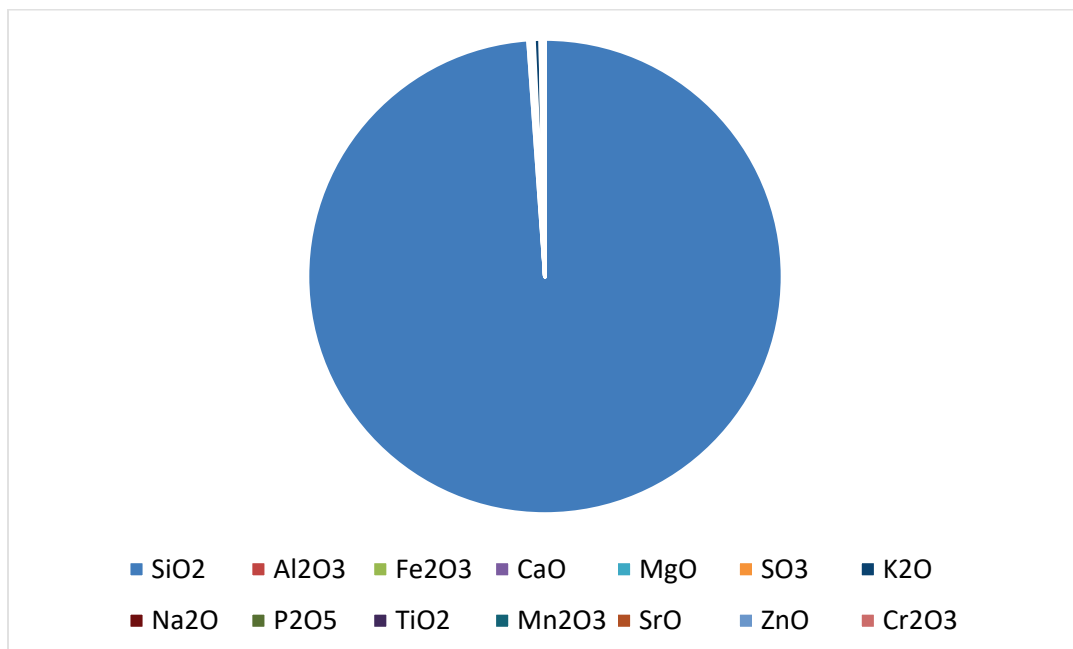


Figure B23. XRF Results of Silica Fume (150050).

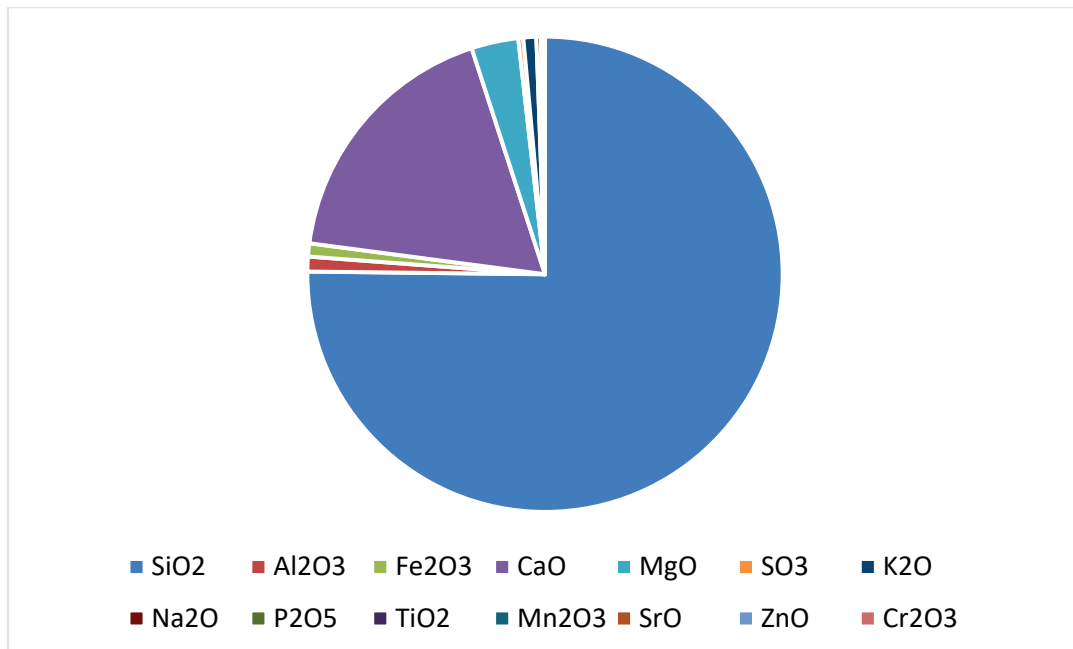


Figure B24. XRF Results of Silica Fume (150051).

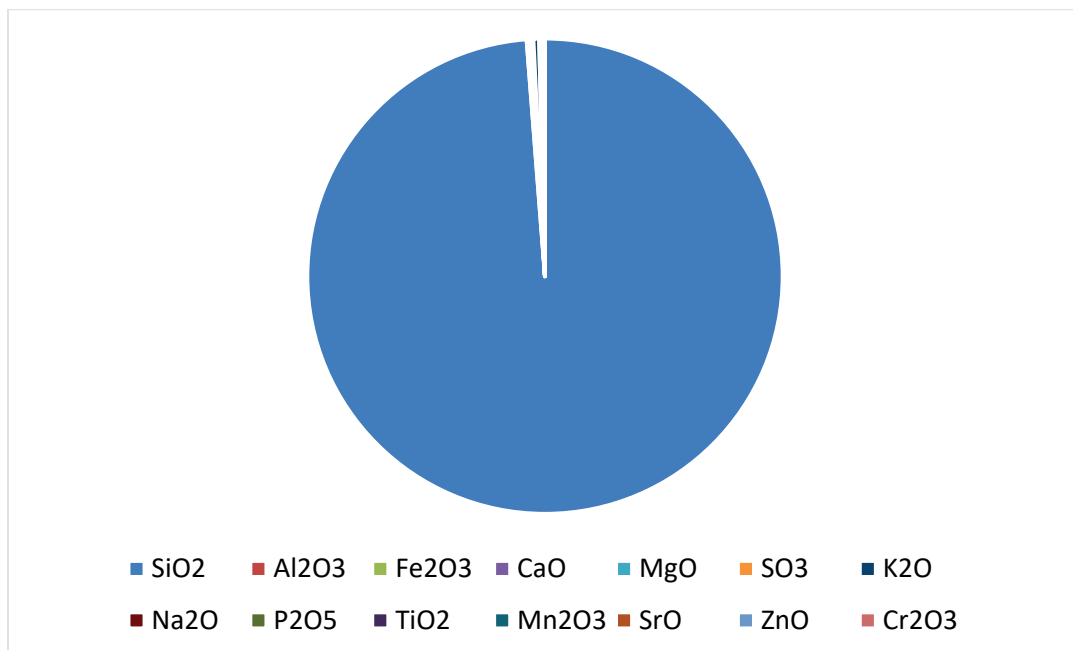


Figure B25. XRF Results of Silica Fume (Ecotech85).

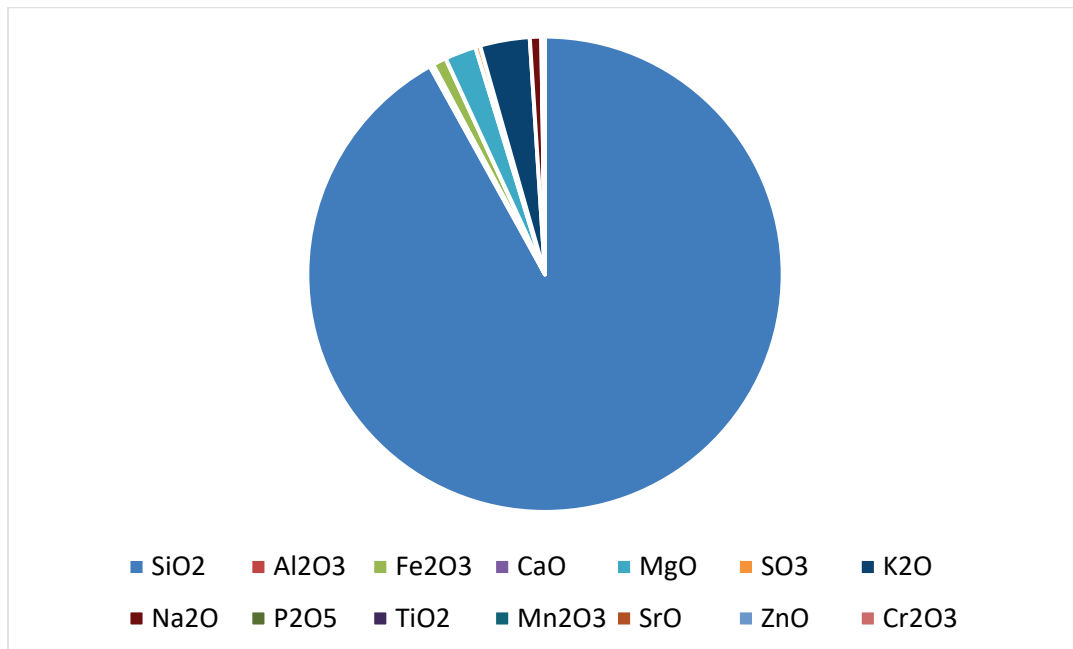


Figure B26. XRF Results of Silica Flour (Sil-co-sil).

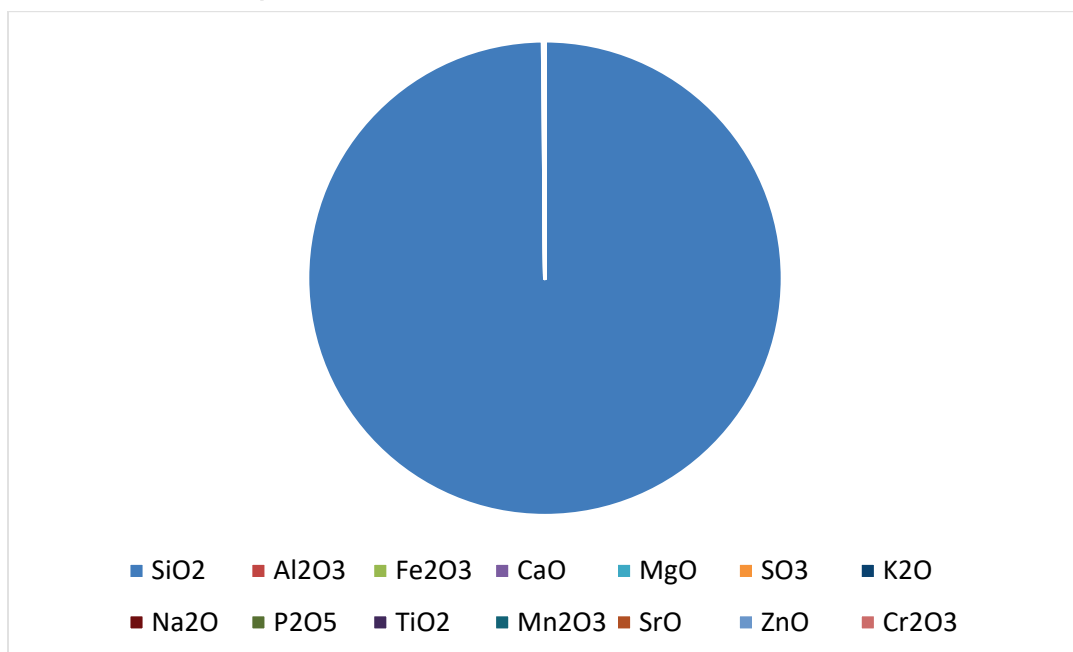
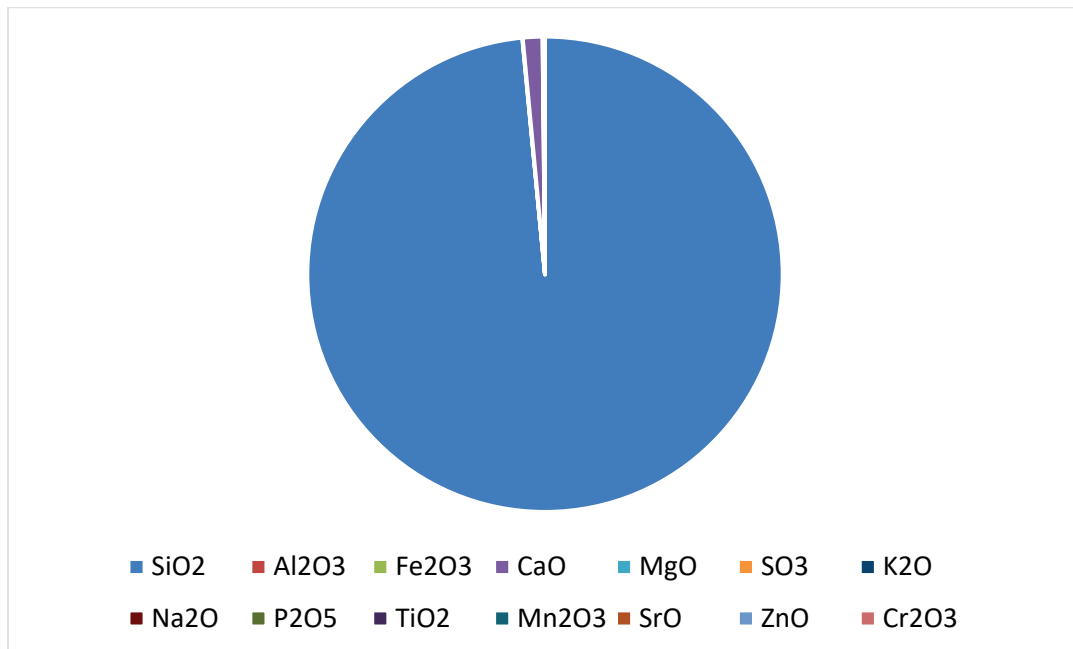


Figure B27. XRF Results of Silica Sand (F50).



Appendix C: X-Ray Diffraction Results

Figure C1. XRD Results of Fly Ash (080159).

Phase ID (3)	Chemical Formula	NA	NR	NP	Wt% (esd)	RIR
Mullite, syn	$Al_{4.59}Si_{1.41}O_{9.7}$	---	29	10	11.2 (4.8)	0.83
Quartz low	SiO_2	---	12	9	7.2 (3.1)	3.02
Hematite	Fe_2O_3	2	11	9	2.5 (1.1)	3.16
Amorphous + Others	SiO_2				79.0 (54.2)	2.50

XRF(Wt%): $Fe_2O_3=2.5\%$, $SiO_2=89.2\%$, $Al_2O_3=8.2\%$

Refinement Converged (R/E=1.31), Round=3, Iter=4, P=33, R=3.97% (E=3.03%, EPS=0.5)

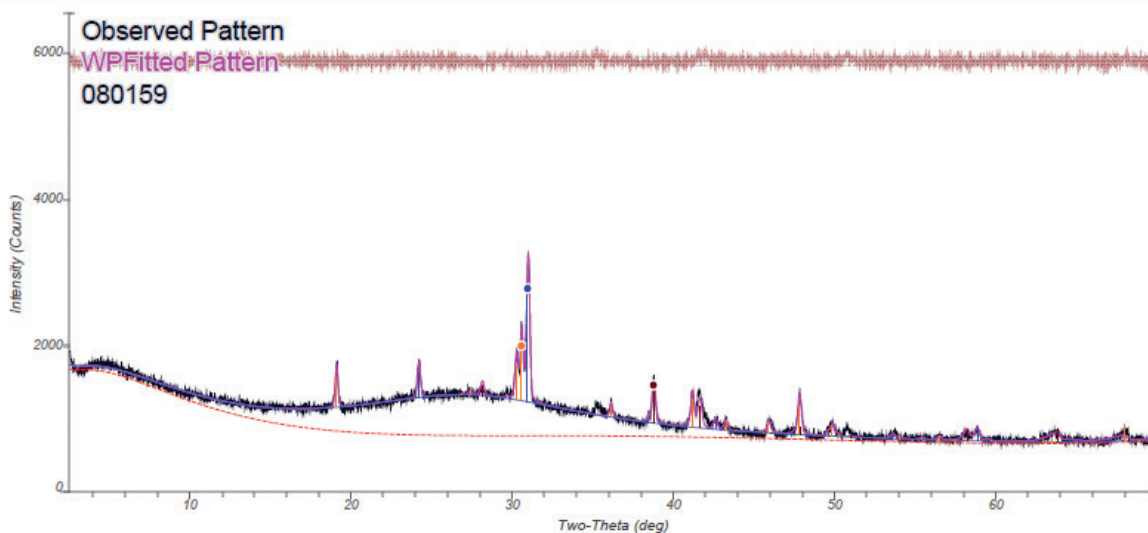
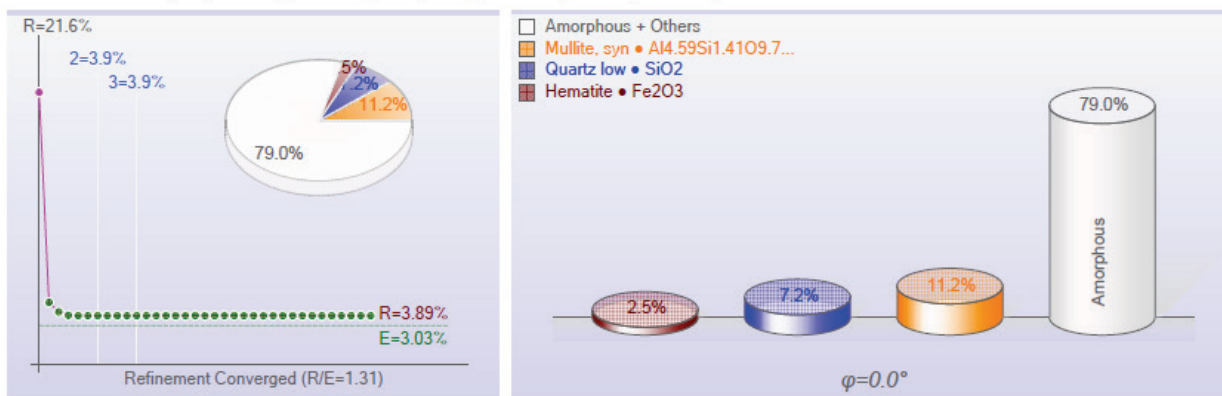


Figure C2. XRD Results of Fly Ash (100015).

Phase ID (4)	Chemical Formula	NA	NR	NP	Wt% (esd)	RIR	
Quartz low	SiO ₂	---	12	9	7.3 (7.4)	3.02	
Mullite, syn	Al _{4.59} Si _{1.41} O _{9.7}	---	29	10	11.6 (11.7)	0.83	
Hematite	Fe ₂ O ₃	2	11	9	5.2 (5.3)	3.16	
Magnetite	Fe ₃ O ₄	---	8	8	6.1 (6.2)	5.13	
Amorphous + Others	SiO ₂				69.9 (123.5)	2.50	

XRF(Wt%): Fe₂O₃=11.5%, SiO₂=80.2%, Al₂O₃=8.5%

Refinement Halted (R/E=1.33), Round=3, Iter=6, P=37, R=3.76% (E=2.82%, EPS=0.5)

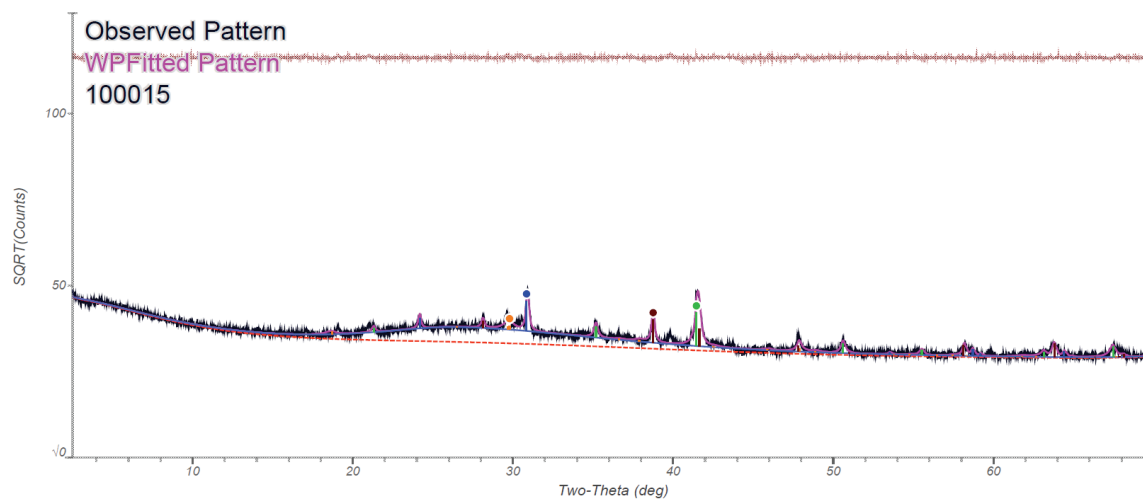
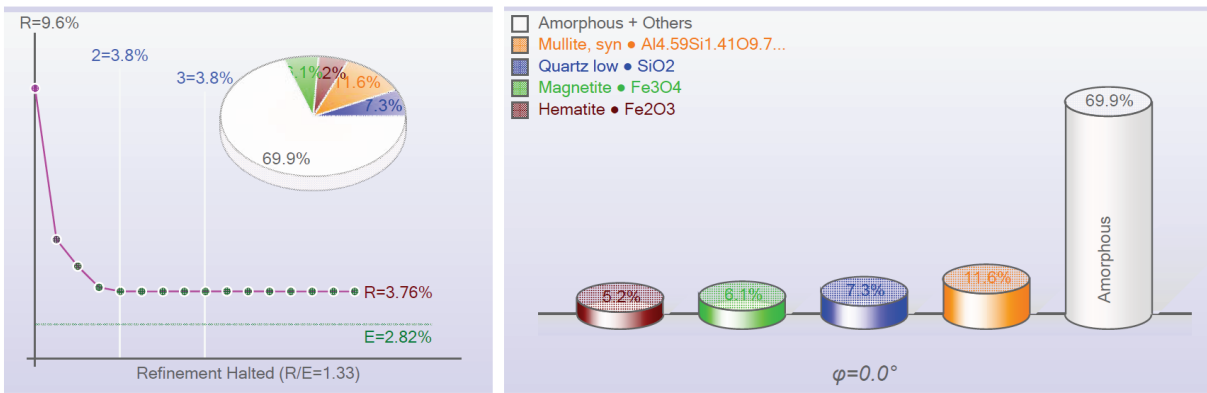


Figure C3. XRD Results of Fly Ash (100069).

Phase ID (4)	Chemical Formula	NA	NR	NP	Wt% (esd)	RIR
Mullite, syn	$\text{Al}_{4.59}\text{Si}_{1.41}\text{O}_{9.7}$	---	29	10	8.6 (3.9)	0.83
Quartz low	SiO_2	---	12	9	16.8 (7.7)	3.02
Magnetite	Fe_3O_4	---	8	8	3.2 (1.5)	5.13
Hematite	Fe_2O_3	2	11	9	5.3 (2.4)	3.16
Amorphous + Others	SiO_2				66.1 (54.5)	2.50

XRF(Wt%): $\text{Fe}_2\text{O}_3=8.6\%$, $\text{SiO}_2=85.2\%$, $\text{Al}_2\text{O}_3=6.3\%$

Refinement Halted (R/E=1.58), Round=3, Iter=6, P=37, R=4.7% (E=2.98%, EPS=0.5)

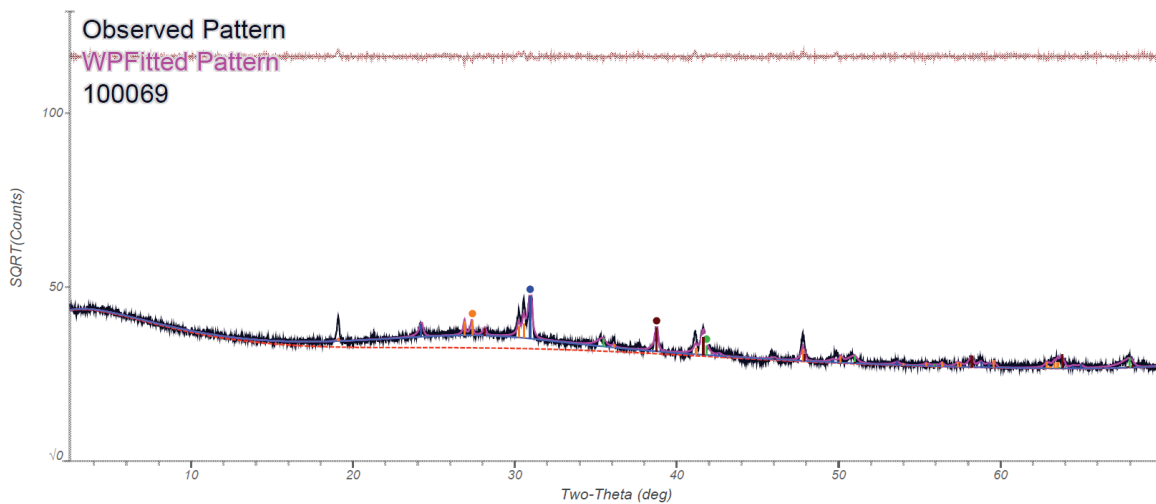
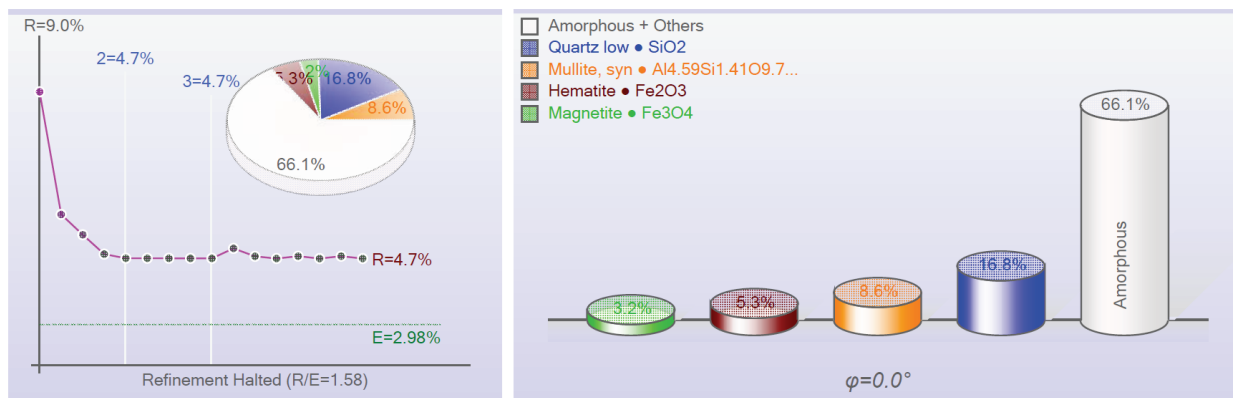


Figure C4. XRD Results of Fly Ash (120013).

Phase ID (4)	Chemical Formula	NA	NR	NP	Wt% (esd)	RIR
Quartz low	SiO ₂	---	12	9	9.7 (3.6)	3.02
Mullite, syn	Al _{4.59} Si _{1.41} O _{9.7}	---	29	10	8.9 (3.5)	0.83
Hematite	Fe ₂ O ₃	2	11	9	3.6 (1.3)	3.16
Magnetite	Fe ₃ O ₄	---	8	8	3.7 (1.4)	5.13
Amorphous + Others	SiO ₂				74.0 (45.6)	2.50

XRF(Wt%): Fe2O3=7.5%, SiO2=86.1%, Al2O3=6.5%

Refinement Halted (R/E=1.32), Round=3, Iter=6, P=37, R=3.85% (E=2.92%, EPS=0.5)

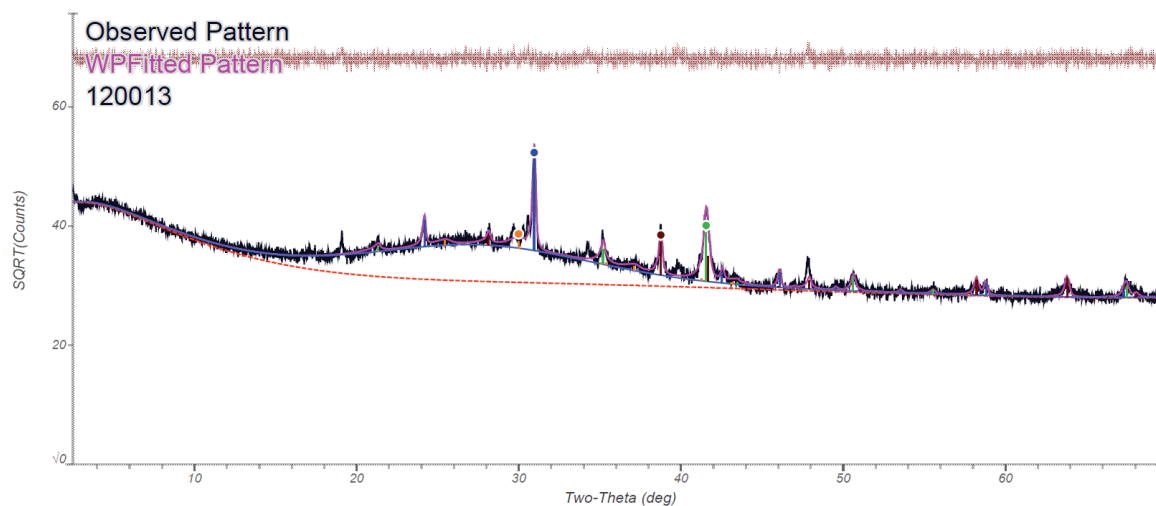
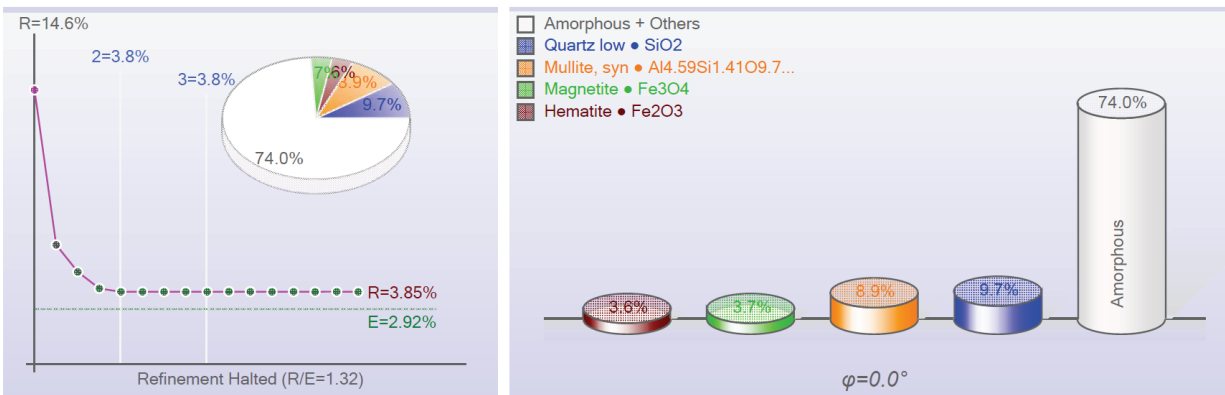


Figure C5. XRD Results of Fly Ash (120083).

Phase ID (3)	Chemical Formula	NA	NR	NP	Wt% (esd)	RIR
Quartz low	SiO ₂	---	12	9	15.2 (3.1)	3.02
Mullite, syn	Al _{4.59} Si _{1.41} O _{9.7}	---	29	*38	4.5 (1.0)	0.83
Hematite	Fe ₂ O ₃	2	11	7	1.3 (0.3)	3.16
Amorphous + Others	SiO ₂				79.0 (26.1)	2.50

XRF(Wt%): Fe₂O₃=1.3%, SiO₂=95.4%, Al₂O₃=3.3%

Refinement Halted (R/E=1.25), Round=3, Iter=8, P=46, R=4.0% (E=3.19%, EPS=0.5)

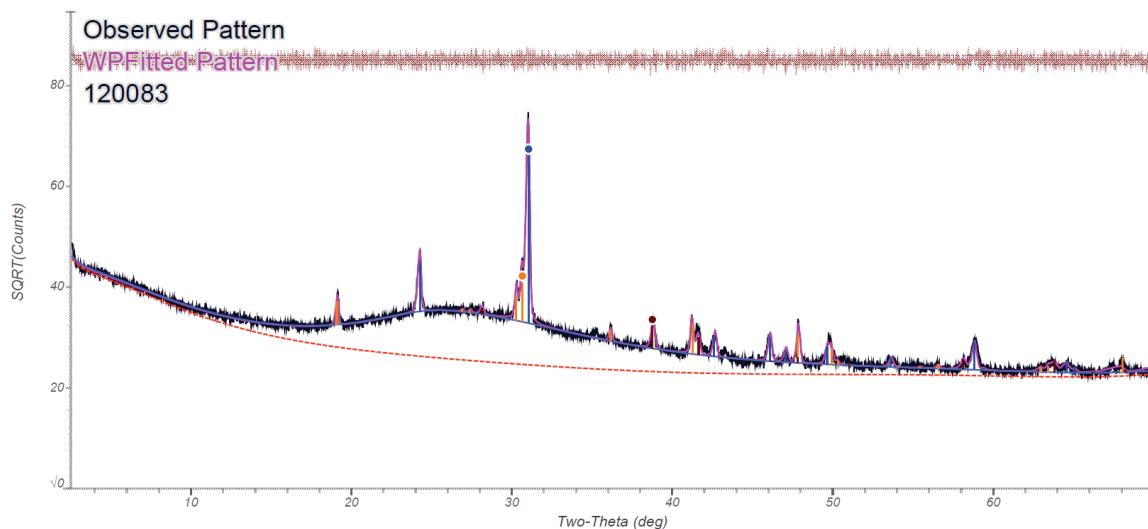
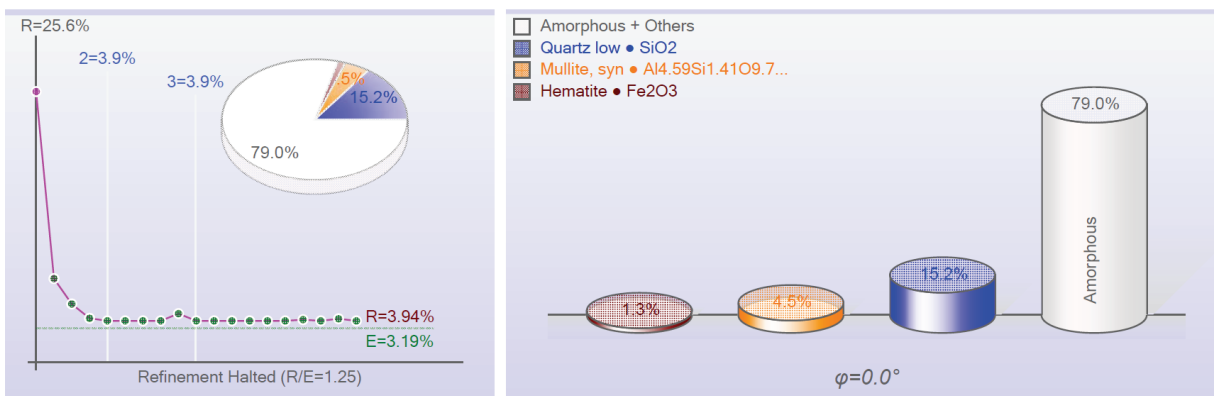


Figure C6. XRD Results of Fly Ash (120104).

Phase ID (3)	Chemical Formula	NA	NR	NP	Wt% (esd)	RIR
Quartz low	SiO ₂	---	12	9	22.1 (2.6)	3.02
Hematite	Fe ₂ O ₃	2	11	7	1.0 (0.2)	3.16
Mullite	Al(Al _{1.272} Si _{0.728} O _{4.864})	---	30	10	8.9 (1.3)	0.85
Amorphous + Others	SiO ₂				67.9 (13.3)	2.50

XRF(Wt%): Fe2O3=1.0%, SiO2=92.5%, Al2O3=6.5%

Refinement Converged (R/E=1.17), Round=3, Iter=3, P=34, R=3.97% (E=3.38%, EPS=0.5)

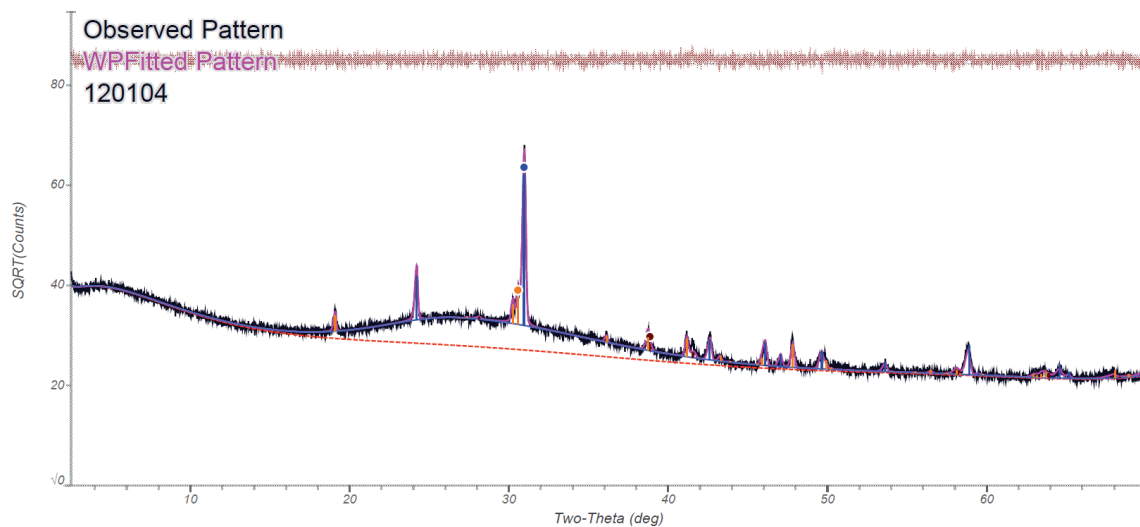
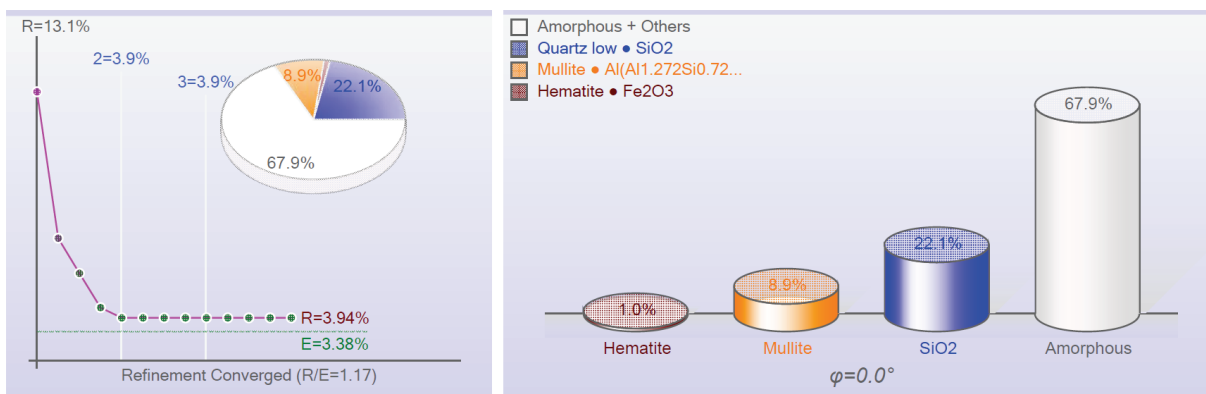


Figure C7. XRD Results of Fly Ash (120222).

Phase ID (3)	Chemical Formula	NA	NR	NP	Wt% (esd)	RIR	
Quartz low	SiO ₂	---	12	9	13.4 (2.6)	3.02	
Mullite	Al(Al _{1.272} Si _{0.728} O _{4.864})	---	30	10	11.2 (3.5)	0.85	
Hematite	Fe ₂ O ₃	2	11	9	3.6 (0.8)	3.16	
Amorphous + Others	SiO ₂				71.8 (17.3)	2.50	

XRF(Wt%): Fe2O3=3.6%, SiO2=88.3%, Al2O3=8.1%

Refinement Converged (R/E=1.27), Round=3, Iter=4, P=34, R=4.31% (E=3.39%, EPS=0.5)

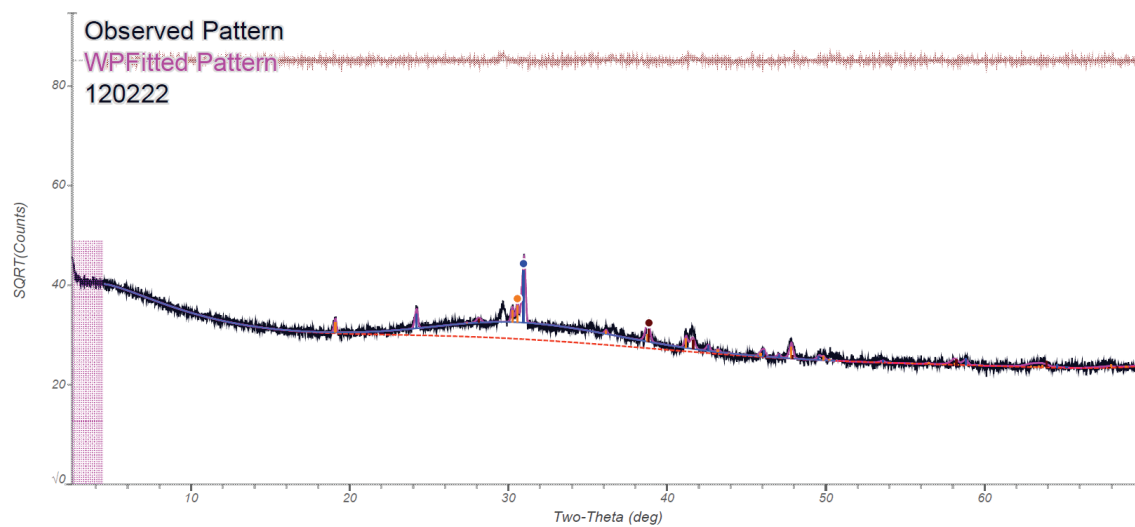
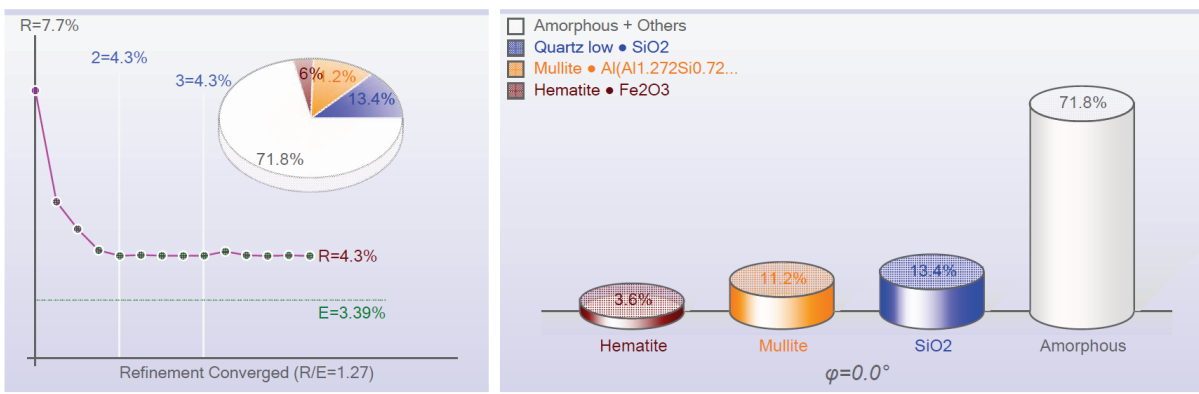


Figure C8. XRD Results of Fly Ash (130032).

Phase ID (3)	Chemical Formula	NA	NR	NP	Wt% (esd)	RIR
Quartz low	SiO ₂	---	12	9	7.9 (5.6)	3.02
Hematite	Fe ₂ O ₃	2	11	9	3.6 (2.5)	3.16
Mullite	Al(Al _{1.272} Si _{0.728} O _{4.864})	---	30	10	11.7 (8.2)	0.85
Amorphous + Others	SiO ₂				76.8 (88.6)	2.50

XRF(Wt%): Fe2O3=3.6%, SiO2=88.0%, Al2O3=8.5%

Refinement Converged (R/E=1.53), Round=3, Iter=6, P=34, R=4.39% (E=2.87%, EPS=0.5)

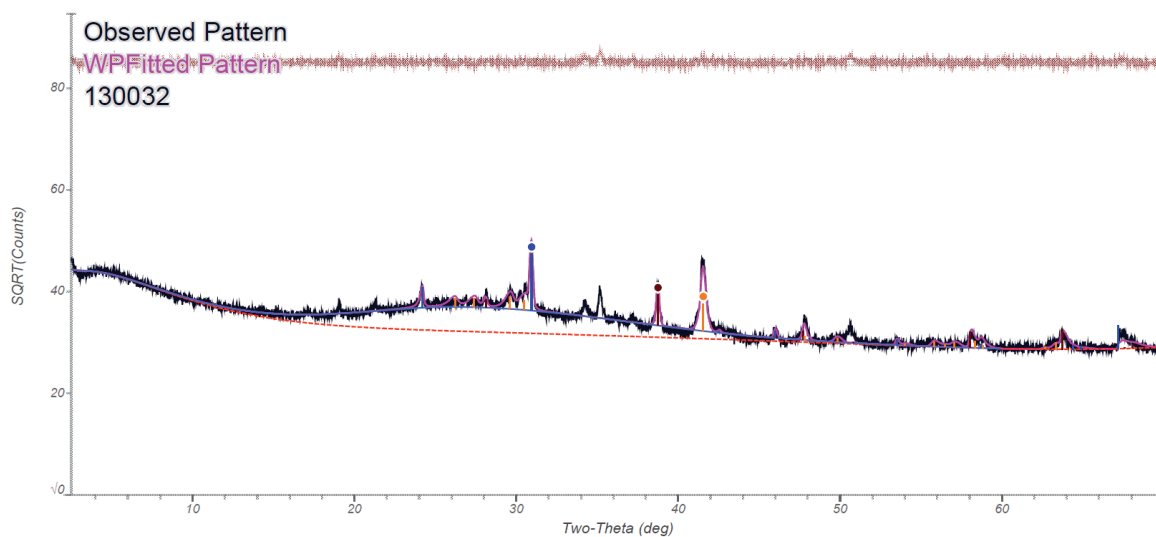
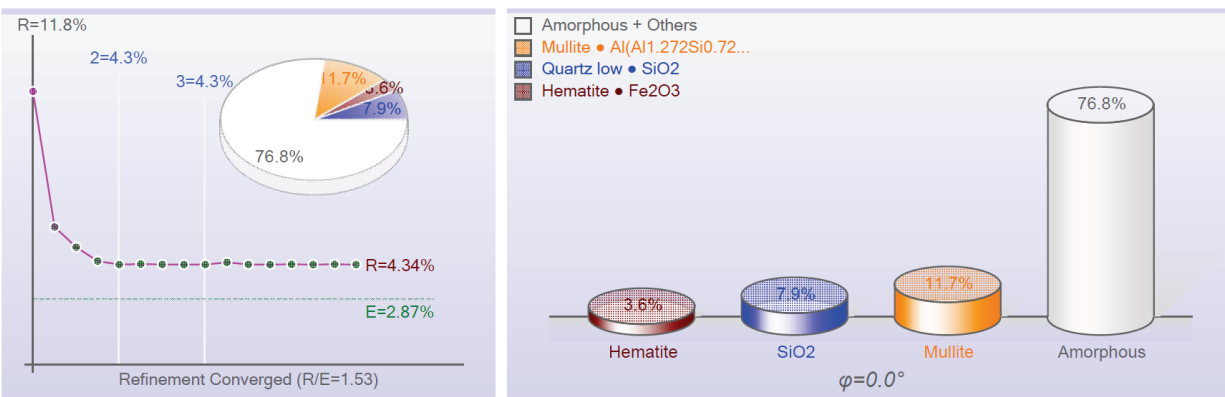


Figure C9. XRD Results of Fly Ash (140022).

Phase ID (4)	Chemical Formula	NA	NR	NP	Wt% (esd)	RIR
Mullite, syn	$\text{Al}_{4.59}\text{Si}_{1.41}\text{O}_{9.7}$	---	29	10	20.9 (2.2)	0.83
Hematite	Fe_2O_3	2	11	9	2.0 (0.3)	3.16
Quartz	SiO_2	---	12	9	17.0 (1.6)	3.07
Maghemite	$\text{Fe}_{1.966}\text{O}_{2.963}$	---	80	9	2.7 (0.4)	2.79
Amorphous + Others	SiO_2				57.5 (10.6)	2.50

XRF(Wt%): $\text{Fe}_2\text{O}_3=4.6\%$, $\text{SiO}_2=80.1\%$, $\text{Al}_2\text{O}_3=15.3\%$

Refinement Converged (R/E=1.29), Round=3, Iter=3, P=36, R=4.31% (E=3.35%, EPS=0.5)

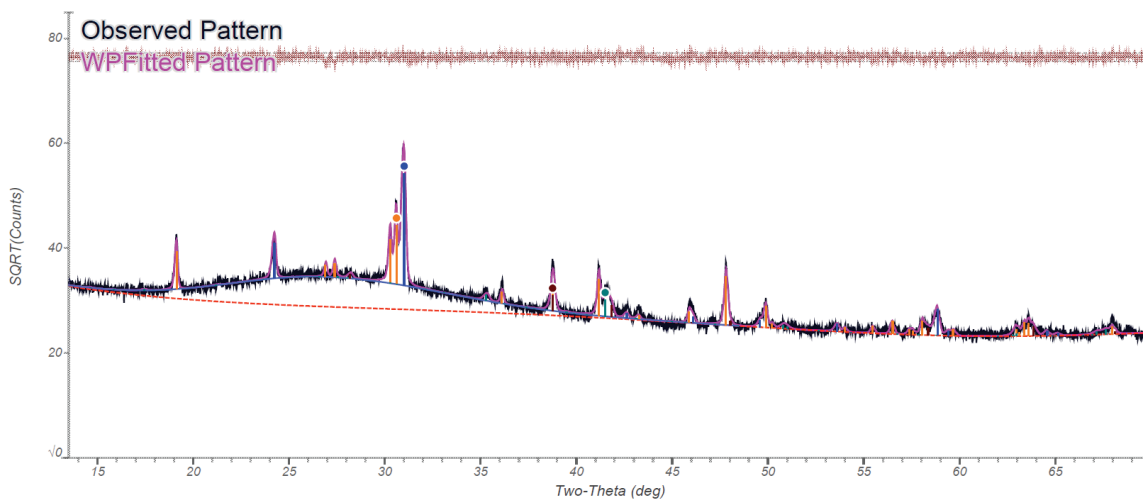
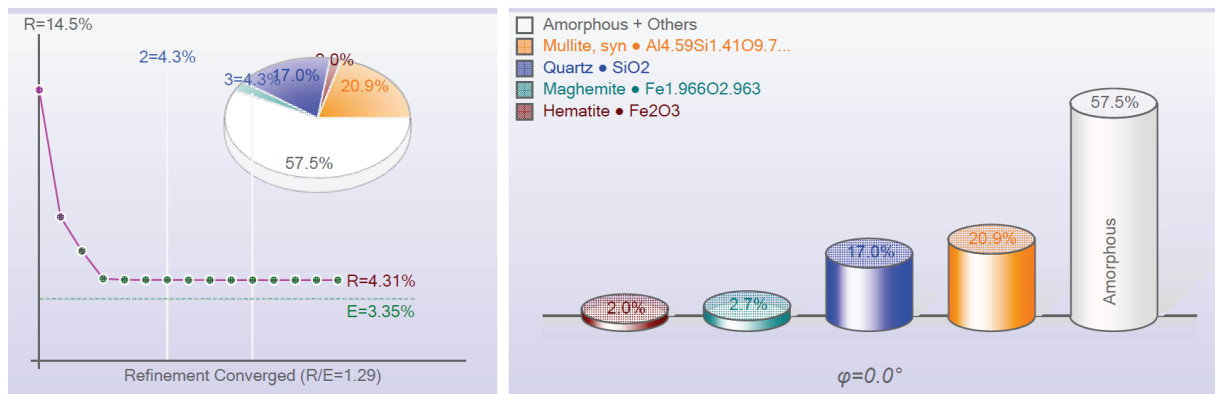


Figure C10. XRD Results of Fly Ash (140054).

Phase ID (4)	Chemical Formula	NA	NR	NP	Wt% (esd)	RIR
Magnetite	Fe_3O_4	---	8	5	0.0 (0.0)	5.13
Quartz low	SiO_2	---	12	9	26.4 (4.5)	3.02
Mullite, syn	$\text{Al}_{4.59}\text{Si}_{1.41}\text{O}_{9.7}$	---	29	10	6.3 (1.2)	0.83
Hematite	Fe_2O_3	2	11	7	1.4 (0.3)	3.16
Amorphous + Others	SiO_2				65.9 (20.4)	2.50

XRF(Wt%): $\text{Fe}_2\text{O}_3=1.4\%$, $\text{SiO}_2=93.9\%$, $\text{Al}_2\text{O}_3=4.6\%$

Refinement Converged (R/E=1.38), Round=3, Iter=6, P=34, R=4.68% (E=3.40%, EPS=0.5)

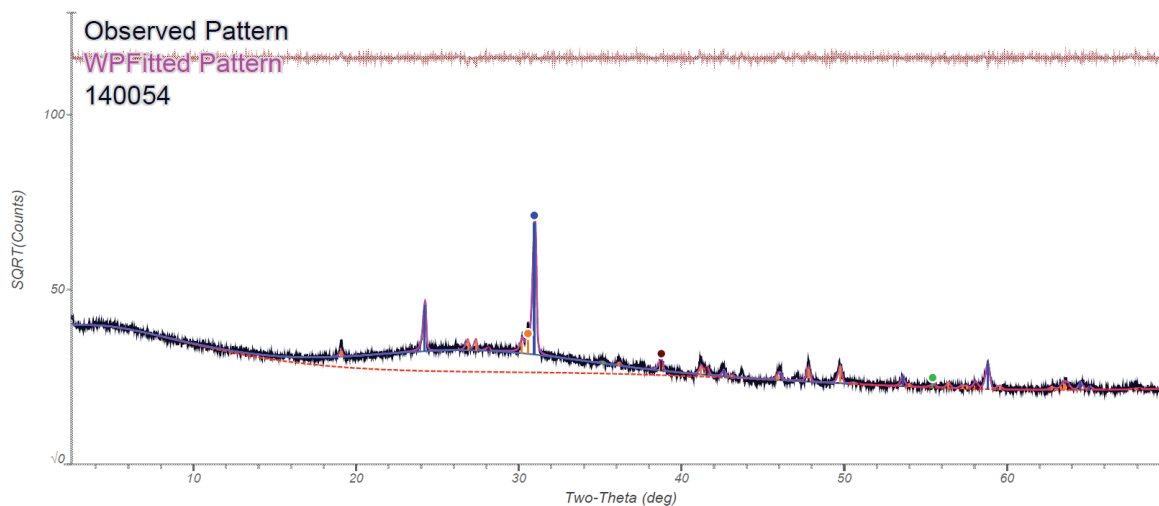
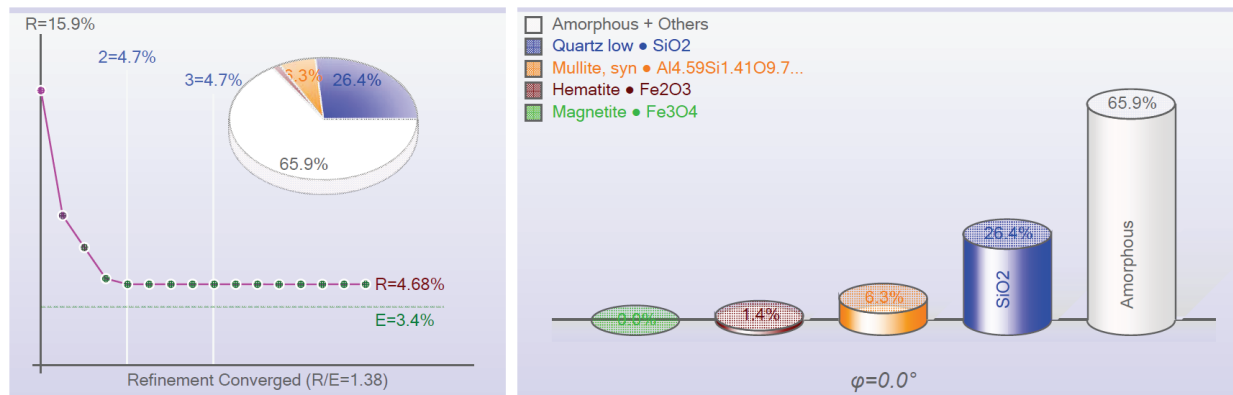


Figure C11. XRD Results of Fly Ash (150108).

Phase ID (4)	Chemical Formula	NA	NR	NP	Wt% (esd)	RIR
Quartz	SiO_2	2	34	9	5.9 (1.0)	4.04
Magnetite, syn	$\text{Fe}^{+2}\text{Fe}_2^{+3}\text{O}_4$	---	7	8	3.1 (0.5)	4.90
Mullite	$\text{Al}_{2.28}\text{Si}_{1.72}\text{O}_{4.86}$	8	31	10	10.0 (1.7)	0.85
Hematite	Fe_2O_3	2	11	9	1.0 (0.2)	3.15
Amorphous + Others	SiO_2				80.0 (21.4)	2.50

XRF(Wt%): Fe2O3=4.2%, SiO2=88.6%, Al2O3=7.3%

Refinement Halted (R/E=1.2), ♦ Round=3, Iter=6, P=37, R=4.23% (E=3.54%, EPS=0.5)

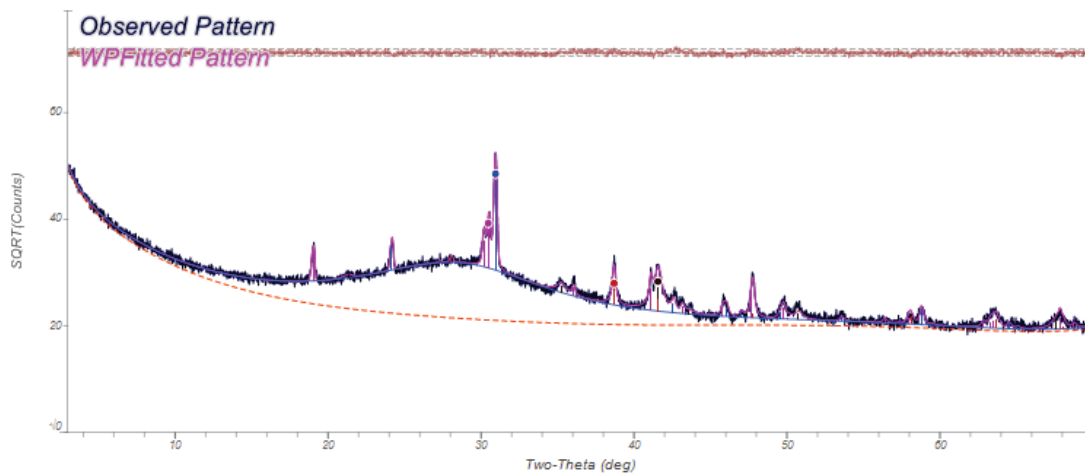
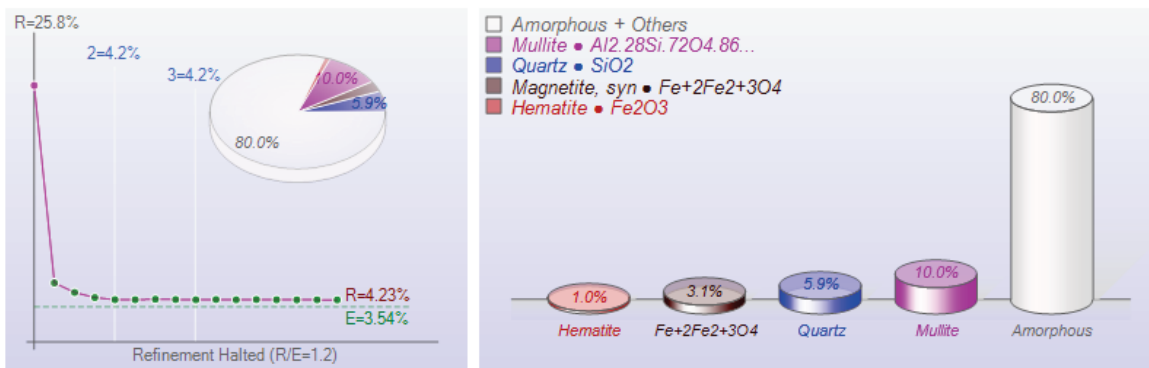


Figure C12. XRD Results of Fly Ash (150109).

Phase ID (3)	Chemical Formula	NA	NR	NP	Wt% (esd)	RIR
Mullite	$Al_{2.28}Si_{7.72}O_{4.86}$	8	31	10	10.8 (2.9)	0.86
Quartz	SiO_2	2	34	9	12.1 (3.2)	4.04
Hematite	Fe_2O_3	2	11	9	1.3 (0.4)	3.18
Amorphous + Others	SiO_2				75.8 (32.8)	2.50

XRF(Wt%): Fe2O3=1.3%, SiO2=90.8%, Al2O3=7.9%

Refinement Converged (R/E=1.26), ♦ Round=3, Iter=3, P=34, R=4.74% (E=3.77%, EPS=0.5)

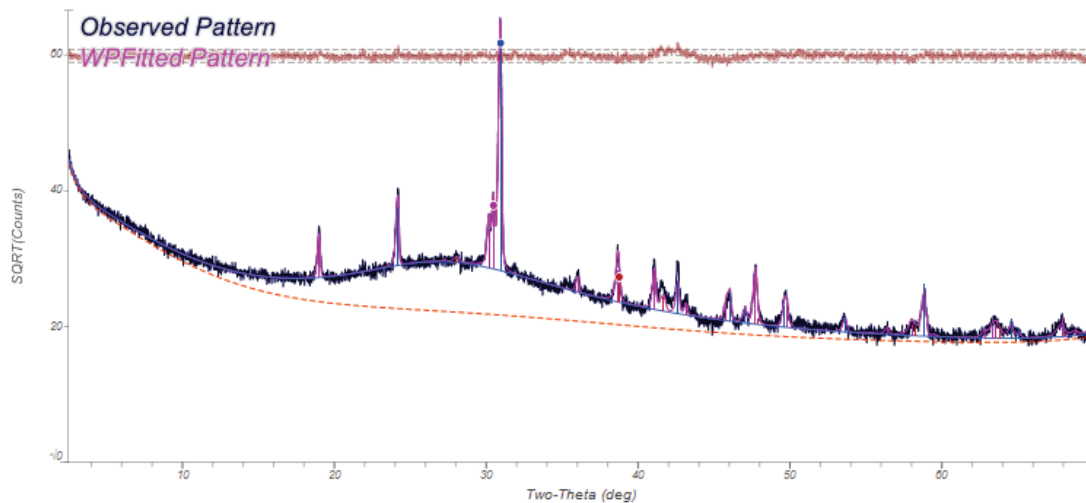
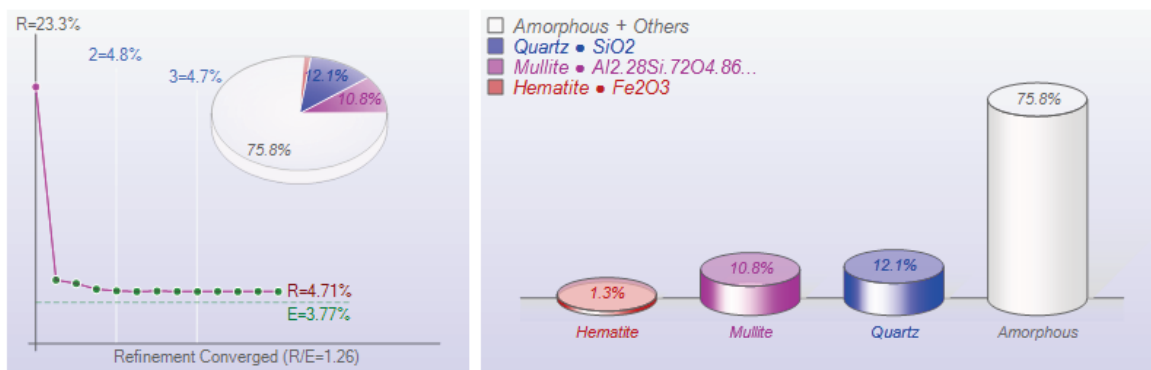


Figure C13. XRD Results of Fly Ash (150110).

Phase ID (7)	Chemical Formula	NA	NR	NP	Wt% (esd)	RIR
Quartz	SiO_2	2	34	9	18.8 (2.4)	4.04
Mullite	$\text{Al}_{2.28}\text{Si}_{7.72}\text{O}_{4.86}$	8	31	10	4.5 (0.7)	0.86
Hematite	Fe_2O_3	2	11	9	1.2 (0.3)	3.18
Magnetite, syn	$\text{Fe}^{+2}\text{Fe}^{+3}\text{O}_4$	---	7	4	0.0 (0.0)	4.90
Anatase	TiO_2	2	14	7	0.3 (0.2)	4.74
Albite (heat-treated)	$\text{Na}(\text{AlSi}_3\text{O}_8)$	---	114	11	0.1 (0.0)	0.64
Iron	Fe	1	2	5	0.4 (0.1)	8.16
Amorphous + Others	SiO_2				74.7 (15.7)	2.50

XRF(Wt%): Fe2O3=1.8%, TiO2=0.3%, SiO2=94.8%, Al2O3=3.3%, Na2O=0.0%

Refinement Halted (R/E=1.21), ♦ Round=3, Iter=6, P=51, R=5.11% (E=4.23%, EPS=0.5)

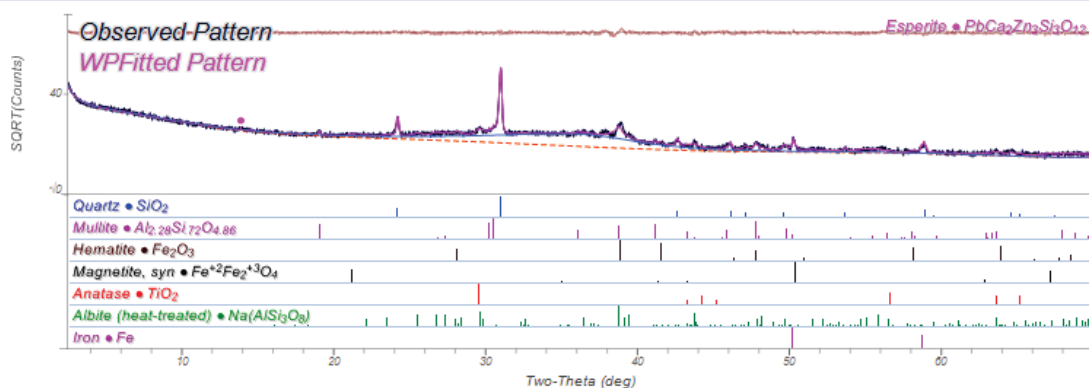
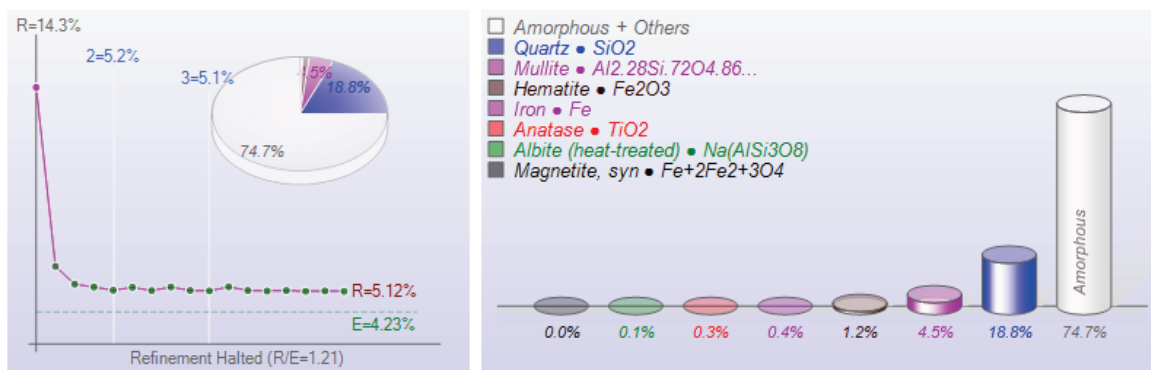


Figure C14. XRD Results of Fly Ash (150111).

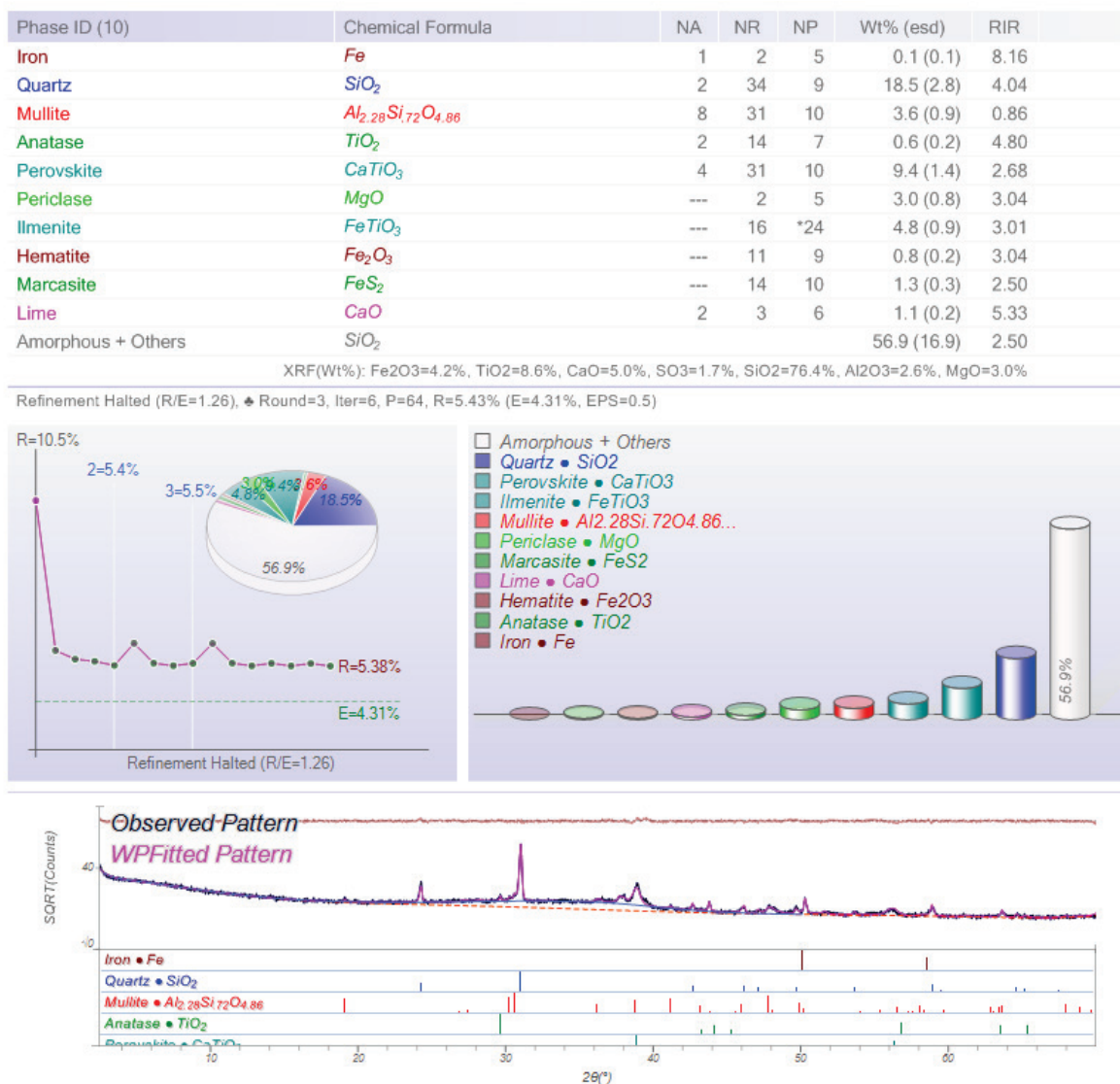


Figure C15. XRD Results of Fly Ash (150112).

Phase ID (8)	Chemical Formula	NA	NR	NP	Wt% (esd)	RIR
Portlandite, syn	Ca(OH)_2	---	8	9	17.7 (2.0)	2.90
Brucite	Mg(OH)_2	---	6	9	13.8 (1.6)	2.12
Anatase, syn	TiO_2	---	8	9	1.3 (0.2)	4.96
Hematite	Fe_2O_3	2	11	9	1.1 (0.2)	3.16
Quartz	SiO_2	---	12	9	15.2 (1.7)	3.07
Periclase, syn	MgO	---	2	5	8.5 (1.0)	1.00
Hatrunite, syn	Ca_3SiO_5	---	72	11	4.4 (0.5)	0.93
Zincite	ZnO	2	10	9	0.0 (0.0)	6.00
Amorphous + Others	SiO_2				38.0 (12.1)	2.50

XRF(Wt%): ZnO=0.0%, Fe₂O₃=1.1%, TiO₂=1.3%, CaO=16.7%, SiO₂=54.3%, MgO=18.0%

Refinement Halted (R/E=1.21), ♣ Round=3, Iter=6, P=54, R=5.26% (E=4.34%, EPS=0.5)

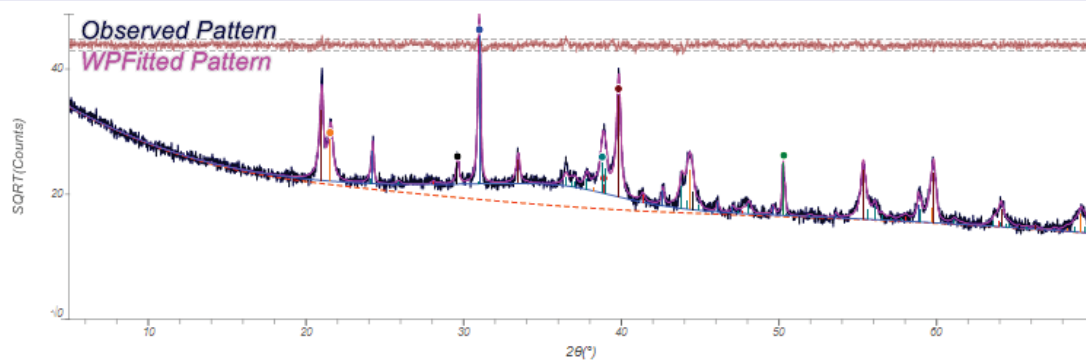
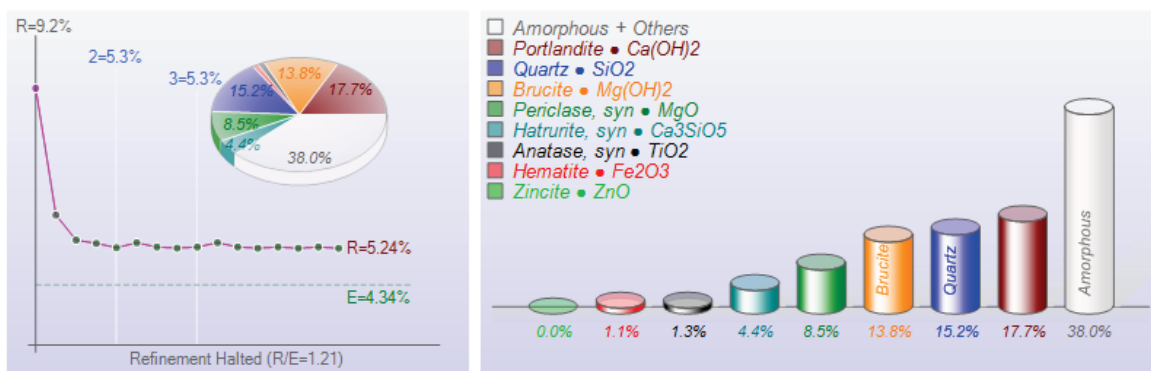
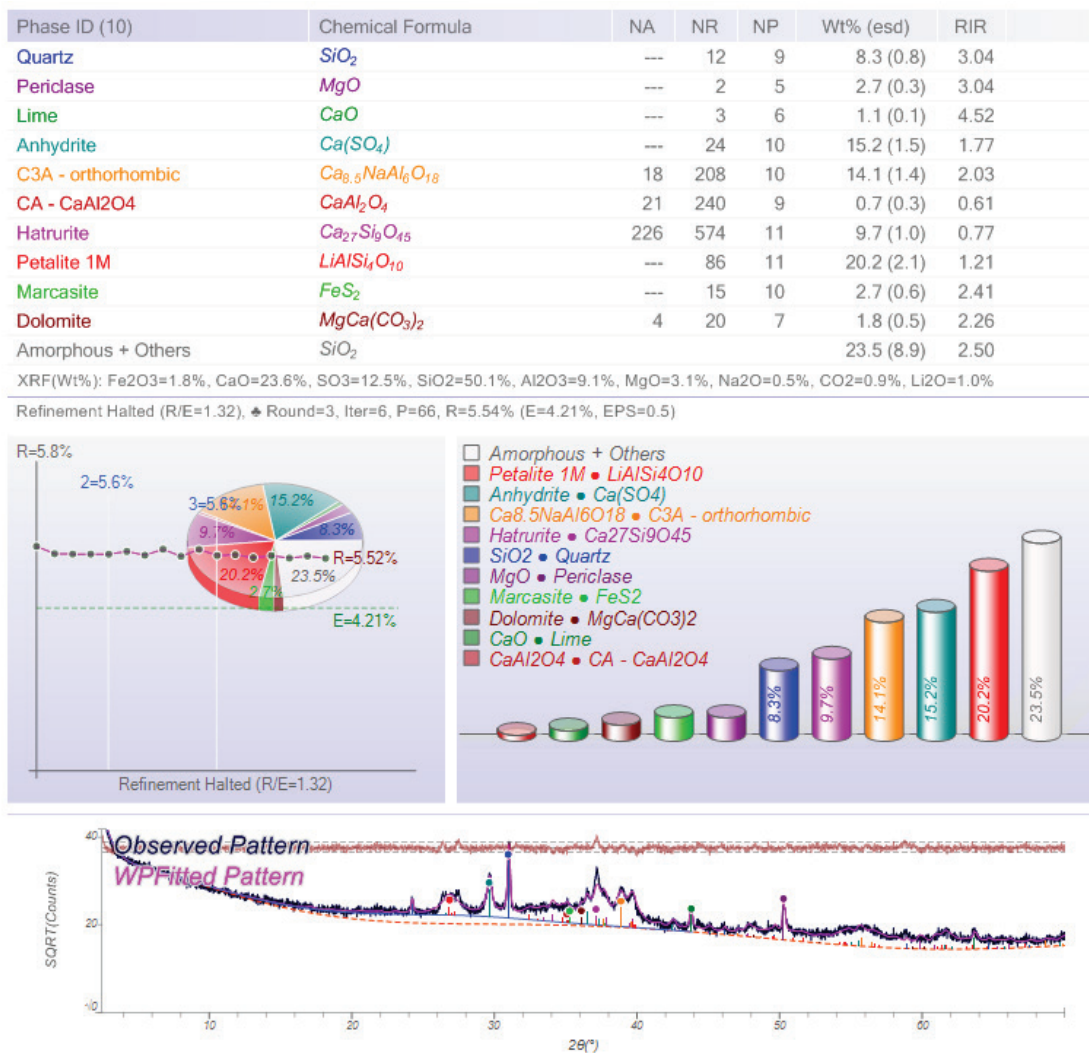


Figure C16. XRD Results of Fly Ash (150113).



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14. ABSTRACT This effort was undertaken as a part of the Materials Modeling for Force Protection (MMFP) Military Engineering 6.2 applied research program. One of the main philosophies of MMFP is the creation of a materials-by-design process for predictive properties and material selection capabilities. This report details the capability improvements to facilitate materials-by-design technologies and the collection and characterization of material data to compile and calibrate an integrated materials-by-design model. The Virtual Cement and Concrete Testing Laboratory (VCCTL) was evaluated for suitability in hydration model generation, and various programs were evaluated for initial virtualization of physical elements. To provide model inputs for particle size analysis, X-ray diffraction (XRD), X-ray fluorescence (XRF), scanning electron microscopy (SEM), and energy-dispersive spectroscopy (EDS) testing were conducted on a wide variety of constituent materials, and characterization data were processed included herein.					
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