

UNIVERSITY OF FORT HARE

OPTIMIZING GEOCHEMICAL SAMPLING SIZES AND QUANTIFYING UNCERTAINTIES FOR ENVIRONMENTAL RISK ASSESSMENT USING ANGLOGOLD-ASHANTI GOLD MINES AS A CASE STUDY

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DECLARATION

I, Elizabeth Chihobvu, declare this dissertation to be my own unaided work. It is being submitted for the Degree of Science at the University of Fort Hare, Alice. It has not been submitted before for any degree or examination in any other University.

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ABSTRACT

Generally, and particularly in South Africa, limited work done on the development of methodologies for sample sizing and quantifying uncertainties in geochemical sampling and analyses. As a result, little trust is placed on the long-term predictions of geochemical modelling for Environmental Risk Assessment (E.R.A). In addition, this leads to the slow approval of mining authorisations, water use licenses and mine closure plans.

This dissertation addresses this deficiency in geochemical sampling and analyses specifically for ERA and proposes two methodologies (i) for quantifying uncertainties in geochemical sampling and analysis as a function of sample size and analyses and (ii) for determining the optimum sample size to ensure data quality.

The statistical analysis approach was adopted as the best method for sample size determination. The approach is based on the premise that the size of the study sample is critical to producing meaningful results. The size of the required samples depends on a number of factors including purpose of the study, available budget, variability of the population being sampled, acceptable errors and confidence level.

The methodology for estimating uncertainty is a fusion of existing methodologies for quantifying measurement uncertainty. The methodology takes a holistic view of the measurement process to include all processes involved in obtaining measurement results as possible uncertainty components. Like the statistical analysis approach, the methodology employs basic statistical principles in estimating the size of uncertainty, associated with a given measurement result. The approach identifies each component of uncertainty; estimates the size of each component and sums the

contribution of each component in order to approximate the overall uncertainty value, associated with a given measurement result.

The two methods were applied to Acid-Base Accounting (ABA) data derived from geochemical assessment for ERA of the West Wits and Vaal River (Ashanti Gold mines) tailings dams undertaken by Pulles and Howard de Lange Inc. on behalf of AngloGold Ltd. The study was aimed at assessing and evaluating the potential of tailings dams in the two mining areas to impact on water quality and implications of this in terms of mine closure and rehabilitation.

Findings from this study show that the number of samples needed is influenced by the purpose of the study, size of the target area, nature and type of material, budget, acceptable error and the confidence level required, among other factors. Acceptable error has an exponential relationship with sample size hence one can minimize error by increasing sample size. While a low value of acceptable error value and high confidence are always desirable, a tradeoff among these competing factors must be found, given the usually limited funds and time.

The findings also demonstrated that uncertainties in geochemical sampling and analysis are unavoidable. They arise from the fact that only a small portion of the population rather than a census is used to derive conclusions about certain characteristics of the target population. This is further augmented by other influential quantities that affect the accuracy of the estimates. Effects such as poor sampling design, inadequate sample size, sample heterogeneity and other factors highly affect data quality and representivity hence measurement uncertainty. Among these factors, those associated with sampling, mainly heterogeneity was found to be the strongest contributing factor toward overall uncertainty. This implies an increased proportion of expenditure should be channeled toward sampling to minimise uncertainty. Uncertainties can be reduced by adopting good sampling practices and increasing sample size, among other methods. It is recommended that more information be made available for proper uncertainty analysis.

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ABBREVIATIONS AND ACRONYMS

ABA	. Acid-Base Accounting
AP	Acid Potential
AMD	Acid Mine Drainage
ANSI	American National Standards Institute
ANOVA	Analysis of Variance
ARD	. Acid Rock Drainage
ASTM	American Society for Testing and Materials
CITAC	Cooperation on International Traceability in Analytical
Chemistry	
CRM	.Certified Reference Material
DBARD	.Database for acid rock drainage
DQO	Data quality objectives
EPA	.Environmental protection agency
EURACHEM	.European chemistry
EUROLAB	European laboratories
ERA	Environmental Risk Assessment
GUM	The Guide to Expression of Uncertainty in Measurement
ICP-ES	Inductively Coupled Plasma Emission Spectroscopy
ICPMS	Inductively Coupled Plasma Mass Spectroscopy
ISO	. International Organization of Standardization
IUPAC	.International Union of Pure and Applied Chemistry
NAG	.Net acid generation
NIST	.National Institute of Standards and Technology
NNP	. Net neutralizing potential
NP	Neutralizing potential
NPR	Neutralization potential ratio
QUAM	Quantifying Uncertainty in Analysis Measurements
QA/QC	Quality assurance and quality control
(R)ANOVA	Robust Analysis of Variance
SD	Standard Deviation
UKAS	United Kingdom Accreditation Service
VIM	.International Vocabulary of Basic and General terms in
Metrology	
XRF	X-ray Fluorescence
XRD	. X-ray Diffraction

1.0 MOTIVATION

Quantification of the uncertainties associated with environmental geochemical prediction as function of sample size and geochemical analyses are always a concern when performing an Environmental Risk Assessment (ERA). This is especially so when applying for mining authorization and/or closure and water use licenses. These are particularly critical when:

- 1. Researchers and consultants compile proposals;
- 2. Mines compile their budgets or develop a term of reference;
- 3. Regulators evaluate and or authorize permits, licenses; and
- 4. Referees and reviewers peer-review a paper or examine a report.

Generally, and particularly in South Africa, there has been limited work done on the development of methodologies, quantifying uncertainties in geochemical sampling and analyses and sample sizing. Uncertainties in geochemical sampling and analyses affect the environmental study data quality. As a result, little trust is placed on the long-term predictions of geochemical modeling. This implies that results have to be validated by long duration monitoring programs thereby imposing an additional financial burden on the mining industry. This in addition leads to the slow approval of mining authorizations, water use licenses and mine closure plans.

The primary reason for this situation is lack of nationally standardized and acceptable procedures for uncertainty analysis especially for geochemical investigations for ERA. This is further exacerbated by lack of:

- Knowledge and application of statistical methods;
- Understanding the complexity of geological, soil and mine materials;
- Experience and knowledge of the significance/importance of data quality control;

- Easily understandable, operable and user-friendly methods specifically for geochemical sampling and analyses; and
- Financial constraints, i.e. limited budgets.

From the above it is clear that there is a definite need for the development of a methodology/protocol that addresses geochemical sampling (sample sizing and representativity), analyses and data quality for application, amongst others, in the ERA industry.

1.1 AIMS

1.1.1 MAIN AIM

The main aim of the project is to establish a methodology for quantification of the uncertainty associated with geochemical prediction for E.R.A as function of geochemical sample size and geochemical analysis for use by researchers, consultants, regulators and the mining industry.

1.1.2 DETAILED AIMS

In addition to the main aim stated above there are subsidiary aims which are to:

- Understand the geochemical sampling and analyses requirements for geochemical prediction for ERA.
- Establish a methodology for determining sample size (number and mass or volume) as well as sample representativity for ERA using a statistical method or formula.
- Apply and test the methodologies established for uncertainty quantification, sample optimization for ERA using the Wits basin as a case study.

1.2 METHODOLOGY

In order to achieve the above-mentioned objectives, the following tasks were performed:

 Conducting a comprehensive and extensive literature survey to review the available methodology for:

- Quantifying uncertainties associated with geochemical prediction for ERA as a function of geochemical analyses; and
- Determining geochemical sample size.
- Assessing the statistical theories, principles, concepts and application for quantifying uncertainties associated with geochemical prediction as a function of sample size and geochemical analyses, sample size determination, analytical quality control, etc.
- Develop a statistical method or formula for the determination of geochemical sample size taking into account factors such as purpose of the ERA, level of confidence required, among other factors,.
- Convert the developed methodology into a simple, user-friendly spreadsheet model or tool that can be utilized to determine and or assess geochemical sampling size.
- Identify parameters applicable to quantify uncertainties associated with prediction for ERA as function of sample size and geochemical analyses and develop a methodology that can be converted into a user-friendly format for use by researchers, consultants, regulators and the mining industry.

The methodologies established will be tested using geochemical and mineralogical data from the Wits Basin as a case study.

1.3 OVERVIEW OF THE CHAPTERS

Chapter 2 provides detailed review of literature on sample size and uncertainty in geochemical sampling and analysis for ERA. The chapter begins with an overview of geochemical sampling and analysis and quality assurance and quality control. Then it is followed by an outline of the basic sampling theory, covering measures of the central tendency mainly mean, measures of dispersion (variance, standard deviation and standard error of the mean), central limit theorem and confidence interval. A literature review on sample size determination and methodologies of quantifying uncertainty in geochemical sampling and analysis follows.

Chapter 3 contains the methodologies of the study. Methods presented are:

- The methodology of determining sample size determination using statistical analysis method. The effects of the various parameters on which the calculation of sample size depend on are discussed first. The parameters include standard deviation, standard error and confidence interval. A procedure for determining sample size is discussed and its application presented.
- The methodology for quantifying uncertainties in geochemical sampling and analysis as a function of sample size and analyses. The method is a synthesis of existing methods for estimating uncertainty including GUM by International Standards Organization (ISO), Quantifying Uncertainty in Analysis Measurements (QUAM) by European Chemistry (EURACHEM), GY's Particulate Sampling Theory, and National Institute of Standards and Technology (NIST).

Chapter 4 presents case study information based on the assessment of the potential of tailings dams and waste rock dumps in the Vaal River and West Wits mining areas to impact on water resources and implications of this in terms of mine closure and rehabilitation. The work was carried out by Pulles and Howard de Lange Company for AngloGold Ashanti Gold Ltd.

Chapter 5 outlines the results of the study based on the application of the methodologies discussed in Chapter 3.

Chapter 6 covers discussion, conclusions and recommendations for the study.

1.4 GENERAL

This dissertation is the part of work done for the WRC project entitled "Geochemical sampling and analyses for environmental risk assessment using the Wits Basin as a case study", sponsored by the Water Research Commission. Mr. William Pulles of Pulles, Howard and de Lange Inc., then Golder Associates is the project leader.

CHAPTER 2: LITERATURE REVIEW

2.0 INTRODUCTION

The main purpose of this chapter is to review literature on sample size, sampling and analysis protocols, and uncertainty in order to meet the project requirements as outlined in Chapter one. The topics reviewed include:

- Sampling and analysis process;
- Quality assurance and quality control in geological investigations;
- Basis statistic theory;
- Methods for determining the sample size;
- Uncertainty and error;
- Uncertainty sources in geochemical sampling and analyses; and
- Theories for quantifying uncertainty.

The references used include:

- Published papers,
- Conference proceedings,
- Books,
- Research reports, and
- Internet web pages.

The chapter begins with an overview of geochemical sampling and analyses process including quality assurance and quality control aspects followed by review of the basic concepts in statistics. Three methods for quantifying uncertainty: Quantifying Uncertainty in Analytical Measurement (QUAM) by European Chemistry Board (EURACHEM) (2000), Pierre Gy's Particulate Sampling theory (1992), and Empirical method based on ANOVA by Dr Michael Ramsey (1997) are reviewed in detail. A comprehensive methodology of quantifying uncertainties is then developed from the reviewed theories. The method is discussed in detail in

Chapter 3.

A review of various literatures on sample size determination follows. Aspects covered under this include factors influencing sample size as well as methodologies for determining sample size. Among the methods discussed are the statistical methods, the volume/mass method and the cost method which relates sample numbers to cost.

2.1 OVERVIEW OF GEOCHEMICAL SAMPLING AND ANALYSIS

In order to address the key issues, stated above (uncertainty and sample representativeness), it was necessary to discuss the geochemical sampling and analysis measurement. While some examples of good sampling practices can now be found (Mason, 1992; Smith *et. al.*, 2004; Dominy and Petersen, 2005), the application of correct procedures is far from universal. Figure 2-1 illustrates the various stages involved in geochemical sampling and analysis.



Figure 2-1: The measurement process in geochemical sampling and analysis

Geochemical sampling and analyses involves the processes of collecting and analysing geological materials such as waste dumps and tailings dams in order to answer questions relating to a geological scenario. Thus, to estimate the concentration of contaminants in mine waste material, an environmentalist collects several samples of waste material from the sampling target. Based on the contaminant concentration, the level of contamination of the whole population is estimated.

2.1.1 STAGES IN GEOCHEMICAL SAMPLING AND ANALYSES PROCESS

The process of geochemical sampling and analyses is multi-faceted with the main processes being sampling, sample preparation, analysis and statistical inference. Under each process are sub and intermediary processes such as transportation and sample preservation among, others but not all are always present.

2.1.2 SAMPLING

As depicted in Figure 2-1, the process begins with definition of the project objectives, and a stage which requires the analyst to clearly explain why the study should be undertaken. Koerner (1996) also encourages a clear definition of sampling objectives in order to collect a representative samples in study of coal stockpiles. Under geochemical investigation for ERA, the study may be carried out to either: (i) assess the impact of temporary stock piles of coal on ground water quality and quantity (ii) monitor pollutants' concentration changes over time (iii) application for mine closure licenses or (iv) compliance with regulations. Basically the problem definition is the preface to the investigation process which affects the outcome of the entire investigation, hence it is crucial that project aims and objectives are clearly defined and explained to all project participants.

Following the study objectives is the definition of the target. A sampling target is the portion of material at a particular time that the sample is intended to represent (EURACHEM / EUROLAB/CITAC / Nordtest, 2006). Under this, questions to be addressed include:

• What is to be sampled?

- How big is the sample?
- Where is the sample located?

Target definition is one of the major items that should be elaborated prior to designing a sampling programme. (EURACHEM/EUROLAB/CITAC/ Nordtest, 2006). In geochemical investigation for ERA, a sampling target may be waste rock dump, tailing dams, temporary stockpile or underground mine.

As cited previously, sample representativity constitutes a fundamental part of the investigation process. Every collected sample is expected to reflect certain properties of the population it is drawn from. The ability to gather samples that are representative depends on a number of factors including sampling design. Subsequent to definition of the target is selection of sampling design, a process which outlines how samples are to be collected from the target. Generally, the more coverage the sampling technique has over the target, the higher the chances of collecting representative samples. The commonly used sampling designs are the probability sampling design (Back, 2001). These include simple random sampling, stratified, cluster and systematic, Figure 2-2. The choice of design depends upon the objectives of the study.





Figure 2-2: Sampling designs: a) Simple random sampling, b) Stratified, c) Systematic sampling

Simple random sampling is a probability based sampling design in which each sample occurring within the sampling target has equal chances of being selected. The main characteristic of this design is that the possibility of introducing bias is limited by the sampling design. Appropriate situations for using the design include:

- Estimating the mean and totals for homogeneous population(Gilbert, 1987);
- Cases where limited information is available about the sampling target.

Stratified sampling divides the target into non-overlapping homogeneous units or strata based on physical or chemical properties, knowledge of site history and or vertical and lateral distribution of the contamination. As a probability sampling design, samples are collected randomly from each stratum. The main aim of undertaking stratification is to minimize the effect of heterogeneity that highly affects the precision of estimates. The number of samples within each stratum is proportional to the relative size of the site and stratum, e.g. if the area of the stratum takes up 30% of the target population then 30% of the total samples would be taken from this stratum. One main advantage of the stratified random sampling is that the increased homogeneity in each strata, increases sampling precision (Back, 2001).

Stratified sampling design is not suitable for tracking peak values (MEND, 4.7.1-1, 1994). Simple random and stratified sampling designs are suitable for sampling of discreet units and they do not take into account spatial continuity and spatial correlation of the material properties (Mason, 1992). For this reason systematic grid sampling design is used. Systematic sampling is a statistically-based method of sampling in which the target is divided into a regular grid pattern before sample extraction. Sample locations are selected at regular intervals following one or more directions. Sample locations must be spaced widely enough. One drawback of the systematic sampling is the loss in precision when the population has a periodic trend that corresponds to the sampling interval (Back, 2001).

Selection of sampling strategy is sample size determination. The decision regarding the number of samples required is determined by the purpose of:

- the study;
- accuracy required;
- sampling design;
- size of the target; and
- The budget.

Sample size determination is further discussed in Section 2.4 of this chapter and in Chapter 4. Akin to sampling design, sample size affects sample representativity, thus the larger the sample size, the higher the representativity. However, caution must exercise since collection of large number of samples does not guarantee representativity.

Another useful step is the determination of sampling technique. An appropriate sampling technique should be used when collecting samples from the site. The technique selected depends on several factors including the type of material whether solid, liquid or gas, purpose of the investigation, budget and the sampling depth required. Other considerations include the texture of the material in the case of solid samples. A good sampling tool is one that preserves the integrity of the sample. Possible factors that may cause alteration of the sample from its original state include contamination from the shallow surface material as the sample is drawn from subsurface, contamination from the sampling tool itself and chemical reaction of the sample on exposure to the atmosphere.

The different collection techniques for different types of materials are listed in Table 2.1:

Technique	Phase	Depth	Degree of sample disturbance during collection
Hand auger	solid	on/near the	High
		surface	
Trenching	solid		moderate
Split spoon	solid	subsurface	low
Vibrating core	solid	subsurface	low
drill			
Shovel	solid	on or near the	moderate
		surface	
Piezometers	liquid	subsurface	-
Bailers	liquid	subsurface	moderate
Surface grab	solid/liquid	on surface or	-
		submerged	
Rotating drill	solid	subsurface	high
Suction	liquid	subsurface	moderate
lysimeter			
Syringe	liquid	-	low
sampler			

Table 2.1: Sampling techniques (MEND Report 4.5.1-1, 1994)

Samples collected for ERA play an important role in decision making and should be augmented by proper sample handling, preservation, and storage. The manner in which the samples are to be handled is detected by the type of sample and intended use. The samples should be handled so as to preserve its integrity. Careful planning and selection of the container for holding the samples is important since these can be a significant source of sample contamination.

Table 2.2 summarises sample handling for water samples prior to analysis

Parameter	Sample container	Preservation method	Maximum recommended storage
рН	Plastic or glass	Analyse immediately	Analyse immediately.
Sulphate	Plastic or glass	Refrigerate	28 days
Cyanide	Plastic or glass	Add NaOH to pH >12,refrigerate	14 days
Metals	Plastic or glass	Add HNO ₃ to pH <2	6 months
Volatile organic compound	Glass	Refrigeration, adding HCl to pH <2	Does not specify
Silica	Plastic	Refrigerate	28 days

Table 2.2: Preservation methods for water analysis (Quevauviller, 2002)

2.1.3 SAMPLE PREPARATION

Sample preparation involves the physical and or chemical preparation processes carried out prior to analysis. It includes (i) sample homogenization; (ii) sieving and drying; (iii) sample compositing; (iv) milling and grinding and (v) riffle splitting. From the bulk field sample (primary sample), a laboratory sample is drawn. In most cases, the laboratory sample does not immediately fit for geochemical analysis and an additional preparation step is carried out to acquire a test portion (Geelhoed, 2005). The preparation method assumed relies on the analysis technique, type of material and objectives of the project. Sample dissolution is an example of one sample preparation technique that can be carried out before analysis.

Care must be taken to avoid sample contamination from the equipment. For instance, it is not advisable to dry and sieve samples that are collected for the determination of volatile contaminants, as it alters the composition of the sample.

The choice of the preparation method adopted is a function of purpose of the study, type of sample, budget, and analytical method among other factors. For heterogeneous material or where the budget is limited, various increments may be composited to form a single composite sample. Elemental analysis of major rock elements by X- Ray Fluorescence (XRF) may require the preparation of a glass bead or pellets depending on what is to be determined. In the case of Inductively Coupled Plasma Mass Spectrometry (ICP MS) analysis a different

preparation method is implemented. Sample dissolution by either acid digestion or water extraction may be employed in preparation of the test portion for ICP MS analysis.

Sample preparation for the test portion should be carried out under strict conditions in order to minimize error. However, it is critical that analysts realize and remember that a measurement is only as good as the preparation that has preceded it (US EPA, 2004).

2.1.4 GEOCHEMICAL AND MINERALOGICAL ANALYSES

After sample preparation, the sample is analysed to determine the analyte (s). Numerous analytical methods are available to qualitatively/quantitatively measure the concentration of varying analyte in geological investigations. The choice of the method is based on the material type, required accuracy, budget and most importantly, the project purpose. The analytical methods selected must be sufficiently sensitive to allow detection of a wide range of elements within the sample media at background levels. Examples of analytical techniques often adopted in geochemical studies for ERA include ICP MS which is used for multi-element analysis, XRF for the analysis of trace and major mineralogy, X-ray Diffraction (XRD) for mineralogy and many others. These methods and their application are summarised in Table 2.3.

Each method selected has to be validated. The validation of an analytical method is a process that demonstrate that the results obtained by the method are reliable and reproducible (Quevauviller, 2002). Addition objectives of method validation are:

- Identify of possible errors;
- Identify differences between methods; and
- Indicate if the method is acceptable for intended purpose.

Method validation studies rely on, precision, accuracy, linearity, detection limit, selectivity, bias and robustness.

Methods	Application	Advantages	Limitations
Atomic Adsorption Spectrometry (AAS)	For elemental analysis of solutions	-It is less sophisticated but provides quality results -fast analysis of an element	-low sensitivity to non metals -individual analysis of elements
ICP MS	Multi element analysis	Allow for rapid determination of many elements in a short period of time -high sensitivity -low detection limit	Elemental interferences Difficult to analyse non volatile samples
XRF	Elemental analysis(major and trace elements) for solids and non volatile liquids	-High sensitivity -minimum sample preparation	-Not sensitive to elements with atomic mass < Na -precision is limited by non uniformity of the sample -most suitable for major element analysis
XRD	Mineralogy of solids	Can distinguish polymorphs, isomers and conformers -less expensive -used for identification of secondary minerals	Detection and sensitivity depend on crystallite size
Microscopic study	Mineral forms, mineralogy	-provide information on mineral forms	
NAA	Elemental analysis of major and trace elements for solid and liquids	Very high sensitivity for many elements	Applicable to solid and liquids only

Table 2.3: Analysis method and their application

2.1.5 STATISTICAL INFERENCES

The last step involves making some statistical inferences about the target population based on the sampling and analytical results. Such inferences are based on the application of statistical assessment and estimation to make reliable conclusions about the conditions existing in the target area.

2.1.6 REPORTING ANALYTICAL RESULT

This is the last and crucial step of the analytical test work. It is through result reporting that the quality of measurements can be assessed by information user. Reports must be prepared according to an agreed procedure and they must accurately reflect the findings of the study. In addition, the result must be expressed in an appropriate number of significant figures and should be accompanied by a statement of uncertainty. Uncertainty is dealt with in detail in

Chapter 3.

2.1.7 IMPORTANCE OF GEOCHEMICAL SAMPLING AND ANALYSES MEASUREMENTS

Accurate geochemical predictions for environmental impact depend on the collection of representative data, i.e. data reflecting the actual conditions on the ground. The rest of the other techniques employed during the predictions depend on sampling. It is therefore important that good sampling and analysis protocols be put in place. Poor sampling and analysis techniques result in poor estimates hence decision errors. Making decisions under uncertainty may lead to serious consequences and heavy financial penalties.

2.2 QUALITY ASSURANCE AND QUALITY CONTROL

The value of geochemical measurements depends upon the level of confidence that can be placed in the results. Critical to geochemical sampling and analyses for ERA are data quality objectives and quality assurance and quality control programs. Since results obtained from sampling and analyses play a crucial role in decision making, emphasis should be placed on the quality of measurement results. The quality of a measurement result and the confidence placed on it depend on strict adherence to QA/QC program.

2.2.1 OBJECTIVES OF A QUALITY ASSURANCE/QUALITY CONTROL (QA/QC) PROGRAM: (Downing, and Mills, 2007)

- Document the procedures and methods of sample collection, preparation, and analysis.
- To give assurance that data generated are indicative of the study site characteristics
- To provide assurance as to the precision and accuracy of data.
- To provide assurance as to the accuracy from using recognized reference standards.
- To provide a chain of custody of samples.
- help to ensure that results are valid and fit for purpose (CITAC/EURACHEM, 2002)
- To provide measurement of the uncertainty in the underlying data.
- prevent the entry of large errors into the database used for geochemical modeling,

It is important for both laboratories and their customers to realise that QA cannot guarantee 100% reliable results due to errors which lead to measurement uncertainty (CITAC/EURACHEM, 2002). The programs should be set up in the early planning stages of the investigation.

2.2.2 CRUCIAL ASPECTS FOR QA/QC PROGRAM

A quality assurance and quality control program for acid rock drainage (ARD) should address the following aspects:

2.2.2.1 Sampling

It is the most vital component of ARD. If the sampling work is not done properly, the mistakes associated with the collected samples will be carried over to the succeeding components of ARD. Inconsistent and inappropriate sampling procedures should therefore be avoided. Factors to be considered under this are as follows:

- Qualification of the personnel;
- Quality control samples (Table 2.4);
- Equipment calibration and decontamination;
- Sample record; and
- Chain of custody.

2.2.2.2 Analyses

Correct and reliable analysis relies on clear and adequate specification of the objectives of the study and strict adherence to good QA/QC practices. Without a clear understanding of the needs for analysis it is difficult to obtain useful results that clearly address the problem in question. Factors to be considered under this are as follows:

- Laboratory selection;
- Laboratory accreditation;
- Staff qualification;
- Instrument calibration and servicing;

- Laboratory quality control samples (Table 2.5);
- Documentation of records; and
- Reporting results.

Table 2.4: Field quality control samples (Van Ee, Blum, and	Starks, 1990)
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ТҮРЕ	FUNCTION
Duplicate sample	Used to demonstrate the reproducibility of the sampling techniques and to test the precision of the overall analytical system (field and laboratory).
Field Blank	Used to check on potential sources of contamination resulting from exposure to the ambient air or from improperly cleaned sampling equipment.
Trip blank	Used to trace contamination introduced during shipment
Split Samples	Used to determine the precision of a laboratory analysis by allowing a comparison of analytical results for two parts of the same sample from the same location.
Spiked Sample	Used to provide a proficiency check on analyte recovery as a function of analyte loss during transport and storage of the collected samples and as a function of the analytical procedures and equipment.
Background Sample	Used to compare site conditions to the surrounding environment

ТҮРЕ	FUNCTION
Certified reference material (CRM)	Used for the calibration of an apparatus and assessment of a measurement method.
Laboratory replicate	Used to evaluate the method precision.
Matrix Spike	Used to establish if the method or procedure is appropriate for the analysis of a particular matrix.
Laboratory blanks	Assess contamination within the laboratory

2.2.2.3 Data Management

Sampling and analyses program will generate a considerable amount of data that will need to be properly managed to facilitate periodic review and revision of the sampling strategy. Data generated during a site investigation should be collated and presented in a logical form to enable the information to be assessed. Database provides a means by which the large volumes collected can be managed and maintained. A good database should contain database should contain all information needed to easily evaluate the quality of data, manage that data, and allow easy reference to hard copy (Long, 2007). There are various types of databases available but the main ones are flat file and relational database. Flat file style database consist of one or more unrelated tables containing considerable records. They are usually ideal for small amounts of data. An example are the spreadsheet such as excel. Relational databases have a much more organized structure for storing data. The tables within the database are linked to one another hence the term relation. Unlike most relational databases, spreadsheet approach allows for more complex and detailed analysis and facilitates the comparison of calculation methods.

2.2.2.4 Data analysis.

Data analysis procedures provided a platform by which errors and flaws within the measurement results can be identified. Data analysis involved all procedures such as statistical technique, plots, applied to describe, summarise, and compare data. It is recommended that all collected data should be analysed to determine any spurious results. Simple plots and statistical analysis may be carried out. Example from the ABA results, one may plot inorganic carbonated NP against total NP to determine the possibility of correlation between carbonate content and total NP.

2.2.2.5 Reporting analytical result

This is the last and crucial step of the analytical test work. It is through result reporting that the quality of measurements can be assessed by information user.

At least the following aspects should be incorporated in the results report:

• Sample ID and laboratory ID;

- Analysis date;
- Sample size;
- Sample (analyte)concentration units;
- Description of the analytical procedure adopted and calibration;
- Quantification and validation procedures;
- Quality control sample results; and
- Statement of uncertainty.

2.3 BASIC STATISTICAL THEORY

The main aim of undertaking geochemical sampling and analysis is to make some inference about the population from which a sample is drawn. It is through the application of basic statistical tools that makes inferences possible. This section covers basic statistics including the following:

- Sampling distribution;
- Central limit theorem; and
- Confidence interval.

2.3.1 SAMPLING DISTRIBUTION

Sampling distributions are theoretical distributions that comprise of infinite number of sample statistics taken from an infinite number of randomly selected samples of a specified sample size. Under this, the main question to be answered is "how close is the value of the statistic to the corresponding parameter of the entire population?" (Lenth, 2001). For example, if we have 60 elements, analysed in a sample from a population and our sample mean \bar{x} is 18.2, we would like to know how far this might be from the mean μ of the entire population.

In most cases, given the difficulty of measuring the population parameters, we often infer the value of the mean μ , variance σ^2 and standard deviation σ of the population from the sample mean \bar{x} , sample variance s², sample standard deviation s, respectively. Thus the sampling distribution of the various sample statistics are used as estimates of the corresponding population parameter.

Although many distributions exist, the often used is that of the mean (Crépin and Johnson, 1993). The mean of the sampling distribution is equal to the sampled mean of the population:-

$$\overline{x} = \mu$$
 2.1

Where

 \bar{x} = the samples mean

 μ = the population mean.

The samples mean \bar{x} can be expressed by the equation below:

$$\bar{x} = \frac{\sum_{i=1}^{n} X_i}{n} \qquad 2.2$$

Variance of sampling distribution of the mean indicates how the values spread around the expected mean value. The sample variance can be expressed as:

$$S^{2} = \frac{1}{n-1} \sum_{i=1}^{n} (X_{i} - \bar{X})^{2} \qquad 2.3$$

Where n is the sample size.

Standard deviation (S) is the spread of values around the average in a single sample:

$$S = \sqrt{\frac{1}{n-1} \sum_{i=1}^{n} (X_i - \bar{X})^2}$$
 2.4

The standard deviation (S) value gives an indication of how the values spread within a given dataset. A large standard deviation (S) is an indication of data values that have high dispersion and wide spread of the numbers and are wide distribution of the numbers from their mean (μ). A small value of S shows that the data values are close to the mean of the data.

For a normally distributed dataset, there is a likelihood that approximately 68% of the scores

are within one standard deviation of the mean (μ) and 95% of the scores are within two standard deviations of the mean μ . Further there is a 99.7% chance that individual measurements will fall within three standard deviations of the mean as shown in Figure 2-3 below:



Figure 2-3: Standard deviation (Wikipedia, 2008)

Standard error of the mean: It is the standard deviation of the sampling distribution of the mean (Lane, 1993-2007). Thus it defines how the sample averages () spreads around the average (μ) of a population. It can be expressed as:

Standard error is also related to sample size (n). Equation 2.5 indicates that the standard error of the mean decreases proportionately with the square root of the sample size. Thus, the sampling distribution of will be more concentrated the larger the sample size. Standard error is also one of the main measures of precision and uncertainty.

2.3.2 CENTRAL LIMIT THEOREM

The basis of this theory is that, for a given distribution with a population mean μ and population variance σ^2 , the sampling distribution of the mean approaches a normal distribution with a mean (μ) and a variance σ^2 as the sample size (n), increases (Lenth, 2001). But how large should the sample size before it is close to normal distribution? Generally this is not an easy question to answer since there is not an exact point where it can be said that the sample

size is large enough to warrant an assumption that the sampling distribution is normal. A sample size of n = 30 is usually recommended as a cut-off point where it assumed that a sampling distribution is approximately normal (Stockburger, 2002)

The concept is employed throughout the geochemical measurement process. For instance it is used in determining the number of samples and quality control situations where single measurements are erratic (Long, 2007). If more measurements are made, the average of the results is much more reliable for decision-making than when few measurements are made.

2.3.3 CONFIDENCE LIMITS OR LEVELS

The confidence levels indicate the level of error or risk one is willing to accept. If n samples are drawn from a population with a mean μ and a standard deviation σ :

75% ($\alpha = 0.25$) confidence limits for sample means = $\mu \pm 1.151 * \sigma/\sqrt{n}$ 95% ($\alpha = 0.05$) confidence limits for sample means = $\mu \pm 1.960 * \sigma/\sqrt{n}$ 99% ($\alpha = 0.01$) confidence limits for sample means = $\mu \pm 2.576 * \sigma/\sqrt{n}$

2.4 SAMPLE SIZE

The objective of sampling is the collection of a representative sample-that is a sample that exhibits the true characteristics of the target population from which the sample is drawn. The collection of representative samples is dependent not only on good sampling design, but also on the collection of the right number of samples to satisfy the requirements of the study.

Sample size determination is a crucial part of the investigation procedures. Collection of too many samples is expensive and wasteful while too few samples may be unrepresentative (Reytblat, 2000). In scientific experiments involving human or animal subjects such as ERA, sample size is a key issue for ethical reasons. An under-sized experiment exposes the subjects to potentially harmful treatments without advancing knowledge while an over-sized experiment, an unnecessary number of subjects are exposed to a potentially harmful treatment, or are denied a potentially beneficial one (Lenth, 2001). It is therefore the investigator's

responsibility to make sure that the optimum number of samples exhibiting the true characteristics of the population or area is collected.

2.4.1 DEFINITION OF SAMPLE SIZE

Townend (2002) defines sample size as the number of measurements or observations made from each population.

2.4.2 FACTORS AFFECTING SAMPLE SIZE

Generally the factors that affect sample size mainly depend on the technique to be used in calculating the number of samples, but at minimum sample size required depends on:

(a) **Purpose of the study**: it is one of the major factors affecting sample size. For a small pilot study, prior to detailed sampling, a few samples may be collected to obtain a rough estimate of the conditions on the ground. For detailed studies, a minimum of 30 samples is recommended in order to capture the properties that exist within the material understudy. For the purpose of Acid Mine Drainage assessment of *in situ* rock, British Columbia Acid Mine Drainage Task Force (1989) recommends a minimum sample number based on the size of a geological unit (MEND, 4.5.1-1, 1994).

(b) Available information about the study target: Available information and statistical methods provide guides to the determination of optimum sample size (Germishuyse, Yibas and Pulles, 2002). In geological studies, such information includes: spatial distribution and geological properties of the material; geotechnical and physical properties and geochemical characteristics. If the sampling target was studied before, the previous information can be used to reduce required number of samples. Depending on the objectives of a study, the previously obtained mean and variance estimate values can be used to calculate sample size for the present study (Taylor and Kuyatt, 1994).
(c) Variability of the geological properties of the geological material to be studied: Basically the more heterogeneous a population is, the larger the sample size required to characterize the effect of variability. Conversely, a homogeneous population requires fewer samples.

(d) Size of the sampling target: If the sampling target is large then more samples are required. On the other hand when the sampling unit is small, fewer samples are generally required.

e) Precision: Generally more samples are required to achieve greater precision.

(e) Cost: There is a direct relationship between cost and sample size. As the number of samples increase, the cost also increases. The relationship between cost and sample size can be demonstrated by the Figure 2-4, Myers (1997). When few samples are collected the cost is generally lower but the risk of collecting unrepresentative samples is high. When more samples are taken the sampling cost usually increases but errors and misclassification cost decrease (Back, 2001). Therefore a balance among competing factors should be sought.



Figure 2-4: The relationship between cost and sample size (Myers, 1997)

2.4.3 METHODS OF DETERMINING SAMPLE SIZE

There are numerous techniques for determining sample size. Some of the techniques are based on application of statistical or mathematical formula while some are based on published tables and others on imitating the sample size from a previous similar study (Israel, 1992). For example, EPA tables for estimating the sample size required listed in Appendix B. One major drawback of using such published tables is that they are mostly based on the assumption that the measured attributes are normally distributed, but when the data for the study do not follow this distribution; the use of the tables may lead to unreliable results.

The use of sample size values from a previous similar study carries the risk of repeating errors that were made in the previous study.

The following sections review methodologies for determining sample size based on the application of statistical or mathematical formula in determining sample size. The methods covered include:

- Statistical analysis method; and
- Volume method.

Statistical methods

A number of schools of thought have come up with different statistical formulas for calculating the number of samples required. Crépin and Johnson (1993) recommended the use of the statistic analysis method based on mean in determining the required sample size. The method can be expressed by the equation below:

$$n = \left(\frac{t_{\alpha/2} \times s}{d}\right)^2 \qquad 2.6$$

Where

n = the required number of samples

 $t_{\alpha/2}$ = the Student's t-value for a chosen level of precision, for example a confidence level of 95 % (α =0.05).

s = sample standard deviation.

d = the variability in mean estimation we are willing to accept critical value, d² is the desired level of precision.

Barth *et.al* (1989) suggested a different approach, in order to determine the minimum number of samples, n, required to achieve a specified precision and confidence level at a defined minimum detectable relative difference, one-sided, one-sample t-test Equation 2.7, or one sided, two sample t- test, Equation 2.8 can be used.

$$n \geq \left[\frac{Z_a + Z_b}{D}\right]^2 + 0.5 Z_a^2$$
 2.7

$$n \geq \left[\frac{Z_a + Z_b}{D}\right]^2 + 0.25 Z_a^2$$
 2.8

Where:

n = number of samples

 Z_a = the percentile of the standard normal distribution such that $P(Z \ge Z_a)$ = infinity

 Z_b is the percentile of the standard normal distribution such that $P(Z \ge Z_b) = infinity$

a = probability of a Type I error

b= probability of a Type II error

D= minimum relative detectable difference/CV

CV= coefficient of variation.

The parameter values are listed in Appendix B.

Huesemann, (1994) and Back, (2001) argue that to determine the number of the samples required, a small priori study area is necessary. From this, a rough estimate of the geochemical and analytical variance can be established hence the necessary number of samples and number of analysis per sample to meet the specified objectives can be calculated. The statistical formula to calculate the sample size can be expressed as:

$$n = \frac{S_c^2}{S_x^2}$$
 2.9

Where

n = number of samples

 S_c and S_x are coefficients of variation (relative standard deviations).

For example, if the priori field study has given $S_c=40\%$ and the sampling objective is $S_x \le 10\%$, the approximate sample numbers required is 16.

Volume Method

The British Columbia Acid Mine Drainage Task Force (1991) developed a curve that can be used as a guideline to ascertain the minimum number of samples required to characterize each geological unit, Figure, 2-5. The curve was developed based on a limited number of sites within British Columbia, Canada as a function of the mass of the geologic unit being sampled (Downing, 2007). The recommended minimum number of samples is given below (MEND, 4.5.1-1, 1994):

$$n = 0.026M^{0.5}$$
 2.10

Where

n = number of samples

M = mass of the geologic unit or lot in tons (M> $6*10^3$ ton).

For a 10^6 ton waste deposit, the minimum number of samples is 26 as shown in Figure 2-5.

The sample volume required for a given study may be determined by using Gy's sampling which can be expressed by the following equation:

$$S^2 = \frac{Cd^3}{m}$$
 2.11



Figure 2-5: Sample size determination using the volume method (MEND, 4.5.1-1, 1994)

Rearranging the equation in order to make m subject of the formula would give (Yang, 2006):

$$m = \frac{Cd^3}{S^2}$$
 2.12

Where;

S is the sample standard deviation of the error

d is the size of the largest particle in the sample

C is a product of

- liberation factor;
- shape factor;
- particle size distribution factor; and
- Composition factor.

The output is a mass needed to achieve a specified sampling error (Ramsey, 1998).

Robertson, Robertson and Kristen (1990) discouraged the use of fixed sample numbers e.g. volume method. Instead he suggested a preliminary sampling plan for determining the best sampling strategy. The following strategies were recommended:

Replicate sampling: it involves extracting several samples at the same location. It can be

adopted when the objective is determination of the mean of a given variable.

Sequential sampling: This can be used when the objective is to reduce variance between samples.

Exceedence driven sampling: the strategy is designed to identify outliers within a dataset. Croarkin (2003) defines an outlier as "an observation that lies an abnormal distance from other values in a random sample from a population". When an outlier has been identified, sampling number is increased at locations where these outliers occur.

Markovian sampling strategy reflects the rise beyond a threshold level determined by previous sampling. The sample size is adjustable and based on "alert" levels. (Robertson, Robertson, and Kirsten, 1990)

Stratified sampling strategy divides the waste rock pile into homogeneous stratum. The strategy under its counterparts provides some statistical predictability.

2.4.4 PROCEDURE FOR SAMPLE SIZE DETERMINATION

Croarkin (2003) proposed the following process for determining sample size:

- Determine the purpose or objectives of the study;
- Build up or find an equation that relates the desired precision to sample size;
- Use previous or prior information to estimate unknown parameters such as variance or mean;
- When stratified design is adopted, determine sample size for each stratum; and The final sample size is scrutinized for practicality.

2.4.5 SUMMARY

The determination of the required sample size is a function of the study objectives, cost and variability. Each of the factors should be taken into account before deciding the number required. The method chosen should satisfy the needs at hand. One major drawback associated with use of statistical methods for determining sample size is the assumption that a random sample was drawn. This assumption brings in some complications especially when a different complex sampling design is adopted, for example-stratified sampling. The use of this design requires one to take into account the variances of sub-populations, strata, or clusters prior to

estimation of the variability in the population as a whole can be made (Israel, 1992).

In addition the use of statistical methods in sample size determination requires a good estimate of the standard deviation. However more often than not, when the sample mean is unknown, so is the sample standard deviation. The mass methods are only applicable where solid materials are to be sampled. It cannot be applied where the sampling target is either a gas or fluid. Studies conducted by Smith and James (1981) have revealed that the mass techniques tend to give an "over-estimate" of mass. In addition, determination of the parameters linked to the calculation of the constant C is difficult.

The cost method, although important does not take into account important aspect such as the required precision, confidence, and prior knowledge of the target. The aspects impact greatly on sample representivity and accuracy of the estimates.

Several methods of sample size determination were discussed in detail. Of all the methods, the method which satisfy the requirement of this study is the statistical analysis approach based on the formula $n = Z_{\alpha/2}^2 * S^2/d^2$. One major advantage of the method is that some of the most important parameters that impact on samples size are directly incorporated in the formula for sample size determination. Such parameters include level of required accuracy, standard deviation which shows dispersion and confidence interval. This importance of this is that the user has control over the input hence have an idea of the output. These factors and their effect on sample size determination are further discussed in Chapter 3.

2.5 UNCERTAINTY

With every measurement including geochemical measurements and calculated result, no matter how carefully they are made, there is always uncertainty. Only measurements involved in counting are accurate (Weber, 2006). In geochemical sampling and analyses, uncertainty arises from the fact that only a portion of the target is sampled instead of the whole population hence the obtained measurement value is one probable value out of a universe of possible values for the population under study. That is if one were to repeat the same measurement several times it is likely that different results will be obtained. Variation between these repeated measurements is attributed to some influential factors resulting from inconsistency of the measurement process. The influential factors are referred to as uncertainty sources or components. Given the inherent variability of measurement, measurement results should be accompanied by an estimate of the uncertainty for the measurement result (Adams, 2002).

2.5.1 **DEFINITIONS**

2.5.1.1 Uncertainty

International Standards Organization (ISO, 1995) defines uncertainty as "a parameter, associated with the result of a measurement that characterizes the dispersion of the values that could reasonably be attributed to the measurand". The "measurand" is the quantity being measured (US EPA, *et.al*, 2004). The parameter may be, for example, the experimental standard deviation of the arithmetic mean. Taylor (1996) defines uncertainty as a measure of the incompleteness of one 's knowledge or information about a quantity whose true value could be established with a perfect measuring device. Thus uncertainty can be summarised as a range of values in which the true value of the measurand is expected to lie with high probability. Related to the concept measurement uncertainty are measurement error, precision and accuracy. ISO (1993) define accuracy as the closeness of agreement between a test result and the accepted reference value. Accuracy cannot be measured, it can only be surmised from secondary measures that reflect the likelihood that an analytical method is accurate (Mason, 1992). The precision is a measure (or an estimate) of the reproducibility (i.e. repeatability) of repeated measurements.

2.5.1.2 Error

Error is defined as the difference between observed value and the true value of the measured quantity (ISO, 1993). It can be expressed by the equation below:

$$E = x_i - x \qquad 2.13$$

Where E is error, x is the true value and x_i is the obtained or observed measurement value.

Pitard (2005) defines error as "variation in measurements, calculations, or observations of a quantity due to mistakes or to uncontrollable factors".

Errors are not "mistakes" but refer to inevitable inaccuracies and imprecision of data resulting from limitations of selecting small representative samples from large volumes of material and from the sensitivity of analytical methods. As errors are cumulative, each step of sampling through to analysis can be identified as a source of error (Downing, 2007).

It consists of two components which are systematic and random errors.

Systematic errors: these errors often lead to bias of the measurement, shifting all measurement in a systematic way so that measurement results vary in a predictable manner. For example, a balance may constantly read mass 0.001g higher because it was poorly calibrated. These errors are consistent and can be difficult to identify and correct (Owen, and Jones, 1990). Systematic error can result in high precision but poor accuracy. Unlike random errors, they cannot be minimized by statistical analysis of repeated measurements. Systematic errors can be estimated using a certified reference material and may be reduced by calibration.

Random errors: these errors are variations between successive measurements made under identical experimental conditions due to limitations of the equipment or techniques used to make the measurement. Unlike, systematic errors, the errors cause measurements to vary in an unpredictable way hence the term "random error" (Owen, and Jones, 1990). Sources of random error include lack of sensitivity of the instrument, noise in the measurement and imprecise definition of object dimensions. In experimental measurements, random error can be reduced by taking sufficient number of measurements (Morgan and Henrion, 1990).

2.5.2 UNCERTAINTY VERSUS ERROR

Although some scholars for example Geelhoed (2005) use the two terms error and uncertainty interchangeably, these terms differ in a number of ways. Uncertainty takes a range of values in which the true value of the quantity being measured is expected to lie, while error is made up of an individual value reflecting how the measured value deviates from the true value, Figure 2-6. In most cases, the true value is unknown. The likely errors for the measurement are

estimated by prescribing a range/bound for these errors as illustrated in Figure 2-6. This collective range of errors is called uncertainty.



Figure 2-6: Relationship between value, error and uncertainty (Leito, 2004)

Example 1

For example an analyst from repeated measurement of mass (g) reported the results below:

150.02	150.00	149.97	149.98	150.03
--------	--------	--------	--------	--------

The mean of the measurement result is 150g. The range of these values is 0.03g (150.03-149.97) that is the difference between the maximum and the minimum value. This is an estimate of the possible range of errors. This can be written as:

Mass=150.00g (mean) $\pm 0.03g$

This means that the analyst is confident that the true mass value lies between 149.97 and 150.03g.

Thus ± 0.03 g reflects uncertainty.

2.5.3 NEED FOR EVALUATING UNCERTAINTY

Many world renowned organizations such as, American National Standard Measurement and Associated Instrument Quality Assurance for Radioassay Laboratories, (ANSIN), ISO, EURACHEM, UKAS, and US EPA recommend that a statement of uncertainty accompany each measurement. Most of these bodies have release several reports and guidelines for evaluating and estimating measurement uncertainty. Thus one aim of reporting a result accompanied by an uncertainty value is to comply with regulations for reporting a scientific measurement result. For example an estimation of the measurement uncertainty is required in ISO 17025 /9 (Magnusson, 2004). A proper acknowledgment of uncertainty associated with a given measurement result improves the credibility of the results. Further communication of uncertainty can lead to more informative decision-making, as the decision makers are able to carefully consider what may be possible and the consequences of such events (Huesemann, 1994).

2.5.4 TYPES OF UNCERTAINTY

Uncertainty can be classified into two broad classes' namely sampling and analytical uncertainty. Sampling uncertainty refers to all possible sources of uncertainty, which may occur prior to analysis. In geochemical investigations, sampling uncertainty is often considered as the dominant source of uncertainty (Ramsey, 1998). This uncertainty is a result of many factors with the major source being heterogeneity. Other causes include improper sample handling and incomplete sampling equipment contamination. Sampling uncertainty can be reduced by increasing the sample size.

Analytical uncertainty results from the limitations of analytical method used during analysis procedure. Common contributors include poor detection limits, analytical bias, lack of precision, and susceptibility to interferences. These factors create an uncertainty interval around the analytical result.

The Triad (2004) recognizes five types of uncertainty associated with measurements in hazardous waste site namely:

- Decision;
- Model;
- Analytical;

- Sampling; and
- Relational uncertainty.

Decision uncertainty is the total uncertainty contribution from political, economic, and public perception factors, along model, analytical, sampling, and relational uncertainties. Model uncertainty refers to the uncertainty associated with the inability of a model to accurately portray the key site features. Relational uncertainty is the uncertainty associated with the relationship between a parameter being measured and the true parameter of interest from a decision-making perspective (Triad, 2004). As for sampling and analytical uncertainty, the principles as discussed in section above apply.

Haimes (1998) recognizes two types of uncertainty namely knowledge and variability uncertainty. Variability uncertainty is linked to heterogeneity of the material being sampled. Knowledge uncertainty results from incomplete knowledge of the model that should be applied. The two are sometimes referred to as aleatory and epistemic uncertainty respectively. The two types are summarised in the Figure 2-7.

This classification of uncertainty has the disadvantage of excluding other possible sources attributed to sampling and analysis. For instance, instrumental effects, which are linked to analysis, also contribute to measurement uncertainty.

Back (2001) divides uncertainty into three classes namely pre-sampling, sampling uncertainty, and post sampling uncertainty. Pre-sampling uncertainty refers to uncertainty that occurs prior to sampling. Uncertainty is introduced during preliminary studies when prior information is used to estimate the value of some parameter. For example, use of standard deviation from previous studies to estimate the number of samples required for the present study. In this process, it is inevitable that the estimated parameter will be associated with some uncertainty that is carried over to the present study.



Figure 2-7: Types of uncertainty (Haimes, 1998)

Sampling uncertainty arises due to the fact that it is practically difficult to obtain samples that are reproducible and of highest quality. Generally field sampling is much larger than uncertainty that is associated with post-sampling activities, such as sample preparation and chemical analysis (Crépin and Johnson, 1993). Post sampling uncertainty is uncertainty that occurs after sampling. Source of post sampling uncertainty include:

- Poor sample handling and transport;
- Contamination of the sample during preparation and sub-sampling; and
- Poor instrument calibration and human error.

Gy (1998) recognizes seven types of errors from both sampling and analyses. These are fundamental, grouping and segregation errors, periodic errors, increment delimitation error, increment extraction error, preparation errors and analytical errors. The most important of these is the fundamental error that is associated with heterogeneity. This error unlike other errors cannot be eliminated but can be reduced homoginising the sample.

2.5.5 SOURCES OF UNCERTAINTY

For geochemical measurements, the identification of the possible sources of uncertainty is not as simple a task. This is due to the complexity of the measurement process. In order to accurately identify the sources of uncertainty with confidence, it is advised to consider each process undertaken during the measurement process as potential source of uncertainty.

Sampling strategy

The sampling strategy defines how the samples are collected from a given sampling target. Under this uncertainty results because of:

- Only limited number of locations within the study population is selected.
- Few samples were collected.
- Strategy for sampling does not capture the extent of the variability that exists within the population being sampled.

Sample preparation

Prior to analysis, all samples undergo preparation. The aim of sample preparation is to produce a manageable sample suitable for laboratory analysis. Sample preparation includes both chemical and physical processes such as acid digestion, drying and or sieving. These processes introduce uncertainty through the following (Pitard, 2005):

- Contamination of the sample;
- Loss of fines during crushing or fine particles may stick to the equipment due to electrostatic forces;
- Change in the physical, chemical, or mineralogical composition of the sample due for example to excessive drying; and
- Poor handling of the sample and sample mix up.

> Analysis

The analytical method itself can be a major source of uncertainty for the measurement results. Geelhoed (2005) defined this uncertainty as the difference between the analysis result and the true value of the analyte concentration in the test portion. This variability component arises from imperfections of the analysis operation. It includes uncertainties associated with:

- Variations in the measurement conditions for example humidity changes affect results especially materials sensitive to humidity;
- Recovery of the analyte from a complex sample matrix;
- Reagent purity;
- Computational effects for example rounding off of figures can lead to inaccuracies in the final result;
- Human errors for example reading a meter or scale consistently high or low; and
- Error that occurs during the publication of these data results.

> Heterogeneity

Heterogeneity expresses the variability inherent in the population or material being sampled in terms of the chemical and physical make-up. This may be particulate size, and levels of contamination. It includes both constitutional and distribution heterogeneity. Constitutional heterogeneity, describes the differences in composition, shape and density between the smallest individual and separable constituents. Distributional heterogeneity further describes the part of the overall heterogeneity stemming from spatial distributions at higher levels (Esbensen, 2005).

It has been identified as the major source of measurement uncertainty in geochemical sampling and uncertainty (Minkkinen and Paakkunainen, 2005).

Gy defines heterogeneity as:

$$h_i = \frac{a_{i-}a_L}{a_L} \times \frac{M_i}{\overline{M_i}}$$
 2.14

Where

 h_i = heterogeneity,

 a_i = value of the process variable in sample i,

 a_L = mean value of the process variable in the sample,

 M_i = the mass of the sample i

\overline{M}_i = the mean of the sample mass

A case in point is when uncertainty is introduced because increments from different locations in the target area will have different expected concentrations of analyte in a heterogeneous material concentration. For measurements subjected to averaging, sample-to-sample variation of the analyte concentration introduces uncertainty to the measurement result.

Instrument

The instrument used for extraction of the sample from the target and analysis may present the weakest link in the measurement process. For most measuring instruments, uncertainty results from:

- Poor detection capabilities of some element, for example XRF cannot detect minor elements with accuracy. The causes of such in-capability are attributed to short counting time and high background interference and long decay time;
- Improper instrument calibration; and
- Manufacturer specifications: most measurement instruments have an accompanying statement of accuracy or tolerance level specified by the manufacturer. This value should be incorporated in uncertainty measurements.

2.5.6 METHODS FOR QUANTIFYING UNCERTAINTY

The section reviews some of the most popular method for quantifying uncertainties in environmental studies. These include:

- Particulate sampling theory by Gy (1992);
- EURACHEM method for Quantifying Uncertainty in Analytical Measurements (QUAM) (2000); and
- ANOVA (empirical approach) by Ramsey (1998).

The section begins by listing the parameters for expressing uncertainty and a detailed discussion of each method follows.

2.5.6.1 Parameters for expressing uncertainty

Table 2.6 highlights some of the parameters used in expressing uncertainty.

Parameter	Formula	
Variance	$S^{2} = \frac{1}{n-1} \sum_{i=1}^{n} (X_{i} - \overline{X})^{2}$	
Standard deviation	$S = \sqrt{\frac{1}{n-1} \sum_{i=1}^{n} (X_i - \bar{X})^2}$	
Standard error	$S_{\overline{x}} = rac{s}{\sqrt{n}}$	
Confidence		
interval	$\overline{X}\pm t_{lpha/2}$	
	Rectangular	<i>a</i> /√3
Probability	Triangular	a/\6
distributions	Normal distribution	$a/\sqrt{2}$
		S

 Table 2.6: Parameters for expressing uncertainty

2.5.6.2 Top Down and Bottom up approach

Ramsey (1998) recognized two broad approaches for quantifying measurement uncertainty, which are bottom up and top down approach. In the bottom up approach, each individual source of uncertainty is identified and quantified separately as standard uncertainty (its standard deviation). The overall uncertainty is found by combining the individual standard uncertainty values for each contributing uncertainty component. An example is the Particulate sampling theory by Gy (1992). The bottom up approach uses already established methodology but requires that each source of uncertainty be identified which may be difficult to achieve. Gy's method is discussed in detail in Section 2.5.6.3.

The top down approach estimates uncertainty from replicate samples or results obtained from inter-laboratory trial. (Ramsey and Argyraki, 1997). The scatters of these measurements are then used to estimate the uncertainty. Example is the empirical method that uses robust analysis of variance (ANOVA) to estimate uncertainty. Using this method, the sources of uncertainty are grouped into two main classes, sampling and analytical uncertainty. One of its weaknesses is that it does not identify all the components contributing to uncertainty. The technique is further discussed in section 2.5.6.5.

2.5.6.3 Particulate Sampling Theory, by Gy

Uncertainty from both sampling and analysis can be quantified using Gy's sampling theory model. The theory is based upon the relationship that exists between the variability of the material, the particle sizes in the material, the distribution of the component of interest, and the size of sample taken (Mason, 1992).

According to Gy (1992);

"A sample is correct when all particles in a randomly chosen sampling unit have the same probability of being selected for inclusion in the sample."

Any deviations from correct sampling will result in sampling bias. The theory classifies error into 10 classes (Figure 2-8). Some of these errors have a variance that can be estimated while others can give an idea whether the bias is positive or negative (Gustavsson, Lagerkvist, and Luthbom, 2005). Most sampling errors, except the preparation errors, are due to the material heterogeneity.

• Global Estimation Error (GEE)

The Global Estimation Error (GEE) is the sum of the total sampling error (TSE) and the total analytical error (TAE) (Minkkinen, 2004). It can be expressed as follows:

$$GEE = TSE + TAE \qquad 2.15$$

• Analytical Error (AE)

Analytical error is error resulting from the analytical technique used. These errors include instrumentation error, calibration error, operator errors, poor detection limits, analytical bias, and any other error linked to the analytical process. Unlike sampling errors, analytical errors are usually well-understood and well-controlled by laboratory quality assurance and quality control procedures. Its contribution to GEE is usually small.

• Total sampling error (TSE)

It is a combination of Preparation Errors (PE) and sampling or selection error (SE) (Mason, 1992). The selection or sampling error (SE) comprise of fundamental error, preparation error, increment extraction and delimitation error, periodic heterogeneity error, long range heterogeneity error and grouping and segregation error as shown in Figure 2-8.

This error can be expressed as follows:

$$TAE = PE + \sum (FE + GE + DE + EE + CE_2 + CE_3)$$
 2.16

Where

PE = Preparation error, FE = Fundamental Error, GE = Grouping and Segregation error, DE = Increment Delimitation Error, EE= Increment Extraction Error, CE₂ = Long Range Heterogeneity Fluctuation Error, CE₃= Periodic Heterogeneity Fluctuation error

• Fundamental Error (FE)

Fundamental error is the main source of sampling error. The main cause of fundamental error is constitutional or compositional heterogeneity of the material being sampled. Sample may occur in terms of the particle size, particle weight, or particle shape. Fundamental error always exists. It is the only error that can be estimated before hand, i.e. prior to performing sampling. Fundamental errors can be reduced by reducing the particle size of the material being sampled. The relationship between FE, mass, and particle size can be expressed as follows (Ramsey, 1998):

$$S^2 = \frac{Cd^3}{m}$$

Where S is the standard deviation of the sampling error, m is the sample mass; d is the size of the largest particles in the sample and C, the constant, is a product of a number of factors.



Figure 2-8: Classification of errors (Gy, 1992)

It can be expressed as follows (EURACHEM/ EUROLAB/CITAC/ Nordtest, 2006):

$$C = f g \beta c \qquad 2.17$$

Where f is the shape factor,

g is the size distribution factor (g =0.25 for wide size distribution and g =1 for uniform particle sizes, β is the liberation factor for materials where the liberated particles are completely liberated, β = 1.

For unliberated material an empirical equation, $\beta = (L/d)^x$

L = Diameter of the congregate

D = Single mineral diameter (librated single mineral)

Where values of x ranging from 0.5 to 1.5 have been suggested and c is the constitution factor and can be estimated if the necessary material properties are available by using (Minkkinen, 2004):

$$c = \frac{\frac{\left(1 - \frac{a_L}{\alpha}\right)}{a_L}}{\alpha} \rho_c + \left(1 - \frac{a_L}{\alpha}\right) \rho_m \qquad 2.18$$

Where

- a_L is the average concentration of the lot
- α *the* concentration of the analyte in the critical particles,
- ρ_{c} *the* density of the critical particles
- ρ_m the density of the matrix.

The parameters are illustrated in Figure 2-9. From the equation it can be seen that doubling the sample mass will reduce variance by a factor of two and uncertainty by a factor of $\sqrt{2}$.





$$C = fg\beta c = f^{*}(1 \text{ to } 0.25)^{*}(1 \text{ to } L/D)^{(0.5 \text{ to } 1.5)^{*}}c$$

where *f* is the shape factor, *g* is the size distribution factor (g = 0.25 for wide size distribution and g = 1 for uniform particle sizes, β is the liberation factor for materials where the liberated particles are completely liberated, $\beta = 1.0$ for unliberated material an empirical equation, $\beta = (L/d)^x$ where values of *x* ranging from 0.5 to 1.5 have been suggested and *c* is the constitution factor and can be estimated if the necessary material properties are available by using Equation 2.19 above.

• Grouping and Segregation Error (GE)

This error results from distribution heterogeneity that occurs within the sample material. It results from non random segregation or grouping of sample elements. This is often attributed, to the effects of gravity which causes for instance, high density particles with high levels of analyte often drop to lower levels of the sample, causing subsamples from the top to be biased low and subsamples from the bottom to be biased high (Gerlach, *et.al*, 2004). The relationship between S² (FE) and S² (GSE) when taking N increments is expressed by (Gustavsson, Lagerkvist, and Luthbom, 2005).

¹ Figure adapted from EURACHEM/ EUROLAB/CITAC/ Nordtest Guide, 2006.

$$S^2(GSE) \approx \frac{S^2(FE)}{N}$$
 2.19

S² (GSE) – Variance of the grouping and segregation error

 S^2 (FE) = Variance of the Fundamental Error

N = number of increments

Grouping and segregation errors can be minimized by collecting as many small increments as possible or by homogenization of the sample (USEPA, 1999).

• Preparation error (PE)

These result from:

- Contamination due to improper procedures or contaminated equipment;
- Losses, e.g. fine particles are lost due to dusting or may stick to the sampling equipment due to electrostatic forces;
- A change in the physical, chemical or mineralogical composition of the sample;
- Human error, e.g. mixing up sample labels; and
- Sabotage and fraud-although rare, possibility of it exist (Back, 2001).

These errors can be eliminated by using correct sampling practices and equipment, as well as operator training.

• Increment extraction error (IXE)

This error is occurs due to incorrect extraction of an increment. The extraction is said to be correct if the particles with their centre of mass inside the defined sample volume get into the sample. Thus a correct sampling tool should include materials that should be part of the sample and exclude those that should not be part of the sample (Mason, 1992). This error can be reduced by using the correct equipment.

• Increment delimitation error (IDE)

This error occurs when the sampler or sampling device delimits portions of material to be sampled. This results in a non uniform probability of each particle to be collected.

The expected bias from delimitation error is given in Equation 2.20 (Gustavsson, Luthbom and Lagerkvist, 2005):

$$E[B] = E[C_s - E[C_c]]$$
 2.20

Where E [B] is the expected bias, E $[C_S]$ expected mean concentration in extracted sample and E $[C_C]$ is the expected mean concentration in the sample. The error can be eliminated by using a proper sampling device.

• Periodic Heterogeneity Fluctuation Error (CE₃)

It is a non random and cyclic error resulting from the fluctuations in the material to be sampled. These periodic fluctuations can be both spatial and temporal. For example variations in rainfall patterns from one season to another may result in cyclic pattern of pollutant concentrations found in the vadose zone (Back, 2001). This error can be minimized by compositing samples correctly (USEPA, 1999)

• Long - range Heterogeneity Fluctuation Error (CE₂)

This error is generated by the heterogeneity resulting from local trends (Mason, 1992). The error is nonrandom. Variance of this error can be quantified using variograms (Back, 2001). This error can be reduced by taking many increments to form the sample.

Advantages of the approach are summarised as follows:

- Largest source of uncertainty can be easily identified.
- It gives a transparent method showing which components of uncertainty have been considered in the summation of uncertainty. The theory is applicable to the sampling of particulate materials and fluids (Borgman *et.al.*, 1996). The theory is generally comprehensive in that it takes into account all sources of error resulting from the process of sampling and analyses. Other theories only cover a small part of the sampling problem but Gy's theory is the only theory of sampling of particulate material that is accepted and undisputed world-wide (Pitard, 1993).

• where prior information is available, the approaches can be less costly than extensive experimental studies (EURACHEM /EUROLAB/ CITAC/ Nordtest Guide, 2006)

Limitations of the approach are summarised as follows:

- The method is too theoretical and therefore difficult to implement. May require detailed prior measurements of the mineralogy, grain size and analyte speciation of the material to be samples (e.g. soil) and how these vary across the target.
- The theory is not appropriate where the sampling target consist of volatile or semi volatile chemicals.
- The estimates for C (constant) are difficult to determine.
- Gy has presented his work in a number of French and English publications but the high complexity has restricted the use of his sampling theory by engineers and scientists. In cases where Gy's published works are encountered, they are difficult to understand even if one is able to translate from French into English.

2.5.6.4 Quantifying Uncertainty in Analytical Measurements (QUAM) by EURACHEM

guide offers guidance for the evaluation of measurement uncertainty particularly arising from chemical analysis. The guide operates within the ISO framework based on "Guide to the expression of Uncertainty in Measurement" (GUM). According to the guide, the process of estimating uncertainty for a given measurement result can be broken down into four steps which are:

Step 1: Specification of the measurand;

Step 2: Identify the uncertainty sources;

Step 3: Quantify the uncertainty components; and

Step 4: Calculate the combined uncertainty.

The steps are summarised in Figure 2-11.

Step 1: Specify the measurand.

Measurand is defined by ISO (1993) as the particular quantity subject to measurement. This step requires the analyst to clearly give a detailed statement regarding the measurand including

its relationship to the input quantities upon which it depends. That is, provide a quantitative expression relating the value of the measurand to the parameters on which it depends. These parameters may be other measurands, quantities which are not directly measured, or constants. Basically the required level of detail for the specification of the measurand depends on a number of factors including the required level of accuracy of the measurement and the available information (Adams, 2002)

Step 2: Identify uncertainty sources

Following the definition of the measurand is identification of uncertainty sources. This step requires that a comprehensive list of all possible sources of uncertainty be compiled. In coming up with the list it is crucial to start with the basic expression used to calculate the measurand. A cause and effect diagram² may the used as to avoid double counting of sources. Figure 2-10 illustrates a cause and effect diagram. Possible sources of uncertainty that may result during sampling and analyses are listed in Figure 2-12.



Figure 2-10: Field Sample, uncertainty Sources (Ingersoll, 2001)

Step 3: Quantifying uncertainty

After sources of uncertainty are identified, an estimate of the likely size of these sources is made. Prior to quantifying the various sources it important to first reconcile the information

² Cause and effect diagram is a graphical representation of the component uncertainty effects that cause measurement uncertainty.(Ingersoll, 2001)

requirements with the available data. If there is not enough data, a plan should be made to obtain the further data requirements

• Uncertainty arising from the various sources can be quantified by determining the combined contribution to the uncertainty on the results from some or all of these sources using method performance data. Performance data is data derived from method validation studies of the used analysis method.

OR

• Evaluating the uncertainty arising from each individual source and then combining them using the laws of uncertainty propagation.

1. Uncertainty estimation based on performance data

The guide presents several ways of estimating uncertainty based on performance data. Only one of these methods will be considered, uncertainty estimation using in house development and validation studies.

2. Uncertainty estimation using in house development and validation studies

Method validation parameters such as precision, bias, linearity, detection limit, robustness and specificity can be used as means of estimating measurement uncertainty. According to the Ellison, Rosslein, and William, (2000) uncertainty estimation from these parameters uses the best available estimate precision which can be obtained from calculating the standard deviation of an overall bias and its uncertainty. A good estimate of precision can be obtained from taking the standard deviation value of replicate analyses performed on the study of several samples. Bias can be estimated by repeated analysis of a relevant CRM.



Figure 2-11: Uncertainty estimation process (Ellison, Rosslein and William, 2000)





3. Quantification of individual components.

When method performance data is unavailable it is recommended that uncertainty sources be evaluated individually. The components can be derived from:

Experimental variation of input variables – the uncertainty associated with variables often arises from random effects which cause variation of results when the experiment is repeated. In this case uncertainty is quantified in terms of the standard deviation of the values.

From standing data – standing data such as calibration certificates often provide information regarding uncertainty.

By modeling from theoretical principles – physical theories which provide good models for effects on result exist. One well known example is the effect of temperature on volume. By establishing the relationship between the two, uncertainties can be calculated (Ellison, Rosslein, and William, 2000).

Using judgment - Not all sources of uncertainty make a significant contribution to measurement uncertainty. Prior to combining uncertainty, the contributing uncertainties must be expressed as standard uncertainties (the standard deviations).

Each source of uncertainty is converted to standard deviation. When an uncertainty component is expressed as a standard deviation it is referred to as standard uncertainty. (ISO, 1993) Rules for converting uncertainty components to standard deviation:

- Where uncertainty components were evaluated experimentally from the dispersion of repeated measurements, the standard deviation of the results is the standard uncertainty
- In the case of contribution to uncertainty in single measurements, the standard uncertainty is the standard deviation.
- For results subjected to averaging, the standard uncertainty is the standard error of the mean.

Step 4. Combining standard uncertainty

There are two main rules that can be followed when combining uncertainty. These rules are based on laws of error propagation (Taylor, 1996).

Rule 1

For models involving a sum or difference of quantities, combined standard uncertainty u_c(y) is:

$$u_c(y(p,q....)) = \sqrt{u(p)^2} + u(q)^2 + \cdots$$
 2.21

Rule 2

Models that involve a product or quotient, combined uncertainty is given by

$$u_c(y) = y \sqrt{\left(\frac{u(p)}{p}\right)^2 + \left(\frac{u(q)}{q}\right)^2 + \dots} 2.22$$

Calculating Expanded uncertainty

The combined uncertainty is multiplied by the coverage factor (k) in order to obtain the expanded uncertainty. Expanded uncertainty provides an interval about the result of a measurement that may be expected to encompass a large fraction of the distribution of values that could be reasonably be attributed to the measurand (ISO, 1993). It can be expressed as:

$$U = k * u_c \qquad 2.23$$

Where U is the expanded uncertainty and k is the coverage factor.

In choosing the coverage factor (k), there are a number of aspects that have to be considered:

- The level of confidence required;
- Knowledge of the underlying distribution; and
- Knowledge of the number of values used to estimate random effects.

Reporting uncertainty

Information required when reporting uncertainty includes:

- Method used to calculate the measurand and the standard uncertainty
- The values and all sources of all corrections for both the calculation and uncertainty analysis.
- List of all the components of uncertainty with full documentation on how each was evaluated.

The procedure below is used for reporting combined standard uncertainty

Results: x (units) e.g. Total lead: 4.37% w/w

Standard uncertainty: 0.005% w/w

Expanded uncertainty is reported as

Result: $(x \pm U)$ units e.g. total lead: $(4.37\pm0.05)\%$ w/w

Advantages of the method can be summarized as follows:

- The method is based on existing techniques which makes it easy to apply.
- It also allows the analyst to see which component is contributing more to uncertainty hence the analyst can see where an adjustment in the sampling or analyses process is required so as to minimize uncertainty.

Limitations of the method can be summarised as follows:

- The definition of the element that is being measured is usually problematic. This is the case with geochemical sampling and analyses where a number of measurements and procedures are carried out.
- Some components of uncertainty are not readily quantifiable.
- More emphasis is place on the contribution from analyses toward measurement uncertainty. As illustrated in Figure 2-1, the measurement process begins by sampling and hence it should also be taken into account when quantifying measurement uncertainty.

2.5.6.5 Analysis of Variance (ANOVA) by Ramsey

The classical statistic method ANOVA (analysis of variance) can be used to disaggregate the sampling uncertainty, analytical uncertainty and geochemical variability. The ANOVA, approach also known as the Empirical approach for quantifying uncertainty is based on the broad class that is top down approach (EURACHEM/ EUROLAB/CITAC/ Nordtest, 2006). Unlike Pierre Gy's Sampling theory, one does not have to identify various sources of uncertainty individually. The uncertainty sources are classed into two broad classes, those

associated with random components, and those associated with the systematic component of uncertainty. These are further subdivided depending on whether they arise from sampling or analysis process.

For one to be able to estimate uncertainty, the analyst has to estimate the sizes of errors (random and systematic) caused by sampling and analysis first. Random errors are expressed as the precision of the method using the standard deviation in units of concentration, or as precision relative to the mean value as a percentage at 95 % confidence. Systematic error is expressed as the bias of the method and is the difference between the mean of a number of measurements by a method and the certified value³ obtained from certified reference material. The bias is expressed in the units of concentration, or again relative to the mean as a percentage.

Sources of uncertainty

There are four main sources of uncertainty. These sources are summarised in Table 2.7.

 Table 2.7 Uncertainty contributions in the empirical approach (EURACHEM/

EUROLAB/ CITAC/ Nordtest, 2	2006)	
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Process	Random (Precision)	Systematic (Bias)			
	Analytical variability (combined	Analytical bias(combined effect of			
Analysis	contribution of random effect)	bias sources)			
	Sampling variability (dominated				
	by heterogeneity and operator	Sampling bias (combined effect of			
Sampling	variations)	selection bias, operator bias etc.)			

Statistical models for estimating uncertainty

In order to design experimental methods to estimate uncertainty using this approach, a statistical model describing the relationship between the measured and true values of analyte concentration is required. If the sources of variation are independent, the measurement variance σ^2_{meas} is given by:

³ This value is used since the true concentration is unknown.

$\sigma_{measurement}^2 = \sigma_{sampling}^2 + \sigma_{analytical}^2$ 2.24

Where $\sigma^2_{\text{sampling}}$ is the between-sample variance on one target (largely due to analyte heterogeneity) and $\sigma^2_{\text{analytical}}$ is the between-analysis variance on one sample.

If statistical estimates of variance (s^2) are used to approximate these parameters, we get

$$s_{measurement}^2 = s_{sampling}^2 + s_{analytical}^2$$
 2.25

The standard uncertainty (u) can be estimated using $s_{measurement}$, which is therefore given by

$$s_{measurement} = \sqrt{s_{sampling}^2 + s_{analytical}^2}$$
 2.26

Estimating uncertainty

According to Ramsey (1998), there are four methods that can be used to estimate uncertainty. These vary in terms of the number of samplers⁴ and the number of protocols. These are:

(a) Using single sampler and single protocol;

(b) Using single sampler and multiple protocols;

(c) Using multiple samplers and single protocol; and

(d) Using multiple samplers and multiple protocols.

These can be summarised in Table 2.8 below: In this document focus will be given to the first method.

Method 1: Single sampler and single protocols

The basis of the method is that the sampling precision and analytical precision can be estimated by taking duplicate samples for some proportion of the sample increments. The duplicate samples are not taken at exactly the same place, but separated by a distance that reflects the separation that might have occurred by a totally independent interpretation of the

⁴ Sampler is the sampling personnel ,that is, a person conducting the sampling

sampling protocol. Duplicate chemical analyses are then made on these duplicate samples as in Figure 2-13.

Method	Method Description	Samplers	Protocols	Component estimated			
number		(persons)		P _{samp}	B _{samp}	Panal	B _{analy}
1	Duplicates	Single	Single	Yes	No	Yes	No
2	Protocols	Single	Multiple	Between protocols		Yes	No
3	CTS	Multiple	Single	Between samplers		Yes	Yes
4	SPT	Multiple	Multiple	Between protocols Y -between samplers		Yes	Yes

Table 2.8: Four empirical methods for estimating combined uncertainty (Ramsey, 1998)

Four methods for estimating uncertainty in measurements ($P_{anal} =$ precision of analytical method, $B_{samp} =$ bias of sampling method, CTS = Collaborative Trial in Sampling, and SPT = Sampling Proficiency Test).



Figure 2-13: Balanced design

Calculating uncertainty and its components

Analysis of variance ANOVA is used to estimate the sampling and analytical precision. Analysis of variance (ANOVA) is a statistical technique for testing differences among group means. The purpose of analysis of variance model ANOVA is to identify the sources of variation and construct the proper tests to compare them. Estimation of the uncertainty derived from the systematic components of both sampling and analysis process, analytical and sampling bias are done separately.

Classical ANOVA method is sensitive to outliers in the data set (Back, 2001) hence Robust ANOVA method is used to calculate uncertainty from random component. Robust ANOVA allows the separation of sampling and analytical variances from the total variance. It is not affected by outlier values which makes it provide reliable results as compared to classical ANOVA method (EURACHEM/ EUROLAB/CITAC/ Nordtest, 2006).

Robust ANOVA is implemented using ROBOCOOP4.EXE program which is a compiled FORTRAN program. ANOVA is not only restricted to this program, there are other statistical packages that are available

Output

The output from the ROBOCOOP.EXE4 program for ANOVA consists of three components of variability for robust analysis of variance. The first 5 lines of results are for classic ANOVA and the last five are for robust ANOVA as shown in Table 2.9. From the table showing the output, it can be seen that 4 parameters have been calculated for robust ANOVA. These are mean, standard deviation, percent variance, and sum of squares.
Table 2.9: Output of ROBCOOP4.EXE	program	(Ramsey,	1998)
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CLASSICAL ANOVA RESULTS						
Mean = 317.79999						
Standard Deviation	(Total) = 240.1923	38				
	Between-target	<u>Sampling</u>	Analys	is		
Sums of Squares	173 8031.9	370075.5	6473.0			
Standard Deviation	197.5	135.4	18.0			
Percentage Variance	67.6	31.8	0.6			
ROBUST ANOVA	RESULTS:					
Mean = 297.30884						
Standard Deviation	(Total) = 218.4876	63				
	Between-target	<u>Sampling</u>	Analysis	Measurement		
Standard Deviation	179.674	123.814	11.144	124.314		
Percentage Variance	67.626	32.114	0.260	32.373		
Relative Uncertainty	-	83.290	7.497	83.626		
(% at 95% confidence	e)					

The robust ANOVA results are used for the calculation of uncertainty. Classical ANOVA results are available for comparison purposes.

Interpretation of the results

- The mean value is the mean of the input data in the same units as input data.
- The sigma value represents estimates of standard deviation for three sources of variance.
- The percent variance expresses each of the three variances as a percentage of the total variance.

The sampling standard deviation $S_{\text{sampling}} = 123.8$ and analysis standard deviation $S_{\text{analysis}} = 11.14$.

The combined uncertainty u is a function of two independent variances and can be expressed as:

$$u_{measurement} = \sqrt{s_{sampling}^2 + s_{analytical}^2} = 124.3$$

Note: the results inside the output of ROBOCOOP 4 above are not linked to any calculations Its expanded uncertainty is:

$$\boldsymbol{U} = \boldsymbol{u} \times \boldsymbol{k} \qquad \qquad 2.27$$

Where k is the coverage factor and the value is 1.96 which gives a 95% confidence interval for the uncertainty.

"The calculated value of the uncertainty applies to measurements made on single samples taken in the survey. If a number samples are taken at any individual location within the site, the uncertainty on the average for that location is the value given by calculating the relative uncertainty divided by \sqrt{n} . This is equal to the standard error on the mean value (s/ \sqrt{n})", Equation 2.5 (Ramsey, 1998, pp 101).

Estimation of sampling and analytical systematic error (bias)

- Estimation of analytical bias can be done through using repeated measurements of the certified reference material (CRM). If the systematic error found using this procedure is insignificant, the uncertainty associated with the systematic error is simply the combination of the standard uncertainty on the CRM value with the standard deviation associated with the systematic error.
- Estimation of sampling systematic error (sampling bias):

(i) Can be done using sampling proficiency test. The procedure enables laboratories to monitor their measurement capability both by comparison to peer laboratories, and over time within their own laboratory; and

(ii) Can be done by collecting paired samples for various targets. After chemical analysis of the two samples, the difference between a pair of results is an estimate of the bias for that target. Repeating the procedure for many distinct targets improves the precision of the estimate.

There are no hard and fast rules concerning how the bias estimates can be incorporated into the uncertainty measurements. The estimate for bias can be expressed as standard uncertainty and then incorporated into the measurement uncertainty.

Limitations of the technique can be summarised as follows:

- The method only estimates the uncertainty resulting from random components and does not include those resulting from systematic errors.
- If fewer duplication samples are used there will be large uncertainties on the estimates of variances.
- If large quantities of dataset are involved the process may be tedious and time consuming.
- The method's approach to sampling bias excludes bias introduced by the sampling equipment and bias inherent in the sample according to Pierre Gy's sampling theory (Back, 2001).

2.6 SUMMARY

Reporting uncertainty of a measurement result provides the means for assessing the reliability of a measurement result. Measurement uncertainty prescribes a range in which the true value of the measurand is expected to lie. As results from sampling and analysis play an important role in decision making, it is crucial that uncertainty in sampling and analyses be evaluated and be part of result reporting. Unreliable results lead to poor decisions. Every procedure undertaken from sampling, sample collection and preparation, analysis and the interpretation of the results should be recognized as potential sources of uncertainty.

As indicated above, three methods of quantifying measurement uncertainty were reviewed in detail. Although the methods vary, one assumption common to all the models is that, measurement uncertainty can be defined statistically and expressions are derived for estimating and combining uncertainties from the various uncertainty components. The commonly used parameter is the standard deviation which defines how values spread.

Both Particulate Sampling Theory and EURACHEM methods are based on the bottom up approach in which one has to identify uncertainty sources, quantify them individually and combine them using some statistical model to obtain the overall uncertainty. This concept is contrary to ANOVA model as suggested by Ramsey (1998) in which the overall uncertainty is determined without necessarily identifying the individual uncertainty components.

The method proposed by EURACHEM provides easy to follow steps for quantifying uncertainty but unlike the Particulate Sampling Theory which is comprehensive, the method emphasizes the effect of analyses. As seen above, the geochemical measurement process is multi-tier process including sampling, sample preparation and analyses. These factors should be considered as potential uncertainty sources.

ANOVA on the other hand places more emphasis on the uncertainty components associated with random error and does not give a clear explanation on how uncertainty associated with systematic error is determined. Due to this lack of clarity regarding quantification of effects associated with systematic error, it is mostly likely that systematic effects are neglected hence an underestimate uncertainty value is provided. In addition, the fact that samples have to be accompanied by duplicates make the method expensive to implement especially taking into account that most project normally conducted with limited financial resource.

The Particulate Sampling Theory as well as ANOVA has been applied in the field of environmental geochemistry, for example in estimating concentration of Cadmium and Phosphorous in topsoil (EURACHEM/EUROLAB/CITAC/Nordtest Guide, 2006 pp.34-83). One disadvantage associated with ANOVA is that although the model is simple to use, it requires independent estimates of uncertainty contribution from sampling and analysis bias. Some components of the Particulate Sampling Theory for example estimates of the four factors within Gy's constant C may be difficult to determine with any degree of certainty.

From the reviewed methods it can be seen that there was a definite need for a more user friendly method which can be applied in geochemical studies as stated in the objectives of this study. A new methodology for quantifying uncertainty in geochemical sampling and analyses was developed and it can be summarised as follows:

• Defining the purpose of the investigation;

- Defining of the sampling target;
- Specification of the measurand;
- Description of the sampling and analysis protocol;
- Identification of the source of uncertainty;
- Quantification the sources of uncertainty;
- Calculating the combined implications of the uncertainty components; and
- Calculating the expanded uncertainty

This new method akin to the previously discussed methods is based on the application of statistics in quantifying uncertainty in geochemical sampling and analysis. The method uses basic statistics parameters that are easy to understand. It is based on simple and straight forward steps that can be easily followed. Where complex sampling or analysing procedure has been used, the method is easily altered to suit the need at hand. The method is further discussed in Chapter 3.

CHAPTER 3: METHODOLOGY

3.1 DETERMINATION OF SAMPLE SIZE

Sample determination is one of the most crucial steps in the geochemical investigation process. Studies of geological materials have revealed that variability exists in terms of mineralization, chemical composition and alteration. Under such conditions, it is essential that an adequate sample size that represents the population or area it was drawn from be collected.

This section provides a detailed discussion of sample size determination using statistical analysis approach. In order to successfully determine optimum sample size needed to satisfy the objectives of a study, several factors need to be considered. These include: the purpose of the investigation, type of the material, cost of sampling, variability of the material and available site information. In addition to these are the required level of acceptable error, confidence interval and standard deviation. These factors and their effects are explored in detail in the following sections.

3.1.1 OVERVIEW OF THE STATISTICAL METHOD

Having a population with a mean " μ " and a standard deviation " σ .", if a sample size n with the values (x₁, x₂,...x_n) is drawn from this population, the sample mean " \overline{x} " is given by :

$$\bar{x} = \frac{\sum_{i=1}^{n} X_i}{n} \qquad 2.2$$

Where \overline{x} (sample mean) is assumed to present the best estimate of μ , and therefore $\mu = \overline{x}$. The difference between μ and \overline{x} is estimation error (d). With given confidence level, the sample mean estimate and estimation error are often reported as follows: $\overline{x} \pm \mathbf{d}$

The estimation error (d) equals:

$$d = T_{\alpha/2} \times \frac{\sigma}{\sqrt{n}}$$
 3.1

T α represents the function of T distribution, which is only related to sample numbers and confidence levels (Table 3.1).

Sample	Confidence levels				
numbers	80%	90%	95%	99%	
5	1.533	2.132	2.776	4.604	
10	1.383	1.833	2.262	3.250	
25	1.318	1.711	2.064	2.797	
30	1.311	1.699	2.045	2.756	
40	1.303	1.684	2.021	2.704	
60	1.296	1.671	2.00	2.660	
120	1.289	1.658	1.980	2.617	
>120	1.282	1.645	1.96	2.576	

 Table 3.1: T values for different sample numbers and confidence levels

Based on the Central Limit Theorem, as long as the sample numbers are reasonably large (usually greater than 30), the sample mean will tend to be normally distributed. Therefore, the estimation error (d) can be expressed as follows:

$$d = Z_{\alpha/2} \times \frac{\sigma}{\sqrt{n}}$$
 3.2

Where

 $Z_{\alpha/2}$ represents the function of normal distribution which related to confidence levels, Table 3.2.

 Table 3.2: Z values for different confidence levels

Confidence levels	75%	85%	95%	99%
Z values	1.15	1.44	1.96	2.57

From the Equation 3.2, it can be seen that for given confidence level, the estimation error is dictated by the standard deviation of the population and the sample numbers. On the other hand, the wider the population spreads, the larger the estimation error is. The bigger the sample size, the smaller the estimation error is. For a given confidence level, the estimation error is determined by the sample size. If one wants to reduce the error, sample size should be increased. However, increasing sample numbers means spending more money. So a balance among competing factors has to be sought.

Rearrange the Equation (3.2), we get following formula:

$$n = \left(\frac{Z_{\alpha/2} \times \sigma}{d}\right)^2 \qquad \qquad 3.3$$

Where

n= sample size, $Z_{\alpha/2}$ is the is known as the critical value the positive Z value that is at the vertical boundary for the area $\alpha/2$ of in the right tail of the standard normal distribution, σ is the population standard deviation, d= the desired level of precision.

This is the basic model for determination of sample size. The formula may be used when the population standard deviation σ is known but most often this value is unknown. Information about σ can be obtained from past experience with the same or similar problem, or can be obtained from pilot study. As a general rule of thumb, if the sample size *n* is greater than 30, σ is replaced by the sample standard deviation *s* and Equation 3.3 becomes (**Reytblat**, 2000-2010):

$$n = \left(\frac{Z_{\alpha/2} \times s}{d}\right)^2 \qquad \qquad 3.4$$

The formula is more convenient way for estimating the sample size for continuous data.

3.1.2 PARAMETERS ON WHICH SAMPLE SIZE DEPEND

In order to illustrate the factors that influence sample size determination, pH results for West Wits tailings dams based on study conducted by Pulles et.al (2003), picked at random were used as shown in Table 3.3.

nII	nII	nTT.	nII	nTT
рН 0.00	рН 8 90	рн 8 10	рН 7.60	рН 8 20
9.00	8.80	8.10 8.40	/.0U 8.10	8.20 8.10
9.00	8.30	8.40	8.10	8.10
8.10	8.40	8.70	8.50	8.20
8.30	8.30	8.60	8.20	8.30
8.20	8.30	8.50	8.20	8.10
8.30	7.90	7.60	8.40	8.50
8.30	8.40	7.50	8.00	8.40
8.30	8.40	8.40	8.50	8.60
7.80	8.30	8.20	7.70	8.90
8.20	8.30	8.10	8.50	8.70
8.20	8.20	8.60	8.80	8.40
8.20	8.40	8.50	8.00	8.50
8.10	8.30	8.50	8.60	8.70
8.00	8.30	8.40	8.70	8.60
8.20	8.30	8.40	8.60	9.00
8.30	8.20	8.20	8.70	8.60
7.50	8.50	7.80	8.40	8.80
7.70	8.50	8.20	8.20	8.60
8.20	8.40	8.30	8.30	8.60
8.40	8.40	7.50	8.40	8.60
8.20	8.30	7.90	8.50	8.10
8.00	8.50	7.90	8.30	9.10
7.60	8.70	8.20	8.50	8.50
8.10	8.50	8.10	8.50	8.50
8.60	8.50	8.30	8.60	8.40
8.70	8.60	8.30	8.50	8.70
8.40	8.40	8.30	8.20	8.80
8.20	8.50	8.30	8.10	8.40
8.50	8.50	8.40	8.20	8.40
8.60	8.60	8.40	8.20	8.70
8.30	8.40	8.40	8.30	8.50
8.20	8.50	8.20	8.00	8.40
8.90	8.50	8.10	8.30	8.30
8.70	8.40	8.30	8.60	8.10
8.50	8.10	8.70	8.60	8.80
8.80	8.40	8.90	8.30	8.50
8.30	8.50	7.20	7.80	8.70
8.50	8.60	8.60	7.70	8.50
8.00	8.00	7.90	8.20	8.20
8.60	8.30	8.20	8.60	8.40
Parameter	Sample Size	Mean	Variance	Standard deviation
Value	200	8.35	0.09	0.30

Table 3.3: pH data (Pulles et. al., 2002)

3.1.2.1 Confidence level

Confidence level as discussed in Chapter 2 is the confidence percentage that specifies how confident one is, that the parameter (mean) lies in the specified interval or range. Based on

Equation 3.4, an increase in confidence level means more samples will be required. This relationship can be illustrated using the dataset from Table 3.3.

The mean (\bar{x}) of the data set is 8.35. This is the estimate of population mean (true mean). If we want to know with 75% confidence level and standard error of 0.02 that 75 out of 100 samples are within the true population mean of 8.35 ± 0.02, sample size (n) required is:

$$n = \left(\frac{1.152 \times 0.30}{0.02}\right)^2 \approx 299$$

Therefore the required sample size is 298.

BUT

If the CI is increased to 95%, the required samples are:

$$n = \left(\frac{1.962 \times 0.30}{0.02}\right)^2 \approx 866$$

The calculated results show that if we want to increase the confidence level, we have to increase the sample numbers, but the relationship is not linear, doubling the sample size does not halve the confidence interval (Israel, 1992).

Table 3.4 can illustrate the relationship.

Table 3.4: Sample numbers for different confidence levels

Confidence level	75%	80%	85%	90%	95%	99%
Sample Size	299	369	467	613	866	1486

3.1.2.2 Standard deviation or population spread

Standard deviation defines how the population is spread. This parameter, like standard error and confidence level also affects sample size. Basically the wider the population spreads; the larger sample numbers needed. The relationship can be demonstrated by the example below: The sample standard deviation of the dataset in Table 3.3:

Given a 95% confidence and the standard error of 0.02, the sample size for dataset (Table 3.3) can be calculated as follows:

$$n = \left(\frac{1.962 \times 0.30}{0.02}\right)^2 \approx 866$$

If the standard deviation is increased to 0.40 with 95% confidence level and standard error of 0.02, the required sample size is:

$$n = \left(\frac{1.962 \times 0.40}{0.02}\right)^2 \approx 1540$$

From the results the spread of the population affect sample size. As the spread increases (standard deviation) more and more samples are required.

3.1.2.3 Standard Error

Equation 4.4 shows that a relationship exists between sample size and the standard error. This relationship is exponential; an increase in sample size is followed by a decrease in standard error. Table 3.5 can illustrate this relationship.

Using the data values in Table 4.3, s = 0.30, C.I = 95% and standard error

a)
$$= 0.02$$

b) $= 0.04$

The required sample size n is:

a)
$$n = \left(\frac{1.962 \times 0.30}{0.02}\right)^2 \approx 866$$

b)
$$n = \left(\frac{1.962 \times 0.30}{0.040}\right)^2 \approx 217$$

Estimation error	0.01	0.02	0.03	0.04	0.05
Sample numbers	3 465	866	385	217	140

Table 3.5: Sample numbers for different estimation error (d)

The results above indicate that a small standard error value is associated with a large sample size. In addition to halve the error, as indicated by the formula, the sample size has to be quadrupled.

3.1.2.4 Cost of taking the samples

Cost helps in determining how precise the estimates should be. If the decision to be made, based on the outcome of sampling activity is valuable, then more samples should be collected; But a large sample size means a higher budget thus any increase in the number of samples is followed by an increase in the total expenditure. Therefore a compromise between competing factors must be reached and one must work out the degree of inaccuracy that they are willing to accept.

3.1.3 PROCEDURE FOR DETERMINING SAMPLE SIZE

3.1.3.1 Specify the purpose

In order to correctly determine sample size, it is essential to clarify the objectives of the investigation. One should ask the question "what do I want to know?" and "what information do I expect to gain?" Only when these questions are addressed can one decide on the number of samples to collect. Study objectives vary and hence sample sizes vary as well.

3.1.3.2 Investigate the parameters

Determination of the population parameters is the next step after definition of the study objectives. In this process, the following questions should be investigated:

- Is the population homogeneous?
- What is its statistical property, mean, standard deviation, median, etc?
- How is its spatial distribution?

The critical parameters such as mean, standard deviation, acceptable error can be acquired from various sources of information such as:

- Previous publications and documents;
- From similar previous studies; and
- Pilot study.

3.1.3.3 Choosing the sampling strategy

Once information has been gathered, the next step is to select a sampling strategy. Care must be taken when choosing the sampling strategy. The analyst should take into account such aspects as spatial continuity, correlation of the geological properties, cost and variability. If the sample material is homogeneous, random sampling strategy may be adopted. If the sample area is heterogeneous, stratified or systematic sampling strategy are preferable. With stratified sampling strategy, the area is divided into several subpopulations and random samples are collected from each stratum and the sample size of each subpopulation is calculated separately. In addition stratified design is more economic in that the number of samples is reduced since the target area is divided into different stratum with each stratum having similar properties. The advantage of systematic grids method over its counterpart is that it takes spatial continuity into account (US EPA, 1999).

3.1.3.4 Determination of the acceptable error

In order to calculate the sample size, we must have the value of the estimation error. The acceptable error is the risk that the analyst or researcher is willing to accept.

The four guidelines can be used to decide the specific value of the error: (Yang, 2006)

- The value should be greater than analysis error;
- The value can be calculated as 5% of the sample mean; and
- The value can be taken as the maximum allowable value for the ERA purpose.

This value can be obtained by referring to previous data or consulting secondary sources of information such as journals. As stated in previous sections, the use of previous data should be done with caution. Only when it can be proved that the project from which previous data is derived was successful can one confidently use the data. It should also be borne in mind that geological situations vary so do geological materials. It is safer to conduct a small prior study and use the obtained estimated to determine acceptable error than to use

data from previous studies.

All in all the key point is that the value of the estimation error should be acceptable or allowable for the ERA purpose.

3.1.3.5 Determine standard deviation

Standard deviation is a critical component of the sample size formula. Although the analyst has less control over variance, it must be incorporated into the design. Methods that can be used to estimate the standard deviation value for sample size determination include use of a pilot study, use of data from previous studies of similar population, estimate assisted by some logical mathematical results (Bartlett, Kotrlik, and Haggins, 2001). Detailed discussion of some of these methods is provided under section 5.6.

3.1.3.6 Specify confidence level

Confidence interval provides a range of values that has a specified probability of containing the parameter being estimated. The frequently used and recommended confidence level is 95% and its alpha value is 0.01 (Yang and Zhao, 2005). The z value associated this confidence level is 1.96. This value can be obtained from standard normal distribution tables. If a 95% confidence level has been chosen it simply means that that 95% of the time, it is believed that the population means lies within this range.

One major factor, affecting the choice of confidence level, is the purpose of the study or investigation. Higher confidence levels for example 99% confidence level are often employed in cases where decisions based on research are critical and errors may cause substantial harm (Bartlett, Kotrlik, and Haggins, 2001). Lower confidence levels for example 75% may be used for instance when identifying trivial differences or other statistical phenomena as an antecedent to further studies.

3.1.3.7 Determine the sample size

When the various parameters necessary to determine sample size have been estimated, the parameters are combined using Equation 3.4.

$$n = \left(\frac{Z_{\alpha/2} \times s}{d}\right)^2 \qquad \qquad 3.4$$

There are two types of sample sizes to be decided on (Yang, 2006):

(1) Minimum sample size: When we use the formula to calculate the sample size we assume that the samples are normally distributed, but based on the Central Limit Theorem the assumption is correct only when the sample numbers are reasonably large. In statistics, 30-50 samples can be taken as reasonably large. Considering the costs of the sampling; 30 is considered as a reasonable minimum sample size.

(2) Maximum sample size: The sample size calculated, based on the formula can be taken as the maximum sample size because it takes all samples as random and independent and give each sample an equal weight.

3.1.3.8 Limitations of the method can be summarised as follows:

- One disadvantage associated with the method is that a "good" estimate of the population standard deviation is necessary. Often this value is unknown.
- The approach is based on the assumption that data is normally distributed.
- The method assumes that a simple random sampling scheme is the method of choice and does not apply when other sampling scheme are schemes of choice.



Figure 3-1: Procedure for determining sample size

3.1.4 SPREAD SHEET FORMAT FOR SAMPLE SIZE DETERMINATION

This section outlines the spreadsheet method of calculating sample size based on the statistical analysis approach. Table 3.6 illustrates the spreadsheet method of sample size. The values and the calculations within the table serve to illustrate only.

	Worksheet for Sample Size Determination						
A	В	С	D	E	F	G	Н
3	Data Carro	C	Samp. STDEV (S)			Conf.Fun. (Ζα/2)
4	Data Source	samp.mean (m)			85%	95%	99%
5	Data A	97.55	26.37		1.44	1.96	2.57
6	Data B	19.10	9.86		1.44	1.96	2.57
7							
8	1	Acceptable Estim	ate Error (r)		Samp	le Size Det	ermination
9	5% of mean	2*(S/√30)	Regulation	Accepted	85%	95%	99%
10	4.88	9.63		4.88	61	112	193
11	0.96	3.60		0.96	221	410	704

Table 3.6: Worksheet for sample size determination

The procedure for determining sample size in spread sheet format is based on the following steps:

Step 1: Calculate the sample mean (m) and the sample standard deviation.

For the sample mean (m), the formula is:

"= AVERAGE (Xij: Ynm)"

Where Xij represents the beginning cell of the data matrix and Ynm the ending cell of the data matrix.

For the sample standard deviation (S), the formula is:

"= STDEV (Xij: Ynm)",

Where Xij represents the beginning cell of the data matrix and Ynm the ending cell of the data matrix.

Step 2: Determine the acceptable estimation error

If the related regulation or protocol presents the guideline for the error, use the guideline; otherwise, choose the lesser between "5% of mean" and " $2xS/\sqrt{30}$ ". The formula is:

"=IF (D11=0, MIN (B11, C11), D11)" (for Data B)

Step 3: Calculate the required sample number

For the confidence level of 85%, the formula is:

"=IF [(D5^2*F5^2/E10^2)>30, D5^2*F5^2/E10^2, 30]" (for Date A) or

"=IF [(D6^2*F6^2/E11^2)>30, D6^2*F6^2/E11^2, 30)]" (for Data B)

For the confidence level of 95%, the formula is:

"=IF [(D5^2*G5^2/E10^2)>30, D5^2*G5^2/E10^2, 30]" (for Date A) or

"=IF [(D6^2*G6^2/E11^2)>30, D6^2*G6^2/E11^2, 30]" (for Data B)

For the confidence level of 95%, the formula is:

"=IF [(D5^2*H5^2/E10^2)>30,D5^2*H5^2/E10^2,30]"(for Date A) or

"=IF [(D6^2*H6^2/E11^2)>30,D6^2*H6^2/E11^2,30]" (for Data B)

3.2 METHODOLOGY FOR QUANTIFYING UNCERTAINTY IN GEOCHEMICAL SAMPLING AND ANALYSES AS A FUNCTION OF SAMPLE SIZE AND ANALYSIS

The results of a measurement are incomplete without an accompanying statement of uncertainty. Measurement uncertainty is one of the main factors that fundamentally impacts data quality and therefore decision-making. In the case of geochemical investigations for ERA decision-making under uncertainty may lead to serious consequences which may result in the loss of public trust and confidence and heavy financial losses.

This section presents a methodology for quantifying uncertainties in geochemical sampling and analysis as a function of sample size and analysis. While many existing methods for estimating uncertainty focus on analysis as the main contributing factor to uncertainty, the method also recognizes the effect and contributions from sampling and sample preparation towards measurement uncertainty. This stems from the fact that the measurement process is a multifaceted process comprising of sampling, sample preparation, and analysis; hence the importance of considering the uncertainty contributions from all procedures involved. The resulting value will be a more realistic estimate of uncertainty, than considering the influence and effect of a single factor.

The methodology for quantifying uncertainty is based on nine steps which are:

- Definition the purpose of the investigation;
- Definition of the sampling target;
- Specification of the measurand;
- Description the sampling and analysis protocol;
- Identification of the source of uncertainty;
- Quantification the sources of uncertainty;
- Calculating the combined implications of the uncertainty components;
- Calculating the expanded uncertainty; and
- Reporting the result.

Various uncertainty components are grouped into two categories, Category A and Category Bbased on the method used to evaluate them. Category A encompasses all sources of uncertainty that can be evaluated using statistical methods while Category B evaluates uncertainty components that cannot be evaluated using statistical methods.

The standard uncertainties are combined to reach a total or overall uncertainty using the laws of error propagation. An expanded uncertainty is calculated to define an interval about the result of a measurement that is expected to encompass a specified fraction of the possible values for the measurand.

3.2.1 THE IMPORTANCE OF QUANTIFYING AND REPORTING UNCERTAINTY

The following points summarise the importance of evaluating and reporting the measurement uncertainty:

- A statement of the uncertainty associated with a result conveys to the customer the 'quality' of the result (CITAC/EURACHEM, 2002);
- Providing the value of uncertainty instills confidence in the result of a measurement and shows the data users that the result can be relied on;
- It leads to better-informed decision-making;
- It allows for the comparison of two measurement results; and
- Information is obtained for improving the method.

3.2.2 PROCEDURES FOR QUANTIFYING UNCERTAINTY

The results of geochemical investigation depend on the collection of good quality and representative samples. However, no matter how careful one is in selecting, preparing and analyzing the sample, the results are always uncertain. In other words, measurement uncertainty is unavoidable. This section provides a detailed discussion of the steps that may be adopted when quantifying uncertainties in geochemical sampling and analyses process. The steps are summarised in Figure 3-2 below.

3.2.2.1 Step 1: Define the objectives of the study

Definition of the study objectives or purpose is the most important and critical step in quantifying uncertainty. It is difficult to arrive at a meaningful estimate of uncertainty without clearly understanding what the objectives of the measurements are. A clear description of the purpose of the investigation provides background on the actual issues addressed by the study. It is therefore crucial that the objectives of the measurement are clearly understood prior to undertaking geochemical measurements for ERA.

An example of sampling objective would be to estimate the mean concentration levels of the element lead elements in water and detecting soil mercury contamination.

3.2.2.2 Step 2: Definition of the sampling target

A sampling target is an area or a portion of material that the sample represents at a particular time. The target may be defined in terms of space (geographic boundaries) or time- that is: when, where and which portion of whole is to be sampled? Thus from a 10 ha of waste dump, one may decide to sample only 3 ha. The 3 ha would be the sampling target.

Questions that should be addressed by the target definition include:

- What type of material is present in the target area?
- Is the material heterogeneous or homogeneous?
- What are its variables? (mean, standard deviation)

Information relating to the sampling target can be obtained from geological reports, historic information and conducting a preliminary study. Whichever method is adopted, the definition should reflect the purpose of study.

3.2.2.3 Step 3: Definition of the measurand

This step requires the analyst to give a clear and detailed statement of the quantity being measured. This quantity may be a directly measured variable (value), or indirectly determined relying on input of other variables to determine the measurand in question. The latter is often associated with a mathematical model which relates the input variables (X_1, X_2, \dots, X_N) to the measurand (output), Y (EAL Task Force, 1999). Most measurements specifically in geochemistry are often associated with the latter in which measurand is determined through the measurement of other variables thus the relationship can be generalized as $Y = f(X_1, X_2, \dots, X_N)$. Such measurements are referred to as multivariate measurements and they call for the development of a mathematical expression which relates the measurand Y to the variables X_1, X_2, \dots, X_N on which it depends. The value of developing such mathematical models lies in their ability to precisely describe how the value of the output quantity depends on the values of the input quantities.



Figure 3-2: Steps in quantifying uncertainty

The step can be summarized as:

- Determine what variables need to be measured to estimate the measurand Y that is the input quantities if the measurand cannot be determined directly.
- Transcribe the actual relationship between the measurand and the input or develop the mathematical model outlining the how the measurand Y relate to the input parameters X₁, X₂.....X_N on which it depends on.

Example 1-measuring rock Acid Potential (AP) of gold waste dump material

The measurand can be defined as assessment of Acid Potential (AP) of gold mine waste material which depends on the input variables: total sulfur content, molecular weight of Calcium Carbonate (CaCO₃) and atomic weight of sulfur.

This relationship between the measurand and the input parameters on which it depends can be expressed as:

$$AP = Total Sulphur (\%) \times 31.25kg CaCO_3 \qquad 3.5$$

3.2.2.4 Step 4: Description of the sampling and analysis protocols followed

This step requires the user to give a detailed outline of the sampling and analytical protocols adopted during the investigation. Sampling and analytical protocols provide details of all the measurement processes that were carried out in order to obtain the measurand. The importance of this step is that the protocol followed would be used as reference especially when determining the source of uncertainty

Sampling protocol

Under sampling protocol, the following aspects should be highlighted:

- The sampling method used and whether it was a stratified or random method of sampling. Any modifications or changes made to the technique should be included.
- The number of samples collected.
- The sampling tool used and spacing or interval.

- Sample labeling and preservation.
- QA/QC used in order to ensure quality.

Many samples get contaminated during this stage hence the manner in which the samples were held should be highlighted.

Analyses protocols

This step encompasses all steps carried out from preparation of the test portion to the analysis itself. These procedures should be stated in detail and the list should include:

- Instrument(s) used;
- Specifications of the analysis methodology;
- Specifications of reagents used;
- How sample(s) were handled;
- Any calculations that were done prior to analysis; and
- QA/QC program adopted to ensure quality.

A write up of the sample preparation procedure should be included in this step.

3.2.2.5 Step 5: Identifying sources of uncertainty

Following the specification of the procedure used, it is necessary to compile a list of the possible sources of uncertainty in the measurement process. Since some sources contribute more to uncertainty than others, it is important to start with major sources, the quantities that have the largest influence on the measurement.

As stated at the beginning of this chapter, most methods of quantifying uncertainty emphasise the contribution of laboratory analysis towards measurement uncertainty but studies by Ramsey (1998) revealed that of all primary procedures undertaken during geochemical investigations, primary sampling contributes more to measurement uncertainty than analysis, Figure 3-3 (Schwedt, 1997). As such, it is crucial that all processes involved in the multi tiered measurement process be considered as potential sources of uncertainty.



Figure 3-3: Degree of error in laboratory sample preparation relative to other activities (US EPA *et.al*, 2004)

Common sources of uncertainty in geochemical investigations

The following sections discuss common sources of uncertainty in geochemical sampling for ERA. It is crucial to begin with the input quantities used in determining the measurand in drawing up the list of the sources of uncertainty (USEPA *et.al*, 2004). Secondary sources of information such as journals, manufacturer's specifications, manuals, and books may be consulted for additional information regarding potential uncertainty sources.

1. Sampling

As stated in the previous section, the sampling process like all processes involved in geochemical measurement contributes towards the uncertainty associated with the value of the measurand. Under this several sources are considered including

(a) Heterogeneity (variability): It is one of the major sources of uncertainty in geochemical sampling and analysis. For most geological materials, for example waste rock dumps, heterogeneity exists in terms of the particle size distribution, mineralisation and alteration and analyte concentrations. The effect of this source of uncertainty can be displayed by variations in terms of the observable analyte concentration among samples collected from different points of the sampling target (the random effect). Most scholars including Pitard (1993) argue that heterogeneity remains the largest uncertainty source

contributing towards the overall uncertainty in geochemical sampling and analyses measurements. This uncertainty source can be minimized by taking many increments or reducing particle size of the sample material.

(b) **Sampling design:** Sampling design introduces uncertainty when a limited number of locations are sampled and the strategy fails to capture the complete extent of the inherent variability which exists within the target.

(c) **Incorrect sampling**: Error is introduced when the size and geometry of the sampling tool used violates the fundamental principle of sampling. According to the principle, a selected sample is structurally accurate when all the units within the lot have equal probability of being selected. Any deviation from the principle would result in bias.

(d) **Sample handling:** The container or equipment used for holding the sample during transport from the field to the laboratory may affect the analyte recovery. Bias is introduced when the walls of the container absorb the analyte of interest or contaminate the sample (or allows the analyte to escape in the case of volatile elements such as mercury and arsenic.

(e) Limited sample mass: The mass of the sample analysed may not be enough to actually capture the actual conditions existing within the sampling target

2. Sample preparation

Sample preparation is another dominant source of uncertainty in geochemical investigations. Under this process uncertainty arises from:

- a. Sample homogenisation;
- b. Incomplete extraction of the analyte from the sample;
- c. Loss of analyte due to over drying of the sample;
- d. Loss of volatiles due to excessive drying or crushing; and
- e. Sample contamination.

3. Environmental factors

Environmental factors such as vibrations, changes in humidity, changes in temperature, and electronic noise, or other effects may affect the measurement process thereby contributing to uncertainty. The factors may affect the functionality of the instrument used to take the measurements, introducing errors as a result. Mass, pH and volume measurements are examples of measurements that are highly influenced by the environment.

4. Instrumental effects

The instrument used for analysis may present the weakest link in the measurement process. Under instrumental effects, uncertainty is introduced by:

- Poor detection capabilities of the analytical instrument: The uncertainty associated with the sensitivity of an analytical device consists of a contribution from the standards used for instrument calibration, a contribution from the curve fitting process and a contribution from the model used to describe the response. For example, XRFs generally fail to detect minor elements with accuracy. Causes of such incapability are linked to short counting time, high back ground interference and long decay time. This source of uncertainty can be minimized by using more sensitive techniques.
- **Improper instrument calibration**: The uncertainty component associated with the calibration will include an uncertainty contribution from the reference materials and an uncertainty contribution from the calibration line fitting.
- **Instrumental drift:** Electronic instruments have readings that drift over time. This source of uncertainty can be significant and should be considered.
- **Manufacturer specifications:** Most measurement instruments have an accompanying statement of accuracy or tolerance level specified by the manufacturer. This value should be incorporated in uncertainty measurements.

5. Analytical Error

For most methods of estimating uncertainty, the assumption is that bias has been corrected and it is negligible. However, even if the bias is zero, it has to be estimated and treated as an uncertainty component. Method bias can be assessed and estimated through the interlaboratory comparisons, certified reference material (CRM), recovery test, and comparison to results from reference method⁵.

Common sources of uncertainty in the laboratory results from:

- Incorrect identification of samples;
- Mis- interpretation of the analytical method;
- Sample contamination;
- Inaccuracy of sample weights, or volumes for example constantly reading a meter or scale consistently high or low;
- Inappropriate sample dissolution/treatment;
- Improper or inappropriate instrumentation /inaccurate measurement; Calculation errors; and Data mix up and incorrect reporting.

These sources are closed associated with human error.

Cause and effect diagrams (fish bone) may be used as convenient and effective means for analysing uncertainty sources. The diagram assists in identifying, exploring, and displaying relevant uncertainty sources and highlights their effect on the measurement result. The procedures for constructing cause and effect diagrams can be summed as follows:

- When the result is based on a mathematical equation, it is recommended to use the parameters in the equation as the main branches for the diagram. If the parameters are unknown, one would consider the main procedures of the measurement process (i.e. sampling, sample preparation and analysis).
- The next step is to consider additional factors that form sub-branches for the main effects (main branches).

When all the sources have been identified, it is necessary to remove any duplication and group causes that are related. Figure 3-4 illustrates a cause and effect diagram for uncertainty sources associated with sampling.

⁵ Note that the list stated and the potential sources of uncertainty stated are not exhaustive. Other methods for estimating method bias exist.





Example II

Common sources of uncertainty in ICP MS measurements include:

- Incomplete dissolution of the analyte;
- Spectral interferences including isobaric interference, molecular interferences and doubly charged ion interference;
- Space charge effects;
- Trace elements such as Be, As and Hg have a high first ionization potential with the result that low temperature plasma will give reduced signal for these elements; and
- The presence of salts such as NaCl leads to reduced sensitivity.

Example III

Potential sources of uncertainty in X-Ray Fluorescence (XRF) analysis results include:

- Calibration of the spectrometer: once a measurement device is selected it must be calibrated. Failure to adjust the instrument to the required standard yields incorrect results;
- Sample in-homogeneity;
- Sample preparation: The sample is liable to contamination during homogenization,

grinding, milling, and polishing. Two commonly used polishing agents - Silicon Carbide (SiC) and Aluminum Oxide (Al_2O_3) - contain elements that are often determined silicon (Si) and aluminum (Al);

- Enhancement effect due to matrix effects;
- Counting errors; and
- Spectral overlaps.

In the case of trace element determination, the uncertainty of the x-ray's peak intensity is also an important factor.

3.2.2.6 Step 6: Quantify the uncertainty components⁶

Following the identification of uncertainty sources is the quantification of the uncertainty sources. For this methodology each contributing source is broken down into its constituent components and the size of uncertainty component estimated. Each of these separate contributions to uncertainty is referred to as an uncertainty component (Ellison, Rosslein and William, 2000). The uncertainty components are then expressed as standard deviation. Standard uncertainty denoted as $u_{(s)}$ is term used when uncertainty component is expressed as a standard deviation.

One major setback presented by tying to identify each uncertainty component and quantifying them is that some of the components are difficult to quantify – for example the effect of the sampling strategy. As important as the factor is, as a potential source of uncertainty, assigning uncertainty contribution in a quantitative way can be a complex task. Such problems are curbed by seeking additional information from literature, conducting additional experiments, or using judgment based on experience in order to effectively model uncertainty component.

⁶ <u>NOTE</u>: Not all sources of uncertainty can be quantified. It is also important to recognize that not all of the components of uncertainty will make a significant contribution to combined uncertainty. A preliminary estimate of the contribution of each component or combination of components to the uncertainty should be made and those that are not significant eliminated.(Ellison, Rosslein and William, 2000, pg 4)

Evaluating uncertainty components

After identification of the possible sources of uncertainty, the next step is to assess the uncertainty sources that are accounted for by the available data prior to estimating the size of each contributing component. Additional sources of information may be consulted to gather information about uncertainty components which are not adequately represented by existing information. In cases where data is absent, it may be necessary to conduct experiments to obtain additional information regarding a particular source of uncertainty.

Basically there are two main categories for evaluating standard uncertainty - category A and B, based on the method used to estimate their numerical values. **Category** A^7 is used to calculate standard uncertainty for estimates obtained by statistical analysis for example. Uncertainty components arising repeated measurements. **Category B** is used to evaluate standard uncertainty for uncertainty components obtained by non-statistical means, for example uncertainty stated by the manufacturer. Expressing uncertainty components quantitatively to standard uncertainty enables all estimates to be in equal form facilitating easy combining.

Category A

The Category A approach to uncertainty estimation assumes that statistical methods can provide reasonable estimates of the measurement uncertainty (Adams, 2002) Often sources that can be evaluated using this method are associated with random error that is experiment measurement results which vary when the measurement is repeated.

An example of a Category A evaluation involves making a series of n independent measurements of a quantity, Q_i and calculating the arithmetic mean and the experimental standard deviation of the mean. The arithmetic mean is regarded as the best estimate of the measurand and standard deviation of the mean provides a good approximation of the uncertainty associated with the mean measurement.

⁷ Category A refers to the method by which the uncertainty estimate was obtained, not the nature of the uncertainty contributor itself (Adams, 2002) this is also the case with category B.

Procedure for estimating Category A uncertainty

a. Identifying and removing outliers from the dataset.

Outliers negatively affect the statistic parameters such as mean and standard deviation. Presence of outliers increases the calculated parameter value especially standard deviation and therefore should be discarded. It is recommended that an inquiry be made regarding the presence of outlier prior to their rejection. The analyst should at least try to understand why they appeared.

Several methods of detecting outliers exist including Grubb's test and Chauvenet's criterion. The procedure for detecting outlier based on the Chauvenet's criterion can be summarised as below (Wikipedia, 2007):

- Calculate the mean and standard deviation;
- Determine the probability that a given data point will be at the value of the suspect data point.
- Using the normal distribution function, multiply this probability by the number of data points taken, remove the point if the result is less than 0.5

b. The measurement results are summed and divided by the number of samples to get the mean.

$$\bar{x} = \frac{\sum_{i=1}^{n} X_i}{n}$$
 1.2

c. Determination of the standard deviation (Equation, 2.4) of the measurements, which

characterizes the variability, or spread, of the observed values.

$$S = \sqrt{\frac{1}{n-1} \sum_{i=1}^{n} (X_i - \bar{X})^2}$$
 2.4

d. Determination of the standard error of the mean (Equation 2.5) which is standard uncertainty of the measurement:

$$u_{\overline{x}} = \frac{s}{\sqrt{n}}$$
 2.5

Where $u_{\bar{x}}$ = standard uncertainty.

Example IV

Seven pH measurements for waste rock yielded the following results:

8.12	8.52	8.4	8.28
8.15	8.44	8.32	
Mean	Standard Deviation	Standard Error	
8.32	0.15	0.06	

The arithmetic mean $P\Box$ for these results:

$$P \Box = 8.32$$

The standard uncertainty of P is

$$u_{\bar{p}} = 0.06$$

The procedure for determining standard uncertainty mentioned above is true for multiple measurements. In the case of single measurements, the uncertainty associated with the results is determined by taking the standard deviation of the measurement.

The advantage of using the standard uncertainty based on the standard error is that it relates the uncertainty to the sample size hence uncertainty can be reduced by increasing the sample size.

Category B

A Category B evaluation is used to estimate the standard uncertainty value for uncertainty sources which cannot be evaluated by statistical analysis as the case with Category A. The successful identification and evaluation of these contributions heavily depends on a detailed

knowledge of the measurement process and the experience of the person making the measurements.

Uncertainty components that can be evaluated by category B are chiefly derived from (EAL Task Force, 1999):

- Specifications of the manufacturer for example the reported calibration uncertainty assigned to reference standards;Previous measurement data;
- Data provided in calibration and other certificates;
- The effects of environmental conditions;
- Uncertainties assigned to reference data taken from handbooks, e.g. uncertainties associated with atomic weight; and
- Experience with behavior and properties of instruments.

The mode in which the uncertainty components which can be evaluated by this category are transformed to standard uncertainty primarily depends on available information regarding the uncertainty component. In some instances, conversions to standard uncertainty solely rely on the knowledge, judgment or experience of the analyst.

The following section discusses some of the possible methods employable in assessing and converting uncertainty components to standard uncertainty.

Method 1: This method calls for the conversion of the stated uncertainty to standard uncertainty by dividing the stated uncertainty by the multiplier (Taylor and Kuyatt, 2004).

Example V

A calibration certificate for a weight reference standard provides a value of 15 000.005g with an associated expanded uncertainty⁸ of 45mg (coverage factor of k=1.96). The standard uncertainty contribution from calibration is:

⁸ Expanded uncertainty = uncertainty*coverage factor (k), (commonly used value for k=1.96 which is equivalent to 95% confidence interval)

u (m) = 45mg (stated uncertainty)/1.96 (coverage factor which is the multiplier) =22.95mg

Method 2: If there are specifications that contain an uncertainty bound for input quantity at a stated confidence interval, the standard uncertainty is calculated by dividing the stated uncertainty by an appropriate factor for the confidence interval.

Example VI

For a given Certified Reference Material (CRM), with a certified value of 14.21 ± 0.2 and a 95% confidence level, the standard uncertainty for the CRM is

$$(u_{\rm crm}) = 0.2/1.96 = 0.10$$

Method 3: Probability Distributions:

A probability distribution is a mathematical function giving the probability that a random variable takes any given value or else belongs to a set of values (UKAS, 2007). There are many probability distributions that can be assumed but in this section, only three distributions are considered- rectangular, triangular, and normal distribution.

1. Rectangular/uniform distribution

Rectangular distribution is assumed when the only information available about the input quantity x are the upper and lower limits $\pm x$ and no other additional information is available. Assuming this probability denotes that true value for the quantity lies anywhere within the specified bound as shown in Figure 3-5.

The standard deviation for distribution is given as:

$$u(a) = s = \frac{a}{\sqrt{3}} \qquad \qquad 3.6$$

Conditions for assuming this probability distribution are as follows:

- 1. Uniform probability that is every value between and including the limits have equal probabilities of occurring.
- 2. There are zero chances of the uncertainty contributor, occurring outside of the limits of the distribution



Figure 3-5: Rectangular distribution (Vetter, 2001)

Example VII

A 50 ml Grade A volumetric flask is certified to within ± 0.25 ml. The standard uncertainty (standard deviation) associated with using the flask is calculated by dividing the given limit (uncertainty) with the rectangular distribution.

$$u_{(vol)} = 0.25 / \sqrt{3}$$

= ± 0.14 ml.

Where $\mathbf{u}_{(vol)}$ is the standard uncertainty associated with volume measurement.

Given a reading of 23.5g on an analytical balance with a manufacturer's specified tolerance of 0.1g the standard uncertainty for the reading is:

$$u_{(mass)} = 0.1/\sqrt{3}$$

= 0.06

Therefore the true value of the measurand lies between 23.4 and 23.6g.
2. Triangular

This distribution is assumed when the information available about the input quantity x are the upper and lower limit $\pm a$, and there is high probability that the quantity x values close to the center of the limits are more probable than values near the limits, Figure 3-6.

The standard uncertainty of the distribution is given as:

$$u(a) = s = \frac{a}{\sqrt{6}} \qquad \qquad 3.7$$

Where u (*a*) is the standard uncertainty associated with quantity *a* S is the standard deviation.



Figure 3-6: Triangular distribution (Vetter, 2001)

3. Normal distribution

The adoption of normal distribution is based on several situations. First, the distribution is assumed when estimates are made from repeated measurements of random varying process where standard uncertainty (u_x) equals the standard deviation (s) of the repeated measurements:

$$u_x = s$$

Second, normal distribution is adopted when the uncertainty associated with a given input is given as a standard deviation, relative standard deviation, and or coefficient of variation with the distribution unspecified.

In cases where the uncertainty is given as a confidence interval $x \pm a$ without specification of

the distribution, a normal distribution can be assumed. The standard uncertainties for the two most common distributions are therefore calculated as:

$$u_x = \frac{a}{1.96} (95\% \text{ confidence interval})$$
 3.8

$$u_x = \frac{a}{2.57} (99\% \text{ confidence interval})$$
 3.9

This distribution is illustrated in Figure 3-7.



Figure 3-7: Normal distribution (Vetter, 2001)

Generally for most analytical tests for geochemical sampling and analysis, uncertainty components that are evaluated using Category B usually contribute less than components that can be evaluated by Category A. As such, quality control data are often employed to provide an estimate of the contribution of analysis to overall uncertainty.

Procedures for estimating standard uncertainty are summarised by Figure 3-8.

3.2.2.7 Step 7: Calculating of combined uncertainty

The combined standard uncertainty is the uncertainty of a result when it is obtained from the values of several other statistically independent quantities (Bronaugh and Heirman, 2004). Although uncertainty components are not errors, the individual components are combined using laws of error propagation model. The combined standard uncertainty, which is usually denoted by u_c , is the uncertainty of a result when obtained from the values of several other

statistically independent quantities.



CC=calibration certificate

Figure 3-8: Flowchart for quantifying uncertainty sources

Laws for combining uncertainty

There are two main rules for combining standard uncertainties (EURACHEM/CITAC, 2002):

Rule 1

In cases involving addition or subtraction of quantities such as:

x=f (w + s + t \pm p....) with w, s, t, and p are uncorrelated, the combined uncertainty is calculated as:

$$u_x = \sqrt{u_w^2 + u_s^2 + u_t^2 + u_p^2 \dots}$$
 3.10

• Rule 2

For cases involving multiplication or division for example x=w*f*h, combined uncertainty is given by

$$u_x = \sqrt{\left[\frac{u(w)}{w}\right]^2 + \left[\frac{u(f)}{f}\right]^2} \qquad 3.11$$

In case of geochemical investigations for ERA the combined uncertainty from the common possible factors can be written as:

$$U(s) = \sqrt{u_h^2 + u_v^2 + u_{rm}^2 + u_{mm}^2}$$

Where u_x^2 is the standard uncertainties from:

 u_h^2 effect of heterogeneity u_v^2 volume measurement u_{rm}^2 Reference material u_{mm}^2 Mass measurement

Example VIII

Given that the pH measurements results for a certain target area with the dams a, b and c have the following standard uncertainties derived from standard deviation of repeated measurement values for each dam, dam a = u (*a*) = 0.13, dam b = u (*b*) =0.15 and dam c = u (*c*) = 0.22.The combined uncertainty for these is calculated by applying rule 1

$$\mathbf{u}_{\mathbf{x}} = \sqrt{u_w^2 + u_s^2 + u_t^2 \dots \dots \dots}$$

 $u_{(pH)} = \sqrt{0.13^2} + 0.15^2 + 0.22^2 = 0.296$

Therefore the combined uncertainty for the pH measurements is $u_{(pH)} = 0.296$

3.2.2.8 Step 8: Calculating the expanded uncertainty

Expanded uncertainty is calculated to indicate how confident one is that the true measurand value lies within a given range of values obtained during the measurement process. It is obtained by multiplying the combined uncertainty by a coverage factor k and can be expressed by Equation 3.12 below:

$$\boldsymbol{U} = \boldsymbol{k} \times \boldsymbol{u}_{c} (\boldsymbol{y}) \qquad \qquad 3.12$$

Where U is the expanded uncertainty, $u_c(y)$ is combined uncertainty, k is the coverage factor.

Usually a coverage factor (k) 9 of 1.92 is recommended when calculating expanded uncertainty (UKAS, 1997). This value provides a 95% confidence level, assuming the distribution is normal. If there is a larger contribution to uncertainty from one source in comparison to other sources, having the value k=1.96 will give a confidence level greater than 95%. When choosing the value of k, the level of confidence required should be considered. Table 3.7 list various coverage factors (k) that can be adopted.

 Table 3.7: Coverage factors

Probability (%)	50	68	75	80	85	90	95	99
Coverage factor (k)	0.68	1	1.15	1.28	1.44	1.65	1.96	2.57

Example IX

Using the results from Example 5, $u_{(pH)} = 0.296$, the expanded uncertainty associated with the combined uncertainty above is expressed as:

U=1.96 * 0.296

=0.580 (95% confidence level)

⁹ Value may be insufficient when their degrees of freedom <6.

3.2.2.9 Step 9: Reporting the uncertainty

Instead of reporting a single estimate, it is necessary to include an interval in which the true measurand value is expected to lie. The uncertainty should be reported in the same units as the results. Usually the expanded uncertainty value obtained in the previous step is used when reporting uncertainty. This is illustrated below:

Result:
$$(y \pm U)$$
 units 3.13

Where y is the measurand and U is the expanded uncertainty obtained from the previous step.

Example X: assuming that the pH result obtained for the previous Example is 8.12, the uncertainty associated with this result is given as:

$$pH = 8.12 \pm 0.59 (7.53 - 8.71)$$

The results should be reported whether positive or negative.

3.2.2.10 Advantages of the method:

Some of the advantages associated with this method are:

- The method is based on existing techniques.
- The guidelines enable the data users to identify the components of uncertainty, estimate the uncertainty associated with each component, and evaluate the contribution to uncertainty of the each components.
- It also allows the analyst to determine which component is contributing more to uncertainty hence the analyst can see where an adjustment in the sampling or analyses process is required so as to minimize uncertainty.

3.2.2.11 Limitations of the technique

Some of the limitations in using the technique are:

• The definition of the element that is being measured is usually

problematic. This is the case with geochemical sampling and analyses where a number of measurements and procedures are carried out.

• Some components of uncertainty are not readily quantifiable.

CHAPTER 4: CASE STUDY INFORMATION

4.0 BACKGROUND

The case study is based on geochemical assessment of tailings dams and waste rock carried out by Pulles Howard and De Lange (PHD) for AngloGold Ashanti Gold Mines Environmental Management. The objective of the assessment was to evaluate the potential of tailings dams and waste rock dumps in the Vaal River and West Wits mining areas to impact on water resources and implications of this in terms of mine closure and rehabilitation.

Gold mining waste is known to contain large quantities of heavy metals found in the host ore and chemical elements used in the extraction process such as cyanide. Under favorable conditions, in the presence of water and oxygen, metals and sulphur react with water and oxygen to produce Acid mine Drainage (AMD).

4.1 METHODOLOGY

In order to fully address the problem in question, PHD developed an in house geochemical assessment programme for ERA. The methodology adopted in undertaking geochemical assessment of tailings dams and waste rock dumps can be summarised by Figure 4-1.

Although all procedures illustrated in the figure above are of importance, for the purpose of this study, the main focus will be on sampling and analyses procedure.



Figure 4-1: Methodology followed during the project (Pulles, et.al, 2003)

4.2 SAMPLING TARGET

The sampling targets were tailings dams situated in the West Wits and Vaal River mining area owned by Anglo Gold (Ashanti), Figure 4-2.



Figure 4-2: Location of Sampled areas; Vaal River and West Wits

4.2.1 WEST WITS MINE

The West Wits tailings consist of three complexes namely:

(a) The Old North Tailings Complex with tailings covering an area of 108ha and containing approximately $24.12 \times 10^6 \text{m}^3$ of tailings.

(b) The New North tailings complex occupying an area of 237 ha and consisting of four dams: 5A, 5B, 7A and 7B.

(c) The South Tailings complex has an estimated volume of tailings of $11.12 \times 10^6 \text{m}^3$ and consists of two dams including an upper dam with an area of 55.1 ha and a lower dam with 73.3 ha (Figure 4-3).

4.2.2 VAAL RIVER MINE

In the case of the Vaal River areas two tailing dams were sampled. The West tailings complex occupy as area of 450ha while the West extension tailings dam has an area of 150 ha. The areas are illustrated in Figure 4-4 below.

4.3 SAMPLING PROTOCOL

The section gives an outline of the procedure adopted in sampling tailings dam.

4.3.1 SAMPLING TOOL

Dutch Augers were used for sampling of tailings.

4.3.2 SAMPLE COLLECTION

The sample collection procedure can be summarised as follows:

- Sample positions were planned at 1 sample point per 10 hectares.
- Positions were marked on a plan and Global Positioning System (GPS) was used to locate each point.
- Samples were collected using a Dutch auger at 50cm intervals.
- At shallow sample sites, samples were taken to a depth of 2.5m and deep holes at 10 meters. Due to the presence of saturated zone, very little was recovered and therefore

no deep holes were drilled.

Collected samples were labeled with each sample labeled three times.
 For some proposed sampling points, the positions were changed due to inaccessibility.

4.4 ANALYTICAL PROTOCOLS USED

4.4.1 SAMPLE COMPOSITING

Prior to analyses, samples were composited. All primary samples collected were composited into two composite groups. Composite samples from each shallow hole were made for ABA and moisture content analyses. Where there was no significant differences in paste pH, samples from each deep hole were made into one composite. Composite samples were further analysed for X- Ray Diffraction (XRD), X- Ray Fluorescence (XRF), water extraction Inductive Coupled Plasma Mass Spectrometry (ICPMS), Aqua Regia ICP MS, particle size distribution and microscopic study (Table 4.1):

Parameter	Analysis technique (s)
Acid Base Accounting (ABA)	Sobek Method
рН	pH meter
Particle size	wet and dry sieving
Mineralogy	X ray diffraction(XRD)
	Microscope
Major and trace minerals	X ray fluorescence (XRF)
Multi-element analysis	Inductively Coupled Plasma Mass
	Spectrometry (ICP MS)

 Table 4.1: Static test conducted

For the purposes of this discussion, only Acid Base Accounting (ABA) will be considered. Other methods are discussed in detail in the main report prepared for AngloGold, Ashanti Mines. All results obtained have been archived in a geochemical database.



Figure 4-3: West Wits tailings Dams (Google Earth, 2007)



Figure 4-4 Vaal River Tailings Dams (Goggle Earth, 2007) 109



Figure 4-5: Sampled points, West Wits tailing dams



Figure 4-6: Sampled points, Vaal River tailing dams

4.4.2 ACID BASE ACCOUNTING (ABA)

Acid base accounting (ABA) is the balance between acid production and acid consumption properties of mine waste (Fey, 2003). The potential for a given rock to generate and neutralize acid is determined by its mineralogical composition. This includes not only the quantitative mineralogical composition, but also individual mineral grain size, shape, texture and spatial relationship with other mineral grains (Mills, 1995).

Components of ABA include:

- Total sulphur;
- Paste pH;
- Acid potential (AP);
- Neutralization potential (NP);
- Net Neutralization potential (NNP); and
- Neutralization potential ratio (NPR).

It is usually the first step for determining AMD potential of mine waste.

Acid Base Accounting was carried out for the first batch of composite sample as illustrated in Appendix C.

> Total sulphur

Total sulphur content is used to predict acid generation potential from geological material if all the sulphur (S) present reacts (Kania, 1998). The most common sulphide minerals that weather to produce AMD include pyrite (FeS₂), marcasite (FeS₂), chalcopyrite (CuFeS₂) and arsenopyrite (FeAsS). The total sulphur (%) was measured using a Leco analyser. The resultant values were used to calculate acid potential.

Acid Potential (AP)

Acid potential is calculated from the product of total sulphur and 31.25kg/t of calcium carbonate (Sobek *et.al.*, 1978). The relationship can be expressed by the equation below:

$$AP = Total Sulphur (\%) \times 31.25 kg/t CaCO_3 \qquad 3.5$$

Neutralization Potential (NP)

The procedure used is based on the Fizz test by Sobek, et.al, 1978. NP is expressed as:

$$NP = \frac{50a\left[x\left(\frac{b}{a}\right)y\right]}{c}$$
 4.1

Where

a is the normality of HCl added in digestion,

b is the normality of NaOH used in the titration,

c is the mass of sample in grams,

x is the volume of HCl added in ml; and

y is the volume of NaOH added in titration.

> Paste pH

The pH was determined using pH meter. Further discussion is conducted in Chapter 5.

Net Neutralization Potential (NNP)

Net Neutralization Potential refers to the calculated difference between NP and AP

$$NNP = NP - AP \qquad 4.2$$

According to the total sulphur (%) and the ratio between NP and AP the material was classified as in Table 4.2.

TYPE I	Potentially acidic forming	Total S (%)>0.25% and NP:AP ratio 1:1 or less
TYPE II	Intermediate	Total S(%)>0.25% and NP:AP ratio 1:1 to 3:1
TYPE III	Non Acid forming	Total S(%)<0.25% and NP:AP ratio 3:1 or greater

Table 4.2: Rock type classification (Pulles et. al., 2003)

4.5 RESULTS

All analyses results have been stored in a geochemical database accompanying this report. The database in Excel spreadsheet format consists of several worksheets highlighting the various analytical procedures carried out. The basic unit of all sheets is an individual sample or sample composite. The structure of the database is as follows:

Common fields

- Location: Large locality for example West Wits mining area
- Small locality, e.g. Dam A
- Coordinates: Latitude and Longitude in Decimal degrees
- Sample ID , e.g. A-3S

Specific Fields

♦ Mineralogy worksheet:

Quartz	Hematite	Calcite	Dolomite	Gypsum	Plagioclase	K- feldspar	Amphibole	Mica	Prrophyllite	Chlorite

✤ Acid Base Accounting (ABA) worksheet

Paste pH	Total S (%)	AP (kg /t)	NP: AP rati o	C- NP (kg /t)	C-NNP (kg/t)	Rock Type total S	Rock type AP:NP	Samplin g depth	Comp osite	Latitude (Decimal degrees)	Longitude (Decimal Degrees)
-------------	----------------	------------------	------------------------	------------------------	-----------------	-------------------------	--------------------	--------------------	---------------	----------------------------------	-----------------------------------

✤ Grain Size worksheet

Sample	750 um	150 um	75um	>75 um	Total	Accuracy	Error	Material
weight(g)	(g)	(g)	(g)	(g)	(g)	(%)	(%)	type

✤ Major Elements worksheet

SiO2	TiO2 Al2	I2O3 Fe2O3	MnO	MgO	CaO	Na2O	K2O	P2O5	Cr2O3	L.O.I	Material Type
------	----------	------------	-----	-----	-----	------	-----	------	-------	-------	---------------

✤ Trace elements

As Ba	Ce	Co	Cr	Cu	Ga	Hf	Mo	Nb	Ni	Pb	Rb	Sc	Sr	Та	Th	U	v	W	Y	Zn	Zr	Material type
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GRAIN SIZE														
<u>Mining Area</u>	Large Locality Complex	Small Locality Dam ID	Sample ID	Sample weight(g)	750 um (g)	150 um (g)	75um (g)	>75 um (g)	Total (g)	Accuracy (%)	Error (%)	Mate typ		
	-	Dam A	A_COMP	500.02	33.25	261.18	179.68	25.42	495.54	99.10	0.90	Tailing		
	Now Newto	Dam B	B_COMP	400.00	57.77	124.06	191.45	28.73	395.82	98.96	1.04	Tailing		
	Compley	Dam C	C_COMP	500.01	26.72	165.28	225.47	38.80	497.96	99.59	0.41	Tailing		
	Complex	Dam D	D_COMP	400.01	39.87	171.95	162.51	25.84	398.21	99.55	0.45	Tailing		
			Complex Mear	450.01	39.40	180.62	189.78	29.70	446.88	99.30	0.70			
WEST WITS	Old North	Dam E,F &G	E-F-G_COMP	500.01	33.25	112.34	290.35	62.42	498.36	99.67	0.33	Tailing		
	Complex	Dam H,I&J	H-J-I_TOP_CC	500.02	57.77	383.57	49.15	9.13	499.62	99.92	0.08	Tailing		
	Complex		Complex Mear	500.02	45.51	247.96	169.75	35.78	498.99	99.80	0.20	Tailing		
		Dam M	M_COMP	500.03	26.72	139.07	280.32	51.84	497.95	99.58	0.42	Tailing		
	South Complex	Dam N	N_COMP	500.03	39.87	226.46	193.83	38.30	498.46	99.69	0.31	Tailing		
			Complex Mear	500.03	33.30	182.77	237.08	45.07	498.21	99.64	0.37			
		Dam A & B	VR-A-B-1_CON	500.13	96.83	219.10	133.65	46.65	496.23	99.22	0.78	Tailing		
		Dam B & D	VD-D-B-2_COI	500.02	55.66	185.01	172.01	85.16	497.84	99.56	0.44	Tailing		
	West Complex	Dam C	VR-C-1_COMF	500.00	30.48	111.44	243.52	119.28	504.72	100.94	-0.94	Tailing		
				Dam C	VR-C-2_COMF	500.04	27.06	130.86	228.56	109.70	496.18	99.23	0.77	Tailing
	-		Complex Mear	500.05	52.51	161.60	194.44	90.20	498.74	99.74	0.26	-		
	West Extension	Dam F	VR-F-1_COMP	500.01	32.97	242.30	175.96	46.73	497.96	99.59	0.41	Tailing		
	complex	Dam F	VR-F-2_COMP	500.40	6.34	185.98	270.32	39.92	502.56	100.43	-0.43	lailing		
VAAL RIVER			Complex Mear	500.21	19.66	214.14	223.14	43.33	500.26	100.01	-0.01			
	10/a aka	waste Dump 3	VRD3-1-COMF	500.00	123.42	277.18	93.19	4.85	498.64	99.73	0.27	ROCK V		
	vvaste rock	Waste Dump 3	VRD3-2-COMF	500.06	134.82	2/8.48	/5.05	11.95	500.30	100.05	-0.05	ROCK W		
	onuih a	waste Dump 3	VRD3-3-COMF	500.00	183.69	258.33	52.93	0.99	495.94	99.19	0.81	ROCK M		
		Wester Dune 1	Complex Mear	500.02	147.31	2/1.33	13.72	5.93	498.29	99.66	0.34	Deals		
		waste Dump 4	VRD4-1-COMF	500.01	181.09	206.15	85.90	26.16	499.30	99.86	U.40	ROCK V		
	vvaste rock	waste Dump 4	VRD4-2-COMF	500.00	161.11	264.21	60.16	12.39	497.87	99.57	0.43	ROCK V		
	oump 4	waste Dump 4	VRD4-3COMP	500.00	130.56	215.36	125.22	27.03	498.17	99.63	0.37	ROCK M		
	-		Complex Mear	500.00	157.59	228.57	90.43	21.86	498.45	99.69	0.40			
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Figure 4-7: An example of the structure of the database worksheet

CHAPTER 5: CASE STUDY RESULTS

5.1 SAMPLE SIZE DETERMINATION

This section serves to illustrate how the proposed methodology can be applied. It also serves to assess if the number of samples collected during the case study were enough for ERA. As such, ABA results for Vaal River dam 3 and West Wits dam N will be used to demonstrate the applicability of the method. The data for the two dams illustrated in Table 5.1. Key parameters for acid rock drainage (ARD) study are the acid potential (AP) and the neutral potential (NP). These parameters are used to illustrate how the proper sample size can be determined.

Location	Sample	Total S (wt	Calcite	AP	NP
Location	ID	%)	(kg/t)	(kg/t)	(kg/t)
	VRD3 A1	0.130	8.75	4.06	8.75
	VRD3 A2	0.277	0.25	8.66	0.25
	VRD3 A3	0.326	0.00	10.19	0.00
	VRD3 A4	0.310	0.75	9.69	0.75
	VRD3 A5	0.207	2.00	6.47	2.00
Waal	VRD3 A6	0.320	0.25	10.00	0.25
V aal Divor	VRD3 P1	0.070	2.00	2.19	2.00
Kiver	VRD3 P2	0.125	0.00	3.91	0.00
	VRD3 P3	0.283	0.00	8.84	0.00
	VRD3 P4	0.127	0.75	3.97	0.75
	VRD3 P7	0.188	0.00	5.88	0.00
	VRD3 P10	0.247	0.00	7.72	0.00
	N-1S	0.789	2.75	24.66	2.75
	N-2S	0.656	7.50	20.50	7.50
West	N-3D	0.691	9.75	21.59	9.75
Wits	N-5S	0.764	7.50	23.88	7.50
	N-6S	0.716	7.00	22.38	7.00
	N-7D	0.652	9.75	20.38	9.75

Table 5.1: ABA data from West Wits and Vaal River Mines (Pulles et al., 2003)

The dataset in the table above were derived from results obtained from the case study of Vaal River and West Wits tailing dams as discussed in Chapter 4. The dataset is only a small

portion of the results that were obtained from the study. Population A represents 12 samples taken from Vaal River ABA results and Population B represents 6 samples extracted from ABA results for West Wits mining area.

For the population A, The sample means $(m_{AP} (A), mNP (A))$ and the sample standard deviations $(S_{AP(A)}, S_{NP(A)})$ are calculated as follows:

$$\begin{split} m_{AP (A)} &= 1/12 \sum x_i = 6.80 \\ m_{NP(A)} &= 1/12 \sum x_j = 1.23 \\ S_{AP (A)} &= SQRT \left[(1/11)^* \sum (x_i - 6.80)^2 \right] = 2.77 \\ S_{NP (A)} &= SQRT \left[(1/11)^* \sum (x_j - 1.23)^2 \right] = 2.48 \end{split}$$

For the population B, the sample means $(m_{AP(B)}, m_{NP(B)})$ and the sample standard deviations $(S_{AP(B)}, S_{NP(B)})$ are calculated as follows:

$$\begin{split} m_{AP(B)} &= 1/6 \sum x_i = 22.23 \\ m_{NP(B)} &= 1/6 \sum x_j = 7.38 \\ S_{AP(B)} &= SQRT \left[(1/5)^* \sum (xi - 22.23)^2 \right] = 1.758 \\ S_{NP(B)} &= SQRT \left[(1/5)^* \sum (xj - 7.38)^2 \right] = 2.563 \end{split}$$

Here we set that the acceptable sampling error (r) of AP and NP equals 1(kg/t) and confidence equals 95%.

Based on the information above, the sample sizes can be calculated as follows:

$$n_{AP (A)} = Z_{\alpha/2}^{2} S_{AP(A)}^{2} / r^{2}$$

$$= 1.96^{2} * 2.77^{2} / 1^{2}$$

$$= 30$$

$$n_{NP (A)} = Z_{\alpha/2}^{2} S_{NP(A)}^{2} / r^{2}$$

$$= 1.96^{2} * 2.48^{2} / 1^{2}$$

$$= 24$$

$$n_{AP (B)} = Z_{\alpha/2}^{2} S_{AP(B)}^{2} / r^{2}$$

$$= 1.96^{2} * 1.758^{2} / 1^{2}$$

$$= 12$$

$$n_{NP (B)} = Z_{\alpha/2}^{2} S_{NP (B)}^{2} / r^{2}$$

$$= 1.96^{2} * 2.563^{2} / 1^{2}$$

$$= 25$$

5.1.1 SUMMARY

Sample size determination is a crucial step in the statistical design of the project. An adequate sample size helps to ensure that the study will yield reliable information which is critical for decision making. Studies involving human subjects should be designed with a large enough sample size so that the risk of making wrong decision is eliminated.

Performing a valid sample size calculation requires estimates of the variability in the data, as well as defining the desired confidence level and the acceptable sampling error. Standard deviation, the often used measure of variability and acceptable sampling error are unknown most of the time. These parameters can be estimated from historic data of similar nature or pilot study. The main risk of using data from previous studies in estimating sample size is the possibility of error propagation. Errors made in the previous study may be carried over to the present study; therefore care is required when using such data.

There is an exponential relationship between sample size and acceptable sampling error. In order to reduce error by a factor of two, the sample size has to be quadrupled hence error can be reduced by increasing sample size. For any given confidence level, the larger the sample size, the smaller the confidence interval

In the case of variability, the more heterogeneous a population, the larger the sample size required to obtain a given level of precision. The more homogeneous a population, the smaller the sample size required.

Although the determination of sample size presents a difficult task, the hurdles can be overcome by drafting a proper sampling plan. It is of great importance that a sampler understands the purpose and uses of the sampling data prior to deciding the number of samples required. As sample size affects representativity and decision making, it is essential that the right sample size be collected.

5.2 QUANTIFYING UNCERTAINTY

In the previous chapter, a methodology for quantifying uncertainty in geochemical sampling and analysis was discussed. This chapter outlines how the methodology can be applied. AcidBase Accounting (ABA) data results from the case study discussed in Chapter 4 with the parameters acid potential (AP), neutralizing potential (NP) and paste pH are used to illustrate the application of the methodology. The input values for the measurements are listed in Appendix D.

Since the measurement results for the various parameters shown in Appendix D are mean concentration values within each tailing dam, derived from averaging individual sample results from varying locations of the target, the standard uncertainty associated with these measurements are evaluated under Category A as discussed in Chapter 3. Equation 2.5 is used to determine the associated standard uncertainty:

$$u_{\overline{x}} = \frac{s}{\sqrt{n}} \qquad 2.5$$

Where

u_x is standard uncertainty,

s is the standard deviation and

n is the sample size.

The standard uncertainty for uncertainty components falling under category B are evaluated using the appropriate probability distribution as discussed above in Chapter 3.

The standard uncertainties are combined using the laws of error propagation stated in Equations 3.10 and 3.11.

Expanded uncertainty is calculated by multiplying the combined standard uncertainty by the coverage factor (k) of 1.96 which gives an approximately 95% confidence level. This value is used since the degrees of freedom are above six for most variables.

The majority of the calculations were carried out in Ms Excel 2003. The formulas for calculation of the various statistics in Excel including mean, count, standard deviation and standard are listed below:

Mean is calculated as:

=AVERAGE (C1: C5)

Where, C1 and C5 are the cell numbers containing the input values Standard deviation as:

=STDEV (number1 [number2]....)

Where STDEV is standard deviation

Standard error of the mean with the formula in Equation 2.5 is calculated as

=STDEV (G5:G11)/SQRT (COUNT (G5:G11)

Where

STDEV is the standard deviation

G5:G11 refers to the location of the values used in the spreadsheet – that is values that are located in column G in rows 5 to 11

SQRT is the square root

COUNT is the sample size

Probability distributions are calculated as below:

=G8/sqrt (number).

The dividing number depends on the probability distribution chosen as discussed previously.

Combined Uncertainty as

E.g. addition rule

=SQRT (SUMSQ (number1 [number2].....)) for rule 1 =SQRT (SUMSQ (D8/D5, D20/D17.....)) for rule 2.

Where, SUMSQ is the sum of squares.

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1	ACID BASE AC	COUNTING	•							
-		Large Locality			Paste	Total S				
	Mining area		Small Locality Dam ID	sample ID			AP(ka/t)	NP:AP ratio	C-NP(kg/t)	C-NNP (kg/t)
2		Complex			рH	(%)			,	,
3	West Wits Tailings			A-1S	8.12	0.541	16.91	0.21	3.50	-13.41
4				A-2D	8.15	0.474	14.81	0.42	6.25	-8.58
5				A-38	8.52	0.537	16.78	0.36	6.00	-10.78
6				A-4S	8.44	0.420	13.13	0.40	5.52	-7.88
7				A-5D	8.40	0.493	15.14	0.45	7.00	-8.41
8				A-6S	8.32	0.505	15.78	0.29	4.50	-11.28
9			Dam A	A-7S	8.28	0.541	16.91	0.34	5.75	-11.16
10				A-Mean	8.32	0.50	15.64	0.35	5.50	-10.21
11				A-Standard Dev	0.15	0.04	1.40	0.08	1.16	2.00
12				A-Sample size	7	7	7	7	7	7
13				A-Standard	-STDE	//E3·E0\/S	OPT/E12)	0.031	0.440	0.756
13				uncertainty	STDE	(LJLJ)/J	[number2]	0.051	0.440	0.750
14				B-1S	SIDE	(number 1	, [numberz])	0.39	5.75	-8.91
15				B-2D	8.62	0.551	17.22	0.39	6.75	-10.47
16				B-3S	8.64	0.332	10.38	0.70	7.25	-3.13
17				B-4S	8.64	0.461	14.41	0.47	6.75	-7.68
18				B-5D	8.52	0.455	14.22	0.39	5.50	-8.72
19			Dam B North Iex	B-6S	8.60	0.752	23.50	0.29	6.75	-16.75
20				B-7S	8.35	0.597	18.66	0.71	0.00	-18.66
21				B-mean	8.51	0.52	16.15	0.48	5.54	-10.61
22		New North Complex		B-standard dev	0.18	0.13	4.16	0.16	2.52	5.38
23				B-Sample size	7	7	7	7	7	7
24				B-Standard Uncertainty	0.067	0.050	1 572	0.062	0.952	2 0 3 3
25				C-18	8,23	0.465	14,53	0.33	5.54	-9.78
26				C-2S	8,20	0.475	14.84	0.37	4.75	-9.34
~~										
• •	ABA TAILINGS	WORKSHEET / I	Mineralogy / Grain size /	Major elemets(XRF)	/ Trace e	eleme <	1111			

Figure 5-1: Spreadsheet format for uncertainty determination

The following sections presents examples showing how the methodology discussed in Chapter 4 can be applied. ABA data from West Wits tailings dams are used.

Example 1 calculates uncertainty associated with Acid Potential (AP) measurement result for West Wits tailings dams,

Example 2 evaluates uncertainty associated with of Neutralizing Potential measurement result for all tailings dams.

Example 3 quantifies uncertainty associated with pH results obtained from paste pH for West Wits tailings dams.

5.2.1 EXAMPLE 1: ACID POTENTIAL (AP)

Step 1: Definition of the purpose of AP determination

Acid Potential is one of the main measurements undertaken in Acid Base Accounting. It was determined in AMD prediction with the aim of assessing the amount of acid producing elements present in waste rock and tailings. Chemical elements contributing to acid production include sulphide minerals such as pyrite, and pyrrhotite. These elements, under favorable conditions, react with oxygen and water leading to the production of acidic discharges which have a negative effect on the environment. For example the oxidations of pyrite (FeS2) produce such acidic discharge as highlighted by the reaction below:

$$2\text{FeS}_2(s) + 7\text{O}_2(g) + 2\text{H}_2\text{O}(l) \rightarrow 2\text{Fe}^{2+}(aq) + 4\text{SO}_4^{2-}(aq) + 4\text{H}^+(aq)$$

Step 2: Definition of the sampling target

The targets are tailing dams occurring in the West Wits mining area. The place is made up of three tailings dams namely: The Old North tailings complex, the New North tailings complex and the South tailings complex. The Old north tailings complex covers an area of 108ha containing an estimated 24.12*10⁶ m³ of tailings. The complex consists of six dams but for this study only information of dam F was provided. The New north complex has 4 dams, A, B, C, and D occupying an area of 237 ha. The South tailings complex consists of 2 dams, M and N. Both dams occupy an area of 125ha.

Step 3: Specification of the measurand

The aim of this step is to write down a clear statement of what is being measured. The measurand in this case is the assessment of the possibility of acid production potential (AP) of mine tailings in various dams .The measurand depends on two parameters which are total sulphur and weighed calcium carbonates. Total sulphur is determined using the method suggested by Sobek, *et.al* (1978), step 4 below. Acid Potential (AP) can be expressed as below:

AP = Sulfur content (%) x 1000 kg/100 x molecular weight of CaCO3/atomic weight of sulfur

$$AP = Total Sulphur \times 31.25 kg CaCO_3 \qquad 3.5$$

Step 4: Description of the sampling and analytical protocol

The goal of this step is to outline the protocol following during sampling and analyses.

Sampling protocol (Sampling procedure followed):

- Sample positions were planned at 1 sample point per 10 hectares;
- Positions were marked on a plan and GPS was used to locate each point;
- Samples were collected using a Dutch auger at an interval of 50cm;
- At shallow sample sites, samples were taken to a depth of 2.5m and deep holes at 10m. Due to the presence of saturated zone, very little was recovered and therefore no deep holes were drilled;
- Collected samples were labeled with each sample being labeled three times;
- Prior to analyses, samples; and
- For some proposed sampling points, the positions were changed due to inaccessibility.

Analytical protocol

The first step in determining AP was to calculate the total sulfur percentage. The total sulfur content was measured using Leco sulphur analyser. In a Leco sulphur analyser, a sample is combusted in a high temperature tube furnace containing pure oxygen. The products of combustion are passed through a moisture trap to the sulfur IR cell where sulfur is measured as sulfur dioxide. Using a calibration curve based on known sulfur compounds, the measured sulfur dioxide is converted to sulfur content per unit weight of sample.

The obtained total sulfur value is multiplied by the molecular weight of calcium carbonate divided by its atomic weight according to a method proposed by Sobek, Equation 3.1. According to Sobek *et.al* (1978), 3.125 g of CaCO₃ is capable of neutralizing the acid produced from 1 g of sulphur (S), in the form of FeS₂ - hence the amount of potential acidity in 1000 tons of overburden could be calculated by multiplying the percent S by 31.25. This argument is based on the equation below (Kania, 1998):

 $FeS_2 + 2CaCO_3 + 3.75O_2 + 1.5H_2O \longrightarrow Fe(OH)_3 + 2SO_4^{-2} + 2Ca^{+2} + CO_2$

Step 5: Identifying the potential sources of uncertainty

This section discusses the various factors which could have contributed to the uncertainty associated with the measurand.

Sampling

Sources of uncertainty associated with sampling include the following:

a) Inconsistency of the sampling strategy: The sampling strategy used when collecting samples introduced certain amount of bias. Initially numerous sampling point were planned but these were changed due to inaccessibility of some of the tailings dams hence they sampling technicians ended up, collecting samples from areas which were accessible only. Sampling of accessible areas violates the main principle of correct sampling which recommends equal selection opportunity for material or units occurring within the target.

b) Heterogeneity (constitutional and distribution): This is the major source of measurement uncertainty. The effect of this source can be displayed by variations in terms of the observable analyte concentration among samples collected from different points of a sampling target. Heterogeneity results from the fact that the distribution of elements in a sampling target is not uniform but varies.

c) Limited number of samples collected: In other dams, very few samples were collected which resulted in high uncertainty values for such dams.

d) **Sample compositing:** As stated in Chapter 4, samples were composited and each composite sample was made up of several increments. Test samples were then extracted from these for laboratory analysis. Although a good practice, sample compositing may result in sample dilution, in which an individual increment with a high analyte concentration is combined with an increment with a low analyte concentration resulting in false test results.

e) Sample collection tool: The Dutch auger used for collection introduced a certain amount of error. According to the rule of thumb for sampling, "all the constituents of a lot to be sampled must be given an equal probability of being selected and preserved as part of the sample" (Pitard, 2005, pp. 56). This rule was violated by the tools used for sampling since the instrument failed to penetrate a depth greater than 2 meters which meant that the material occurring above this depth could not be sampled.

Analysis including sample preparation

In addition to the effects of mass, temperature, and calibration of Leco analyser, the AP measurement result has an uncertainty contribution possibly resulting from:

- Incomplete combustion of the sample in Leco Analyser which gives an underestimate of the total sulfur content;
- Formation of sulphur dioxide and trioxide: The formation of Sulphur dioxide and sulphur trioxide during combustion results in inaccurate and imprecise results in measuring the total sulphur concentration;
- The molar mass of calcium carbonate CaCO₃ and atomic weight of sulphur. The uncertainty associated with atomic weight of the elements involved in the calculation of AP is listed in Table 5.2; and
- Mass measurements for CaCO₃ depend on the calibration of the balance used and the effects of resolution.

Element	Atomic weight	Quoted uncertainty ¹⁰
Ca	40.078	±0.004
С	12.0107	±0.0008
0	15.9994	±0.0003
S	32.065	±0.0005

Table 5.2: Atomic weight and their associated uncertainty (IUPAC, 2006)

Various sources of uncertainty are highlighted in the cause and effect diagram illustrated in Figure 5-2 below.



Figure 5-2: Cause and effect diagram for Acid Potential

¹⁰ The tabulated value should be applicable to any normal material. Note that for some geological specimen, the uncertainty may exceed the stated uncertainty value.(IUPAC, 2006)

Step 6: Quantifying Uncertainty

The aim of this step is to quantify uncertainty sources. It should be noted that due to limited information, the size of some sources of uncertainty mentioned in the previous section will not be estimated.

a. Variation between locations (effect of heterogeneity)

Samples taken from different parts of the sampling target exhibit variations in terms of the mean concentration values of acid producing elements, shown in Appendix D. As stated previously, the sampling target for the case study consists of 7 dams. Within each dam, several increments from different points of the dam were combined resulting in numerous composite samples. These composite samples were treated and analysed separately. For example, the composite sample with the ID A-1S is made up of 5 increments, Appendices C and D.

The mean of the single results constitutes the measurement result in agreement with the specification of the measurand, for dam A the final AP measurement value is derived from the average the results for composite samples A-1S, A-2D, A-3S, A-4S, A-5S, A-6S and A-7S. The calculation procedure is true for all of the seven dams within the area.¹¹

Since the measurement results for each dam is based on averaging all obtained sample values obtained as illustrated in the accompanying geochemical database, standard uncertainty associated with each dam's measurement results is therefore calculated using Equation 3.1. The results are listed in Table 5.3.

¹¹ Calculations were carried out in Excel workbook accompanying this document.

Dam ID	Mean	Standard Deviation	Sample Size	Standard uncertainty u _(dam) (kg/t)
Α	15.64	1.4	7	0.53
В	16.15	4.16	7	1.57
С	14.03	1.24	5	0.55
D	17.26	1.14	5	0.51
F	12.42	1.37	4	0.97
Μ	21.98	4.21	3	2.43
Ν	22.23	1.76	6	0.65
Combined standard uncertainty				3.25

Table 5.3: Descriptive statistics and standard uncertainties for AP measurement

Thus the uncertainty contribution from the effect so heterogeneity in AP measurements = 3.25 kg/t.

b. Molar mass of CaCO₃

Another source of uncertainty that could be identified is contribution from the molar weights of the various elements Ca, C, O and S. The standard uncertainties were calculated by dividing the quoted uncertainty by $\sqrt{3}$ (rectangular distribution). The results are listed in Table 5.4.

Table 5.4: Standard uncertainties for Ca, C and O

Elements	Atomic weight	Quoted uncertainty	Standard uncertainty
Ca	40.078	±0.004	0.00231
С	12.0107	±0.0008	0.00046
0	15.9994	±0.0003	0.00017
Combined standard un	0.00236kg/t		

Step 7: Combined uncertainties for AP

The combined uncertainty can be achieved by variance addition of the standard uncertainties from the various factors as below:

$$u_{c(AP)} = \sqrt{0.53^2 + 1.57^2 + 0.55^2 + 0.51^2 + 0.97^2 + 2.43^2 + 0.65^2} = 3.25 kg/t$$

The contribution from the atomic weight is minor when compared with that of variations between locations and it is therefore discarded. Results are illustrated in Table 5.5.

Dam ID	Mean (kg/t)	Standard Deviation	Sample Size	Standard uncertainty (kg/t)
Α	15.64	1.40	7	0.53
В	16.15	4.16	7	1.57
С	14.94	1.24	5	0.55
D	17.256	1.14	5	0.51
F	12.42	1.68	3	0.97
Μ	21.98	4.21	3	2.43
Ν	21.75	1.45	5	0.65
	17.16		U _c (between Location)	3.25

Table 5.5: Combined uncertainty from AP measurements West Wits tailing Dams

Step 8: Calculation of the expanded uncertainty

Expanded uncertainty is Uc (AP) is obtained by multiplying the combined standard uncertainty with the coverage factor of k = 1.96 (approximately 95% confidence level). The expanded uncertainty is:

$$U_{AP} = 1.96 \times 3.25 \, kg/t = 6.7 \, kg/t$$

Step 9: Reporting the result

The AP measurements for The West Wits tailings is

17.16 ± 6.70 kg/t at 95% confidence level

The value 17.16 represents the mean of all seven dams. The results above, show that the true concentration of AP elements lies somewhere between 10.66 and 23.66 kg/t.

Contribution of various dams is summarised by Figure 5-3 below.



Figure 5-3: Standard uncertainty contributions for AP measurements

5.2.2 EXAMPLE 2: NEUTRALISATION POTENTIAL (NP)

Step 1: Definition of the purpose of NP determination

Neutralization Potential is another important measurement carried out in ABA. The main aim of undertaking NP measurements was to assess the amount of acid neutralizing elements present within the tailings. That is assessing the amount of chemical elements capable of neutralizing acid producing elements. Acid neutralizing elements include carbonate minerals such as calcite and dolomite. When their concentration is larger than that of AP elements then the risk of AMD is minimal.

Step 2: Definition of the sampling target

The targets are tailing dams occurring in the West Wits mining area. The place is made up of three tailings dams namely: The Old North tailings complex, the New North tailings complex and the South tailings complex. The Old north tailings complex covers an area of 108ha containing an estimated $24.12*10^6$ m³ of tailings. The complex consists of six dams but for this study only information of dam F was provided. The New north complex has 4 dams, A, B, C, and D occupying an area of 237 ha. The South tailings complex consists of 2 dams, M and N. Both dams occupy an area of 125ha.

Step 3: Specification of the measurand

The measurand is the neutralization potential of tailings within West Wits area. NP can be expressed by the following equation:

$$NP = \frac{50a\left[x\left(\frac{b}{a}\right)y\right]}{c} \qquad \qquad 3.2$$

Where:-

a is the normality of HCl added in digestion,

b is the normality of NaOH used in the titration,

c is the mass of sample in grams,

x is the volume of HCl added in ml and

y is the volume of NaOH added in titration.

Step 4: Description of the sampling and analysis protocols

This step requires the analyst to give a detailed outline of the sampling and analytical procedure followed during measurement.

Sampling protocol

The sampling protocol adopted was discussed in Step 5 of Example 1.

Analytical protocol

The analytical protocol used is based on the methodology for NP measurement by Sobek et.

al., (1978). The procedure is outlined below.

- 1. Place approximately 0.5g of sample (less than 60 mesh) on a piece of aluminum foil.
- 2. Add one or two drops of 1:3 HCl to the sample. Presence of CaCO₃ is indicated by the bubbling or audible fizz
- 3. Rate the fizz in step 2 as shown in Table 6.5. The objective of the fizz test is to provide sufficient acid to complete the reaction with the acid.
- 4. Weigh $2.0g \pm 0.1g$ of sample into a 250ml beaker.
- 5. Add HCl as indicated in Table 5.6.
- 6. Heat nearly to boiling point, mixing occasionally until reaction is complete. Reaction is complete when all the gas (CO₂) has evolved.
- 7. Bring the volume to 125ml by adding distilled water.
- 8. Heat the contents of the beaker to boiling point for a minute. Let it cool.
- 9. Titrate using 0.1N NaOH and pH meter to pH 7.00.
- 10. Calculate NP.

HCl				
Fizz rating	ML	Normality		
None	20	0.1		
Slight	40	0.1		
Moderate	40	0.5		
Strong	80	0.5		

Table 5.6: Fizz ratings (Sobek et al, 1978)

Step 5: Identification of the potential sources of uncertainty

The aim of this step is to list all the possible sources of uncertainty for each of the parameters that affect the value of the measurand. A cause and effect diagram is also included.

Sampling

Sources of uncertainty associated with sampling include the sampling strategy, heterogeneity of the target, sample compositing, and use of unsuitable sampling tools. These sources were discussed in detailed in Example 1 under Step 5.

Analysis - including sample preparation:

a) Sample effect

During acid digestion, some minerals that co-exist with acid neutralizing elements may be digested at the same time as the carbonate minerals interfering with the whole analysis process. For example Kania (1998) found that the presence of siderite (FeCO₃) within the sample to be analysed may increase acidity and makes it difficult to hold the final end point of titration.

b) Mass Measurement

Assuming the measurements were carried out using a tarred scale with an accuracy of $\pm 0.1g^{12}$ common sources of uncertainty associated with using the measurement instrument include:

- Repeatability,
- Digital resolution (readability), and
- Sensitivity and linearity of the balance.

c) Volume measurements

The volume measurements are subject to three major uncertainty sources. These are:

 $^{^{12}}$ Most scales have an accuracy of +-0.1g.
- a) Variations in filling and reading the volume,
- b) Uncertainty in the certified internal volume of the beaker used, and
- c) Variation of the beaker and solution temperature from the temperature at which the volume of the beaker was calibrated.

In addition to the above stated factors affecting volume, the volume of NaOH for titration suffers from the uncertainty related to detection of end point for the titration process.

d) Molar Mass

NaOH and HCI have an uncertainty associated with their relative atomic masses. These are listed in Table 5.7.

Elements	Atomic weight	Quoted uncertainty (IUPAC, 2006)	Standard uncertainty
Na	22.98976928	± 0.0000002	1.1547E-08
0	15.9994	±0.0003	0.00017
Н	1.00794	± 0.00007	4.04145E-05
Cl	Cl 35.453		0.001155
Н	1.00794	±0.00007	4.04145E-05

Table 5.7: Atomic weights and their associated uncertainty

e) Normality of NaOH

The volume of NaOH that is titrated has a normality of 0.1 ± 0.005 .

Various sources of uncertainty are highlighted in the cause and effect diagram illustrated in Figure 5-4 below.

Step 6: Quantifying uncertainty sources

a. Variation NP between locations (Effect of Heterogeneity)

Samples taken from different locations of the dams show variation in terms of NP values as shown in Appendix D. The standard uncertainty of the measurement results (mean value) are calculated from Equation 3.1. Results locations are listed in Table 5.8.



Figure 5-4: Cause and effect diagram for uncertainty components for NP

Dam ID	Mean	Standard Deviation	Sample Size	Standard uncertainty
Α	5.50	1.16	7	0.44
В	5.54	2.52	7	0.95
С	5.96	1.73	5	0.78
D	6.15	1.94	5	0.87
F	6.30	0.92	3	0.53
Μ	9.42	1.23	3	0.71
Ν	8.30	1.34	5	0.60
	6.74			
			U _{c (between} Location)	1.90

Table 5.8: Standard uncertainties for NP measurement

b. Molar Masses of NaOH and HCl

The standard uncertainties associated with molar masses of the NaOH and HCl are calculated using the rectangular distribution (Equation 5.1). No other information was given with reference to the atomic masses.

$$u(a) = s = \frac{a}{\sqrt{3}} \qquad \qquad 3.6$$

The standard uncertainties are listed in Table 5.9.

Elements	Standard uncertainty, u _(mm)
Na	1.1547E-08
0	0.00017
Н	4.04145E-05
Cl	0.001155
Н	4.04145E-05
Combined uncertainty (u _{c mm})	$\sqrt{.1547E-08^2+0.00017^2+4.04145E-}$ $05^2+0.001155^2+4.04145E-05^2=$ 0.001168842

Table 5.9: Standard uncertainties associated with molar weights

c. Uncertainty of the laboratory glassware used

A volumetric beaker of 125ml was used during the analysis. Research of literature indicates that a 125ml beaker usually has an accuracy of $\pm 5\%$ which is 0.05ml.The standard uncertainty associated with the beaker is therefore:

$$u_{vol} = 0.05/\sqrt{3} = 0.03ml$$

d. Normality of NaOH

Because there is no additional information about the uncertainty value, a rectangular distribution is assumed. The uncertainty associated with NaOH normality is:

$$u_{normality} = 0.005 / \sqrt{3} = 0.002$$

e. Mass Measurement of the sample

The uncertainty associated with weighing 2.0g of the sample is

$$u_{mass} = 0.1/\sqrt{3} = 0.06g$$

A rectangular distribution was assumed.

Step 7: Calculate the combined uncertainty

The combined uncertainty is calculated as below:

 $u_c(NP) = \sqrt{uc(mm)2 + u(vol)2 + u(mass)2 + u(normality)2 + uc (btwn Loc)2}$

$$u_{c(NP)} = \sqrt{1.90^2 + 0.03^2 + 0.06^2} = 1.90 \ kg/t$$

Therefore the combined uncertainty u_c (NP) for NP West Wits measurements is **1.90 kg/t**. The contribution to uncertainty from molar mass and normality are negligible therefore ignored.

Step 8: Calculating the expanded uncertainty

The expanded uncertainty Uc ($_{NP}$) is obtained by multiplying the combined standard uncertainty with the coverage factor of k= 1.96 (approximately 95% confidence level).

The expanded uncertainty (Table 5.10) is

$$U_{AP} = 1.96 \times 1.90 \ kg/t = 3.72 \ kg/t$$

Dam ID	Mean	Standard Deviation	Sample Size	Standard uncertainty
Α	5.50	1.16	7	0.44
В	5.54	2.52	7	0.95
С	5.96	1.73	5	0.78
D	6.15	1.94	5	0.87
F	6.30	0.92	3	0.53
Μ	9.42	1.23	3	0.71
Ν	8.30	1.34	5	0.60
			u _c (btwn Loc)	1.90
			u _c (Vol)	0.03
			u _c (mass)	0.06
			u _c (NP)	1.90
			U	3.72

Table 5.10: Expanded uncertainty for NP measurements West Wits tailings

Reporting the result

Thus the amount of Neutralizing Potential within West Wits tailing dams is

6.74 ± 3.72 kg/t

Where the stated uncertainty is calculated using a coverage factor of 1.96 (95% confidence interval), 6.74 is the mean of all dams.

Figure 5-5 is a plot of uncertainties of various dams.



Figure 5-5: Standard uncertainty contribution for NP measurements

5.2.3 EXAMPLE 3: PASTE pH MEASUREMENTS

Ophardt (2003) defines pH as the negative logarithm of the hydrogen ion concentration. It is measured on a pH scale containing values ranging from 0 to 14. On the pH scale, a reading above 7 indicates alkalinity and below 7 is an indication of acidity (Table 5.11).

Table 5.11: pH scale

pH < 7	Acidic
pH = 7	Neutral
pH >7	Basic

For example the concentration of H^+ in a solution is 10^{-3} the pH is 3, acidic

Step 1: Defining the purpose of pH determination

The main aim for measuring the pH was to determine the concentration of hydrogen ions H^+ , in the tailings within the target area. If the hydronium ion concentration exceeds that of the hydroxide ion concentration the material is said to be acidic. The reverse is true for basic materials which have a pH of above 7 at 25^oC. The value obtained will be used in conjunction with AP and NP to assess and make a conclusion regarding the ARD potential generation of mine tailings.

Step 2: Definition of the sampling target

The targets are tailing dams occurring in the West Wits mining area. This area is made up of three tailings dams namely The Old North tailings complex, The New North tailings complex and The South tailings complex. The Old north tailings complex occupies an area of 108 ha containing an estimated 24.12*10⁶ m³ of tailings. The complex consists of six dams but for this study only information on dam F was provided. The New north complex has 4 dams, A, B, C, and D occupying an area of 237 ha. The South tailings complex consists of 2 dams, M and N. Both dams occupy an area of 125ha.

Step 3:Specification of the measurand

The measurand is the concentration of the amount of hydrogen ions (H^+) in mine tailings. pH is equal to the negative log of H^+ , Equation 5.1 (Prichard, 2003).

$$pH = -log_{10}a_{H+}$$
 5.1

Step 4: Description of the sampling and analysis protocols followed

The aim of this step is to discuss the measurement procedure in paste pH.

Sampling protocol

The sampling protocol followed was discussed in Step 4 of Example 1.

Analytical protocol

The analytical procedure used is based on the method for measuring paste pH suggested by Sobek *et.al*, 1978. The steps can be summarised as below:

- The pH meter is calibrated for test temperature using pH buffer solutions and standard procedures according to meter instructions.
- 10.0 g of sample is weighed into a beaker and approximately 5 ml of distilled water is added to sample.
- The sample is stirred with a spatula to form thin paste, adding more water or sample to keep sample at saturation point. Paste should just slide off spatula easily.
- pH electrode is carefully placed into paste and move about to ensure contact between paste and electrode. Care must be exercised to avoid impact and scratching of electrode.

• Record the pH of the paste.

The procedures are summarised in Figure 5-6:



Figure 5-6: Procedures involved in pH measurement

Step 5: Identifying potential sources of uncertainty

a. Sampling

Sources of uncertainty associated with sampling include the sampling strategy, heterogeneity, sample compositing and sampling tools used. The sources were discussed in detail under Step 5 of Example 1.

b. Analysis including sample preparation

Most uncertainty sources for pH result are linked to pH meter which is used for in obtaining the measurement. Potential sources of uncertainty include:

- a. Soluble salts: The presence of neutral salts affects pH readings by influencing ionic activities and in turn gives rise to activity errors.
- b. Uncertainty of the certified pH values for standard buffer solution.

- c. Drifting: Drifting of the pH reading occurs if the glass electrode is insufficiently cleaned between samples. It also occurs when due to absorption of CO_2 from the air or alkali is leached from the glass.
- d. Buffer solution: if the buffer solution does not have the exact pH value attributed to it, all the readings on the pH meter will be in error by the amount by which the pH of the buffer differs from its reputed value.
- e. Temperature: The temperature at which the pH measurement is carried out may differ from the temperature at which the pH value for the standard buffer is valid. This usually results in a larger uncertainty contribution from temperature.
- f. Reading the meter: figures may be rounded off
- g. The pH meter only gives an accurate measurement for solutions between -5 and 60° C. In addition, other contributing uncertainty sources include:
 - The mass of the sample- factors contributing to uncertainty are listed in step 6 of example 1.
 - The volume of the distilled water and sample contained in the beaker is subject uncertainty sources as listed in step 5 of Example 2.

These sources are listed in Figure 5-7.



Figure 5-7: Cause and effect diagram for pH measurements

Step 6: Quantifying uncertainty components

Analysis

The uncertainty contribution from analysis was found to be 0.15. Therefore the standard uncertainty is:

u_c (analysis) = 0.15

Variations of pH between locations due to the effect of heterogeneity

The effect of heterogeneity was discussed in detail under step 5 of example 1.Results showing the standard uncertainties resulting from variations between locations due to heterogeneity are listed in Table 5.12.

Dam ID	Mean	Standard Deviation	Sample Size	Standard uncertainty
Α	8.32	0.15	7	0.06
В	8.51	0.18	7	0.07
С	8.31	0.21	5	0.09
D	8.50	0.25	5	0.11
F	8.50	0.07	3	0.04
Μ	8.22	0.24	3	0.14
Ν	8.42	0.27	5	0.12
			u _c (btw Loc)	0.25

Table 5.12: Standard uncertainties for the tailing dams

Step 7: Calculating the combined uncertainty

The combined uncertainty associated with pH measurements is:

$$u_c(pH) = \sqrt{0.25^2 + 0.15^2} = 0.29$$

The spread sheet calculations are listed in Table 5.13.

Dam ID	Mean	Standard Deviation	Sample Size	Standard uncertainty
A	8.32	0.15	7	0.06
В	8.51	0.18	7	0.07
С	8.31	0.21	5	0.09
D	8.50	0.25	5	0.11
F	8.50	0.07	3	0.04
М	8.22	0.24	3	0.14
Ν	8.42	0.27	5	0.12
	8.40		u _c (btw Loc)	0.25
			u _c (analysis)	0.15
			u _c (pH)	0.29

 Table 5.13: Combined uncertainty for pH measurements.

Step 8: Calculating the expanded uncertainty

Expanded uncertainty (Table, 5.14) is calculated by multiplying the combined uncertainty value by a coverage factor of k = 1.96;

$$U_c(pH) = 1.96 \times 0.29 = 0.57$$

Dam ID	Mean	Standard Deviation	Sample Size	Standard uncertainty
А	8.32	0.15	7	0.06
В	8.51	0.18	7	0.07
С	8.31	0.21	5	0.09
D	8.50	0.25	5	0.11
F	8.50	0.07	3	0.04
М	8.22	0.24	3	0.14
Ν	8.42	0.27	5	0.12
	8.40		u _c (btw Loc)	0.25
			u _c (analysis)	0.15
			u _c (pH)	0.30
			U(pH)	0.57

 Table 5.14: Expanded uncertainty for pH measurements

Step 9: Reporting the result

Thus the pH (>7) for West Wits tailings is:

8.40 \pm 0.57 (95% confidence interval)

Where the stated uncertainty is calculated using a coverage factor of 1.96.

Uncertainties in pH measurements are displayed in Figure 5-8 below:



Figure 5-8: Uncertainties in pH measurements

5.3 SUMMARY

The chapter outlined the application of the methodology of sample size determination and a methodology of quantifying uncertainties in geochemical sampling and analyses discussed in Chapter 3. The parameters of ABA, acid potential, neutralising potential and paste pH from the case study of AngloGold Ashanti Mine, presented in Chapter 4 were used as example of how the methods can be applied. Calculations were carried out in Excel spread sheet accompanying this document. The observable trends within these obtained results are discussed in subsequent chapter.

CHAPTER 6: DISCUSSION, CONCLUSION AND RECOMMENDATIONS

6.0 INTRODUCTION

This chapter presents discussion of the findings of the study, conclusions and recommendations. The chapter sets out to investigate if uncertainty in geochemical sampling and analyses can be quantified as a function of sample size and analyses. Although this study was carried out under limited time frame and resources some trends were observed and hence some conclusions were reached.

6.1 DISCUSSION

6.1.1 SAMPLE SIZE DETERMINATION

Chapter 4 outlined the statistical analysis technique for sample size determination. The principle behind this technique is that "the size of study sample is critical to producing meaningful results" (Germishuye, Yibas, and Pulles, 2002).

The approach is based on two main assumptions, which are:

- (i) It is assumed that the selection of samples is random and unbiased; and
- (ii) The parameters used to calculate sample size are assumed to be normally distributed.

These conditions should be met before the application of the method

In order to determine a representative sample size using the approach, knowledge of the distribution of the variables is required. Additional factors on which sample size depend on include purpose of the study, confidence interval, spread of the population, homogeneity and the estimation error, among other important factors. Failure to recognize the impact of these factors on sample size may lead to poor results and hence poor decisions.

Adoption of the approach requires a good estimate of the variability and acceptable error. Unfortunately, there is no way of knowing for sure how variable the population is until after a study has been carried out (Townend, 2002). This information can be acquired from:

(i) Previous research of similar nature

If for example, paste pH has been used for geochemical research of a given mine residue deposit and the research was successful, the standard deviation value could be calculated from the data used in that specific research. This can be justified by the fact that similar populations may differ in their means but they may differ slightly in terms of spread (standard deviation) (Chou, 1989). This same concept may be used to estimate acceptable error. However it should be borne in mind that the distributions of variables for example mine residue deposits is governed by complex geochemical and other factors inherent to that specific geological and mining environment hence they vary.

(ii) **Double sampling**

Double sampling involves sampling in phases. Results obtained from the first phase, mean, variance or standard deviation are used to determine the sample size for the next phase.

(iii) Use pilot study results

If no information about the population to be studied is available, a preliminary or pilot study can be carried out. A pilot study aids in establishing procedures, understanding and protecting against things that can go wrong, and to obtain variable estimates needed in determining sample size.

6.1.1.1 Interpretation of results

The results of the exercise indicate a positive correlation between sample size and sample standard deviation. As the population variability, represented sample standard deviation s increases, sample size n follows suit. This effect can be demonstrated by comparison of Neutralization Potential (NP) results for population A to NP results for population B. Close examination of the NP measurement results for population A and B shows that results for population A vary significantly from population B. Such variations in NP measurements results may be attributed to un-uniform distribution of neutralizing elements in population A in order to

capture all patterns existing in the target area. In the case of population B which exhibited uniformity in terms of the distribution of neutralizing elements, fewer samples were required.

The results also highlight an exponential relationship between sample size (n) and acceptable error (d). The impact of estimation error on sample size can be demonstrated by taking AP results for population A. By selecting measurement results at random to create different sample sizes and calculating error associated with each as illustrated in Table 6.1, it can be seen that a low acceptable or estimation error is associated with large sample size. Figure 6-1 illustrates this relationship. The observable trend in this figure is, as d value gets smaller, sample size increases significantly.

Sample size (n)	Mean (m) (kg/t)	Standard deviation (s)	Error (d) (kg/t)
12	6.8	2.8	2.2
10	7.4	2.7	2.3
9	6.7	2.9	2.8
6	6.3	3.0	3.6
3	7.1	2.9	5.0

Table 6.1: Relationship between Sample size and Error

The results from Chapter 5 also highlight the positive correlation between sample size and confidence level. Any increase in confidence level requires that more samples are collected. The problem with this is that an increase in sample size entails high expenditure. Therefore a tradeoff among the competing factors must be reached, and one needs to work out the degree of confidence and acceptable error that they are willing to accept.

Based on the results above, for population A, 30 samples are acceptable for AP and 24 samples are acceptable for NP; for population B, 12 samples is acceptable for AP and 25 samples is acceptable for NP. However, according to our minimum sample constraint, at least 30 samples for each population should be collected.



Figure 6-1: Relationship between sample size and error

6.1.1.2 Methods of reducing the required sample size

The relationship between sample size and the various parameters, on which it depends on can be used as means of cutting on the required sample size and therefore cost. For example one may increase the allowable or acceptable error in order to minimize the number samples require and therefore cost. One major risk associated with increasing acceptable sampling error is the possibly of reducing the quality of results and representativity which may lead to poor decisions.

Decreasing material variability within population being sampled is another method that can be adopted in order to reduce the required number of samples. Methods that can be adopted include sample compositing. Mason (1992) recommends taking a large number of small increments and combining them to form a single composite sample which is submitted to the laboratory for analysis.

The use of parameters derived from a previous study of similar nature is an effective means of minimizing the required sample size. One may use the standard deviation or acceptable error from a previous research, which is equivalent to using the same value of n to conduct the similar research. It must be borne in mind that the accuracy of sample size calculation relies heavily on the accuracy of estimates of the parameters used in the calculation.

6.1.2 QUANTIFYING UNCERTAINTY

The methodology for quantifying uncertainty discussed in Chapter 5 follows existing methods for quantifying uncertainty. Such methods include those proposed by ISO, EURACHEM, UKAS and NIST, among other institutions. The approach uses a predefined model, which identifies each potential source of uncertainty, making estimates of each component, and sums them in order to make an overall estimate of uncertainty. Prior to combining uncertainty sources, sources are first converted to standard uncertainty (standard deviation) and then combined using the laws of uncertainty propagation. Unlike other techniques which use complicated procedures in estimating uncertainty, the methodology uses easy to follow steps for estimating measurement uncertainty. Furthermore, it employs simple statistical applications which are simple to comprehend.

The ability to identify individual uncertainty components and quantify their contribution facilitates comparisons and determination of the uncertainty component with the largest contribution to the overall uncertainty. Based on this information, investigations can be carried out to determine the main cause and measures may be put in place to prevent recurrences. In addition, such assessments may lead to the development of new methods designed to minimize uncertainty.

6.1.2.1 Can uncertainty be quantified as a function of sample size and analyses?

This section interrogates if uncertainty can be quantified as function of sample size and analyses. As demonstrated by in Equation 2.5, a relationship exists between uncertainty and sample size. This relationship as previously stated in the preceding chapters is exponential; an increase in either sample size or uncertainty is followed by a decrease of another parameter. This relationship can be illustrated by the following examples:

As an illustration, Acid Potential uncertainty results for West Wits tailing dams (Figure 6-2) shows that dam M has the largest standard uncertainty value compared to the rest of the dams. The dam contributed 34% towards the overall uncertainty as illustrated in below. Dam B trails behind, contributing only 22% toward overall uncertainty. Together, the two dams contributed more than 50% towards the overall uncertainty. On the other hand, dam A and D contributed 7% each.



Figure 6-2: AP uncertainty: West Wits tailings

Such variation in uncertainty contribution from these dams is attributed to the differences in sample sizes. Dams A, C, D and N with sample sizes 7, 5, 5 and 6 have uncertainty values of \pm -0.61 which is generally lower than uncertainty values for Dams F and M with sample sizes of less than 5.

Based on this relationship one has to quadruple sample size in order to reduce uncertainty by a factor of two, *ceteris paribus*. This can be illustrated as follows:

Standard deviation for dam M = 4.21If the initial 3 sample (sample size) are multiplied by 4 (quadrupling the sample size) a new sample size of 12 obtained.

Therefore standard uncertainty u_M is:

$$u_M = \frac{s}{\sqrt{n}} = \frac{4.21}{\sqrt{12}} = 1.215$$

This trend is also visible in pH results for West Wits. Figure 6-3 highlights the sample size and standard uncertainty plot for pH values from West Wits tailings dams. A comparison of dam A, B, C and F results with dam M supports the exponential relationship. A high uncertainty value is associated with small sample size.



Figure 6-3: Sample size and uncertainty for pH measurements West Wits

In addition if the original sample values listed in Appendix D are selected at random to form several sample sizes as shown in Table 6.2, the calculations show a remarkable decrease in uncertainty value with increase in sample size.

Sample size (n)	Standard dev (s)	Uncertainty (u _x)
35	0.21	0.04
28	0.23	0.04
18	0.22	0.05
13	0.22	0.06
9	0.26	0.09
6	0.30	0.12
3	0.28	0.16

 Table 6.2: Change in uncertainty at varying sample sizes

Plotting the two variables against each other, uncertainty against sample size produces a graph that gradually decreases as that number of samples increases, Figure 6-4.

It is therefore evident that a lower uncertainty value is associated with large sample sizes. The reason behind the association is that, as the sample size increases the value of the estimates moves closer to the actual value, based on the central limit theorem. Although it is difficult to tell with certainty whether the actual value lies within the stated range, the equation for determining uncertainty above, Equation 2.5, provides at least 68% chance that the actual concentration value lies somewhere within the stipulated range. "Even if the true value lies outside this range, it will probably not be far outside" (Townend, 2002).



Figure 6-4: Uncertainty vs. Sample size

On the contrary, the Neutralizing Potential for West Wits dams disobeys the trend. A comparison of dam B and dam F shows that, although the sample size for dam B is larger than dam F, dam B has a larger uncertainty value (Figure 6-5). These uncertainty results indicated a different trend independent of the known relationship between sample size and uncertainty. Such a trend can be explained in terms of selection bias as well as heterogeneity.



Figure 6-5: Sample size and standard uncertainty for NP measurements



Figure 6-6: pH-uncertainty component contributions

Figure 6-6 shows various uncertainty contributions for pH results, West Wits. The figure indicates a 19% contribution to uncertainty from analysis. This percentage is significantly smaller compared to the contributions from sources linked to sampling which have a total contribution of 81%. In the case of AP and NP of West Wits uncertainty contribution from analysis was extremely low hence they were deemed insignificant.

While both sample size and analysis are crucial aspects in geochemical measurement process and contribute to uncertainty, from the results above, it is evident that it is not sufficient to quantify uncertainty solely based on the effect of sample size and analyses alone. This shows that uncertainty in geochemical investigations is interplay of many factors and therefore cannot be quantified solely based sample size. In addition to the effect of sample size other effecting factors associated with sampling for example heterogeneity, poor sample preparation and flaws in the analytical procedure should also be considered and their effect acknowledged, thus Taylor, (1996) stated:

"The total uncertainty is the sum of the contributions from random uncertainty and systematic uncertainty. Taking more samples can reduce the random uncertainty but not the systematic component of uncertainty which is independent of the number of samples. The total uncertainty includes not only sampling uncertainty but also other types of uncertainty."

As highlighted in Chapter 2, the measurement process in geochemical investigations is a multifaceted process hence all processes involved must be considered as potential uncertainty sources.

6.2 CONCLUSIONS

Chapter 3 provided practical methods for sample size determination and quantifying uncertainties in geochemical sampling and analyses and their applicability. This section outlines the conclusions reached and challenges faced.

6.2.1 SAMPLE SIZE DETERMINATION

- Sample size determination is dependent on, the purpose of the study, cost, acceptable error (risk that one is willing to accept), variability of the population (standard deviation) and confidence level required.
- An understanding of the geology as-well as the distribution of pollutants is crucial in geochemical investigation for ERA.
- The sample size chosen determines the accuracy and representativity of the results and hence decision making. In order to achieve high precision of our estimate, a large sample size is required.
- Various parameter or variables required in calculating sample size n, Equation 3.1 may be derived from either historic datasets or conducting pilot study.
- Use of information from a previous study of similar nature is crucial especially when background information of the population to be studied is not available.
- A large sample size is required for a heterogeneous material, while fewer samples are need for homogeneous material.

6.2.2 QUANTIFYING UNCERTAINTY

Conclusions reached are as follows:

 Both sampling and analyses contributed to the overall uncertainty associated with geochemical measurements for ERA. It is therefore crucial to consider all process carried during geochemical measurement as potential uncertainty source;

- The results also demonstrated the dominant contribution of sampling and related factors toward overall uncertainty;
- Uncertainties in geochemical measurement generally decrease with increases in sample size hence it can be concluded that uncertainty can be reduced by increasing the sample size;
- While there are many sources of uncertainty, heterogeneity of the material being sampled remained the most significant source of uncertainty as observed from variation in the geochemical measurements for the different parameters measured;
- In addition it can be seen from the examples that the effect of material heterogeneity cannot be reduced by increasing sample size as highlighted by tailings dams which had a large sample size but have high uncertainty values;
- No matter how carefully geological materials are selected, homogenized, they always vary and there is always uncertainty; and
- Not all sources of uncertainty are quantifiable. Although some of these sources are difficult to quantify and some insignificant, their effect is nevertheless essential to understand and control.

Uncertainties in geochemical measurements are unavoidable but can be minimised by the following:

• Implementing Quality assurance and quality control program and adoption of good sampling practices

The quality and reliability of geochemical studies result sorely depend on the quality of field sampling and analytical procedures. A good QA/QC program should be put in place in order to minimize inconsistency and uncertainties in geochemical data. QC samples, such as external standards, duplicates and blanks, should be used in order to monitor in the processes of the sampling and the geochemical analysis.

Good sampling and analysis practices such as the use of clean gloves, decontamination of sampling devices and sampling containers are important in order to minimize contamination of the samples. Sampling techniques such as random sampling can be used to minimise sampling bias. Proper calibration of analytical instruments and sampling equipment is crucial.

• Increasing sample size

In ideal circumstances, an increase in sample size would lead to a decrease in uncertainty. This is due to the fact that as the sample size increases, the estimate of a population parameter, for example sample mean, will move closer to the true population mean, thus increasing the precision and accuracy of the estimate.

• Decreasing the particle size

Another method of minimising uncertainty is to decrease the particle size of the sample. Decreasing the particle size of the sample by grinding reduces the effect of heterogeneity which in most cases if not dealt with would result in large uncertainty.

• Use of equipment made from inert material

The equipment used during the measurement process may introduce errors through cross contamination of the sample from the fabric the equipment is made from. Such contamination may occur during sample collection, preservation, preparation (grinding and milling) and analysis. In addition to decontamination of the equipment, the use of equipment made from inert material is recommended. Such materials include glass, ceramics and stainless steel.

• Sample compositing

One commonly used method for homogenization is sample compositing. Sample compositing refers to the physical mixing of samples to form composites. It is an effective way to reduce inter-sample variance caused by the heterogeneity. Mixing samples from different locations is also an effective means of reducing cost since it is only the composite sample that is analysed. Although a good practice, compositing samples may result in loss of information and sensitivity due to dilution of the samples. Therefore great caution should be taken when deciding on whether to composite samples or not.

Improving the sampling strategy

In order to improve the precision of the results, in addition to increasing the sample size, one may work towards improving the sample design. Stratified ensures proper coverage unlike in simple random sampling where no specific order is followed. This minimizes the chances of some portions of the target being missed or left unrepresented. The methods can also be combined in order to ensure coverage and therefore sample representivity.

• Establishing the site history

The history of the site should be known before carrying out the investigations. This involves having a list of all the previous activities including the processes done. Site history may be obtained from previous work performed in the area or local authorities such as municipalities. Knowledge of the site history assists in determining the suitable sampling design and strategy which in turn helps in obtaining representative samples and therefore aids in the minimization of errors and uncertainty.

• Improving quality of measurements

Since results obtained from geochemical sampling and analyses are used for long term prediction, it is important that measurements made in the past be compatible with those made in the future. This facilitates the monitoring of changes in quality and establishing whether the quality has improved or deteriorated over time. In addition it is also essential that a more reliable method of archiving or storing data be put in place in order to monitor quality.

6.2.3 CHALLENGES

One major drawback was lack of information and resources. This was mainly attributed to the fact that the author did not undertake any practical work to fully address the requirement of the project. This is further exacerbated by limited literature on the subject and lack of appreciation the concept of "measurement uncertainty" in the field of geology and environmental sciences.

Identifying possible sources of uncertainty was a difficult task especially considering the fact that the author was not involved in undertaking the geochemical investigation discussed in the case study information presented in Chapter 4.

Although the author managed to identify possible uncertainty sources for the three examples given above, quantifying them was a challenge. Some potential sources of uncertainty were identified and listed but were not further quantified due to limited information.

As previously stated, the project was generally short, a study of this nature required practical work in order to make sound conclusions.

The assessment was general and more qualitative than quantitative due to limited information and resources. More data was required in order to accurately delineate the size of each component contributing towards uncertainty.

6.3 **RECOMMENDATIONS**

This section provides recommendations. These can be summarised as follows:

- It is recommended that in order to actually explore the practicality and applicability of the discussed method for uncertainty analysis and sample size determination, a detailed study coupled with extensive field data collection is required.
- Prior understanding of the concepts of geochemical sampling and analyses for ERA as well as sample size and uncertainty is necessary when embarking on studies of this nature.
- A study of this caliber require a proper sampling and analyses plan specifying clearly the objectives of the study and how sampling and analyses tasks will be executed. The drafted plan should be accompanied by a QA/QC program.
- Since the concept of measurement uncertainty is not well known especially among Geoscience practitioners as evidenced by limited literature on this subject, more research should be conducted in this field in order to develop a universally accepted and user friendly method for quantifying uncertainty in geochemical investigations and the Geoscience field in general.
- As indicated above, it is therefore crucial that the sample planner seeks advice where possible from experts such as statisticians, sampling technician, decision makers as well as analytical chemist during the planning phase of the project.
- It is recommended that either the combined standard uncertainty or an expanded uncertainty be reported with every result obtained from a geochemical investigation in

order to comply with requirements of ISO and other renounce bodies responsible for standardising laboratory practices.

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APPENDICES

APPENDIX A: EPA SAMPLE SIZE DETERMINATION

NUMBER OF SAMPLES REQUIRED IN ONE SIDED ONE SAMPLE t-TEST TO ACHIEVE MINIMUM DETECTABLE RELATIVE DIFFERENCE AT A CONFIDENCE LEVE (1-α) AND POWER OF (1-β), (Barth *et.al*, 1989)

Coefficient of Variation (%)	Power	Confidence Level	Minimum Detectable Relative Difference (%)					
	(%)	(%)	5	10	20	30	40	
10	95	99	66	19	7	5	4	
		95	45	13	Ś	3	3	
		90	36	10	3	2	2	
		80	26	7	2	2	1	
	90		55	16	6	5	4	
		95	36	10	4	3	2	
		90	28	8	3	2	2	
		80	19	5	2	1	1	
	80	99	43	13	6	4	4	
		95	27	8	3	3	2	
		90	19	6	2	2	2	
		80	12	4	2	1	1	
15	95	99	145	39	12	7	5	
		9 5	99	26	8	5	3	
		90	78	21	6	3	3	
		80	57	15	4	2	2	
	90	99	120	32	11	6	- 5	
		95	79	21	7	4	3	
		90	60	16	5	3	2	
		80	41	11	3	2	ī	
	80	99	94	26	9	6	- 5	
		95	58	16	5	3	3	
		90	42	11	4	2	2	
		80	26	7	2	2	1	
20	95	99	256	66	19	10	7	
		95	175	45	13	9	5	
		90	138	36	10	5	3	
		80	100	26	7	4	2	
	90	99	211	55	16	9	6	
		95	139	36	10	6	4	
		90	107	28	8	4	3	
		80	73	19	5	3	2	
	80	99	164	43	13	8	6	
		9 5	101	27	8	5	3	
		90	73	19	6	3	2	
		80	46	12	4	2	2	
Coefficient of Variation	Power	Confidence Level		Minir Relat	num Detec ive Differe (%)	table		
--------------------------------	-------	---------------------	-------------	----------------	---------------------------------	----------	--------	
(%)	(%)	(%) г	6	10		20		
	05	00	207	10	20	30	40	
2	93	99	277	102	20	14	9 6	
		90	216	55	15		5	
		90 90	155	40	13	<i>'</i>	2	
	90		320		- 24	12		
	30	95	272	70	10	12	6	
		90	166	42	12	6	4	
		80	114	20	8	4	3	
	80	99	254	66	19	10	7	
	00	95	156	41	12	6	4	
		90	114	30	8	4	3	
		80	72	19	5	3	ž	
30	95	99	571	145	39	19	12	
50		95	391	99	26	13		
		90	310	78	21	10	6	
		80	223	57	15	7	4	
	90	99	472	120	32	16	11	
		95	310	79	21	10	7	
		90	238	61	16	. 8	5	
		80	163	41	11	5	3	
	80	99	364	84	26	13	9	
		95	224	58	16	8	5	
		90	164	42	11	6	4	
		80	103	26	7	4	2	
35	95	99	775	196	42	25	15	
		95	532	134	35	17	10	
		90	421	106	28	13	8	
		80	304	77	20	9	6	
	90	99	641	163	43	21	13	
		95	421	107	28	14	8	
		90	3 23	82	21	10	6	
		80	222	56	15	7	4	
	80	99	495	126	34	17	11	
		95	305	78	21	10	7	
		90	222	57	15	7	5	
		80	140	36	10	5	3	

APPENDIX B: QUALITY CONTROL SAMPLES (Van Ee, Blum, and Starks, 1990)

Procedure	Description
Field Blank	A sample container filled with distilled deionized (DDI) Water, exposed during sampling and then analysed to detect accidental or incidental contamination.
Sample Bank Rinsate	A sample (last rinse of DDI water) of DDI water passed over the sample preparation apparatus, after cleaning, to check for residual contamination.
Field Rinsate	A sample (last rinse of DDI water) of DDI water passed over the sampling apparatus after cleaning to check for residual contamination.
Calibration Check Standard	A standard material to check instrument calibration.
Spiked Sample	A separate aliquot of the soil sample having an appropriate standard reference material added to check for soil and extract matrix effects on recovery.
Total Recoverable	A second aliquot of the sample which is analysed by a more rigorous method to check the efficacy of the protocol method.
Laboratory Control Standard	A sample of a soil standard carried through the analytical procedure to determine overall method bias.
Triplicate Samples (Splits)	The prepared sample is split into three portions to provide blind duplicates for the analytical laboratory and a third replicate for the referee laboratory to determine interlaboratory precision.
Duplicate Sample	An additional sample taken near the field sample to determine total within-batch measurement error.
Field Audit	A sample of well-characterized soil that is taken into the field with the sampling crew, sent through the sample bank to the laboratory with the field samples to detect bias in the entire measurement process and to determine batch to batch variability.
External Laboratory Audit	A sample of well-characterized soil sent directly to the laboratory for analysis. The analyte concentrations are unknown to the laboratory. This type of sample is used to estimate laboratory bias and batch-to- batch variability. It may also be used for external quality control of the laboratory.

APPENDIX C: SAMPLE RECORDS PRIOR TO ANALYSES

Table C1. Sampling records of tailings dams at West Wits - continued

		Field D	ata					Laboratory	Anal	yses	_			_		_
Complex	Dam	Sample No	Field colour description	Depth (m)	Colour	Paste pH	Composite for ABA	Composite for detailed analysis	XRD	XRF	м	SA	AI	wi	PSD	CL
		A-ISA A-ISB A-ISC A-ISD	grey with yellow inclusions grey wet clay grey in color grey	0.5 1 1.5 2	Charcoai Dark grey Grey Dark grey	7.6 8.1 8.5 8.2	A-15	R								
		A-2DA A-2DB A-2DC A-2DD A-2DD	grey (yellow inc) stiff grey silt grey grey to light grey	0.5 1 1.5 2	Dark grey Grey Dark grey Dark grey	8.4 8.0 5.4 8.5 7.7	A-2D									
New North Complex		A-3SA A-3SB A-3SC A-3SD A-3SE	silt grey grey grey moist grey moist grey	0.5 1 1.5 2 2.5	Dark grey Dark grey Charcoal Dark grey Dark grey	8.5 8.8 8.0 8.6 8.7	A-38									
	7B	A-4SA A-4SB A-4SC A-4SD A-4SE	grey grey grey grey grey	0.5 1 1.5 2 2.5	Dark grey Dark grey Charcoal Dark grey Grey	8.6 8.7 8.4 8.2 8.3	A-45	A-COMP	1	0	1	0	1	1	1	
		A-5SA A-5SB A-5SC A-5SD	grey grey grey grey	0.5 1 1.5 2	Dark grey Dark grey Dark greenish yellow Dark grey	8.4 8.5 8.3 8.5	A-SS									
		A-55E A-65 A A-65 B A-65 C A-65 D A-65 E	grey moist grey grey grey moist grey moist grey	2.5 0.5 1 1.5 2 2.5	Charcoal Grey Dark grey Charcoal Dark grey	8.5 8.5 8.2 8.1 8.2	A-6S	-								
		A-7SA A-7SB A-7SC A-7SD	grey grey grey	0.5 1 1.5 2	Greenish yellow Dark grey Dark grey Deep	8.2 8.3 8.0 8.3	A-75									

M - Microscopy; SA - Surface area; Al = Aquategia + ICP-MS; Wi = Water extract + ICP-MS; PSD = Particle size distribution; CL = column leach

		Field D	ata		1000			Laboratory	Anal	yses		_				_
Complex	Dam	Sample No	Field colour description	Depth (m)	Colour	Paste pH	Composite for ABA	Composite for detailed analysis	XRD	XRF	м	SA	Ai	wı	PSD	CI
		B-1SA	grey	0.5	Dark grey	8.2										
		B-1SB	grey	I	Dark grey	8.1										
		-	Constanting of the second	16	Greenish		B.IS			1.1						
		B-15C	tight grey to	1.5	yenow	0.4	0.10	8		1.1	1.3			1. 1		
		B-1SD	yellowish	2	Dark grey	8.3										
		B-1SE	silty grey	2.5	Dark grey	8.1			3		0.1			i i		1
					Dark											
		B 36 A	Kabi area	0.5	groenish	85				1.0	11			1.3		
		B-25R	orev	1	Dark grey	8.4	B-25									1
	X 6	B-2SC	grev	1.5	Dark grey	8.6		1							1.5	1
		B-2SD	moist grey	2	Dark grey	8.9			I .							
		B-2SE	moist grey	2.5	Dark grey	8.7			1	10	1.1			1.1		
		B-3SA	grey	0.5	Dark grey	8.4		1	1					1.1	10.1	1
	1 1	B-3SB	grey	1	Charcoal	8.5	1.65229.5	1		1.8						L
		B-3SC	moist grey	1.5	Dark grey	8.7	B-3S	1	1	1.18			1			1
		B-3SD	moist grey	2	Dark grey	8.6						1	I			1
		B-3SE	grey	2.5	Dark grey	9.0		4								L
		B-4SA	grey	0.5	Charcoal	8,6		1	1			1	1			
	N 3	B-4SB	grey	1	Dark grey	8.8		1	1			1				L
	111	B-4SC	light grey	1.5	Dark grey	8.6	8-45								100	L
	1.1	B-4SD	moist grey	2	Dark grey	8.6		1	1							L
	1 8	B-4SE	moist grey	2.5	Dark grey	8.6		-								
10000		B-5DA	grey	0.5	Grey	8.1			L .	1						
New	74	B-SDB	moist clay	1	Dark grey	9.1		B-COMP	1	0	1	1	1	1	1	
Complea		B-SDC	light grey to	15	Dark grey	8.5				1			1			L
	1.3	10000	1	1.44	Greenish	1			1				1		1.1	
		B-5DD	grey	2	yellow	8.5	1		1	1						
		B-5DE	moist grey	2.5	Dark grey	8.4		-	1					1		L
		B-6SA	grey	0.5	Dark grey	8.7				1						
	11 1	B-6SB	silty grey	1	Dark grey	8.8	2									L
		B-6SC	grey	1.5	Dark grey	8.4	B-6S		1							1
		B.6SD	orev	2	vellow	8.4		1		1						L
	14 8	B-6SE	ercy	2.5	Dark grey	8.7	Law			1						
	10	B-7SA	silty erey	0.5	Dark grey	8.5	1	1	1							
			1.500.00.0	10331/	Dark	2005			1							ł
			(1 8	greenish	1				1			1	1		L
		B-/58	grey	1 1 6	Greek	83	B-75		1				1	1	1	
		D-750	grey and	1.0	Citey	0.0					1		1	1		Ł
		B-7SD	moist	2	Dark grey	8.1	1	1		1	1		1	1		Ŀ
		B-7SE	grey	2.5	Dark grey	8.1	-			1	1		1		1	
		Coperation of			Light	ante:	1									1
		B-7SF		3	brown	8.1	1			1						
		B 760		25	Light	84			1							1
		5-730	20	3.5	Dark	0.4	B-75-2		1							
		Pastaro		355	greenish	1 mars	1									
		B-7SH		4	yellow	8.7		1								
					greenish	S. Same	1									
		B.151		4.5	vellow	86	1	1		1			12-	1		

Table C1. Sampling records of tailings dams at West Wits - continued

M = Microscopy; SA = Surface area; AI = Aquaregia + ICP-MS; WI = Water extract + ICP-MS; PSD = Particle size distribution; CL = column leach

		Field D	ata			a		Laboratory	Anal	yses						
Complex	Dam	Sample No	Field colour description	Depth (m)	Colour	Paste pH	Composite for ABA	Composite for detailed analysis	XRD	XRF	м	SA	AI	wı	PSD	CL
		C-5DA C-5DB C-5DC C-5DD C-5DE C-5DF C-5DG C-5DG	silty grey grey to light grey grey grey and moist grey	0.5 1 1.5 2 2.5 3 3.5 4	Dark grey ND Dark grey Medium brown Charcoal	8.2 ND 8.5 7.6 7.5 8.4 8.2 8.1	C-5D									
		C-3SA C-3SB C-3SC C-3SD C-3SE	grey silty grey grey grey grey	0.5 1 1.5 2 2.5	ND ND Dark grey ND Dark grey	ND ND 8.6 ND 8.5	C-3S									
New North Complex	5B	C-4DA C-4DB C-4DC C-4DD C-4DD	grey grey light grey to yellowish grey moist grey	0.5 1 1.5 2 2.5	Light brown Brown ND ND Charcoal	4.5 4.4 ND ND 8.5	C-4D	C-COMP	1	0	1	0	1	1	1	
		C-2SA C-2SB C-2SC C-2SD C-2SE	light grey grey grey moist grey grey	0.5 1 1.5 2 2.5	Dark grey Dark grey ND Dark grey C15-4	8.4 8.4 ND 8.2 7.8	C-28									
		C-1SA C-1SB C-1SC C-1SD C-1SE	light grey moist and grey grey grey grey	0.5 1 1.5 2 2.5	Dark grey Dark grey ND ND Dark grey	8.2 8.3 ND ND 8.2	C-1S									

Table C1. Sampling records of tailings dams at West Wits - continued

M = Microscopy; SA = Surface area; AI = Aquaregia + ICP-MS; WI = Water extract + ICP-MS; PSD = Particle size distribution; CL = column leach

		Field D	ata					Laboratory	Analyses					-	
Complex	Dam	Sample No	Field colour description	Depth (m)	Colour	Paste pH	Composite for ABA	Composite for detailed analysis	XRDXRF	м	SA	AI	wı	PSD	CL
		D-1SA D-1SB D-1SC D-1SD D-1SD D-1SE	BLeà BLeà BLeà BLeà	0.5 1 1.5 2 2.5	Dark grey Dark grey ND ND Dark grey	8.2 8.0 ND ND 8.6	D-15	-2							
	2	D-3SA D-3SB D-3SC D-3SD D-3SE	grey light grey to yellowish grey grey	0.5 1 1.5 2 2.5	Dark grey Dark grey Dark grey Deep yellow Charcoal	8.3 8.3 8.3 8.4 8.5	D-3S								
New North Complex	SA	D-2SA D-2SB D-2SC D-2SD D-2SE	grey grey light grey to yellowish silt grey	0.5 1 1.5 2 2.5	Dark grey Dark grey ND ND Dark grey	8.3 8.4 ND ND 8.3	D-2S	D-COMP	1 0	1	0	1	1	1	
		D-5DA D-5DB D-5DC D-5DD D-5DE	द्वारुप्र इरुप्र इरुप्र	0.5 1 1.5 2 2.5	Dark grey Dark grey Dark grey ND Dark grey	8.8 8.9 8.7 ND 8.6	D-5D								
		D-4DA D-4DB D-4DC D-4DD D-4DE	light grey dark grey light grey grey	0.5 1 1.5 2 2.5	Dark grey Dark grey ND ND Dark grey	8.7 8.8 ND ND 8.9	D-4D								

Table C1. Sampling records of tailings dams at West Wits - continued

M = Microscopy; SA = Surface area; AI = Aquaregia + ICP-MS; WI = Water extract + ICP-MS; PSD = Particle size distribution; CL = column leach

		Field D	ata					Laboratory	Anal	yses			_			
Complex	Dam	Sample No	Field colour description	Depth (m)	Colour	Paste pH	Composite for ABA	Composite for detailed analysis	XRD	XRF	м	SA	AI	wı	PSD	c
		E-ISA	grey	0.5	Deep yellow Medium	7.5										
		E-1SB	grey	1	brown	7.9										
	1 1	E-1SC	light grey	1.5	Yellow	7.9	E-IS									
		E LOS	grey to	-	Deep	0.7										
		6-150	grev to	-	Deep	0.2										L
		E-ISE	yellowish	2.5	yellow	8.1										
	4	E-2S A	grey	0.5	Deep yellow Deep	8.3										
		E-25 B	light grey	1	yellow	8.3										
		E AR C	Vield and	12	Deep	8.2	E-28		1							
		E-23 C	light grey to	1.2	yenow	0.3			1							L
		E-2S D	yellowish	2	Yellow	8.4			ł.							
		F-25 F	light grey to	2.5	Deep	8.4			1							
	\vdash	Lo and to	Tenomian	Ard I	Dark	0.1	-	1	1	1.1						
		F-1D A	light grey	0.5	beige	8,4			1							
		F-IDB	light grey	1	Yellow	8.2			1		1					
		EIDC	light ores	15	Deep	8.2			1			1.1				
		1 - 467 50	infline fires	1146	Greenish					2						
	1	F-IDD		2	yellow	8.3			1							Ľ
		F-IDE	yellowish	2.5	Yellow	8.4										L
		F-IDE	yellowish	2.7	ND	ND		f -	1							L
		EIDE	light yellow	3	Vellow	85	F-ID-I		1							
Old	1 1	1-101	wins Brea	1.	Light	0.5	ALCONCE:	E-F-G-	1.	h., -	L.	1.1		100		1
omplex		F-IDG	grey	3.5	yellow	8.5		COMP	1.	1.	Ľ.	1	Ľ.	1.1		
-124280-020		FaiDH	OTPY	4	Deep	8.4			1							L
		F-IDI	grey	4.5	ND	ND			1							L
		100-000	grey &	11275	10555.5	1925										Ľ
		F-1D1	yellowish	4.9	ND	ND			1							Ŀ
		F-IDJ	irrev	5	Deep	8.6			1							L
		F-IDK	grey	55	ND	ND			1							L
	30				Dark			1	1							
		F-1D K	dark grey colour	6	greenish yellow Dark	8,5										
		E IDI	13-221	22	greenish	1	F-1D-2	1	ł.							L
		F-IDL	grey	0.0	Dark	5.5		1							1	L
		F-ID M	grey	7	beige	8.7			1						1	Ľ
		697742755		1	Medium	-			1			1.8				
		F-IDN		7.5	brown	8.5		-	1			1 3				L
		F-4S A	silt grey light grey to	0.5	yellow	8.3										
		F-4SB	colour	1	Yellow	8.4										1
					Deep		F-45									
		F-48 C	grey	1.5	yellow	8.4	1.000.000.0									
		F-4S D	grey	2	vellow Dark	8,5										
	1	EASE		3.5	greenish	9.5			1	1						

Table C1. Sampling records of tailings dams at West Wits

M = Microscopy, SA = Surface area; AI = Aquaregra + ICP-MS, WI = Water extract + ICP-MS; PSD = Particle size distribution, CL = column leach

		Field D	ats		Lungar		North Constant	Laboratory	Analy	vses		(1) (1) (2)				
Complex	Dam	Sample No	Field colour description	Depth (m)	Colour	Paste pH	Composite for ABA	Composite for detailed analysis	XRD	XRF	м	SA	AI	wī	PSD	CL
		F-25 A	light grey light	0.5	Light brown	8.2			Π							
		F-25 B	□ellowish colour	1	Light brown Light	8.3	F-25	14								
		F-2S C	light grey	1.5	brown	8.3									1.1	
		F-25 D	grey silt	2	Yellow	8.3										
	1	E.38 E	orev silt	25	Deep	84										
	3	F-15 A	light grey	0.5	Yellow	82		1					Ċ.	<u>1</u> .		
		1-33 A	ugur grey	4.4	Deep	1.200										
		F-3S B	grey	1	yellow	8.3										
		F-35 C	light grey	1.5	Greenish yellow	8.3	F-3S									
		F-3S D	light grey	2	Greenish yellow	8.4										
		F-3S E	moist)	2.5	yellow	8.4	1 1								1.2	
Old		G-2SA	grey	0.5	Greenish yellow Dark greenish	7.9		F-F-G-								
North Complex		G-2SB	grey	1	yellow Dark	8.3		COMP	1	I	1	1	1	1	1	1
		G-2SC	light grey	1.5	yellow Dark	8.3	G-25-1									
		G-2SD	grey	2	greenish yellow Dark	8.4									R I	
		G-2SE	grey	2.5	yellow	8.3		1. I								
	6	G-2SF		3	Light brown Medium	8.8	6.25.2									
	1	G-2SG		3.5	brown	9.0	0-20-2									
		G-2SG		3.8	brown	9.0										
		G-ISA	light grey	0.5	Deep yellow	8.1										
		G-ISB	yellowish	1	yellow	8.3		li l								1
		G-1SC	light grey to yellowish	1.5	Deep yellow	8.2							1			
		G-1SD	light grey to yellowish	2	Greenish yellow	8.3										
8		G-1SE	light grey to yellowish	2.5	Deep yellow	8.3	G-15	-								

Table C1. Sampling records of tailings dams at West Wits - continued

M - Microscopy, SA = Serface area; AI = Aquaregis + ICP-MS; WI = Water extract + ICP-MS; PSD = Particle size distribution; CL = column leach

		Field D	ata					Laboratory	Ana	yses				_		
Complex	Dam	Sample No	Field colour description	Depth (m)	Colour	Paste pH	Composite for ABA	Composite for detailed analysis	XRD	XRF	м	SA	AI	wi	PSD	CI
		H-ISA	yellow color with orange inclusions	0.5	Yellow	4.3										
i d		H-ISB	yellow	1	Deep yellow	4.2					1					
	1	H-1SC	yellowish grey with	1.5	yellow	8.1										
	•	H-ISD	yellowish inclusions	2	Deep yellow	3.1	H-1\$		1 2 2 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2							
		H-ISE	yellowish silt	2.5	Yellow	3.0	6	0								
	а С	H-ISF		3	yellow Dark	4.3					100					
		H-ISG		3.5	greenish yellow	3.7			88							
		H -1\$I		4	greenish yellow	8.3										
		1-15A	yellow silt	0.5	Greenish yellow	4.2										
		I-1SB	yellow silt	1	Greenish yellow Dark	4.2	I-1S-1									
Old		I-ISC	yellowish color	1.5	greenish yellow	3.7		H-I-J_top-	,				4		San I	
Complex		I-1SD	yellowish color with grey inclusions	2	Deep yellow	7.8		comp			ġ.	ľ		1.50		
		I-1SE	moist light yellowish	2.5	Medium brown	8.2	1-S-2									
		I-1SF		3	brown Medium	8.2			13							
		1-1SG	vellowieh	3.5	brown	8.2										
		J-1SA	and moist yellowish	0.5	yellow	4.2		3								
		J-1SB	mixture yellow and	1	yellow	4.6	J-18-1									
	1	J-1SC	dark grey mixture	1.5	Greenish yellow	5.9										
		1-ISD	dark grey yeilow and	2	Greenish yellow	8.1		5								10.00
		J-1SE	dark grey mixture	2.5	Greenish yellow	8.3	J-1S-2									
		J-1SF		3	Light brown	8.2										

Table C1. Sampling records of tailings dams at West Wits - continued

		Field D	ata		0.00-000			Laboratory	Asal	yses	_					_
Complex	Dam	Sample No	Field colour description	Depth (m)	Colour	Paste pH	Composite for ABA	Composite for detailed analysis	XRD	XRF	м	SA	AI	wı	PSD	CI.
		N-ISA	grey	0.5	Dark grey	8.5										
		N-ISB	stiff grey color	1	Dark groy Dark	8.7										
	Low	N-ISC	grey	1.5	greenish yellow	8.4	N-IS	N-COMP	1	0	1	1	1	1	1	
	er	N-ISD	grey grey with light vellowish	2	Charcoal	8.2	1210 845	Are Art Ale			10000					
		N-ISE	inclusions	2.5	Brown	8.2	1				_					_
	51	M-4DA	silty grey	0.5	Dark grey	7.7								11		
	11	M-4DB	silty grey	1	Dark grey	7.8						6				
		M-4DC	grey	1.5	Dark grey	72	M-4D	0 0					6.5	1.		
		M-4DD	moist and grey in color	2	Dark grey	8,6										
		M-4DE	grey	2.5	Charcoal	8.6								1.3		
South		M-5SA	grey with yellow inclusions	0.5	Dark grey	8.0							11///2-5			
Complex		M-5SB	grey	1	Dark grey	7.9	1995532									
	11	M-5SC	moist grey	1.5	Dark grey	8.5	M-5S									
	Upp	M-5SD	dark grey and stiff erev and	2	Dark grey	83		M-COMP	1	0	ī	0	1	ĩ	1	
	er	M-5SE	moist	2.5	Charcoal	8.5		0		n,	1	10000	(any	1.0	1.825	
		M-6SA	silty grey	0,5	Dark grey	8.0										
		M-6SB	grey	1	Dark grey	8.6										
		M-6SC	silty grey	1.5	Dark grey	8.3				E C						1
	1	M-6SD	moist grey	2	Charcoal	8.2				E d						
		M-6SE	moist grey	2.5	Dark grey Dark greenish	8.6	M-6S									
		M-6SF		3	yellow Dark	8.4										
		M-6SG		3.5	yellow	8.7										
		M-6SH		4	yellow	8.8										

Table C1. Sampling records of tailings dams at West Wits - continued

_		Field D	ata					Laboratory	Anal	vses			_	_	_	_
Complex	Dam	Sample No	Field colour description	Depth (m)	Colour	Paste pH	Composite for ABA	Composite for detailed analysis	XRD	XRF	м	SA	AI	wı	PSD	CI
		N-7DA	grey with brownish staff		Light bown	7.5										
		N-7DB	yellowish color		Light bown	7.7		108								
		N-7DC	grey clay		bown	8.2										
		N-7DD	very miost arev clay		Medium	8.4										
		10.055	6-92		Medium		N-7D									
	100	N-7DE	grey	÷	Dark	8.2					. 1					Ł
		N-7DF		3	greenish yellow	8.0										
		N-7DG		3.5	Greenish yellow Dark	7.6										
		N-7DH		4	greenish yellow	8.1										
	11				Dark											
		N-6SA	grey	0.5	greenish yellow Dark	8.6								-		
		N-6SB	grey	1	greenish yellow	8.7					1					
		N-65C	grey and very moist	1.5	Greenish yellow	8.4	N-6S									
South Complex	Low er	N-6SD	grey	2	Greenish yellow Dark	8.2		N-COMP	1	0	1	1	1	i	ì	
	11	N-6SE	grey	2.5	yellow	8.5	1									
		N-55 A	grey with light yellowish inclusion	0.5	Medium brown	8.6					2 2 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1					
		N-5S B	silty grey	1	ND	ND	N-55									
		N-5S C	grey	1.5	brown	8.3					1.1		81			
		N-5S D	grey	2	Medium	8.2										
		N 20 P	very moist		Medium											
		N-35 E	grey	2.5	Medium	8.9					1		83			
		N-3DA	grey	0.5	brown	8.7										
		N-3DB	grey	1	C28-5	8.5					6 U		1.3			
		N-3DC	grey	1.5	Brown	8.8	N-3D				1					
		N-3DD	grey	2	brown	8.4					1		1.4			
		N-3DE	grey	2.5	Brown	8.1					1					
		N-2SA	stiff grey clay	0.5	Brown	8.7										
		N-2SB	grey clay	1	brown	8.9	Hat									
		N-2SC	grey and moist	1.5	Brown	8.3	Pe-25				1.1					
		N-2SD	grev	2	E17-7	8.6										
		N-2SF	grou	25	Brown	8.8							11			

Table C1. Sampling records of tailings dams at West Wits - continued

Location	Dam	ID	Paste pH	AP(kg/t)	C- NP(kg/t)	Latitude	Longitude
		A-1S	8.12	16.91	3.50	-26.4327	27.3482
		A-2D	8.15	14.81	6.25	-26.4324	27.3511
		A-3S	8.52	16.78	6.00	-26.4344	27.3499
	Α	A-4S	8.44	13.13	5.52	-26.4365	27.3485
		A-5D	8.40	15.14	7.00	-26.4371	27.3508
		A-6S	8.32	15.78	4.50	-26.4398	27.3491
		A-7S	8.28	16.91	5.75	-26.4391	27.3523
		B-1S	8.18	14.66	5.75	-26.4318	27.3544
		B-2D	8.62	17.22	6.75	-26.4316	27.3563
		B-3S	8.64	10.38	7.25	-26.4343	27.3550
	В	B-4S	8.64	14.41	6.75	-26.4343	27.3574
		B-5D	8.52	14.22	5.50	-26.4363	27.3564
		B-6S	8.60	23.50	6.75	-26.4386	27.3563
		B-7S	8.35	18.66	0.00	-26.4381	27.3591
		C-1S	8.23	14.53	5.54	-26.4298	27.3610
		C-2S	8.20	14.84	4.75	-26.4317	27.3609
West Wite Tailings	С	C-3S	8.55	14.22	5.50	-26.4329	27.3627
Dams		C-4D	8.50	17.09	9.00	-26.4307	27.3267
		C-5D	8.07	14.03	5.00	-26.4303	27.3646
		D-1S	8.26	18.78	5.75	-26.4277	27.3654
		D-2S	8.33	15.72	7.50	-26.4274	27.3632
	D	D-3S	8.36	17.5	4.25	-26.4256	27.3649
		D-4D	8.80	17.59	8.75	-26.4261	27.3665
		D-5D	8.75	16.69	4.50	-26.4245	27.3682
		F-1S	8.43	10.66	6.91	-26.4228	27.3763
	F	F-2S	8.30	14.00	5.25	-26.4256	27.3773
		F-3D	8.32	12.59	6.75	-26.4235	27.3801
		M-4D	7.98	17.63	8.00	-26.4531	27.4053
	Μ	M-5S	8.24	22.28	10.25	-26.4508	27.4073
		M-6S	8.45	26.03	10.00	-26.4505	27.4053
		N-2S	8.66	20.5	7.50	-26.4557	27.4096
		N-3S	8.50	21.59	9.75	-26.4542	27.4120
	Ν	N-5S	8.50	23.88	7.50	-26.4516	27.4144
		N-6S	8.48	22.38	7.00	-26.4557	27.4170
		N-7D	7.96	20.38	9.75	-26.4497	27.4159
			-				
	Α	VR-A-1S	3.98	11.22	0.00	-26.9353	26.6773
		VR-A-4D	6.10	31.53	2.50	-26.9440	26.6739
		VR-B-1S	3.99	31.09	1.25	-26.9384	26.6792
		VR-B-2S	4.01	39.69	1.25	-26.9384	26.6830
		VR-B-3S	4.07	16.39	2.50	-26.9415	26.6792
Vaal River Tailing	В	VR-B-4D	3.79	16.22	0.00	-26.9415	26.6830
Dams		VR-B-5D	3.72	16.94	0.00	-26.9415	26.6863
		VR-B-6S	4.04	14.13	5.75	-26.9446	26.6757
		VR-B-7S	4.26	17.84	3.50	-26.9446	26.6792
		VR-B-8S	3.87	10.47	6.00	-26.9446	26.6830
	C	VR-C-1D	6.83	20.19	5.00	-26.9322	26.6863
	C	VR-C-2S	6.28	27.38	5.50	-26.9322	26.6900
		VR-C-3S	6.55	26.16	6.50	-26.9352	26.6830

APPENDIX D: INPUT DATA (ABA) USED IN VARIOUS EXAMPLES

	VR-C-4S	6.72	32.75	5.50	-26.9352	26.6863
	VR-C-5D	6.85	36.25	7.75	-26.9362	26.6892
	VR-C-6S	7.18	27.97	5.00	-26.9384	26.6863
	VR-C-7S	7.17	25.03	7.00	-26.9384	26.6900
	VR-C-8S	7.20	36.84	5.25	-26.9415	26.6900
F	VR-F-1S	7.4	41.25	6.75	-26.9290	26.7077
	VR-F-2S	7.45	26.75	7.25	-26.9330	26.7108
	VR-F-4S	7.63	24.72	4.00	-26.9322	26.7108
	VR-F-5D	7.65	25.78	4.50	-26.9331	26.7084
	VR-F-6S	7.78	25.03	4.50	-26.9352	26.7045
	VR-F-8S	8.05	31.41	6.25	-26.9352	26.7108
	VR-F-9D	7.87	23.97	3.75	-26.9372	26.7108
	VR-F-					
	10S	7.76	21.47	4.75	-26.9384	26.7076