Investigation of the Variability in the Results of the NZ Vibrating Hammer Compaction Test

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Dedicated to my parents Huda Abu Neama and Wasfy Shahin
ABSTRACT

The New Zealand vibrating hammer compaction test procedure has been proven to provide inconsistent results. As supported by the Opus investigation, repeatability and reproducibility values of the New Zealand Standard for the vibrating hammer compaction test method are higher than those values found in standards both in America and the United Kingdom.

The research examined the variability in the vibrating hammer compaction test results. Two approaches were implemented to achieve a sound and scientific understanding of the variability associated with the test results. Firstly, repeated testing of the vibrating hammer compaction test was conducted under constant conditions to determine the natural variability of the test. Secondly, X-ray diffraction tests were conducted to verify the homogeneity of the source aggregate being used for testing.

Results have confirmed that the variability is significantly large considering the tests have been conducted under constant test conditions. Under these conditions, factors that could possibly affect the reliability of the test results have been kept the same throughout testing. The natural variability in the source aggregate explains approximately 30% of the observed variation in the Opus Interlaboratory study. As evident in the results, the amount of compactive effort applied to the sample during compaction determines the degree of Dry Density achieved.

X-ray diffraction results have shown that there are some differences within the aggregate in terms of physical properties and mineral constituents. However, it is unknown to what extent, if any, these differences contributed to the variation in the vibrating hammer compaction test results.

Future research is recommended in areas such as the amount of contribution that segregation and degradation has on the variation of the results. Additional testing should be done on aggregates passing the 19 mm sieve to observe whether oversized particles have the effect of interlocking and interfering with compaction.
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# Abbreviations and Acronyms

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<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>AADT</td>
<td>Annual Average Daily Traffic</td>
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<tr>
<td>AM</td>
<td>Asphalt Mix</td>
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<tr>
<td>ASTM</td>
<td>American Society for testing and materials</td>
</tr>
<tr>
<td>BSI</td>
<td>British Standard Institution</td>
</tr>
<tr>
<td>CBR</td>
<td>California Bearing Ratio</td>
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<tr>
<td>CETANZ</td>
<td>Civil Engineering Testing Association of New Zealand</td>
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<tr>
<td>CI</td>
<td>Clay Index</td>
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<tr>
<td>CoV</td>
<td>Coefficient of Variation</td>
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<tr>
<td>CS</td>
<td>Chip Seal</td>
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<tr>
<td>DD</td>
<td>Dry Density</td>
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<tr>
<td>ESA</td>
<td>Equivalent Standard Axles</td>
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<tr>
<td>MDD</td>
<td>Maximum Dry Density</td>
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<td>NZ</td>
<td>New Zealand</td>
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<td>NZS</td>
<td>New Zealand Standard</td>
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<td>NZTA</td>
<td>New Zealand Transport Agency</td>
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<tr>
<td>OWC</td>
<td>Optimum Water Content</td>
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<td>PI</td>
<td>Plasticity Index</td>
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<td>PSD</td>
<td>Particle Size Distribution</td>
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<td>SE</td>
<td>Sand Equivalent</td>
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<tr>
<td>TNZ</td>
<td>Transit New Zealand</td>
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<tr>
<td>UK</td>
<td>United Kingdom</td>
</tr>
<tr>
<td>UoA</td>
<td>University of Auckland</td>
</tr>
<tr>
<td>USA</td>
<td>United States of America</td>
</tr>
<tr>
<td>WC</td>
<td>Water Content</td>
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<tr>
<td>XRD</td>
<td>X-ray Diffraction</td>
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<td>ZAV</td>
<td>Zero Air Voids</td>
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Chapter 1. INTRODUCTION

1.1 Research Motivation

Vibratory hammer compaction is a relatively new approach in aggregate laboratory testing. Originally, the test had been developed specifically for the compaction of granular aggregates. The reason for the development of this test was that the Standard Proctor compaction method was deemed an ineffective approach to compacting materials of a granular nature. The issue with the Standard Proctor test method was that the granular material would often break down and displace when struck by the impact rammer (Farrar, 2000; Felt, 1968). However, the method which was adopted by the New Zealand Standards authority (NZS) for the vibrating hammer compaction test was also deemed inadequate due to its inconsistent nature. Consequently, Opus Ltd conducted an inter-laboratory investigation to identify the degree of variability in the vibrating hammer compaction test results where the study found repeatability and reproducibility values of the New Zealand vibrating hammer test method to be significantly higher than those specified in the British and American Standards. Therefore, it is evident by the repeatability and reproducibility values in the Opus investigation that the New Zealand test produces inconsistent results and therefore may need revision and/or minor alterations (New Zealand Standards, 1986b; Opus International Consultants Limited, 2008). The variation and unreliability in the results of the test is not only apparent in New Zealand (NZ) but also by countries abroad (British Standards Institution, 1980; Opus International Consultants Limited, 2008).

The purpose of laboratory compaction is to determine the Maximum Dry Density (MDD) and corresponding Optimum Water Content (OWC) so that these values could be targeted in field compaction. However, previous research has shown that the laboratory vibrating hammer compaction test adopted in NZ produces unreliable and significantly variable results. Therefore, research is essential to identify the reasons for the variability and where possible to better control those variables.

This research, kindly funded by the New Zealand Transport Authority (NZTA) through a Roading New Zealand project, will undertake laboratory testing facilitated by Stevensons
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Ltd. It will consider one source aggregate material (Greywacke) to determine the minimum statistical variation associated with the test method adopted in NZ.

1.2 Research Objectives

The primary objective of this research was to determine a sound and scientific understanding of the variability in the results of the NZ Vibrating Hammer Compaction Test (New Zealand Standards, 1986b) through rigorous testing and provide conclusions and recommendations based on the results obtained. Essentially, there are two factors which can influence the vibrating hammer compaction test results. The first factor is the natural variability in the properties and mineral constituents of the aggregate, where samples taken from the same aggregate source tested under the same test conditions can yield different results. The second factor being the test conditions of the experiment (such as hammer type and age, mould size and technician experience) (Wilson & Shamseldin, 2010). The research focuses on identifying the natural variability in the test procedure. Thus, although aggregate quality control and natural aggregate variability will be briefly tested, a large portion of the research will be focused on conducting the vibrating hammer compaction test to note its variability.

1.3 Research Methodology Overview

The research methodology was split into three main phases.

Phase 1  **Review of Literature** – A comprehensive review of appropriate literature available.

Phase 2  **Testing** – The testing phase was further split into three stages, namely:

- **Quality Control Testing** – Strength and durability tests were carried out on the source aggregate to ensure a certain level of quality was maintained.
- **Vibrating Hammer Compaction Testing** – This is the main stage of the testing phase. Conducting the vibrating hammer test a sufficient amount of times to obtain viable data to provide statistical analysis on the results.
• X-ray Diffraction testing – Perform X-ray Diffraction tests to determine any variability in the properties and mineral composition of the source aggregate.

Phase 3 **Results Analysis and Conclusions** – Evaluation and analysis of results obtained from the testing phase to provide conclusions and recommendations.

### 1.4 Organisation of the Thesis

The structure of the thesis follows from a literature review of existing studies through to testing, analysis and presentation of results obtained. This Section provides an overview of the subsequent chapters of the thesis.

Chapter 2 provides an introduction to the theory of compaction. The chapter begins by defining the pavement structure followed by an outline of the compaction process including the importance of compaction. The compaction curve is then described, where the two variables which make up the curve, the Water Content and Dry Density, are explained. The types of laboratory compaction used today are then reviewed. Field compaction is then briefly outlined. Specifications relating to the compaction of unbound pavement layers in New Zealand are then discussed, followed by a summary of the chapter.

Chapter 3 describes the typical aggregates used in New Zealand for the basecourse layer in pavement construction. A brief history of the types of aggregates sourced for basecourse is initially introduced in the chapter. Aggregate quality control tests are then described followed by a review of the X-ray Diffraction analysis used for identifying the variability in the properties and mineral constituents of the source aggregate. The effect of an aggregates’ grading on its performance is then discussed, followed by a summary of the chapter.

Chapter 4 reviews and describes the possible factors which could contribute to the variation of the vibrating hammer compaction test results. It also describes the ruggedness test, which is used in identifying factors which have a major effect on the variation of a test experiment. A case study which is relevant to this research is then discussed. The chapter is then summarised.
Chapter 5 provides a review of the various vibrating hammer compaction test standard methodologies available, including those from the United States of America (USA) and United Kingdom (UK) test standards. In this chapter, the New Zealand Standard for the vibrating hammer compaction test method is compared to international standards which have adopted a slightly different approach to using the vibrating hammer for compaction testing. Evaluation of each section of the standard is conducted to find any major differences between these standards.

Chapter 6 describes the research methodology that has been adopted to successfully and efficiently perform the research. The research tasks are first identified, followed by a detailed description of the methodology of each of the three different types of tests performed.

Chapter 7 presents and discusses the results obtained from the various tests conducted in the research. It begins by a brief introduction to the chapter, followed by discussion of the quality control test results obtained for the source aggregate. Chapter 7 then discusses the statistically analysed results of the vibrating hammer compaction tests. And finally the results of the X-ray diffraction tests are discussed.

Chapter 8 provides the conclusions drawn from the research and the recommendations for important issues within the vibrating hammer compaction topic which require further research.
Chapter 2. **COMPACTION OF PAVEMENT MATERIALS**

2.1 Pavement Structure

2.1.1 Introduction

Pavements are primarily designed to provide a stiff surface that serves a specific transportation need. The surface stresses by the tyre pressure on the pavement must be transferred to the subgrade (underlying materials beneath the pavement) with the least amount of unrecoverable deformation strain. The function of a pavement is thus to reduce and distribute the surface stresses to an acceptable level at the subgrade. This basic function must be conducted under different seasonal and environmental conditions and without permanent deformation or cracking. The initiation of any of these distress conditions would reduce the functionality of the pavement (Huang, 1993; Thom, 2008).

The key mechanism used in the transfer and reduction of loads in a pavement is the use of layers of decreasing strength from top to bottom. The different layers distribute the load, thus, decreasing its intensity with depth, and resulting in subgrade stresses being much less than stresses on the surface. These lower stresses at the subgrade ensure it does not undergo excessive deformations (Papagiannakis & Masad, 2008).

There are two basic pavement types but with a number of variations to each type. The two types used in the roading industry are flexible and rigid pavements. Theoretically, flexible pavements transfer uniform stresses throughout the layers but deflections are non uniform. Conversely for a rigid pavement, the transferred stresses are not uniform but the deflections are uniform. In practice, the stress and deflection distributions throughout the flexible and rigid pavements depend on the relative stiffness of the top layers to the underlying granular layers (Huang, 1993; Thom, 2008).

2.1.2 Flexible Pavements

The cross-section of a generic flexible pavement is shown in Figure 2-1. The layered system consists of high grade materials on the top where the intensity of stress is high and
inferior materials at the bottom where the stress intensity is low. Starting from the top, the pavement consists of a wearing surface, basecourse, subbase and subgrade (Thom, 2008).

![Cross-section of Flexible Pavement](image)

**Figure 2-1: Cross-section of Flexible Pavement (Papagiannakis & Masad, 2008)**

The wearing surface is the top course of a flexible pavement, sometimes called the surface course. Within NZ, the wearing surface is usually made from an Asphalt mix (AM) layer or Chipseal (CS). The CS surface is better known as the ‘surface dressing’ in many parts of the world such as the United Kingdom. It is considered the dominant surfacing type in NZ and comprises of a uniformly sized stone chips embedded onto a 1 to 2mm layer of sprayed bitumen (Transit New Zealand et al., 2005). The wearing surface must be tough to resist distortion under traffic and provide a smooth and skid-resistant riding surface. It must also be impermeable to protect the entire pavement and subgrade from the weakening effect of water (Thom, 2008).

The basecourse layer sits directly below the surface layer, as shown in Figure 2-1, and helps provide additional load distribution where the imposed load on the surface course is spread over a bigger area of the road base. Because it lies directly below the wearing surface layer, it experiences the second highest intensity of stresses. The depth of the basecourse layer largely depends on the California Bearing Ratio (CBR), a measure of the bearing strength, of the layers below it (Subbase and subgrade layers). It also depends on the traffic characteristics of the road being built such as the Annual Average Daily Traffic (AADT) and Equivalent Standard Axles (ESA). The basecourse layers are usually constructed in NZ from unbound aggregates. As will be discussed in Section 2.2.1, unbound aggregates are essentially free-bound particles with no cohesion. Due to this fact, compaction becomes a key role in achieving the desired strength of the basecourse
layer as it compresses the free-bound particles tightly together and provides confinement as shown in Figure 2-3. In New Zealand, typical basecourse aggregates require a 95% minimum relative compaction to achieve the desired strength. Thus, aggregates used for the basecourse layer must be compacted effectively to prolong the pavement’s design life and to avoid pavement damage such as rutting (Henning, 2008a; Hoffman, 2008; Transit New Zealand, 2005b).

The basecourse layer also serves to contribute to frost resistance and shrinkage and swelling control. If water reaches the subgrade, it may cause it to shrink or swell; the basecourse layer serves as a surcharge load imposed on top of the subgrade reducing the amount of swelling or shrinkage taking place. The basecourse layer also contributes to drainage. As water enters the pavement structure through cracks and joints, an open-graded basecourse layer can carry this water away to the road side (Huang, 1993).

The subbase is the layer of material beneath the basecourse and usually consists of larger sized crushed aggregate. This material has better engineering properties, such as modulus of elasticity, than the subgrade layer thus resulting in a higher bearing capacity; however, it is lower in quality than the basecourse layer above it. It is important to note that the use of two layers, basecourse and subbase, consisting of aggregates is for economic reasons. The basecourse is the stiffer layer because it uses higher quality aggregates and because the stress intensity decreases further down the layers, lower quality aggregates can be used thus forming the subbase (Papagiannakis & Masad, 2008; Thom, 2008).

The final layer in the pavement is the subgrade and can consist of the local in-situ soil or compacted fill imported from different locations. If the in-situ soil is used, the top layer of soil is usually scarified, sometimes undercut and the replaced fill is then compacted to the desired density and Optimum Water Content (OWC) (Huang, 1993; Thom, 2008).

The thickness of each layer varies with the type of axle loading, available materials and expected pavement design life. The expected design life is the number of years the pavement is expected to provide an adequate service with the expected ESA before it requires asset intervention and a major rehabilitation is required.
2.1.3 Rigid Pavements

In contrast to flexible pavements, rigid pavements can be placed either directly on the prepared subgrade or on a single layer of granular or stabilised material. Figure 2-2 shows a cross-section of a generic rigid pavement structure. As is the case with flexible pavements, the subgrade is often scarified, blended and compacted to the desired density. Above this layer is the basecourse which has the same function and is from the same material as the basecourse layer in the flexible pavement. The top layer is constructed from either unreinforced or reinforced concrete and acts as an impervious layer that reduces water ingress. It also provides a skid-resistant smooth surface on which vehicles can operate. The disadvantage of using concrete is that it cracks under thermal stresses. To counter the effect of shrinkage, transverse contraction joints are built into the pavement. Load transfer devices, such as dowel bars, are placed in the joints to minimise deflections and reduce stresses near the edges of the slabs (Papagiannakis & Masad, 2008).

![Figure 2-2: Cross-section of Rigid Pavement (Papagiannakis & Masad, 2008)](image)

2.2 Compaction Process

The compaction process (also known as densification) is defined as the removal of air voids from the material by application of mechanical energy with zero or minimal change in the water content. Compaction should not be confused with soil consolidation, which is void reduction in saturated soils over a length of time due to the expulsion of water. Compaction modifies and enhances the engineering properties of the material. Compacted materials display higher strength, lower permeability and lower compressibility (Drnevich...
et al., 2007). Carson (2004) defines compaction as “the method of mechanically increasing the density of soil”. Carson (2004) best explains this by the use of a diagram as shown in Figure 2-3.

![Figure 2-3: Soil Structure Before and After Compaction (Carson, 2004)](image)

The expulsion of air during the compaction process causes both the density and unit weight of the material to increase. Although similar, these two parameters are different and are commonly confused between each other. Density is the amount of mass per unit volume and is expressed as kg/m³ (kilograms per meter cubed) using the symbol \( \rho \). On the other hand, unit weight refers to the measure of weight per unit volume and is expressed as kN/m³ (kilo Newtons per meter cubed) using the symbol \( \gamma \) (Drnevich et al., 2007).

The degree of compaction is expressed in terms of the dry density (\( \rho_d \)). The reason for expressing the degree of compaction by the dry density and not the bulk density (\( \rho \)) is that the bulk density contains water, and water offers no strength. Hence, the performance of a compacted material can be best expressed as the amount of dry soil solids per unit volume (Drnevich et al., 2007). The formula used to calculate the dry Density is as follows:

\[
\rho_d = \frac{100\rho}{(100 + w)}
\]

where:

- \( \rho_d \) = Dry Density of Soil (t/m³)
- \( \rho \) = Bulk Density of Soil (t/m³)
- \( w \) = Water Content expressed as a decimal number
However, it is important to note that Dry Density (DD) is not a direct measure of material properties, that is, different material with the same or similar DD will not exhibit the same or similar engineering properties (Drnevich et al., 2007).

### 2.2.1 Importance of Compaction

Compaction is an imperative process during the pavement construction phase to ensuring the desired performance levels from the pavement are achieved (Christopher et al., 2006). Budhu (2000) mentions that compaction is one of the most popular techniques of aggregate property improvement. Some of these improvements include:

- The increase in shear strength of the aggregate.
- The decrease in compressibility; reducing the potential of excessive long term settlement of fills and soils.
- The reduction in permeability; restraining flow of water through the compacted basecourse layer.
- The general decrease in void ratio; this helps prevent water from being withheld by the basecourse layer, thus maintaining strength and stiffness properties.
- Achieving a state of increased unit weight.

These aggregate property enhancements are vital to prolonging the life of the pavement. To achieve an optimum degree of compaction, Maximum Dry Density (MDD) must be reached. The MDD largely depends on the Water Content (WC) of the aggregate. Thus, preliminary laboratory compaction tests are conducted on the sample to obtain the MDD value and corresponding Optimum Water Content (OWC) value. Subsequent to obtaining these values from laboratory testing, optimum field compaction can be targeted according to these values.

In addition, New Zealand pavements are fatigue structures that are predominantly (>90%) unbound granular aggregates. These types of aggregates and pavements rely heavily upon reaching design compaction levels to be able to withstand an adequate design life of repeated traffic cycles. Thus, achieving the desired compaction levels is immensely important to getting the expected design life out of the total pavement structure (Black, 2009; Henning, 2008a).
Unbound aggregates are a skeleton of particles with principally no cohesion to keep these particles stuck together. Thus, there must be a significantly high degree of confinement of these particles to achieve the desired stiffness of the unbound granular layer. Confinement of these particles is provided by the horizontal stresses that arise as a result of compaction and subsequent traffic loading (New Zealand Institute of Highway Technology, 2000).

Contractors are required to determine the laboratory Maximum Dry Density (MDD) and the Optimum Water Content (OWC) by using the Vibrating Hammer Compaction method specified in the NZS 4402: Test 4.1.3. This test will set the target dry density that the contractor must achieve when compaction occurs on the field (Frobel & Moulding, 2006; Transit New Zealand, 2005b).

### 2.3 Compaction Curve

#### 2.3.1 Introduction

The Water Content (WC) at which a material is compacted defines the degree of compaction achieved. The WC is usually expressed in percentage and is defined as the ratio of the mass of the water to the mass of the solids in an aggregate.

Defined as the mass per unit volume, Dry Density (DD) is considered an important property of engineering stabilisation. The symbol used to denote Dry Density is $\rho_d$ and is expressed in terms of kg/m³ (or t/m³). The degree of density is defined by how loosely or closely the particles are packed.

#### 2.3.2 Water Content – Dry Density Relationship

Subsequent to obtaining laboratory compaction results, the OWC at which the MDD occurs is determined from the graph of plotted results. Theoretically, the results obtained from the laboratory vibrating hammer compaction test produces a bell-shaped curve. The peak of the curve is defined as the Maximum Dry Density (MDD), the WC at which this MDD occurs is better known as the Optimum Water Content (OWC).

The Dry Density and Water Content relationship varies depending on the type of material being compacted. According to the British Standard of the vibrating hammer compaction test “BS EN 13286 – part 4 Test methods for laboratory reference density and water content – Vibrating hammer” aggregate compaction curves can take three different forms.
as shown in Figure 2-4. The first two forms are discussed in the next two Sections to come where the convex downwards curve appears when cohesive material is compacted, the convex upwards curve shown in Figure 2-4 occurs when cohesionless material is compacted. Because this research is testing a cohesionless unbound granular aggregate, more focus will be given to the curve produced by cohesionless material. The flat curve, where the material dry density is independent and insensitive to the water content, is rare and does not occur often and hence will not be discussed (BS EN 13286 - 4, 2003).

![Different Forms of Compaction Curves](image)

**Figure 2-4: Different Forms of Compaction Curves (BS EN 13286 - 4, 2003)**

**Cohesive Material**

An optimal degree of compaction will only be reached when the Optimum Water Content (OWC) has been achieved. This is because compactive forces are resisted by the friction between the material’s particles. The water available in the voids helps reduce this friction. For every aggregate type there exists an Optimum Water Content and a Maximum Dry Density as illustrated in Figure 2-5. It however, must be noted that the curve shown below is a typical curve based on a compacted cohesive material. As will be discussed in the next Section, cohesionless graded materials exhibit a slightly different curve.

From Figure 2-5, it can be seen that water contents less than the OWC provide an increase in dry density if the water content is increased, this is where the increase in water acts as a lubricant and helps reduce friction between particles. However, at water contents above the OWC, the increase in water prevents the expulsion of air and/or water and hence a decrease in dry density is observed (Drnevich et al., 2007).
Investigation of the Variability in the Results of the NZ Vibrating Hammer Compaction Test

Figure 2-5: Typical Compaction Curve for Cohesive Material (Drnevich et al., 2007)

The Zero Air Voids (ZAV) line shown in Figure 2-5 is a property of the aggregate that is dependent on the Solid Density $q_s$ of the soil. The ZAV represents the soil when it is fully saturated, i.e., the voids in the soil are completely filled with water (no air). It can be calculated using the following formula (New Zealand Standards, 1986a):

$$Q_{ZAV} = \frac{1}{\frac{1}{q_s} + \frac{w}{100q_w}} W$$  \hspace{1cm} (2.2)

where:

- $Q_{ZAV}$ = Dry Density at saturation (at Zero Air Voids) (t/m$^3$)
- $q_w$ = Density of Water (t/m$^3$)
- $q_s$ = Solid Density of Soil Particles (t/m$^3$)
- $w$ = Water Content (%)

Cohesionless Material

Most of the granular materials behave differently during compaction and hence exhibit an unusual compaction curve than that observed for cohesive soils. For these materials, the Maximum Dry Density (MDD) is at either dry conditions (0% WC) or near saturation, while lower dry density values are obtained at intermediate WC’s (Bergeson et al., 1998; Forssblad, 1981; Hilf, 1991; Parsons, 1992; Pike, 1972). An example of a typical granular soil compaction curve is shown in Figure 2-6.
Forssblad (1981) explains that effective compaction at oven-dry conditions works well for graded materials with as much as 30% fines. In contrast, Brandl (2001) reported that even though the Maximum Dry Density may occur at oven-dry conditions, the MDD should nevertheless be chosen at its other peak where the Optimum Water Content lies near saturation, he supports his argument by explaining that if the material was to be compacted at oven-dry conditions it would favour long-term grain rearrangement and hence differential deformation.

On the other hand, a compacted layer of material where the MDD and corresponding OWC were achieved on the saturation curve can exhibit ‘sponge-like’ behaviour. Hence, Brandl’s (2001) report suggests that if the MDD and corresponding OWC lie on the saturation curve, a slightly lower Water content should be chosen as an OWC to prevent the ‘sponge-like’ behaviour of the layer.

![Compaction Curve for Granular Material](image)

**Figure 2-6: Typical Compaction Curve for Granular Material (Drnevich et al., 2007)**

The low dry densities that occur in the compaction curve of granular material at intermediate Water Contents (as seen in Figure 2-6) is explained by a phenomenon known as bulking (Hilf, 1991); the range of Water Contents at which this phenomenon occurs are called bulking Water Contents. Capillary stresses which exist under low Water Contents cause bulking. Tension stresses are formed in partially saturated material where a curved surface develops at the air-water boundary. The tension stresses (and water
available within the material) help keep the particles in place and resist the compactive effort applied on the sample (USBR, 1990). Thus, this phenomenon only occurs at intermediate Water Contents because the tensile stresses do not exist at completely dry conditions, and begin to reduce as the aggregate starts to saturate; allowing for higher compactive effort and therefore effective compaction (Bergeson et al., 1998).

Engineering judgement should be exercised when selecting the OWC, to achieve the appropriate MDD. This is because, as mentioned earlier, sometimes a curve would indicate that, at completely dry conditions, the OWC and corresponding MDD have been achieved, however these values should not be chosen due to the fact that a compacted soil that is too dry favours long term grain rearrangement and hence differential deformation. Also on the other hand, if the optimum appears to be on the saturation line (ZAV) then the optimum should be selected a little below this value, because at saturation level the compacted layer would be rather spongy.

2.3.3 Compaction Suitability

A various number of laboratory compaction tests have been developed to suit the different aggregate types that exist. These different aggregate types require different methods of compaction in order to be compacted correctly to an optimum level. This is because different aggregates behave differently under the application of loads. The main two material types are cohesionless and cohesive materials. This research deals with the compaction of cohesionless granular graded materials. Thus, it is important to analyse the different methods of compaction and determine the advantages and/or disadvantages of these methods particularly when compacting granular graded materials.

2.4 Laboratory Compaction

This Section discusses the different methods of compaction and discusses the advantages and/or disadvantages of each technique relative to granular soils. Compaction processes in a laboratory can be classified under five categories (Luxford, 1975), namely:

- Impact Compaction
- Static Compaction
- Kneading Compaction
- Vibratory Compaction, and
2.4.1 Impact Compaction

Developed originally by Proctor in 1933 to aid in earth dam stabilisation (Proctor, 1933), impact compaction is one of the most widely used compaction techniques today. The Proctor test (named after its developer) basically involves dropping a hammer of a known weight from a set height onto the sample. The test is relatively easy and cheap to perform, however some disadvantages exist within the test (Luxford, 1975).

Felt (1968) reports that the impact compaction test is not suitable for cohesionless material containing sands and/or course graded crushed stones or similar material possessing inherent angular stability. Felt (1968) further explains that the test is unworkable with cohesionless material due to a number of factors; firstly, because there is no confinement on the sample, the cohesionless particles easily displace when struck by the rammer. Second, the impact force is considered small and limited when compacting cohesionless soils. Third, the mould restraint and friction between the particles oppose the requirement of the particles packing closer together by moving horizontally.

Furthermore, repetitive ramming degrades the sample. Reports by Hoover, Kumar and Best (1970), and Dunlap (1966) confirm that impact compaction does not produce satisfactory results when compacting cohesionless granular materials due to degradation of the sample. Strikes produced by the impact hammer tend to “break down” the course granular material otherwise known as degradation (Farrar, 2000; Felt, 1968). Degradation tends to increase and become more of a problem as the percentage of coarse aggregates is increased in graded materials (Johnson & Sallberg, 1960).

An article by Sherwood (1970) showed that reproducibility of the impact compaction test is unsatisfactory for compaction control purposes. However, the degree of reproducibility was considered acceptable for design purposes. Hence, it could be argued that this is unacceptable for testing purposes.

In addition, another problem in using the Proctor test on granular materials is that it is very difficult to get a flat surface of the specimen by levelling the top of the mould for testing measurements (Strohm et al., 1967).
A new procedure was later adopted by the New Zealand Standards (NZS 4402 Test 4.1.2 Heavy compaction) to account for the improvements in technology in field compaction equipment. The ‘Heavy’ compaction test which is also usually referred to as the ‘Modified Proctor’ compaction test is largely based on the standard Proctor test with a few minor changes such as a heavier rammer that is dropped from a greater height and the compaction is done in five layers rather than three as in the standard Proctor test (Hausmann, 1990). The differences between the two tests are provided in Table 2-1.

**Table 2-1: Comparison of the Standard and Modified Proctor Compaction Test Methods – Reproduced from (Brandl, 2001)**

<table>
<thead>
<tr>
<th>Detail</th>
<th>Standard Compaction</th>
<th>Modified Compaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mould Volume (cm³)</td>
<td>1000</td>
<td>1000</td>
</tr>
<tr>
<td>Mould diameter (mm)</td>
<td>105</td>
<td>105</td>
</tr>
<tr>
<td>Mould height (mm)</td>
<td>115.5</td>
<td>115.5</td>
</tr>
<tr>
<td>Rammer diameter (mm)</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>Rammer drop height (mm)</td>
<td>300</td>
<td>450</td>
</tr>
<tr>
<td>Rammer mass (kg)</td>
<td>2.7</td>
<td>4.9</td>
</tr>
<tr>
<td>Number of layers</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>Blows per layer</td>
<td>25</td>
<td>25</td>
</tr>
<tr>
<td>Energy input (kJ/m³)</td>
<td>596</td>
<td>2703</td>
</tr>
</tbody>
</table>

The modified compaction test was still deemed unsatisfactory for use on granular materials. Due to the same problem mentioned earlier for the standard Proctor, the cohesionless nature of unbound granular material causes the particles to simply displace under each strike of the rammer. The granular materials move under each successive rammer blow, however not much actual compaction or densification occurs. Thus, in order for effective compaction of granular material to take place, confinement of the particles is vital to prevent the displacement of particles (Luxford, 1975).

In addition to the standard and modified Proctor methods of compaction, another impact compaction method, known as the Marshall Hammer Compaction Test was introduced specifically for compacting dense graded aggregates, although this method was never actually implemented. In short, the Marshall Hammer test uses a slightly bigger hammer (in diameter) and a smaller mould than the standard and modified proctor tests. These
changes allow for increased confinement for the sample preventing it (to a certain extent) from displacing freely and actually compacting. This test was found to produce Maximum Dry Densities of graded aggregates which were much more achievable in the field than the standard and modified compaction methods (Roberts, 1976).

2.4.2 Static Compaction

Static compaction involves compressing a pre-weighed specimen in a cylindrical mould by placing it in a compression testing machine. Compression forces are progressively increased until the Maximum Dry Density is reached (Hausmann, 1990). However, because of the way the test is done, particle orientation is likely to be different from that achieved in the field since the field technique of compaction is not simulated in any way in this test. A report conducted by Johnson and Sallberg (1962) showed a few factors that influence the test, these include:

- In order to prevent segregation graded granular aggregates must be placed into the mould very carefully.
- Long periods of static load application onto the sample results in expulsion of water producing a Maximum Dry Density at unrealistic water contents.

2.4.3 Kneading Compaction

Inspired by the kneading action produced by the sheepsfoot roller (see Section 2.5) in field compaction, the kneading compaction laboratory test was developed. Similar to the sheepsfoot roller in the field, the laboratory compaction efforts on the sample are gradually built up then gradually released.

The development of an automatic kneading compactor by Dodd and Dunlop (1971) showed that the kneading compaction method is not suitable for the compaction of granular materials such as sand; it was observed that surface deformation occurred under the compactor foot and that compaction results were unsatisfactory. Significantly higher Dry Density (DD) values were achieved at lower Water Contents (WC) by vibratory compaction.
2.4.4 Vibratory Compaction

Compaction of granular soils and aggregates is often confounded by the lack of an appropriate test method. Other methods of compaction such as impact compaction have been deemed unsuitable to compact these types of soils due to their cohesionless nature. Thus, new methods such as vibratory compaction have been developed in an attempt to better compact these types of soils. Since field compaction equipment use vibrations to compact aggregates effectively, vibratory compaction yields a better correlation between field and laboratory results. Compaction by vibratory means can be achieved in two ways as illustrated in Figure 2-7.

![Figure 2-7: Methods of Vibratory Compaction](image)

The difference between the two procedures is that the vibrating table method (Figure 2-7b) places a static surcharge load on top of the sample contained within a mould and applies continuous vertical vibrations from the bottom. In contrast, the vibrating hammer method (Figure 2-7a) utilizes a vibrating hammer which is placed on top of the sample contained within the mould and applies vibratory forces for a specific set time from the top. The vibrating hammer method is considered better due to the fact that it better simulates field compaction (Drnevich et al., 2007). Since this research is concerned with the vibrating hammer compaction test, focus will be given to this method.

Initially designed for heavy duty demolition work, vibrating hammers were later utilised for soil compaction. Being considered the most suitable for the compaction of granular
soils, the vibrating hammer compaction method provides the required confinement granular aggregates need in order for effective compaction. In this method, compaction occurs by vibration, which means the specimen is compacted thoroughly throughout its depth (Luxford, 1975).

Since its development, extensive research has been carried out on the vibrating hammer test not to only ensure its validity, but to also seek its acceptance by international standards Authorities such as the American Society for Testing and Materials (ASTM), British Standards Institution (BSI) and New Zealand Standards (NZS).

The first to perform a thorough investigation and research on the use of the vibrating hammer compaction test was Parsons (1964). Where the focus was on five different factors affecting the test:

- Type of hammer and tamper size used
- Magnitude of static load applied
- Period of operation of hammer
- Size and shape of mould, and
- Voltage supplied to hammer.

The results in Parsons’ (1964) investigation led to the adoption of the vibratory hammer compaction test by the British Standards Institution “BS EN 13286 – 4:2003 Unbound and hydraulically bound mixtures – Part 4: Test methods for laboratory reference density and water content – Vibrating Hammer” (BS EN 13286 - 4, 2003; Luxford, 1975). ASTM approved the vibrating hammer test in December 2007 as a result of research conducted by Drnevich, Porchaska and Evans, where a comprehensive investigation was conducted by them at the University of Purdue in Indiana regarding the vibrating hammer test and its reliability (IHS, 2010). Thus, the test is slowly being recognised worldwide, however due to some of its uncertainties (such as its repeatability and reproducibility) some parts of the world (such as Australia) continue to use the Standard Proctor compaction method for granular materials.

2.4.5 Gyratory Compaction

Gyratory compaction is the result of research and studies conducted by the U.S Army corps of Engineers and the Texas Transportation Institute (Ping et al., 2003). The
Compaction method has shown great promise for compacting granular base course aggregates. Because sample preparation is done in one layer, segregation and stratification are prevented (Luxford, 1975).

There are conflicting reports on the degradation that occurs in gyratory compaction. Some authors have observed very little or no degradation in the sample whilst others have stated significant amounts of degradation has been observed (Luxford, 1975).

### 2.5 Field Compaction

It is important to understand the process and theory behind field compaction in order to devise a satisfactory laboratory experimental compaction procedure which replicates and realistically represents actual compaction in the field.

Because of the advancements and improvements in field compaction equipment in more recent years, the MDD and corresponding OWC can be reached in the field at much lower values than those obtained in the laboratory (as shown in Figure 2-8). The improved heavy rollers could achieve MDD at much lower Water Contents than those specified in the laboratory. However, this could lead to degradation of the material being compacted and so it is imperative that very heavy compactors are not used in the field. A balanced relationship between laboratory compaction and field compaction should be established.

As can be seen in Figure 2-9, the compactive effort greatly affects the MDD achieved, thus, reasonable field equipment that are not too powerful should be specified depending on the compactive effort used in the laboratory.

The NZ specification for the compaction of unbound pavement layers, which will be discussed in greater detail in Section 2.6, specifies that the Maximum Dry Density for field compaction is the higher of the maximum laboratory dry density and the pleatau density at Optimum Water Content (OWC). This requirement ensures that the maximum laboratory dry density is the minimum requirement achieved on the field and therefore prevents the need for the TNZ B/2 specification to specify restrictions on the allowable weight of compaction rollers to be used on the field (Transit New Zealand, 2005a). The TNZ B/2 also specifies “A maximum number of tonnes mass per meter of roll width has been retained to give some guidance on when rollers are likely to significantly change the gradation of TNZ M/4 basecourse materials” (Transit New Zealand, 2005a)
Figure 2-8: Effect of Compaction Effort on the Compaction Curve – Laboratory and Field (Ping et al., 2003)

Figure 2-9: Effect of Compaction Effort on the Compaction Curve - Different Hammers (Ping et al., 2003)

Figure 2-10 displays two different types of rollers used in field compaction. The sheepsfoot roller (Figure 2-10a) as the name suggests resembles a “sheepsfoot” and is used first during the compaction process. It compacts the material from the bottom up.
The smooth roller (Figure 2-10b) on the other hand, compacts the layer from the top down and is considered the final stage of compaction which is usually applied to the layer after the sheepfoot roller.

![Sheepsfoot Roller and Smooth Roller](image)

**Figure 2-10: Field Compaction Rollers**

### 2.6 Compaction of Unbound Pavement Layers in NZ

#### 2.6.1 Introduction

The New Zealand Transport Agency (NZTA), formerly Transit New Zealand (TNZ), has implemented specifications for the construction of unbound granular pavement layers in New Zealand. The applicable specification for compaction is referred to as the TNZ B/2 and includes a guideline for compaction of these types of pavement layers.

#### 2.6.2 Compaction Criteria

TNZ B/2 specifies that compaction should be undertaken in the minimum number of passes of compaction field equipment. The contractor’s responsibility involves conducting the New Zealand laboratory vibrating hammer compaction test on a sample representative of the material used in the field to find the MDD and corresponding OWC (Transit New Zealand, 2005b).

Once the MDD and corresponding OWC values are known, compaction can then take place. During the field compaction process, TNZ B/2 specifies that the contractor monitor Dry Density levels by undertaking the Plateau density test using a nuclear density meter.
shown in Figure 2-11. This meter is a sophisticated piece of equipment which measures the Dry Density at the location it is placed.

![Nuclear Density Meter](image)

**Figure 2-11: Nuclear Density Meter**

In order to achieve a satisfactory degree of compaction, the TNZ B/2 standard specifies a requirement which the compacted pavement layer must comply with in order to be deemed an acceptable level of compaction. These requirements are given in Table 2-2.

<table>
<thead>
<tr>
<th>Parameter Values</th>
<th>Basecourse Pavement Layer, % of MDD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean Value</td>
<td>≥98</td>
</tr>
<tr>
<td>Minimum Value</td>
<td>≥95</td>
</tr>
</tbody>
</table>

These values given in Table 2-2 are believed to be achievable in the field. However, this largely depends on the power produced by the field compaction equipment. Thus, it is vital that the contractor utilises appropriate equipment depending on the nature and size of the pavement being compacted. The choice is usually dependent on the strength of aggregate, and layer thickness (Transit New Zealand, 2005a).
2.7 Summary

This chapter discussed the theory of compaction and its vital role in achieving stable pavement structures with an adequate design life. Pavement structures in New Zealand are predominantly constructed from unbound granular material. These types of material rely heavily upon reaching optimum compaction to be able to withstand repeated traffic loadings. Thus, effective compaction of these types of materials is imperative to constructing a pavement structure that will meet the expected design life criterion.

Different types of materials behave differently under the application of load due to their natural characteristics and properties. To effectively compact a particular type of material, an optimum method should be selected which best suits that type of material. The vibrating hammer compaction test was deemed the best approach to compacting cohesionless granular material. Although this research is concerned with the laboratory vibrating hammer compaction test, field compaction was discussed due to the fact that grasping an understanding of field compaction is imperative to devising a method in the laboratory which realistically reflects compaction on the field.

The NZTA has implemented a specification for the construction of unbound granular layers in pavement structures, which includes a guideline to effective compaction. The specification includes compaction criteria that must be met to ensure proper compaction levels have been met. In addition to these requirements, basecourse material used for the construction of these unbound granular layers must also comply with a set of requirements specified in the TNZ M/4 specification. The next chapter discusses these basecourse aggregates in greater detail.
Chapter 3. NEW ZEALAND BASECOURSE AGGREGATES

3.1 Introduction

Most aggregates in New Zealand are sourced from Greywacke and Volcanic rocks. Within these two categories falls a range of different rock types, each having a unique matrix of properties which are defined by the minerals and other constituents contained (and their arrangement) in these rocks. The roading industry in New Zealand is a major consumer of aggregates, using approximately 24 million tonnes per year on New Zealand’s roading network. Failure to meet MDD reduces the pavement’s stability and strength, consequently reducing its life expectancy leading to expensive rehabilitation works. Because aggregates are a non-renewable source, they must be used sparingly and effectively. There are very limited high quality aggregates available in New Zealand with many already exhausted. Thus, optimum compaction levels must be met to avoid the inefficient use of these valuable limited resources (Black, 2009).

New Zealand aggregates are only about 150 million years old; this is considered geologically young as opposed to aggregates internationally, where in some areas (such as North America and Australia), Greywacke rocks are about 1 billion years old. An aggregate’s age can influence its homogeneity, geological constituents, physical strength and response to application of loading. Relatively young aggregates have not been exposed to geological metamorphism, where pressure and heating over long geological time periods modify the rock source physical and chemical properties. Therefore, these aggregates tend to be much more heterogeneous than significantly older aggregates. Geologically heterogeneous aggregates are inconsistent; the aggregate does not behave in the same manner throughout. Thus, heterogeneous rocks are undesirable for civil engineering purposes due to their unpredictable behaviour (Black, 2009).

For example a sample from a 1000 kg batch of heterogeneous aggregate may have an MDD of 2.24 kg/m³ in a laboratory test, however because the laboratory test only uses about a 5kg portion, the MDD value of 2.24 kg/m³ may not be representative of the full
1000kg batch. Furthermore, if the aggregate is heterogeneous, then different parts of the quarry can produce significantly different results.

### 3.2 Aggregate Property Tests

#### 3.2.1 TNZ Basecourse Specifications

TNZ M/4 refers to a standard specification for basecourse use in flexible granular pavements. The specification states that the aggregate shall be of high quality to be used on NZ road pavements as a road base (Henning, 2008a). Typical aggregates used on heavily trafficked roads such as State Highways are usually crushed from sources accepted as a regional basecourse aggregate (Transit New Zealand, 2006b).

The aggregate being used in this research is predominately Greywacke and is sourced from an accepted region as stated in the TNZ M/4 acceptable regional basecourse table (Transit New Zealand, 2006b). It is quarried and crushed (All Passing 40 mm) by Stevensons Ltd, south of Auckland.

Aggregates must be tested using standardised procedures to maintain quality of the aggregates being used as basecourse layers in New Zealand pavements. Quality control is sometimes referred to as quality assurance and is defined as providing a product or service that will satisfy certain requirements for quality (Geological Society Engineering Geology, 2001). The NZTA has specified a specification (TNZ M/4 2006 – Specification for basecourse Aggregate) which includes a quality control procedure where the aggregate must undergo a set of tests to ensure its performance is up to an acceptable level (Black, 2009; Transit New Zealand, 2006b). As will be discussed in the Section 3.2.2, quality control tests are split into ‘Source’ and ‘Production’ tests.

#### 3.2.2 Source and Production Properties

The specification set out by NZTA for basecourse aggregates (TNZ M/4:2006) distinguishes between source and production properties. It is thought that the ‘source’ properties are those which are inherent properties of the rock and should not change or differ significantly over time, tests such as crushing resistance, weathering resistance fall under this category. On the other hand, ‘production’ properties refer to properties which are solely influenced by the production process of the rock, and are known to
significantly change over time. Particle-size distribution (PSD), Broken face Content and Sand Equivalent (SE) are classified as ‘production’ property tests (Black, 2009; Transit New Zealand, 2006a).

This distinction made between ‘source’ and ‘production’ properties may have been valid at the time the TNZ M/4 specification was created, however due to the advancements in quarry equipment and technology, all property tests including ‘source’ properties (which are initially thought to be governed only by the inherent characteristics of the aggregate) can now be controlled and manipulated through the processing method a quarry undertakes (Ellis, 2010). This fact is also supported by Black (2009) where it is reported that “all aggregate properties are dependent on the processing methodology used to produce them” (Black, 2009).

Black (2009) reports a test that has been carried out where a range of aggregates sourced from the same rock have been processed in a quarry in different ways and levels in an aim to observe if any source properties would be affected by the different levels and methods of processing. The results revealed that there was a significant variation in ‘source’ properties between the aggregates meaning that the different methods and levels of processing do affect the ‘source’ properties of an aggregate. Hence, all of the standard aggregate properties tests are, to some extent, influenced by the processing method a quarry undertakes.

Based on the original NZTA distinction made between ‘source’ and ‘production’ properties, the flow chart shown in Figure 3-1 shows the TNZ M/4 quality control procedure in accepting an aggregate for use as a basecourse layer.
Figure 3-1: Quality Control Flow Chart for Basecourse Aggregate (Transit New Zealand, 2006b)
3.2.3 Source Property Tests

‘Source’ properties define the properties at which the aggregate is sourced from. Theoretically speaking, these are not dependant on the processing method a quarry undertakes; rather they rely on the basic inherent properties (such as crushing and weathering resistance) of the source these aggregates are extracted from. A brief description of each of the source property tests performed on the aggregate is provided in the following Sections (Transit New Zealand, 2006a).

Because source properties for a reasonably homogenous quarry do not change significantly over time, they shall be sampled and tested (using all source property tests such as Crushing Resistance, Weathering Quality and CBR) at least once every 10,000m³ (Transit New Zealand, 2006b).

**Crushing Resistance**

The purpose of the Crushing Resistance Test is to indicate the strength and likelihood of attrition of the aggregate. In the aggregate industry the strength of a rock is defined by the stress at which the material begins to fail. Hence, the Crushing Resistance test involves applying a specified load to the aggregate and consequently measuring the amount of fines it produces (Black, 2009; New Zealand Standards, 1991g; Transit New Zealand, 2006a).

**Weathering Quality Index**

The aggregate being used as a basecourse layer must not degrade under environmental changes, hence, it must meet minimum weathering quality index criterion.

The weathering quality test assesses the degree of the aggregate’s ability to resist the effects of environmental changes such as wetting, drying, heating and cooling. The test attempts to represent natural adverse weather conditions where aggregates are exposed to the combined agencies of wetting and drying and heating and cooling (New Zealand Standards, 1991h; Transit New Zealand, 2006a).

**California Bearing Ratio**

The California Bearing Ratio (CBR) is a penetration test that was developed in California by the California State Highway Department, the test assesses the aggregate’s bearing
capacity and compares to that of a high quality crushed stone. The test is used to evaluate the strength of an aggregate (HEICO, 2010; New Zealand Standards, 1991i).

The TNZ M/4 aggregate to be used as a basecourse layer must attain a minimum required CBR of 80% (Henning, 2008b; Transit New Zealand, 2006b).

3.2.4 Production Properties

The production properties of an aggregate are defined by how the aggregate is processed in a quarry. Different quarries process aggregates in different ways and hence the TNZ M/4 specification for basecourse aggregates specifies a number of tests (such as Sand equivalent, Broken Face content and Particle size distribution etc.) that an aggregate must be tested for to ensure that a satisfactory quality assurance is met. The following Sections identify and give a brief description about the production property tests. The number of samples of each of the production tests to be performed depends on the lot size as shown in Table 3-1.

Table 3-1: Minimum Sampling Rate for 'Production' Property Tests – Reproduced from (Transit New Zealand, 2006b)

<table>
<thead>
<tr>
<th>Lot Size</th>
<th>Number of Samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>From</td>
<td>To</td>
</tr>
<tr>
<td>1 m³</td>
<td>400 m³</td>
</tr>
<tr>
<td>400 m³</td>
<td>1500 m³</td>
</tr>
<tr>
<td>1500 m³</td>
<td>4000 m³</td>
</tr>
</tbody>
</table>

If a lot size exceeds 4000m³ then the number of additional tests performed should be at the rate of one per 1000m³

**Sand Equivalent**

The Sand Equivalent test (SE) measures the relative amounts of silt or clay size particles in granular soils indicating its cleanness.

Black (2009) argues that although the SE test replaces two other tests (Plasticity Index and Clay Index), these tests are not directly comparable, studies of the relationship between the three tests for a range of different aggregate types indicate that there is no strong correlation between these three tests. The study also revealed that the three tests
lack the ability to determine the presence of moisture sensitive fine particles. The presence of fine clay particles in the aggregate has a deleterious effect on the aggregates permeability (Black, 2009; New Zealand Standards, 1991f; Transit New Zealand, 2006a).

**Clay Index**

The Clay Index (CI) test is basically a methylene blue titration test that is used to “...estimate the percentage of expansive clay minerals in natural fines or rock powders” (New Zealand Standards, 1991b). Black (2009) defines the test as a measure of the surface area of fraction fines in the aggregate by determining how much methylene blue can be adsorbed on the surface of the aggregate fines.

**Plasticity Index**

The Plasticity Index (PI) is the difference between the plastic and liquid limits. The test determines the PI of fine fractions of an aggregate. It is heavily criticised as the results are subjective and are dependent on the experience of the lab technician (Black, 2009; New Zealand Standards, 1991c).

**Broken Face Content**

The Broken Face Content test determines how many ‘broken faces’ an aggregate fraction of a test sample has. In order for an acceptable level of performance an aggregate must have a number of broken faces when crushed. This helps increase the strength and interlock forces of the particles in the aggregate (New Zealand Standards, 1991d).

**Particle Size Distribution**

The Particle Size Distribution (PSD) method (also known as aggregate grading) is a simple sieving test which can be performed wet or dry (wet sieving is the preferred method). The PSD must conform to the envelope limits (upper and lower). The test provides an assessment of how well the material may mix and compact, thus providing the interstitial strength of an unbound granular material (Black, 2009; New Zealand Standards, 1991e; Transit New Zealand, 2006a).

### 3.2.5 Concerns about the Property Tests in New Zealand

Most of the property tests that have been mentioned above were developed in Europe and North America to help predict those countries’ aggregate performance. Since then, these tests have been adopted with minor modifications to suit New Zealand aggregates.
However, Black (2009) argues that these tests have been developed specifically for Europe and North America’s aggregates which are considered old continental rocks. 69% of America’s crushed stone aggregate production is carbonate, and Europe produces approximately 60% of carbonate aggregates. In contrast, New Zealand produces around 70% of greywacke aggregate where the remainder is derived from young volcanic rocks. Thus, there are major differences in the natural properties of Europe/North America’s aggregates and New Zealand’s geologically young aggregates. This follows that “the tests that have been developed specifically for old continental rocks may not be an effective predictive tool on our geologically young aggregates” (Black, 2009; Lowe et al., 2010).

Most of these property tests are measuring more than one physical property at one time, because these properties cannot be independently controlled, it is quite difficult to interpret results. The physical strength of an aggregate for example (i.e. the crushing resistance) has an effect on more than one property test; it governs (to some extent) whether or not the aggregate will conform with other tests such as PSD, weathering quality and SE (Black, 2009).

There is also growing criticism regarding the weathering resistance test, as it is thought that the test does not reflect real environmental weathering conditions. The test offers a poor prediction of the weathering of the aggregate in service, however, the weathering resistance test is better than many inadequate tests and gives an indication of the aggregates weathering performance (Black, 2009; Transit New Zealand, 2006a).

### 3.3 Aggregate Mineral Composition Testing

The mineral composition of an aggregate defines its performance and behaviour under the application of load. Aggregates containing different mineral constituents could be contributing to the variation in the vibratory hammer compaction test results. Testing of the aggregate’s mineral composition is therefore imperative to achieving a sound understanding of the variability in the vibratory hammer compaction test results.

X-ray Diffraction (XRD) is a test method used to understand the mineral composition of an aggregate. Wavelength X-rays are introduced to a powdered sample of the aggregate, where the reflections are then recorded and the data analysed to calculate the inter-atomic spacing between each mineral layer. These inter-atomic spacings provide unique patterns
which allow the identification of the minerals present within the aggregate (Lowe et al., 2010).

Clay minerals that are most commonly found in NZ rocks are chlorite, illite and smectite. Of particular concern, is the smectite clay mineral which is classed as a swelling clay. Swelling is defined as the volume change in the aggregate due to the absorption of water. The effect of swelling clays is problematic to the durability and overall performance of the aggregate (Lowe et al., 2010).

To test for the presence of these expansive minerals, two X-ray diffractions tests are conducted on the sample. The first sample being untreated and is tested in its natural state, this sample does not identify any swelling clays. Ethylene glycol is added to the second sample to allow for the identification of the expansive swelling clay minerals during X-ray diffraction testing (Lowe et al., 2010).

3.4 The effect of grading on performance of basecourse aggregate

Land Transport New Zealand (now NZTA, as of 2008) has carried out a research investigation into the effect of the Particle Size Distribution (PSD) on the performance of aggregates. The PSD also referred to as the grading of an aggregate plays an important role in aggregate behaviour under an imposed load. A dense graded material refers to a material in which each particle size fits closely into the space left between bigger particle sizes within the grading. Uniformly graded aggregates refer to those aggregates which predominately contain one particle size (Arnold et al., 2007).

Grading envelope curves are specified in the TNZ M/4 basecourse specification to ensure that dense grading is achieved. Grading curves are established by plotting the diameter of particles (in mm) on a negative log scale, against the mass percentage of the material smaller than that diameter. Formula 3-1 is used to work out the grading curve of a particular aggregate.
\[ p = 100 \left( \frac{d}{D} \right)^n \] 

(3-1)

where:

\( p \) = Percent passing sieve size \( d \)

\( D \) = Maximum particle size and

\( n \) = known as Talbot’s exponent \( n \) value, it is an integer which has a common range of 0.3 (fine grading) and 0.6 (coarse grading)

It is hence, a simple and convenient way to describe the grading of an aggregate by the use of Talbot’s exponent \( n \)-value, where \( n \)-values > 0.5 refer to course graded aggregates and \( n \)-values < 0.5 are fine graded aggregates (Arnold et al., 2007).

The study involved testing a similar material to that used in this research, the TNZ M/4 AP40, however sourced from a different quarry located in the south island. This material was referred to as “Material 1” in the study. “Material 7” was kept anonymous in the research and hence will not be discussed as the source of the material is unknown.

![Figure 3-2: Effect of Talbot’s Grading Exponent n on Rutting Performance for Material 1 and 7 (Arnold et al., 2007)](image)

Figure 3-2 shows the observed trend for Material 1 under dry and wet conditions. It can be seen that in dry conditions, PSD’s with fairly fine particles outperform PSD’s with fairly fine particles at wet conditions. Literature reviewed in this study also supported that dry materials with a high fines content can help reduce permanent deformation. On the
other hand, as Talbot’s n-value increases (i.e. as the particle size distribution increases within an aggregate) from approximately 0.4 to 0.8, Material 1 (wet) begins to outperform the Material 1 (dry) as shown in Figure 3-2. The report concluded from the findings that testing on the TNZ M/4 AP40 at gradings with a Talbot’s n-value of 0.3, 0.4, 0.55 and 0.8 showed that the best performance with the least rutting observed in wet conditions was at the n-value of 0.8 (coarse graded). While the best performance ratings obtained at dry conditions was observed at the n-value of 0.3 (fine graded) (Arnold et al., 2007).

In conclusion, a balanced aggregate grading is recommended for the basecourse layer, where the particle size distribution is slightly balanced in terms of fine aggregate to coarse aggregates. This will allow for optimum performance under dry and wet conditions.

3.5 Summary

This chapter discussed the basecourse aggregates used in New Zealand. NZTA implemented a specification known as the TNZ M/4 which sets out the requirements for a basecourse aggregate to be used in New Zealand pavements. The TNZ M/4 includes a set of tests to ensure quality of the aggregate is met before it is used as a basecourse layer. Each of these tests were discussed in this chapter. A test procedure known as the X-ray Diffraction test, which allows for the determination of mineral constituents of the aggregates, is also discussed. The X-ray Diffraction test is hoped to determine the natural variability of the aggregate being used in testing. While other sources of variation which could be contributed by test conditions are discussed in greater detail in the next chapter.

The effect of grading on the performance of the pavement was also discussed. It was found through a literature search that aggregates with a relatively coarse grading outperform finer grading aggregates when wet. In contrast, fine grading aggregates outperform coarser aggregates in dry conditions. Thus, a balanced gradation must be achieved in order to obtain an aggregate which can perform well under wet and dry conditions.

In addition to the grading, several other factors can significantly affect the performance and consequently the results of the vibrating hammer compaction test. These are discussed in the next chapter.
Chapter 4. **VARIATION IN VIBRATING HAMMER COMPACTION TESTS**

4.1 Introduction

Growing concern over the repeatability and reproducibility of the laboratory-based vibrating hammer compaction test has been expressed from the time the test was adopted by the New Zealand Standards (NZS). The test is known to produce inconsistent and significantly variable results (Opus International Consultants Limited, 2008). These laboratory results are used as benchmark values on the field. Thus, it is important that these values are reliable and accurate (Frobel & Moulding, 2006). However, it was clear from field experience that contractors often could not reach the target Dry Density specified by the laboratory test.

To address the problem of varying results, NZTA requested an inter-laboratory (round robin) study be carried out to investigate the influence of a range of test factors on the reproducibility and repeatability of the New Zealand vibrating hammer compaction test (Opus International Consultants Limited, 2008).

The vibrating hammer compaction test is used by several countries such as USA, Britain and New Zealand. Unlike New Zealand, the USA and Britain have designed distinct test methods for both unbound granular materials and cohesive material (ASTM D 7382 - 08, 2008; BS EN 13286 - 4, 2003). In the case of the New Zealand Standard, a generic test method is used for all material types. However, the standard does mention that the vibrating hammer compaction test is particularly suitable for granular material (New Zealand Standards, 1986b). It seems intuitive that different materials which have different physical properties, mineral composition and behaviour under application of load should not be tested using the same test method.

It is worth noting that the UK initially had only one test method for all types of material. However, after experiencing the same problems that are currently experienced in NZ, the UK authorities introduced an additional standard specifically targeted for graded granular materials (BS 5835, 1980).
4.2 Possible Causes of Variation

Inconsistency in the results of an experiment, which has been repeated a number of times in the same manner and under the same conditions, could be due to a number of factors. This Section discusses the possible causes of variation in the results of an experiment and particularly in the vibrating hammer compaction test. The possible factors contributing to the variability of a test result are split into two categories as shown in Table 4-1 (ASTM E 177 - 10, 2010).

**Table 4-1: Potential Factors Affecting Variability in Vibrating Hammer Compaction Test Results**

<table>
<thead>
<tr>
<th>General sources of variability</th>
<th>Sources of variability specific to the vibrating hammer test</th>
</tr>
</thead>
<tbody>
<tr>
<td>Operator/Technician</td>
<td>Hammer</td>
</tr>
<tr>
<td>Calibration of Apparatus</td>
<td>Segregation</td>
</tr>
<tr>
<td>Environmental Conditions</td>
<td>Degradation</td>
</tr>
<tr>
<td>Test Sample</td>
<td>Mould Type and Size</td>
</tr>
<tr>
<td>Time</td>
<td>Oversized Particles</td>
</tr>
</tbody>
</table>

It is important to investigate all the sources of variability and quantify their effects on the results in an attempt to substantially reduce or eliminate their contribution. The factors illustrated in Table 4-1 are discussed in more detail in Sections 4.2.1 and 4.2.2.

4.2.1 General sources of variability in a laboratory-based test

General sources which may contribute to the variation in an experiment’s results are those which arise in any experiment conducted. Such variations include different technicians conducting the test, the duration of the test and the environment at which the test was done. This Section identifies these sources and describes their potential significance to the variation in an experiment’s results.

**Operator/Technician**

Variability among different operators/technicians carrying out the same test can be significant. It is therefore vital that a test method is written in a very clear and concise manner to avoid confusion and serious differences in interpretation by various operators.
It is also important that technicians follow the test method closely and accurately. However, no matter how clear and concise a test method is, different operators have different techniques in conducting an experiment. Human error such as reaction time, colour sensitivity, scale reading and interpolation differs from person to person, which in turn could affect the nature of the test results (ASTM E 177 - 10, 2010).

The level of experience and familiarity of the test method by technicians also contributes to the variability in test results. Experienced technicians are aware of common faults and mistakes within a test method. Therefore, the level of uncertainty in a test conducted by an experienced technician is much lower than that for an amateur technician who is unfamiliar with the test.

**Calibration of Apparatus**

Improper calibration of apparatus, different levels of tolerances and uncertainties can contribute to the variation in test results. The test methods should provide information on the frequency at which an equipment must be recalibrated (ASTM E 177 - 10, 2010).

**Environment**

Material properties are sensitive and can be easily influenced by environmental effects such as temperature, humidity, atmospheric pressure and contaminants. Although it is common for a test method to specify the environmental conditions for testing, these conditions cannot be perfectly controlled within and between laboratories. A margin of error must be incorporated in test methods relating to the inevitable variability which will occur due to environmental effects (ASTM E 177 - 10, 2010).

**Test Sample**

A bulk of material should be checked for quality periodically through property testing because it is unlikely that the material is homogenous throughout. As discussed in Section 3.1 aggregates, particularly in New Zealand, are geologically young materials that have not been exposed to geological metamorphism. This means that these materials may be heterogeneous and may have varying mineral constituents. The differences in mineral compositions (and other properties) can yield varying test results (Black, 2009).

X-ray diffraction and property tests should be conducted periodically on the material to ensure its uniformity throughout a quarry source. This helps eliminate any variations
caused by the non-uniformity of the material (Lowe et al., 2010). The test specimen must also be prepared and tested in the same manner every time to avoid inconsistencies in sample preparation. If sample storage conditions are specified within a test standard, these should be followed and kept constant for all samples being tested (ASTM E 177 - 10, 2010).

**Time**

Time can influence each of the factors mentioned above; the longer the period between two or more test results, the less likely the changes in the factors mentioned above will stay at a minimum, and therefore would increase observed differences in test results. The degree of control exercised by a laboratory over the above factors will govern the amount of variation due to time (ASTM E 177 - 10, 2010).

The duration of each test can also contribute to the variation in test results. Tests must all be conducted from the same source in roughly the same amount of time to avoid uncertainties due to settlement of water in a material for example. In addition, the vibrating hammer compaction test samples require a curing period after being wetted to the required level of Water Content. This curing period ensures that the test sample has thoroughly been soaked in water to establish equilibrium. It is important to ensure that all samples being tested are given equal periods of curing time (New Zealand Standards, 1986b).

### 4.2.2 Sources of variability specific to vibrating hammer test

Within the vibrating hammer compaction test, there are a number of factors which could possibly affect the reliability of this test method. These factors include segregation and degradation of the material, mould shape and size, type and age of hammer and any oversized particles present during compaction. A discussion of each factor follows.

**Hammer**

It is no surprise that the type of hammer could have a significant impact on the results produced. Hammers with different power and frequency ratings yield different results. A round robin study carried out by Opus (2008) proved that laboratories which used hammers with low power ratings generally produced lower Dry Densities. Those which used hammers with power ratings in the higher end of the spectrum yielded notably
higher Dry Densities, this will be discussed in greater detail in Section 4.4 of this chapter (Opus International Consultants Limited, 2008).

The calibration test for acceptable hammers that is incorporated in the New Zealand Standard test method for the vibrating hammer compaction test (NZS 4402 Test 4.1.3) is highly criticised. It involves compacting Leighton Buzzard Silica Sand at a specified Water Content, and only has a minimum requirement and not a maximum. Because an upper limit does not exist, bigger and more powerful hammers can be used to achieve higher Dry Densities. However, powerful hammers can affect and damage the nature of the material (degradation) (Frobel & Moulding, 2006; New Zealand Standards, 1986b).

**Degradation**

As discussed in Section 3.4, the grading of a material influences its characteristics and performance, therefore it is vital to ensure that the specified gradation does not change during compaction. Degradation, which is a phenomenon opposite to gradation, is defined as the breakdown of aggregate particles in smaller sized fragments. It can occur during compaction due to the compactive effort. As mentioned above, high compactive efforts can damage the nature of the aggregate and cause it to breakdown. Because an aggregate is carefully graded before its intended use, the breakdown of particles will affect the gradation of the aggregate and in turn the performance of this aggregate (Luxford, 1975).

**Segregation**

Segregation is defined as the non-uniform distribution of the coarse and fine particles within an aggregate. The tendency of fine particles separating from the larger coarse particles creates an inevitable scenario where segregation will occur, especially in the vibrating hammer compaction test where segregation becomes most pronounced. Segregation occurs either during the compaction process or when placing the specimen into the compaction mould. Thus, in order to keep segregation to a minimum, it is essential that the placement of the specimen into the mould is done very carefully and in a consistent manner. Although segregation during compaction is inevitable, it can be substantially reduced through the application of adequate surcharge weight on the top surface of the test specimen. The imposed weight on the sample helps prevent movement of finer particles away from coarse particles and therefore helps minimise segregation (Panarese, 1972). It seems that as a result of excessive segregation, a British Standard for the vibrating hammer compaction test method (BSI 5835 “Compactibility test for graded
aggregates” has repeatedly increased its surcharge specification in the vibrating hammer compaction test over the years. Two amendments (amendment 1 in 1983 and amendment 2 in 1987) have been introduced to the original specification to increase the surcharge weight from 350 N to 450 N (BS 5835, 1980).

In order for an aggregate to perform adequately, its gradation must be uniform throughout its depth; however during compaction, the vibrating action causes the finer particles within the aggregate to roll down to the bottom outer edges of the mould. When the finer particles of the aggregate accumulate in one layer during compaction on the field, the permeability of the aggregate can be grossly influenced (Luxford, 1975).

Unfortunately, a standard qualitative method for measuring the amount of segregation (and degradation) which takes place during compaction does not exist. Researchers in the field of compaction studies state that segregation should be visually analysed. However, some suggest (without any experimental development yet), that it is possible to measure the degree of segregation by extruding out thin sections of a compacted sample and carrying out gradation analysis on these sections (Luxford, 1975).

**Mould shape and size**

Mould size and shape can influence the reliability in the results of the vibrating hammer compaction test. The mould diameter size selected can influence results depending on the maximum particle size available in the sample being compacted. According to Drnevich et. al. (2007) research has found that the Maximum Dry Density (MDD) is reached in moulds six to eight times the maximum particle size available in the sample. The NZ vibrating hammer compaction test standard does not conform to this theory. It allows for particles of maximum size of 37.5 mm to be compacted in a 152 ± 0.5 mm. The diameter of the specified NZ mould is only four times the maximum permissible particle size (New Zealand Standards, 1986b). Research conducted by Bishop and Green (1965) suggests that the height to diameter ratio should not be less than 2 to 1. The NZ vibrating hammer compaction test standard (NZS 4402 Test 4.1.3 “New Zealand vibrating hammer compaction test”) specifies a mould size such that the height to diameter ratio is approximately 1 to 1.
**Oversized particles**

The NZS 4402 Test 4.1.3 allows for up to 37.5 mm particles in a 152 ± 0.5 mm diameter mould, international standards such as the USA does not allow for particles of this size in such a small mould. As will be discussed in Section 5.4, the USA standard employs two methods, each specified for different maximum allowable particle sizes available within the sample. Drnevich et. al. (2007) suggest that particles retained on the 19mm sieve that are compacted in a 152 mm diameter can lead to interlocking of particles which can ultimately interfere with the compaction process.

### 4.3 Ruggedness Test

#### 4.3.1 Introduction

The purpose of the Ruggedness, also known as robustness, test is described in the ASTM standard as the identification of “...those factors that strongly influence the measurements provided by a specific test method and to estimate how closely those factors need to be controlled” (ASTM E 1169 - 07, 2007). The ruggedness test was initially introduced to avoid facing problems in inter-laboratory tests and to identify the potential factors responsible for inconsistency of results. Thus, the ruggedness test is part of the validation phase of the development of a standard test method; it is recommended that the ruggedness test precedes an inter-laboratory (round robin) study (ASTM E 1169 - 07, 2007; Massart et al., 2006).

#### 4.3.2 Concept of the Ruggedness test

The ruggedness test is a planned experiment where environmental and test factors are intentionally varied in order to record the effects on the test results of such variation. It requires making systematic changes in the test factors which are believed to have potential effect on the results, and then observing the subsequent effects of these changes on the end results of a test method. The steps involved in conducting the ruggedness test are briefly identified (ASTM E 1169 - 07, 2007):

1. Identification of relevant factors
2. Selection of levels for each factor (two realistic extremes for each factor, usually a high and low extreme)
3. Display the treatment combinations in a cyclic shifter order
4. Execution of runs arranged in a random order
5. Statistical analysis to determine the effect of factors on the test method results, and
6. Possible revision of test method if needed.

4.3.3 Suitability to vibrating hammer test

The ruggedness test was initially incorporated in the research proposal as part of this study. It is important to determine which factors have a significant effect on the vibrating hammer compaction test results (Wilson & Shamseldin, 2010). However, before identifying these factors using the ruggedness test, it is vital that an understanding of the natural variability of the test is achieved.

4.4 Inter-laboratory (Round Robin) Study

4.4.1 Introduction

The build-up of uncertainty in the results of the vibrating hammer compaction test in New Zealand was agreed upon at a Civil Engineering Testing Association of New Zealand (CETANZ) meeting in 2008. At the meeting, the New Zealand vibrating hammer compaction test (NZS4402:1986 Test 4.1.3) was discussed and it was agreed by participants that an inter-laboratory study be carried out between current laboratories conducting the test to establish actual repeatability and reproducibility values of the test. Thus, this Section reviews the inter-laboratory study that has been conducted by Opus international Consultants Ltd., Central Laboratories (Opus International Consultants Limited, 2008).

In order for a full rework or replacement of the current test standard for the vibrating hammer compaction, the repeatability and reproducibility values must be evidently higher than those values obtained by previous inter-laboratory studies conducted by the UK and USA.

The round robin study conducted by Opus consisted of thirty three laboratories and was aimed at determining the effect of a range of different equipment variables on the reproducibility and repeatability values of the New Zealand vibrating hammer compaction test. As a requirement set out by ASTM to determine repeatability values,
testing was to be carried out by the same technician in each lab (ASTM E 691-09, 2009).

Two materials were tested; TNZ M/4AP40 and GAP40. These were tested at two water contents; 2.0% and 6.0% for the TNZ M/4AP40 (denoted by TNZ2 and TNZ6 respectively) and 1.0% and 4.0% for the GAP40 (denoted by GAP1 and GAP4 respectively). The determination of water contents after compaction was carried out as specified in the NZS 4407 Test 3.1 “The Water Content of aggregate” (New Zealand Standards, 1991a).

Results from five laboratories were excluded from the data analysis from the study simply because these results varied significantly from the rest of the data. All laboratories used identical samples to conduct the test. Therefore, this variability by the five laboratories could be attributed largely to the inconsistency in the operating procedures.

4.4.2 Discussion of Results

The inter-laboratory study conducted by Opus produced some important results which helped in setting the objectives for this research. This Section analyses and discusses these results in relation to the objectives set in this research.

**Repeatability and Reproducibility**

The precision of an experiment is defined by the “closeness of agreements between independent test results obtained under stipulated conditions” (ASTM E 691-09, 2009). Precision is measured by two parameters; **Repeatability** and **Reproducibility**.

**Repeatability** is the precision of a test method where the independent test results are obtained using the same equipment and conducted by the same technician in the same laboratory. Hence, repeatability ensures the reliability of a test method under constant conditions (ASTM E 691-09, 2009).

The precision of a test method where the results are obtained following the same test method, in different laboratories with different technicians and equipment is measured by the reproducibility value of that test method (ASTM E 691-09, 2009).

The repeatability and reproducibility values were calculated in the report produced by Opus and are shown in Table 4-2.
Table 4-2: Variability Parameters of the Dry Density – Reproduced from (Opus International Consultants Limited, 2008)

<table>
<thead>
<tr>
<th>Material</th>
<th>Average Dry Density t/m³</th>
<th>Repeatability, r t/m³</th>
<th>Reproducibility, R t/m³</th>
</tr>
</thead>
<tbody>
<tr>
<td>TNZ2</td>
<td>2.175</td>
<td>0.078</td>
<td>0.151</td>
</tr>
<tr>
<td>TNZ6</td>
<td>2.242</td>
<td>0.078</td>
<td>0.212</td>
</tr>
<tr>
<td>GAP1</td>
<td>2.232</td>
<td>0.048</td>
<td>0.125</td>
</tr>
<tr>
<td>GAP4</td>
<td>2.222</td>
<td>0.046</td>
<td>0.133</td>
</tr>
</tbody>
</table>

Repeatability values of the TNZ material are identical (0.078 t/m³) and are higher than repeatability values for the GAP material (0.048 and 0.046 t/m³). Hence, for the vibrating hammer compaction test method, the type of material does influence test repeatability. There appears to be high inter-laboratory factors contributing to the variance in reproducibility values, particularly in the TNZ6 material. This could be due to the differences in the method in which technicians are conducting the test. In addition, significant amount of water loss has been observed by some laboratories during the compaction process particularly for the TNZ6 material (discussed in more detail in the following Section) (Opus International Consultants Limited, 2008).

A similar precision study was carried out in 1988 by the British Standards Institution (BS EN 13286-4 2003 Part 4 Test methods for laboratory reference density and water content – Vibrating hammer) where 12 laboratories took part. The British Standard states a reproducibility value, R of 0.054 t/m³ and a repeatability value, r of 0.033 t/m³ for a gravel subbase material (BS EN 13286 - 4, 2003).

In addition, the vibrating hammer compaction test method in the ASTM standards (ASTM D 7382 -08 Standard Test Methods for Determination of Maximum Dry Unit Weight and Water Content Range for Effective Compaction of Granular Soils Using a Vibrating Hammer) states a repeatability value, r of 0.05 t/m³. The ASTM D 7382 does not mention what type of material this repeatability value is based on. Reproducibility studies have not yet been completed by this standard (ASTM D 7382 - 08, 2008).

A comparison of the repeatability and reproducibility values of the three different standards is provided in Table 4-3.
Table 4-3: Comparison of the Repeatability and Reproducibility Values

<table>
<thead>
<tr>
<th>Material</th>
<th>Repeatability, r t/m³</th>
<th>Reproducibility, R t/m³</th>
</tr>
</thead>
<tbody>
<tr>
<td>GAP1 (NZ)</td>
<td>0.048</td>
<td>0.125</td>
</tr>
<tr>
<td>GAP4 (NZ)</td>
<td>0.046</td>
<td>0.133</td>
</tr>
<tr>
<td>TNZ1 (NZ)</td>
<td>0.078</td>
<td>0.151</td>
</tr>
<tr>
<td>TNZ6 (NZ)</td>
<td>0.078</td>
<td>0.212</td>
</tr>
<tr>
<td>Gravel Subbase (UK)</td>
<td>0.033</td>
<td>0.054</td>
</tr>
<tr>
<td>Unknown (USA)</td>
<td>0.05</td>
<td>N/A</td>
</tr>
</tbody>
</table>

The New Zealand repeatability results are of the same order as the ASTM D 7382 repeatability value (approximately 0.05 for GAP samples and 0.08 for TNZ samples, compared to 0.05 for USA ASTM). Reproducibility values for the New Zealand precision study (0.13 t/m³ to 0.21 t/m³) appear to be significantly higher (up to four times) than the UK value of 0.054 t/m³ (Opus International Consultants Limited, 2008).

However, it is important to note that the values produced by the UK and USA studies are not directly comparable to the NZ values. This is due to a number of reasons:

- The materials used in each study (USA, UK, and NZ) are quite different.
- Water Content ranges used for each study could have been different (the USA and UK studies do not specify over which Water Contents the tests were undertaken).
- Each study adopted their own test methodology of the vibrating hammer test, which vary slightly (as will be discussed in chapter 5).

Thus, although the values cannot be directly compared, UK and USA values can be used as an indication of likely repeatability and reproducibility values for this type of test (Opus International Consultants Limited, 2008).

Effect of Water Content on Variability of Dry Density Results

The water content at which a material is compacted determines the degree of compaction achieved. Hence, Opus (2008) investigated if water content was a major factor in causing
the significant variability evident in the test method results. Two graphs (for the two materials) were plotted to analyse this. One of them is reproduced in Figure 4-1.

From Figure 4-1 the Opus (2008) report explains that a significant effect of water content on dry density can be seen particularly for the TNZ6 material. The $R^2$ value of this particular material (TNZ6) denotes approximately 7% of variation can be explained by water content variation, assuming the WC-DD relationship is linear. However, since it is known that this relationship is non-linear (parabolic), this is not a valid assumption.

Thus, the Opus report concludes that although WC is known to affect the attained DD, Figure 4-1 indicates that the variability due to the between-laboratory factors overwhelm this effect (Opus International Consultants Limited, 2008).

**Effect of Hammer Power Rating on Variability of Dry Density Results**

The degree of compaction (Dry Density), among other factors is dependent on the compactive effort applied. Thus, the hammer input power rating has significant influence on the variation of the vibrating hammer compaction test results. This is due to the fact that hammers with high input power ratings apply a greater compactive effort on the specimen during compaction than a hammer with a relatively lower input power rating.
Opus (2008) investigated the hammer input power rating effect on dry density by plotting a graph of DD (t/m³) against Hammer input power (watts). Figure 4-2 is an example of one of the graphs analysed by the Opus (2008) report, as can be seen, there is a general trend for DD to increase as the hammer power input Rating increases.

**Table 4-4: Predicted Increases in Dry Density When Changing from an 1140 W Vibrating Hammer to a 1700 W Vibrating Hammer – Reproduced from (Opus International Consultants Limited, 2008)**

<table>
<thead>
<tr>
<th>Material</th>
<th>Dry Density Increase with Hammer Power, t/m³</th>
</tr>
</thead>
<tbody>
<tr>
<td>TNZ2</td>
<td>0.045</td>
</tr>
<tr>
<td>TNZ6</td>
<td>0.074</td>
</tr>
<tr>
<td>GAP1</td>
<td>0.038</td>
</tr>
<tr>
<td>GAP 4</td>
<td>0.053</td>
</tr>
</tbody>
</table>

Based on the equations shown in Figure 4-1 and other graphs plotted in the Opus (2008) report for the other materials, Table 4-4 was produced, which shows the predicted DD increase when changing from an 1140 W vibrating hammer to a 1700 W hammer. These
increases appear to be quite significant, considering the repeatability and reproducibility values shown in Table 4-2 are quite low.

**Calibration of Hammer**

It is a requirement that the hammer used for compaction testing be properly maintained. To ensure a satisfactory level of compactive effort is applied on the specimen, a test method for hammer calibration is incorporated in Note (5) of the NZS 4402:1986 Test 4.1.3 (New Zealand Standards, 1986b).

Leighton Buzzard sand passing a 600µm sieve with a permissible water content of 2.5 ± 0.5% is to be compacted according to the standard’s test methodology. The hammer is considered satisfactory if the DD of the sample exceeds 1.74 t/m³ (New Zealand Standards, 1986b).

Criticism of the hammer calibration method has been expressed. The calibration method only has a minimum requirement and of the hammer power input rating not a maximum. This can lead to potential variation in the results. In addition, because a maximum limit of the hammer power input rating does not exist, degradation (as discussed in chapter 2) can further contribute to the variation in results (Frobel & Moulding, 2006).

Luxford (1975) reports an investigation carried out by Parsons on the hammer type and power rating stating that care should be taken when selecting a suitable hammer for laboratory compaction. Although hammers with a high power rating achieve corresponding high DD values, this can lead to degradation and damage of the nature of the material which contributes to additional variation in results obtained.

In Opus’ (2008) investigation, seventeen of the thirty three laboratories supplied calibration results based on the NZS 4402:1986 test 4.1.3 method. Two DD values of the seventeen laboratories were below the minimum limit of 1.74 t/m³. Both of these laboratories used hammers of the same model with a power input of 1140 W, which are near the bottom of the power input rating range of the hammers used in the study.

Although the NZS 4402 Test 4.1.3 standard does not specify the hammer power rating, it recommends hammers with ratings of 600 to 1200 W with the following comment “Hammers with ratings of 600 to 1200 W power consumption have been found to be satisfactory” (New Zealand Standards, 1986b). This however, may reflect the power
ratings of many hammers at the time the test method standard was created, with the advancements in technology, vibrating hammer power ratings have increased significantly. As there is no maximum limit on the hammer power output, most laboratories use hammers slightly higher than the recommended range stated in the NZS 4402 Test 4.1.3.

4.4.3 Conclusions Drawn From the Study

The conclusions drawn from the Inter-laboratory study conducted by Opus were as follows:

- The hammer calibration method may need revision. An attempt should be made to perhaps incorporate a maximum limit of vibrating power input rating.
- NZ reproducibility values appear to be two to four times larger than those stated in the British Standard. This is quite significant and may justify a rework and/or revision of the current test standard.
- Hammers with high input power ratings tend to provide higher dry density values. Although this only accounts for roughly 10% of the variability.
- Five laboratory results out of the thirty three laboratories have been excluded from data analysis due to the extent of variability observed in these five results. It is evident by these five laboratories that the way in which the vibrating hammer compaction test method is conducted can significantly influence the reliability of the results obtained.

4.5 Summary

This chapter discussed the possible causes of variation due to the test conditions. The precision of a test experiment is affected by the experiment’s factors such as uncertainties in apparatus used, technician level of experience and the environmental conditions.

The chapter discussed the concern expressed over the reliability of the vibrating hammer compaction test results and identified the possible causes of this variation. Sources of variability in a test experiment were described briefly.

A similar study that has been conducted by Opus in the past focused on the between-laboratory variability of the test showed that the variation amongst different laboratories in New Zealand is significant. Factors such as different equipment, different levels of
technician experience and general laboratory environmental conditions can affect between-laboratory studies.
Chapter 5. REVIEW OF METHODOLOGIES

5.1 Introduction

Laboratory compaction by an electric vibrating hammer has spread widely since its development in the early 1970’s. To date, the test method has been adopted by various Standards’ Institutions worldwide including the New Zealand Standards institution (NZ), British Standards Institution (UK) and ASTM (USA). Although the fundamentals of the test method are identical, variations in the test method between each institution exists. These variations are based on the institution’s knowledge and extensive research carried out regarding the test. Hence, an important step to understanding the variability in the results of the New Zealand test method is to compare New Zealand’s version of the vibrating hammer test method to USA and UK’s versions of the test. Any apparent differences in the NZ standard, which the American and/or British standards do not concur with, could be contributing factors to the variation in test results observed in NZ.

The comparison process should help point out any major differences in the New Zealand vibrating hammer test. Whether these differences contribute significantly to the variation in results or not can be investigated once testing is conducted and results are analysed.

Due to the fact the New Zealand Standards authority works closely with the Australian Standards authority, it was hoped to include the Australian Standards method of vibrating hammer in the comparison. Unfortunately the Australian Standards has not adopted a vibrating hammer compaction test method to this day. Australia uses either the Standard Proctor or the Modified Proctor methods for the compaction of cohesionless soils (SAI AS 1289.5.1.1, 2003; SAI AS 1289.5.2.1, 2003).

The test method adopted by NZS for the vibrating hammer compaction test is described in the NZS 4402 Test 4.1.3 titled “Determination of the Dry Density/Water Content relationship – New Zealand Vibrating hammer compaction test” (New Zealand Standards, 1986b).

The American Society for testing and materials (ASTM) have also adopted a standard for the vibrating hammer compaction test described in D 7382-08 titled “Standard tests methods for determination of Maximum Dry Unit Weight and Water Content range for
effective compaction of granular soils using a vibrating hammer”; the standard in itself is divided into two methods to account for the difference in materials being compacted (ASTM D 7382 - 08, 2008).

The British Standards Institution (BSI), however, has adopted 3 different standards to accommodate for a suitable compaction method for every type of material. Initially the BS 1377:1975 Part 4.3 titled “determination of the dry density/moisture content relationship of granular soil (vibrating hammer method)” test method, which is considered a purely British Standard, was developed and published by BSI. However, the BS 1377 brought rise to another standard known as the BS 5835 Part 1: in 1980 titled “Recommendations for testing of aggregates Part 1 – Compactibility test for graded aggregates”, due to the fact that the BS 1377 test method was deemed unreliable when applied to aggregates that are commonly used for road sub-bases and base materials. It is important to note though, that the BS 5835 did not supersede BS 1377 because although BS 1377 was deemed unreliable when applied to graded aggregates, the test method was still applicable to other types of materials and soils. Since this research is dealing with graded aggregates, the comparison of standards will include the BS 5835 and not the BS 1377. The third British Standard for the vibrating hammer compaction test is the BS EN 13286-4:2003 Part 4 “Test methods for laboratory reference density and water content – vibrating hammer” which was originally a European Standard that was later adopted by BSI as a British Standard. As will be discussed in later Sections of this chapter, the type of material (and particle size) being compacted governs which one of these British standards should be used (BS 5835, 1980; BS EN 13286 - 4, 2003).

This chapter will examine the main sections common to the three standards and discuss the differences found.

5.2 Scope

The NZS 4402 Test 4.1.3 test method determines the dry density when soil passing a 37.5mm sieve is compacted by a vibrating hammer over a range of water contents, including that which provides the Maximum Dry Density (MDD).

The ASTM D 7382 test method determines the dry unit weight of granular soils specifically, by compaction using the vibrating hammer. The standard is divided into two
methods (Method A and Method B), each is created based on percentage of maximum particle size present in the aggregate.

- **Method A** – Applies to material passing a 19.0 mm sieve and containing up to 35% of the total dry mass passing a 75 µm sieve.
- **Method B** – Applies to material passing a 50 mm sieve and containing up to 35% of the total dry mass passing a 75 µm sieve.

If however, the material contains 30% or less of its mass retained on the 19.0 mm sieve Method A can still be used by applying a correction procedure specified in Practice D 4718. As will be discussed in the procedure Section of this chapter, Method B is a lengthy and more complicated method. Thus, for ease of operations, it is highly recommended to use Method A, unless Method B is required due to the gradation of the aggregate not meeting the above requirement of no more than 30% of the aggregates’ mass retained on the 19.0mm sieve. The aggregate being used in this research does not comply with method A, thus, method B will be analysed for comparison purposes.

The BS EN 13286 test method also determines the relationship between the dry density and water content by the process of compaction using a vibrating hammer. It applies to mixtures containing less than 30% of their mass retained on the 20 mm sieve. It does not apply to aggregate mixtures containing more than 10% of its mass retained on the 40 mm sieve. However, the standard includes an annex that has a different procedure designed specifically for mixtures with particles that do not conform to the 20 mm sieve requirement.

The BS 5835 allows for aggregates with particles smaller than 37.5 mm, any material retained on the 37.5 mm sieve is removed and discarded.
5.3 Apparatus

Table 5-1 identifies the apparatus used in the various test methods and their associated dimensions.

<table>
<thead>
<tr>
<th>Apparatus</th>
<th>Standard</th>
<th>NZ</th>
<th>BS 5835</th>
<th>BS EN 13286</th>
<th>ASTM D 7382</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mould</td>
<td>Diameter (mm)</td>
<td>152 ± 0.5</td>
<td>150 to 150.08</td>
<td>149.8 to 150.5</td>
<td>279.4 ± 1.1</td>
</tr>
<tr>
<td>Height (mm)</td>
<td>125 to 127</td>
<td>NA</td>
<td>NA</td>
<td>230.9 ± 0.5</td>
<td></td>
</tr>
<tr>
<td>Perforation</td>
<td>Base</td>
<td>Side</td>
<td>Side</td>
<td>Non perforated</td>
<td></td>
</tr>
<tr>
<td>Hammer</td>
<td>Frequency (Hz)</td>
<td>25 to 60</td>
<td>N/A</td>
<td>33</td>
<td>53 to 58</td>
</tr>
<tr>
<td></td>
<td>Power (W)</td>
<td>600 to 1700 W</td>
<td>N/A</td>
<td>900</td>
<td>N/A</td>
</tr>
<tr>
<td>Frame</td>
<td>Surcharge Load (N)</td>
<td>350 ± 50</td>
<td>N/A</td>
<td>640 ± 10</td>
<td>285 to 570</td>
</tr>
</tbody>
</table>

5.4 Sample Preparation

Proper sample preparation is vital to achieving correct and accurate end results. All the standards being reviewed suggest that samples being tested be kept in a cool dry place and well away from direct sunlight, to help minimize the problem of condensation and water loss from the material.

Some differences in the way samples are prepared do exist among the standards being reviewed, more specifically the curing period after wetting up the samples. The NZS 4402 Test 4.1.3 specifies a period of at least 16 hours before any testing is done on the sample and advises that for some soils such as heavy clays, an even longer period is required to establish equilibrium. ASTM D 7382 on the other hand specifies a short soaking period of only half an hour. Both British standards (BS EN 13286 and BS 5835) suggest a 12 hour period to allow for thorough wetting.

As discussed earlier, each standard specifies a limit on the maximum particle size allowed in the material being compacted. Preparation of samples involves sieving out any
oversized particles present in the material. The NZS 4402 Test 4.1.3 specifies that any material retained on the 37.5 mm sieve is to be removed and discarded from the sample. The NZS 4402 Test 4.1.3 does not specify an additional procedure for materials over the maximum particle size limit of 37.5 mm. Method B of the ASTM D 7382 standard applies to those aggregates which pass the 50.0 mm sieve. On the other hand, the BS EN 13286 provides two separate methods. The first method is applied for aggregates which contain less than 30% by mass retained on the 20.0 mm sieve, and the other method is applied for those aggregates which do not conform to this requirement. The BS 5835 is similar to the NZS 4402 Test 4.1.3 in that it only adopts one method to compact any material passing the 37.5 mm sieve, any material retained on this sieve is weighed and noted down.

It can be seen by the sample preparation review; both the NZS 4402 Test 4.1.3 and the BS 5835 contain one broad method which applies to material having a wide range of particle sizes. In terms of other standards such as the ASTM D 7382 and BS EN 13286, particles over the 19.0 – 20.0 mm range are considered oversized and hence require a special method to be compacted properly and effectively.

5.5 Procedure

The most critical step which can significantly affect the precision of results is the procedure in which the vibrating hammer compaction test is conducted. Each standard sets out a different procedure in terms of the number of layers the sample is compacted, the time period of hammer operation and the different parameters measured during the test.

The NZS 4402 Test 4.1.3 achieves compaction in two layers to ensure effective compaction throughout the depth of the sample. The first layer which approximately half-fills the mould is compacted by the vibrating hammer for 180 ±5 seconds. The hammer is then removed and an additional layer of the sample is added, enough to protrude into the extension collar of the mould. The sample is then compacted by the vibrating hammer for a further 180 ± 5 seconds. The mass of the mould and sample, and the height of the sample are two of the main measurements which are recorded during the test for calculation purposes.
The ASTM D 7382 standard, as mentioned earlier, specifies two methods – Method A and Method B. This research deals with an aggregate which contains more than 30% by mass retained on the 19.0 mm sieve, this is a case which only conforms to method B of the ASTM D 7382 standard. Hence, only Method B will be discussed and used for comparison. The ASTM D 7382 Method B uses a large 279.4 mm diameter mould to prevent interlocking of particles during compaction. Since the permissible maximum particle size in this method is around 50 mm, a small 152.4 mm diameter mould may cause these large particles to interlock against each other and the inner walls of the mould which consequently interferes with effective compaction of the sample. Method B specifies that the sample be compacted in 3 layers. The specified tamper used cannot cover the full surface of this large mould of 279.4 mm diameter; thus each layer is compacted in 8 locations, as shown in Figure 5-1, for 52 ± 5 seconds at each location.

The BS EN 13286 test standard also specifies two methods as mentioned earlier. The method that is suitable for the material used in this research specifies that compaction be done in one single layer. The hammer is applied to the sample for a period of 180 ± 5 seconds.

![Diagram](image.png)

**Figure 5-1: Sequence of 152 mm Tamper Positions in 279.4 mm Mould (ASTM D 7382 - 08, 2008)**
5.6 Summary

This chapter reviewed the New Zealand Standard for the vibrating hammer compaction test with international standards. The United Kingdom and United States of America employ similar test methodologies for the compaction of cohesionless material. As a part of the investigation in the variability of the results in the New Zealand vibrating hammer compaction test, it was suggested to compare this standard with international standards.

The comparison proved that differences do exist with the New Zealand Standard. The testing and analysis of results will help in determining whether these differences in the New Zealand Standard are the reason for the variability in the results.

Important differences that exist in the New Zealand Standard include:

- The NZS allows for particles up to 37.5 mm to be compacted in a 152 mm diameter mould. Whereas the USA and UK standards specify special methods for particles larger than 19 mm.

- Procedural differences such as compacting the sample in two layers in the NZS rather than 3 layers as in the USA method, or a single layer as in the UK method.

- Curing time subsequent to wetting the samples differs in each standard. The NZS specifies a curing time of at least 16 hours. While the USA standard specifies only 30 minutes. The United Kingdom standard specifies a curing period of 12 hours.
Chapter 6. **ADOPTED RESEARCH METHODOLOGY**

6.1 Introduction

This chapter describes the methodology adopted to conduct this research and perform testing in order to achieve the specified objectives stated in Section 1.2. Two different tests were conducted to achieve a sound and scientific understanding of the variability in the results of the New Zealand vibrating hammer compaction test. Each test focused on one aspect of the two possible contributing factors, these are:

- **Variability Due to Vibrating Hammer Compaction Test Conditions** – The test methodology used is largely based on the NZS 4402:1986 Test 4.1.3 “New Zealand vibrating hammer compaction test” with a few minor modifications applied to help reduce inconsistencies in the results.

- **Variability Due to Aggregate Property Variation** – The X-ray diffraction method was utilised to provide results (such as aggregate property and mineral constituents) regarding the material being used in the test.

In addition to these two tests, quality control tests were also performed to ensure that the aggregate being used in this research maintained an acceptable level of quality.

6.2 Research Tasks

To meet the primary objective of determining an understanding of the variability in the results of the vibrating hammer test the following research tasks were implemented.

1. **Grasp a Deeper Understanding of the Problem** – The first step was to obtain a better and much deeper understanding of the problem at hand. This was done in a number of ways:
   - A review of existing literature available.
   - Seeking advice of personnel who have experience and have dealt with the test in question.
- Analysis of raw data provided by Opus from similar research conducted in the past.

2. **Quality Control Tests** – Conduct aggregate property tests such as strength and durability tests on the source aggregate to ensure an acceptable level of quality is maintained.

3. **XRD Analysis** – Perform X-ray Diffraction analysis on the aggregate being used in this research to determine if there was any major variability in the properties and mineral composition of the aggregate that could explain any significant variation from one test result to another.

4. **Adopt an Experimental Methodology for the Vibrating Hammer Compaction Test** – Perform a thorough comparison of the New Zealand Standard for the vibrating Hammer Compaction Test (NZS 4402 : 1986 Test 4.1.3 New Zealand vibrating hammer compaction test) with other international standards such as the British Standard (BS EN 13286-4:2003 Part 4: Test methods for laboratory reference density and water content – Vibrating hammer) and American Standard (ASTM D 7382 – 07 Standard test methods for determination of maximum dry unit weight and water content range for effective compaction of granular soils using a Vibrating Hammer). Adopt a modified experimental procedure based on the comparison of these standards to use for testing.

5. **Calibration of Apparatus** – Perform necessary calibration tests on all apparatus used (such as hammer, scales, moulds, timers, straight edges etc.) to control variability due to equipment.

6. **Hammer Power Output Test** – Devise a test which measures the power output of the vibrating hammer and use it to measure the output before any compaction tests were conducted and then after compaction testing was completed.

7. **Perform Vibrating Hammer Compaction Tests** – Conduct multiple vibrating hammer compaction tests at various Water Contents until a sufficient number of
tests were reached to enable reliable statistical analysis of the results. Sub-tasks in this step included:

- Keeping as many test factors constant (such as hammer type, mould size etc.), to determine the minimum natural variability of the test method.
- Conduct the vibrating hammer compaction test using a more powerful hammer than the one initially used and compare results between the two hammers.
- Conduct a Standard Proctor compaction test on the same aggregate and compare results with the vibratory hammer compaction test results.
- Conduct a Modified (heavy) Proctor compaction test and compare the results with the vibrating hammer compaction test results.

8. Evaluation and Analysis of Test Results – Statistically analyse test results obtained by utilising the statistical analysis software SPSS. Manipulate test results to graphically show trends and correlations.

9. Presentation and Conclusion of Findings – Report findings and provide conclusions and recommendations based on the analysis of results.

6.3 Quality Control

Quality control tests were carried out to ensure the aggregate being tested was up to the specified acceptable standard to be used as a basecourse aggregate. This phase was carried out before any compaction testing took place. It was also hoped to carry out these tests regularly as the compaction tests were underway to maintain a certain standard of quality, however due to the length of time it takes to conduct these tests and considering the time constraints for this research, this was not possible. The brief outline of each quality control test is described in Section 3.2.

6.4 Vibrating Hammer Compaction Test Methodology

6.4.1 Introduction

The adopted test method is largely based on the New Zealand Standard method (NZS 4402:1986 test 4.1.3 “New Zealand vibrating hammer compaction test.”). It was
developed progressively throughout the practice runs of testing. Initially, it was decided to adhere to the NZS 4402 Test 4.1.3 as much as possible and carry out the steps stated in the NZS 4402 Test 4.1.3 precisely. The reason for following the NZS 4402 Test 4.1.3 closely was to investigate the variation and suggest a revision of the current test method. However, during practice tests carried out by the researcher, a few minor modifications were suggested to keep variation to a minimum and enable consistent testing.

The degree of accuracy of data analysis relies heavily on the amount of data available. For research projects of this nature, where variation in results for a particular test method is investigated, repeating the test to acquire a considerable amount of data is desirable. However, due to time constraints, it was advised that the test be conducted forty times, which is considered an acceptable number of repeats to ensure that the statistical data analysis is reliable.

The adopted experimental procedure used to carry out the laboratory vibrating hammer compaction test is described below.

6.4.2 Scope

This test method is used for determining the dry density of granular soils, passing a 37.5mm sieve, by the use of a vibratory hammer over a range of Water Contents. The Dry Density will then be used to calculate the MDD and OWC.

6.4.3 Apparatus

The apparatus used to conduct the tests are described below.

Mould

The New Zealand vibrating hammer compaction test standard specifies an allowable internal diameter of 152 mm ± 0.5 mm and an adequate depth to provide a specimen height of 125 to 127 mm.

The NZS 4402 Test 4.1.3 also suggested that perforations be present in the base of the mould, however, consequent to practice tests conducted using the perforated base mould, it was recommended that a non-perforated base mould be used for actual testing due to the significant amount of water escaping through these perforations. As discussed in Section 2.1, compaction is defined as the expulsion of air voids from the aggregate by the
application of mechanical energy with zero or minimal water loss. Because the water present in the aggregate sample contributes to the samples total mass, loss of water can lead to a decrease in mass and hence a decrease in the dry density. Thus, it is important that water does not escape during compaction as this may lead to inconsistencies in the results.

To keep inconsistencies in results to a minimum, a new mould was constructed for the purpose of this test. The new mould was used solely for the purpose of this research, while Stevensons Laboratory Ltd used a different mould for their regular commercial compaction testing. Figure 6-1 shows the mould used for testing. It is recommended that the mould has ‘guide’ lines as shown in Figure 6-1 to give the technician an idea of how much material to fill the mould in each layer.

![Figure 6-1: Compaction Mould used for Testing](image)

**Vibrating Hammer**

The specifications of the two hammers used for the research are provided in Table 6-1. Several laboratories from the Opus study have used the Kango 950K hammer which has an input power of 1,700 W. It was recommended for the purposes of this research that a less powerful hammer be used to avoid issues of degradation and damage of the nature of the aggregate. The majority of testing (forty tests) was carried out using the Metabo KHE75; however an additional five tests were conducted using the Kango 950K to observe the differences in Dry Density values produced by these two different hammers. The power input spectrum from the Opus Inter-laboratory study varied from 750 W to
1,700 W. Thus, the Metabo KHE75 approximately falls in the middle of this spectrum. Therefore, the compactive effort provided by the Metabo hammer is neither too high nor too low, this will consequently help in obtaining average dry density values that are not disadvantaged the by application of significantly low or high compactive efforts.

Unfortunately, the hammer calibration method incorporated into the NZ vibrating hammer compaction test in the New Zealand Standards, is a heavily criticised and broad method which only specifies a minimum limit on the permissible hammers used in the test (Frobel & Moulding, 2006). Therefore, although the standard does not specify an upper limit, a hammer with an appropriate level of power input must be selected, due to the fact that significantly more powerful hammers can easily degrade the sample being compacted, leading to additional problems in the variation of results.

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Metabo KHE75</td>
<td>1,150</td>
<td>690</td>
<td>10</td>
<td>6.4</td>
</tr>
<tr>
<td>Kango 950K</td>
<td>1,700</td>
<td>850</td>
<td>7 - 27</td>
<td>11.8</td>
</tr>
</tbody>
</table>

The Metabo KHE75 and Kango 950K vibrating hammers are shown in Figure 6-2. The tampers used in each hammer are of similar dimensions and weight and are designed as per the specifications stated in the NZ vibrating hammer compaction test standard, where the diameter of the tamper is 145 mm and its weight does not exceed 3 kg. As shown in Figure 6-2, the Kango hammer is much larger than the Metabo hammer. These two hammers impose different surcharge loads on the sample due to the differences in their weight as shown in Table 6-1. The differences in imposed loads can affect the consistency of the test results. The Kango hammer is much heavier at 11.8 kg than the Metabo hammer which is only 6.4 kg. However, combined with the surcharge load in the hammer frame, both of these hammers are within the NZS 4402 Test 4.1.3 imposed load specification limits of 30 to 40 kg (350 ± 50 N) (New Zealand Standards, 1986b).
Other equipment

Associated apparatus that are needed for conducting the tests include:

- A balance that is readable and accurate to 10 grams
- A timer that is readable and accurate to 1 second
- Trays with various sizes ranging from 600mm x 500mm x 80mm to smaller sized trays of roughly 300mm x 300mm x 80mm
- Heavy grade plastic bags
- Commercial drying oven capable of heating up to 110°C, and
- A soundproof cabinet is recommended to conduct the test in.
6.4.4 Procedure

The procedure is outlined in Figure 6-3 and further described in the following paragraphs.

Referring to the main procedural steps in Figure 6-3:

Step A1 The various aggregate sample bulk of particle sizes which make up the aggregate grading are oven dried overnight (approximately 12 hours) and then allowed to cool.

Step A2 The sample recipe (known as grading) is then prepared. For each compaction curve, seven samples are used to cover a specified range of water contents (3.5% to 6.5% in 0.5% increases). Each sample is weighed to achieve approximately 5.5 kg. In contrast to the suggestion of riffling or
quartering as specified in the New Zealand Standard for the vibrating hammer compaction test, the adopted procedure makes each sample grading from scratch to help minimise the effect of segregation.

Step A3  The seven samples which are prepared according to the specified grading are then wetted to the chosen range of Water Contents (which are 3.5%, 4.0%, 4.5%, 5.0%, 5.5%, 6.0%, and 6.5%).

Step A4  Each wetted sample is then placed in a heavy grade plastic bag, which is sealed to reduce air space between the sample and the bag to minimize the problem of condensation. The samples are left to cure overnight in a cool place away from direct sunlight for a recommended period of 16 hours. Note that it is important to stay consistent in this step by giving all samples the same amount of time for curing.

Step B1  Ensure that the apparatus assembly is perfectly clean and dry. Determine to the nearest 0.5 mm the average internal diameter of the mould and record.

Step B2  Place a straight-edge across the top of the surface of the mould and measure the depth from the straight-edge to the bottom of the mould using a steel ruler. Take at least six readings around the mould and calculate the mean height and record.

Step B3  Weigh the mould to the nearest 10 g and record.

Step C1  The mould is then placed onto the base of the loading frame with the vibrating hammer drawn aside to allow free access to the mould.

Step C2  Empty one sample into a tray and thoroughly mix to help minimize segregation, and scoop enough of the material to half fill the mould when compacted (the first guide line shown in the mould in Figure 6-1 represents the ‘half fill’ mark). It is important to take extra care when scooping the sample into the mould to ensure segregation is kept to a minimum.

Step C3  Place the vibrating hammer with the tamper inside the mould so that the vibrating hammer is in position for operation. Operate the vibrating hammer for 180 seconds. Then remove the vibrating hammer and tamper from the mould.

Step C4  Add another layer of the aggregate sample, ensuring to scoop enough
material into the mould so that when compacted, the specimen just protrudes the second guide line in the mould. Repeat Step C3.

Step C5 Remove the mould from the loading frame and clean, with a dry cloth, any sample particles from the outside of the mould.

Step D1 Adopt a consistent measurement approach to measure the height of the sample in the mould. That is to say, measure the height of every sample in approximately the same six locations every time, to avoid inconsistency in results. Place a straight edge across the top surface of the mould and measure, to the nearest 0.5 mm, the depth from the straight-edge to the surface of the specimen. Take at least six readings from six different locations around the mould and record.

Step D2 Weigh the mould complete with the specimen to the nearest 10 g and record.

Step D3 Remove the compacted specimen from the mould and place it in a small pre-weighed tray. Immediately take a portion over the full height of the specimen and determine the Water Content as specified in the NZS 4402 Test 3.1 “The Water Content of Aggregate” (New Zealand Standards, 1991a).

Step D4 Perform calculations to determine Water Content – Dry Density curve as specified in the NZS 4402 test 4.1.3 “New Zealand vibrating hammer compaction test” (New Zealand Standards, 1986b).

6.5 X-ray Diffraction Test Methodology

6.5.1 Sample Preparation

The source aggregate was examined and split into three different types. Each type represented aggregates with a similar physical appearance and particle size. Figure 6-4 shows the three different types.
Two types of samples are usually required for the X-ray diffraction test. Each type requires a slightly different method in sample preparation. These methods will be discussed below.

**Bulk or Random Orientation Sample**

This type of sample is used to identify the mineral constituents of the aggregate and also to determine the proportion of minerals in the aggregate. The methodology used in preparation of this type of sample is as follows:

1. The aggregate which contains coarse chips is reduced in grain size by crushing it in a Rocklabs steel ring mill.
2. The Rocklabs steel ring mill should not be used to powder the sample as this may damage and distort the mineral grains in the sample. Instead, the sand sized sample is hand powdered in a pestle and mortar.
3. The powder is then loosely packed into an aluminium holder. Care should be taken when packing the powdered sample, it should not be pressed as this may risk damaging and orienting the mineral grains in the sample.
4. Once packed into the aluminium holder, the sample is then ready to be inserted into the X-ray diffractometer.

**Oriented Sample**

These samples are used to determine the nature of the clay minerals. The sample preparation procedure is as follows:

1. The powdered bulk sample can be used for this method.
2. Approximately 1.5 ml of the powdered sample is deposited into a plastic test tube.
3. The tube is then filled with distilled water and hand shaken until the sample is completely dispersed in the distilled water.
4. The tube is then left to settle for approximately 20 – 30 minutes. Subsequently, a portion is drawn off from the top 2 ml of the test tube and carefully deposited onto a glass slide and allowed to spread over an area of around 20 mm in diameter.
5. The glass slide is then air dried in a dust-free environment to allow the clay particles in the sample to sediment onto the glass slide.
6. The oriented sample is X-rayed three times. Initially as an untreated sample. Followed by another X-ray on the now glycolated sample. And finally the glycolated sample is heated at 550°C for at least an hour before it is X-rayed for the last time. Performing the X-ray on the glycolated sample allows for the identification of zeolites present in the aggregate which usually cause volume changes such as swelling. Swelling is an unwanted phenomenon in aggregate properties as it negatively affects the performance of the aggregate. The final X-ray on the heated sample reveals the collapsed dehydrated basal layer of the clays.
6.5.2 Experiment Set-up

The set-up of the experiment is shown in Figure 6-5. The goniometer is computer controlled and data is obtained by using the XRD software. After inserting the sample slide into the diffractometer, a printout is outputted from the computer.
Chapter 7. **TEST RESULTS AND DISCUSSION**

### 7.1 Aggregate Property Test Results

Results of the conducted quality control tests will be presented and their significance discussed in this Chapter. A number of different tests have been carried out to identify the physical properties and mineral composition of the aggregate being used for testing. The analysis of these test results is imperative to better understand the level of quality of the source aggregate being used for testing. The classification of the aggregate property tests will be similar to that in Section 3.2.2 where tests fall under “Source Property Tests” or “Production Property Tests”, depending on the nature of the test.

#### 7.1.1 Source Property Tests

**Crushing Resistance**

The TNZ M/4 basecourse specification states that the test should done under a load of 130 kN where fines passing the 2.36 mm sieve as a result of the load application must be less than 10% to deem the aggregate of acceptable strength. The test returned a result of 0.5% of fines passing the 2.36 mm sieve under the specified 130 kN load. This result indicates that the source aggregate being used for testing is of high strength quality, however during the vibrating hammer compaction tests, slight degradation to the aggregate was visually observed.

**Weather Quality**

Figure 7-1 shows the results obtained from the weathering quality tests conducted. The graph displays the different indices based on the cleanness value and percentage of sample material retained on the 4.75 mm sieve. The TNZ M/4 specification for basecourse aggregates allows weathering quality values of AA, AB, AC, BA, BB or CA. Thus, the results obtained for weathering quality are ‘AA’ as shown by the highlighted cell in Figure 7-1. The aggregate displays an extremely high resistance to environmental effects. In comparison with the specification, the aggregate obtained the highest permissible weathering quality index of ‘AA’ with a cleanness value of 98 with 98% of particles being retained on the 4.75mm sieve. Before the oven drying process, the bulk
material used for the vibrating hammer compaction test was left outside due to space constraints in the laboratory. Thus, weathering of the material was a concern, however the results shown here indicate that the material is highly resistant to environmental effects (such as wetting, drying, heating and cooling). Hence, these results imply that environmental effects on the source aggregate that has been left outside should not be a concern as the aggregate appears to be highly resistant to the environmental effects. In saying that, concern has been expressed regarding the weathering quality test, as it is believed that it does not reflect real environmental weathering conditions. Weathering effects should nevertheless be controlled as best as possible in laboratory testing to help reduce inconsistencies in results.

![Image of weathering quality results](image)

**Figure 7-1: Weathering Quality Results**

*California Bearing Ratio*

Table 7-1 summarises the results obtained from the California Bearing Ratio test and also the specifications that the results must meet in order to pass the test. As can be seen the minimum permissible CBR for a material being used as a basecourse layer is 80%, the result obtained is well above that at 275%. The CBR test provides an idea of the strength of the material being tested and in this case the source aggregate is considered high quality having a relatively high strength characteristic.
Table 7-1: California Bearing Ratio Results

<table>
<thead>
<tr>
<th>Detail</th>
<th>Results</th>
<th>Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Period of Soaking (days)</td>
<td>4</td>
<td>Minimum 4</td>
</tr>
<tr>
<td>Compacted Dry Density (t/m³)</td>
<td>2.24</td>
<td>-</td>
</tr>
<tr>
<td>Compacted Water Content (%)</td>
<td>5.0</td>
<td>-</td>
</tr>
<tr>
<td>Soaked Water Content (%)</td>
<td>5.9</td>
<td>-</td>
</tr>
<tr>
<td>Rate of Penetration (mm/min)</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Depth CBR Recorded (mm)</td>
<td>5.0</td>
<td>0</td>
</tr>
<tr>
<td>California Bearing Ratio CBR</td>
<td>275%</td>
<td>Minimum 80%</td>
</tr>
</tbody>
</table>

7.1.2 Production Property Tests

Sand Equivalent

The TNZ M/4 specification for basecourse aggregates specifies that the Sand Equivalent Index achieved must be above 40 for the aggregate to be compliant. The result obtained from the test was 51, considering the minimum limit is 40, this result is deemed above satisfactory.

Clay Index

The fraction of the aggregate passing the 75 µm sieve shall have a clay index value of less than 3 in order to be compliant to the TNZ M/4 specification for basecourse aggregates. The result obtained from this test method is 0.7; this is relatively low compared to the specification value.

Plasticity Index

The plasticity index of the fraction of aggregate passing the 425µm sieve shall not be greater than five according to the TNZ M/4 specification. The result obtained for this test is five. The result is barely complies with the limit specified. However, the TNZ M/4 specification states that in order for compliance, the aggregate being tested shall pass at least one of the three tests (Sand Equivalent, Clay Index or Plasticity Index). Thus, even though the result barely passes the specification stated, it has already complied with the other two tests mentioned above.
Broken Face Content

Table 7-2 summarises the results obtained from the Broken Face Content test. The results achieved are 100% for all fractions exceeding the specified 70% minimum stated in the TNZ M/4 specification. 100% Broken face content for all particle fractions is considered very good. The angularity and broken faces of particles helps to define its performance when compacted. As these irregularly shaped aggregates are compacted, they provide a much stiffer structure than aggregates with a lower broken face content.

<table>
<thead>
<tr>
<th>Sieve Fraction (mm)</th>
<th>Results (%)</th>
<th>Specification (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;37.5</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>19 – 37.5</td>
<td>100</td>
<td>Minimum 70</td>
</tr>
<tr>
<td>9.5 – 19</td>
<td>100</td>
<td>Minimum 70</td>
</tr>
<tr>
<td>4.75 – 9.5</td>
<td>100</td>
<td>Minimum 70</td>
</tr>
</tbody>
</table>

Particle Size Distribution

The particle size distribution is sometimes referred to as the grading of the aggregate. Grading has a major effect on the performance of the aggregate. Figure 7-2 shows the grading of the aggregate used for testing as well as the upper and lower limits specified by the TNZ M/4 basecourse specification. The dotted line (which represents the grading of the aggregate) lies between the upper and lower specified limits. Table 7-3 provides more detailed information regarding the results obtained. As can be seen, the results are always within the limits specified at each aggregate fraction. Keeping a constant grading/particle size distribution throughout the vibrating hammer compaction tests was a vital step to ensuring the consistency in end results.
Figure 7-2: Source Aggregate Particle Size Distribution

Table 7-3: Shape Control of Source Aggregate

<table>
<thead>
<tr>
<th>Fraction (mm)</th>
<th>Result (mm)</th>
<th>Specification (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>19 – 4.75</td>
<td>33</td>
<td>28 – 48</td>
</tr>
<tr>
<td>9.5 – 2.36</td>
<td>21</td>
<td>14 – 34</td>
</tr>
<tr>
<td>4.75 – 1.18</td>
<td>20</td>
<td>7 – 27</td>
</tr>
<tr>
<td>2.36 – 0.600</td>
<td>16</td>
<td>6 – 22</td>
</tr>
<tr>
<td>1.18 – 0.300</td>
<td>12</td>
<td>5 – 19</td>
</tr>
<tr>
<td>0.600 – 0.150</td>
<td>8</td>
<td>2 – 14</td>
</tr>
</tbody>
</table>
7.2 Vibrating hammer compaction test results

7.2.1 Data Selection

Statistics were used to analyse and determine the extent of variation in the results obtained from testing. Data was analysed in various ways in an attempt to better understand the causes of variation. In addition to this research’s data, the raw data of the Opus inter-laboratory investigation was also included in the analysis to better understand and compare results.

Although forty vibrating hammer compaction tests have been conducted, initial analysis of the data has shown that the first ten test results exhibited a considerably larger variation than the rest of the data as can be seen in Figure 7-3. Consequently, a decision was made to exclude the first ten test results from the data analysis. The exclusion of the first ten tests can be justified by the fact that the researcher was initially inexperienced and unfamiliar with the test. After approximately ten tests, results show that the researcher began to gain confidence and progressively improved whilst initially conducting the test.

![Figure 7-3: Contribution (by Percentage) of each Group of 10 Tests to the Total Variation](image)

Figure 7-3 shows that Tests 1 to 10 contribute approximately 40% of the total variation of the test. This is an unrealistically high contribution; the inexperienced researcher is more
likely to make errors than a technician who is experienced. Thus, the inclusion of the first ten tests in the analysis could falsely represent the actual variation present within the test method. Tests 11 to 40 all seem to be contributing a relatively equal amount of variation indicating the true and natural variation of the test method.

### 7.2.2 Overall Variation – Scatter / Box and Whisker Plot

The natural variation in the vibrating hammer compaction test is shown in Figure 7-4. The 2.0% WC data from the Opus Round Robin study have been included in Figure 7-4 for comparison purposes. The data appears to follow the general trend shown in Figure 2-6 where the Dry Density exhibits an initial peak at dry conditions (in this case at 2.0% WC since data for 0% WC was not available). This was then followed by a decrease in the Dry Density at intermediate WC levels and finally a gradual increase again in the Dry Density as the WC increases until saturation (Zero Air Voids) creating another peak at approximately 6.5% WC. The phenomenon of bulking appears to be taking place as supported in the literature review in Section 2.3.2.

This variability in the aggregate source is quite large considering that as many factors that could possibly affect the test have been kept constant. Thus, the variation, which is represented by the spread of the points at each Water Content shown in Figure 7-4 represents the natural variation of the New Zealand vibrating hammer test method. Natural sources such as the test method itself and/or the natural variability in the aggregate properties are to be held responsible for the observed variability.

The variability (excluding that from the Opus data at 2.0% WC) appears to be increasing as the WC increases. This can be explained by the increase in the amount of ‘splashing’ of water and material as the WC increased. As the sample gets wetter it becomes difficult to compact without some loss of material and/or water. This loss contributes to a decrease in mass and hence a variation in results. The New Zealand Standard for the vibrating hammer compaction test method specifies mould and tamper sizes such that there is a gap between the tamper and inner walls of the mould that is approximately 3.5 mm. Although the reason for this gap is not mentioned in the New Zealand Standard, it could simply be the specified gap prevents rubbing between the tamper and inner walls of the mould which can cause interference to the compaction process. However, the issue with the gap
between the tamper and inner walls of the mould is that it allows material and/or water to escape during compaction as shown in Figure 7-6.
Investigation of the Variability in the Results of the NZ Vibrating Hammer Compaction Test

Figure 7-4: Variation of Vibration Hammer Compaction Test
Figure 7-5: Natural Variability in Dry Density at each Water Content
The apparatus used in the British Standard on the other hand, completely surrounds and contains the sample in place to prevent any loss of material and/or water during compaction. The tamper (otherwise known as anvil in the BS) is in contact with the inner walls of the mould and does not allow any gaps and provides full confinement to the sample. A cross sectional diagram of the apparatus used in the BS is shown in Figure 7-7; a similar apparatus should be considered in the New Zealand Standard as even a minor loss of material and/or water can cause significant variation in the end result due to the change in mass.

**Figure 7-6: Effect of the Gap between the Tamper and the Mould on the Loss of Water and/or Material**

Whilst it is difficult to measure, phenomena such as segregation and degradation were visually noted during compaction. It is unknown whether segregation and degradation contributed to the variation observed in the results and if so to what extent. This should be investigated in future research. According to past research segregation is inevitable during the compaction process, however as mentioned in Section 4.2.2 it can be substantially reduced by applying an adequate amount of surcharge weight.

The large variability in the Opus data at 2.0% WC is mostly contributed to the nature of the study. The Opus round robin study involved thirty three different laboratories and therefore the larger variation can be explained by the different technicians and equipment used by each laboratory.
As can be seen from Figure 7-4, some points at 6.5% WC have crossed the Zero Air Voids (ZAV) line which is theoretically impossible. This could be due to human error during the compaction test.

![Figure 7-4 Cross-section of Mould and Tamper Apparatus used in BS 5835 (BS 5835, 1980)](image)

Although the range of variability in DD only appears to be quite small (approximately 0.15 t/m³) minor uncertainties in the DD is largely pronounced in the degree of compaction achieved.

The Dry Density variability at each Water Content is represented by a box and whisker plot in Figure 7-5. The medians (and means, as are shown in Table 7-4) are gradually increasing as the WC increases. This follows the same trend presented in Figure 2-6 for cohesionless material, where a relatively high increase in DD from intermediate WC to high WC levels is noted. Followed by a minor increase in DD where the curve begins to level off before reaching saturation.

The median at 5.5% is slightly lower than its preceding WC; this however, can be explained by the large variability exhibited at this WC which seems to have caused the median to reduce. The mean on the other hand, follows the general trend, where it is larger than its preceding WC. The variability observed at 5.5% WC is slightly larger than
that of any other WC value. Although the reason behind this particularly large variability at 5.5% WC is unknown, it could be due to a number of factors such as errors in test procedures and recording of measurements at that particular WC. However, as seen in Table 7-4 the coefficient of variation at 5.5% WC is not significantly larger than that for 6.0% and 6.5% Water Contents.

The spread of data at each Water Content shown in Figure 7-5 could be due to interlocking of particles during the compaction process. The NZS standard for the vibrating hammer compaction test method allows for particles of 37.5 mm to be compacted in a 152 mm diameter mould. International standards such as the ASTM standard for the vibrating hammer compaction test, consider these particles ‘oversized’ and specify that for compaction of an aggregate with these oversized particles, a much larger mould should be used. Moreover, the ASTM D 7382 specification states that particles retained on the 19 mm sieve should be compacted in a 279.4 mm mould (ASTM D 7382 - 08, 2008). The compaction of an aggregate which contains 37.5 mm particles in a 152 mm mould could lead to the interlocking of these oversized angular particles. Interference to the compaction process can occur once these particles interlock against each other and the inner walls of the mould.

7.2.3 Comparison of Dry Density at 6% Water Content – Opus and UoA

In the Opus Round Robin study, the same TNZ M/4 AP40 material as was tested in this research at Water Contents of 2% and 6% was used. The UoA research however, tested the TNZ M/4 material across a wider range of Water Contents. To observe the variability between studies, a DD box and whisker graph has been plotted for both 6% Water Contents (Figure 7-8).

As would be expected, the variability from the Opus Round Robin test is much greater due to the fact that the study involved testing across a various number of laboratories rather than just one laboratory as in the UoA study. As discussed in chapter 4, the same test being conducted in different laboratories can yield significantly variable results due to the different equipment being used, different technicians and variations in test methodologies used in conducting the test.
If it is assumed that the variability observed in the UoA 6% WC is explained by the natural variability present within the test method and natural variability in the source aggregate, Figure 7-8 shows that an approximated 30% (ratio of the size of the UoA box-plot to the size of the Opus box-plot) of the variability can be contributed to the natural variability of the test method/source aggregate; this means factors such as unreliability in test methods, and natural variability of the material are to be held accountable for 30% of the total variation. The other 70% of the variation can be attributed to the variability in different equipment used to conduct the test, the different technicians, and in general between-lab variation.

The median in the UoA 6% WC is slightly higher than that obtained in the Opus 6% WC, this could be due to the types of hammers used in both studies. The UoA research conducted all tests by using the same hammer which had an input power of 1,150 W, while the Opus study involved using a variety of different hammers with input powers ranging from 750 W to 1700 W. Lower end power input hammers produce relatively low
Dry Density values and hence hammers used in the Opus study with a relatively low input power could be the reason for the decrease in the median in the results.

It is important to note that the majority of residual Water Contents of the Opus Study at 6% were considerably lower than 6%; in fact the average of residual water contents at 6% was a low 4.9%. There has been considerable water loss during compaction in the Opus study, although the UoA study has experienced problems with water loss during compaction, an acceptable variance level of 0.3% was put in place, that is to say if the sample had lost more than 0.2% of its water content during compaction then that result was rejected a new sample compacted instead. Opus did not have a similar restriction in place, and thus part of the variation experienced in the Opus data can be explained by this highly variable residual Water Content.

### 7.2.4 Numerical Comparison of data

Table 7-4 provides statistical results based on the analysis of the raw test data. The number of samples analysed from the UoA data was 30 samples at each WC after the first initial ten tests were removed. The Opus data on the other hand, had tested 66 samples at each WC.

By comparing the statistical range of values in Table 7-4, it can be seen that, as would be expected, the ranges for the Opus data are significantly larger than any range obtained for the UoA data. However, the comparison of the range value has its limits due to the fact that it is defined as the difference between the maximum and minimum value, where outliers in the data can influence this statistic significantly.

The standard deviation and Coefficient of Variation (CoV) provide a much more genuine and accurate representation of the degree of variability in the data. Thus, by comparing the standard deviation, it can be seen that the Opus data has values larger than that of the UoA. The standard deviation in the UoA data appears to increase gradually as the WC increases, however at 5.5% WC, the standard deviation is larger than WC values subsequent to it. This was not expected from the results as it was thought that variation tended to increase as the WC increased due to the fact that the wetter the sample gets the more the problem of water splashing/loss of fine material during compaction was pronounced. The CoV follows a very similar trend to the standard deviation in that there is a gradual increase in CoV as the WC increases, with the same exception observed in
the standard deviation at 5.5% WC where the CoV is slightly larger than its subsequent WC values.

The mean values on the other hand, confirm the general trend (with no exceptions at 5.5% WC) of a gradual increase in DD from intermediate levels of WC to high levels of WC followed by a slow levelling off of the curve at approximately saturation levels (6.5% WC).

The repeatability values shown in Table 7-4 for the UoA results (0.07 to 0.13 t/m³) are quite large when compared to other repeatability studies by ASTM D 7382 and BS EN 13286 where repeatability values are 0.033 t/m³ and 0.05 t/m³ respectively. Since repeatability values are based on tests conducted in the same laboratory, by the same technician, using the same apparatus, a high repeatability value suggests a natural variation in the test procedure or the aggregate being tested.

It was concluded in the Opus Interlaboratory study that the repeatability values vary with the material being tested but are not significantly affected by the target water content. Repeatability values for the TNZ material tested in the Opus study at 2% and 6% WC were both 0.078 t/m³ (Opus International Consultants Limited, 2008). On the other hand, the UoA results shown in Table 7-4 suggest otherwise, where repeatability values increase as the Water Content increases.
<table>
<thead>
<tr>
<th>Water Content (%)</th>
<th>Number of Samples</th>
<th>Range</th>
<th>Minimum</th>
<th>Maximum</th>
<th>Mean</th>
<th>Standard Deviation</th>
<th>Skewness</th>
<th>Coefficient of Variation (CoV)</th>
<th>Repeatability</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Opus</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>66</td>
<td>0.290</td>
<td>2.020</td>
<td>2.310</td>
<td>2.177</td>
<td>0.059</td>
<td>-0.040</td>
<td>0.027</td>
<td>N/A</td>
</tr>
<tr>
<td>6</td>
<td>66</td>
<td>0.770</td>
<td>1.930</td>
<td>2.700</td>
<td>2.248</td>
<td>0.113</td>
<td>1.289</td>
<td>0.050</td>
<td>N/A</td>
</tr>
<tr>
<td><strong>UoA</strong></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.5</td>
<td>30</td>
<td>0.080</td>
<td>2.118</td>
<td>2.198</td>
<td>2.159</td>
<td>0.026</td>
<td>-0.139</td>
<td>0.012</td>
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<tr>
<td>4.0</td>
<td>30</td>
<td>0.119</td>
<td>2.114</td>
<td>2.233</td>
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<td>0.031</td>
<td>0.020</td>
<td>0.014</td>
<td>0.085</td>
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<tr>
<td>4.5</td>
<td>30</td>
<td>0.128</td>
<td>2.126</td>
<td>2.254</td>
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<tr>
<td>5.0</td>
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<td>2.290</td>
<td>2.226</td>
<td>0.031</td>
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<td>0.086</td>
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<td>2.307</td>
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<tr>
<td>6.0</td>
<td>30</td>
<td>0.155</td>
<td>2.168</td>
<td>2.323</td>
<td>2.255</td>
<td>0.042</td>
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<td>0.019</td>
<td>0.116</td>
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<tr>
<td>6.5</td>
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<td>2.182</td>
<td>2.330</td>
<td>2.259</td>
<td>0.043</td>
<td>-0.140</td>
<td>0.019</td>
<td>0.120</td>
</tr>
</tbody>
</table>
7.2.5 Sample Size needed to Achieve Target Confidence Interval

In Figure 7-9 a graph is shown that represents the number of samples required to be tested in order to achieve a certain confidence limit. As would be expected, targeting a high allowable error such as 10% yields a relatively low number of samples to be tested. On the other hand, as the allowable error target decreases, that is to say as the precision target of the test increases the number of samples rapidly increases following an exponential behaviour.

**Figure 7-9: Number of Samples needed to be Tested to Achieve a Desired Precision**

All WC curves follow the same general trend with little difference in the quantity of samples to be tested at each allowable error. As discussed previously, since the variability at 5.5% WC is the largest, the curve experiences the most number of samples required to be tested out of all other Water Contents at each allowable error. The rest of the curves follow the logical trend of an increase in variation as the WC increases and therefore an increase in the number of samples as the WC increases at each allowable error.

The curves shown in Figure 7-9 are based on the CoV values of each WC, thus, the number of samples that should be tested to attain a certain target of precision in the test is based on the UoA data. The number of samples to achieve only 90% precision in the test
stands quite large at 50 samples. Hence, the variation associated with this test is significant and a revision of the test method should be considered with further research.

7.2.6 Scatter of Opus Data with Respect to UoA 95% Confidence Interval

![Opus Dry Density at 6% WC and 95% Confidence Limits of UoA 6% WC](image)

**Figure 7-10: Scatter plot of Opus 6% WC with Regards to 95% Confidence Interval of UoA 6% WC**

Figure 7-10 shows the scatter of the Opus data with reference to the upper and lower 95% confidence limits of the UoA data.

As expected only a small portion of the Opus data falls within the upper and lower 95% confidence limits of the UoA data. This proves that between-laboratory testing creates a much higher variation than testing within a single laboratory. The variation observed by the Opus data is mainly contributed to factors such as the differences in technicians conducting the test, different hammers and moulds used. Although all technicians follow the same test standard, each has their own ways of conducting the test according to their interpretation of the test standard. Thus, this puts an emphasis on the importance of the
clarity of a test standard to avoid different interpretations by different laboratory technicians.

This graph once more emphasises the degree of variability of the test if carried out in different labs rather than just one lab. Only roughly around 10% of the Opus data falls within the limits. The limits are based on the data obtained from the testing in this research where 30 tests have been conducted under exactly the same conditions, thus, the reason for the small gap between the two limits.

7.2.7 Effect of Different Compactive Efforts on Compaction

In addition to the thirty vibrating hammer compaction tests conducted, Standard Proctor and Modified (heavy) Proctor tests were carried out along with an additional five vibrating hammer compaction tests conducted with a much more powerful hammer than the one initially used, as shown in Figure 7-11. These tests were carried out for comparison purposes to observe the differences when conducting compaction in different ways.

An average of every 5 tests from the total thirty tests of the initial hammer was taken. The more powerful hammer (Kango 950K) has an input power of 1700 Watts which is 550 Watts more than the initial Metabo hammer used. The two hammers have passed the calibration test specified in the NZS 4402 Test 4.1.3 test method. Yet both have produced significantly different results as can be seen in Figure 7-11. The Kango-based tests produced higher DD values as expected, as it is the more powerful hammer. There has been growing criticism about the calibration test within the NZS vibrating hammer test method due to the fact that it only specifies a minimum limit over the power of hammer and not a maximum. Therefore, significantly more powerful hammers can be used for the test since the test standard does not enforce a maximum limit. These powerful hammers however, can easily cause degradation of the sample. A way forward to this problem is to develop a correlation between the powers produced by field compactors and laboratory hammers used for compaction. This will ensure the MDD obtained in the laboratory better predicts the compaction achieved in the field at the same OWC specified by the laboratory.

The results shown in Figure 7-11 align with what has been discussed in the literature review. The application of a higher compactive effort yields higher Dry Density values as
shown in Figure 7-11. The Standard Proctor method applies the least amount of compactive effort onto the sample and hence has yielded the lowest Dry Density results. This is followed by the Modified Proctor method, which as discussed in Section 2.4.1 applies a greater amount of compactive effort than the Standard Proctor; this is evident by the Dry Density values achieved which are higher than that for the Standard Proctor. The vibrating hammer compaction test method is known to apply a greater compactive effort than both of the Proctor tests; this is evident by the Metabo Dry Density values shown in Figure 7-11. The Kango hammer is known to produce an even greater compactive effort than that of the Metabo due to its powerful 1700 W power input. Evidently the Dry Density values achieved by the Kango hammer are much higher than the Metabo hammer.

The curves produced by the Metabo hammer appear to be erratic, while the Kango hammer curves are considerably more consistent. This could be due to the fact that the Metabo hammer is not delivering a sufficient amount of compactive effort on the sample during compaction and hence not achieving the optimum Dry Density results. During the testing phase at the Stevensons laboratory, there was a transition from using the Metabo hammer to using the Kango hammer for the vibrating hammer compaction tests. This change simply came about due to the fact that the Kango hammer produces much more consistent results as opposed to the erratic curves produced by the Metabo hammer as can be seen in Figure 7-11.
Chapter 7. Test Results and Discussion

Figure 7-11: Effects of Different Compactive Efforts on Compaction
7.3 X-ray Diffraction Test Results

7.3.1 Bulk Sample Results

Figure 7-12 and Figure 7-13 show two of the various graphs obtained from the XRD tests. The graphs show the minerals found in two types of the source aggregate. As explained in
Section 6.5, the aggregate was classified into 3 types. Samples were grouped based on their similar physical features. The two graphs shown in Figures 7-12 and 7-13 represent XRD tests for Types B and C. The amount of minerals available within each type is represented by the curves shown in Figure 7-12 and Figure 7-13. As can be seen, the quantity of quartz is much higher in Type C than Type B, and conversely the quantity of Feldspar in Type B is higher than type C. The amount of clays present in rock types B and C also differs. Chlorite seems to be generally larger in quantity in rock type B, while illite is present in larger amounts in rock Type C.

Thus, there are differences within the same source aggregate; however it is unknown to what extent, these differences contribute to the variation of the vibratory hammer compaction results. A correlation must be investigated between the differences in mineral composition and the variation in the vibrating hammer compaction test to see if these differences do play a dominant role in the variability in results or not.

7.3.2 Oriented Sample Results

The oriented sample was X-rayed twice. Once untreated and again when it has been glycolated to note if there are any volume changes/swelling occurring in the aggregate due to zeolites (swelling clays). However, as can be seen from Figure 7-14 and Figure 7-15, the peaks appear to be relatively equal with the exception of the beginning of the curve where the untreated peak is at 400 counts and the glycolated peak is at roughly 320 counts.

Figure 7-14: Output of X-ray Diffraction Oriented Sample A – Glycolated
7.4 Power Output Test

7.4.1 Introduction

As per the research tasks, a test measuring the power output of the vibrating hammer was developed. The test was to be conducted twice, once before the vibrating hammer compaction testing phase started and again after the phase was completed. The purpose of conducting the same test twice was to note any differences in the results between the two identical tests. Due to the fact that these commercial vibrating hammers are heavy duty and usually last several years, an investigation of the ageing of these hammers on the contribution of the significant variability within the test is essential. The Opus Inter-laboratory study provided information on the age of hammers in each laboratory. Based on this information, the average age of hammers in NZ laboratories is approximately 8.5 years old.

Theoretically speaking, the vibrating hammer degrades and loses its full power as it gets older. This reduction in power could be contributing to the variation in the vibrating hammer compaction test results.

Figure 7-16 presents the experiment set up and identifies the components and equipment used in the test to allow for the effective measurement of the hammer’s power output.


7.4.2 Experiment Set-up

![Experiment Set-up Diagram]

**Figure 7-16: Power Output Experiment Set-up**

The purpose of the test was to measure the displacement (using the portal gauge shown in Figure 7-16) produced by the hammer when a certain amount of weight is imposed on the tamper. The displacement which is measured by the portal gauge was expected to resemble a wavelength where it will oscillate about the x axis. The portal gauge which is supported by two rods as shown in Figure 7-16; one rod that is embedded into a wooden block and the other rod which is attached to the tamper to record vertical displacements.

7.4.3 Problems with the test

Unfortunately the test did not produce understandable test results as was initially hoped. The problem faced during the test was that the portal gauge was not very sensitive to register every movement created by the vibrating hammer. As the hammer was operating, movement of the tamper was too rapid for the portal gauge to record and thus the output obtained gave a false impression of what was actually happening in reality.
Another challenging problem faced was keeping the weights attached to the tamper. The use of clamps allowed for hammer operation for approximately 30 seconds, thereafter the clamps began to loosen and the weights would fall off. Due to time constraints, it was not possible to redevelop the test to turn it into a viable one. However, it is hoped that future research investigates the possibilities of the development of a new and feasible test.
Chapter 8. **CONCLUSIONS AND RECOMMENDATIONS**

8.1 Conclusions

The results of this research have proven that the natural variability associated with the New Zealand vibrating hammer compaction test is a lot higher than international experience has shown. Repeatability values stated in the USA and UK standards for the vibrating hammer compaction test method are 0.05 t/m³ and 0.033 t/m³ respectively. In comparison to this study’s repeatability value of 0.086 for 5% Water Content, UoA repeatability values are significantly larger than those obtained in international standards. This means that either the New Zealand vibrating hammer compaction test procedure is less precise or the source aggregate being used is more variable than those procedures and materials being used in the USA and UK.

The natural variation within the source aggregate used for testing in this research explains approximately 30% of the total variation observed by the Opus Round Robin Inter-laboratory tests. The other 70% can be explained by the between-laboratory differences in equipment and technician level of experience.

Due to suspected interlocking of oversized particles during compaction, the New Zealand Standard for the vibrating hammer compaction test should specify that vibratory hammer compaction tests be conducted on aggregates passing the 19 mm sieve. Material retained on the 19 mm sieve should be discarded.

The NZS 4402 Test 4.1.3 allows for particles of 37.5 mm to be compacted in a 152 mm diameter mould. International standards such as the USA standard consider these particles ‘oversized’ and state that these particles should not be compacted in such a small mould due to boundary effects. The USA standard for the vibrating hammer compaction test (ASTM D 7382 - 08, 2008) specifies that compaction of aggregates with particles retained on the 19 mm sieve should be compacted in a 279.4 mm diameter mould. Compacting particles passing the 37.5 mm sieve in a 152 mm diameter mould can lead to the interlocking of these oversized angular particles during the compaction process. Consequently, the interlocking of these particles interferes with effective compaction.
The amount of splashing of water and/or loss of material during the vibratory hammer compaction test increased as the Water Content of the sample increased. The loss of water and/or material during compaction has a direct effect on the dry density achieved. Thus, as the loss of water and/or material increases, the variability also increases. The apparatus specified in the NZS 4402 Test 4.1.3, particularly the mould and tamper size, cause a greater of the loss of water and/or material. The mould and tamper sizes are specified such that there is a permissible gap present between the tamper and inner walls of the mould, the loss of water and material is escaping through this gap.

The Coefficient of Variation of the 30 test samples ranged from 0.012 at 3.5% Water Content to 0.019 at 6.5% Water Content. The Coefficient of Variation at 5.5% Water Content is particularly larger than any other Water Content. The Dry Density values experienced at this Water Content are quite variable. This could be attributed to the amount of water/material lost during compaction.

A comparison between the previous Opus research and the UoA data demonstrated that, as expected, the variability in the Opus data is significantly larger than the UoA data. This can be explained by the different approaches and objectives of each study. The Opus investigation conducted the vibrating hammer compaction test in thirty three different laboratories to investigate the between-laboratory variation of the test method. Clearly, this variation would be much larger than conducting repeat tests of the vibrating hammer compaction test within the same laboratory as is the case for the UoA data. Different technicians conducting the same experiment using different apparatus induced a much larger variation in test results than test results obtained from an experiment conducted under the same technician and same apparatus.

The initial ten tests have been excluded from the data analysis as it was found these increase variability due to the researcher becoming familiar with the test and gaining experience. The first ten test results have shown to contribute approximately 40% to the total variation in the test results, that is approximately double the contribution of any other set of ten tests in the data. The exclusion of these first ten tests is justified by the fact that the researcher was initially inexperienced and unfamiliar with the test method.

As the target confidence interval of the test increases, the number of samples to be tested also increases. In addition, due to the fact that the variability increases as the Water
The number of samples increases even further as the target confidence interval increases.

As expected, very few points of the Opus 6% WC data fall within the 95% confidence interval limits of the UoA 6% WC. This proves that the variability associated with the Opus results is far greater than that in the UoA results.

Different methods of compaction yield different results. The results in Figure 7-11 show that as the compactive effort is increased, higher Dry Density values were achieved. The Standard Proctor test method achieved the lowest Dry Density values, followed by the Modified Proctor test method. The modified (heavy) Proctor test method is largely based on the standard Proctor method but utilises a significantly higher compactive effort. The Metabo hammer of the vibrating hammer compaction method achieved higher Dry Density values than the Standard and Modified Proctor methods. Lastly, the Kango hammer of the vibrating hammer compaction method achieved the highest Dry Density values. The Kango hammer is a much more powerful hammer than the Metabo (producing a higher compactive effort) and seems to produce a much clearer Water Content – Dry Density relationship.

The Metabo hammer curves (1-30) are noticeably erratic, while the Kango curves are relatively more consistent. This could be due to the fact that the Metabo hammer is not delivering sufficient compactive effort onto the sample to achieve a consistent behaviour from the aggregate.

The hammer calibration method incorporated in the New Zealand Standard requires updating, as although it specifies a minimum limit on the hammer power spectrum, a maximum limit does not exist. Furthermore, the minimum limit appears to be quite low, as a hammer with apparent insufficient power can pass the calibration test. However, these hammers’ power may not be sufficient to effectively compact the sample.

Analysis of production and source tests of the aggregate has shown that the aggregated used in the testing phase of this research is of high quality.
8.2 Summary of Conclusions

This research investigated the variability in the results of the New Zealand vibrating hammer compaction test. Based on the comprehensive review of literature and the analysis of results, the investigation has found the following:

- The natural variability within an aggregate explains approximately 30% of the variation observed by the Opus Inter-laboratory study.
- The compactive effort applied to the sample during compaction governs the degree of Dry Density achieved.
- Repeatability values obtained are significantly larger than those stated by similar studies conducted in the USA and UK.
- The New Zealand vibrating hammer compaction mould of 152 mm nominal diameter should only use a maximum particle size of 19 mm due to the suspected effect of interlocking of particles.
- The mould and tamper apparatus specified in the New Zealand Standard for the vibrating hammer compaction test should be reconsidered due to the significant loss of water/material during compaction through the permissible gap between the tamper and inner walls of the mould.
- The between-laboratory variability as observed in the Opus investigation, is much larger than that observed in the within a single laboratory variation in the UoA study.
- Compaction curves produced by the Metabo hammer appear to be erratic, while the Kango hammer curves are much smoother and consistent. The Metabo hammer appears to be delivering an insufficient amount of compactive effort on the sample.
- The laboratory vibrating hammer compaction test for aggregates produces a much flatter and less definable shape of curve. It is sometimes difficult to determine the Optimum Water Content and Maximum Dry Density values are for field compaction.
8.3 Recommendations and Future Research

The extent of segregation and degradation occurring during the vibrating hammer compaction test should be further investigated. Research is required to determine to what extent these two phenomena have on the affect of the precision of the test results.

The development and implementation of a hammer power output measurement test is required. The test will help determine whether the hammer output decreases as the hammer gets older, and whether the decrease in hammer power output has an effect on the DD results achieved.

It is recommended that similar apparatus to that used in the BS 5835 for the mould and tamper, be adopted and incorporated in the NZS for the vibrating hammer compaction test method (New Zealand Standards, 1986b). This will help in the prevention of water and/or material loss during compaction through the permitted gap in the current New Zealand Standard test method.

Vibrating hammer compaction testing should be conducted on samples with particles passing the 19.0 mm sieve to observe whether the variability has slightly decreased or not. If the variability decreases then interlocking of particles can be determined to be interfering with compaction and consequently, affecting the reliability of the test results.
REFERENCES


