Yucca Mountain Site Characterization Project

Bulk and Mechanical Properties of the Paintbrush Tuff Recovered from Boreholes UE25 NRG-2, 2A, 2B, and 3: Data Report

P. J. Boyd, R. H. Price, R. J. Martin, J. S. Noel

Prepared by
Sandia National Laboratories
Albuquerque, New Mexico 87185 and Livermore, California 94550
for the United States Department of Energy
under Contract DE-AC04-94AL85000

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BULK AND MECHANICAL PROPERTIES OF THE PAINTBRUSH TUFF
RECOVERED FROM BOREHOLES UE25 NRG-2, 2A, 2B, AND 3:
DATA REPORT

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ABSTRACT

Experimental results are presented for bulk and mechanical properties measurements on specimens of the Paintbrush tuff recovered from boreholes UE25 NRG-2, 2A, 2B, and 3, at Yucca Mountain, Nevada. Measurements have been performed on Timber Mountain tuff and two thermal/mechanical units, TCw, and Ptn of the Paintbrush tuff. On each specimen the following bulk properties have been reported: dry bulk density, saturated bulk density, average grain density, and porosity. Unconfined compression to failure, confined compression to failure, and indirect tensile strength tests were performed on selected specimens recovered from the boreholes. In addition, compressional and shear wave velocities were measured on specimens designated for unconfined compression and confined compression experiments. Measurements were conducted at room temperature on nominally water saturated specimens. The nominal strain rate for the fracture experiments was $10^{-5}\text{s}^{-1}$.
This report was prepared under the Yucca Mountain Site Characterization Project WBS number 1.2.3.2.7.1.3. The planning documents that guided this work activity are Site Characterization Plan Section 8.3.1.15.1.3; Study Plan SP-8.3.1.15.1.3, Revision 0; and Work Agreement WA-0090. The information and data documented in this report are qualified and may be used in licensing the repository.
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## CONTENTS

<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>ABSTRACT</td>
<td>i</td>
</tr>
<tr>
<td>CONTENTS</td>
<td>iii</td>
</tr>
<tr>
<td>1.0 INTRODUCTION</td>
<td>1</td>
</tr>
<tr>
<td>2.0 EXPERIMENTAL PROCEDURE</td>
<td>6</td>
</tr>
<tr>
<td>2.1 Sample Preparation</td>
<td>6</td>
</tr>
<tr>
<td>2.2 CT Scans of Each Specimen Tested in Unconfined Compression</td>
<td>8</td>
</tr>
<tr>
<td>2.3 Drying, Saturation, Bulk Density, Average Grain Density, and Porosity</td>
<td>8</td>
</tr>
<tr>
<td>2.3.1 Procedure for Drying a Specimen</td>
<td>8</td>
</tr>
<tr>
<td>2.3.2 Procedure for Water Saturation</td>
<td>9</td>
</tr>
<tr>
<td>2.3.3 Average Grain Density Measurement Using Water Pycnometry</td>
<td>11</td>
</tr>
<tr>
<td>2.3.4 Dry Bulk Density, Saturated Bulk Density, and Porosity</td>
<td>14</td>
</tr>
<tr>
<td>2.4 Compressional and Shear Wave Velocity Measurements</td>
<td>14</td>
</tr>
<tr>
<td>2.4.1 Detailed Procedures for Compressional and Shear Wave Velocity Measurements</td>
<td>18</td>
</tr>
<tr>
<td>2.5 Unconfined Compression to Failure</td>
<td>19</td>
</tr>
<tr>
<td>2.5.1 Experimental Procedures For Unconfined Compression Tests</td>
<td>22</td>
</tr>
<tr>
<td>2.6 Confined Compression to Failure</td>
<td>25</td>
</tr>
<tr>
<td>2.7 Indirect Tensile Strength Tests</td>
<td>26</td>
</tr>
<tr>
<td>2.7.1 Experimental Procedures for Indirect Tensile Strength Tests</td>
<td>26</td>
</tr>
<tr>
<td>3.0 RESULTS</td>
<td>29</td>
</tr>
<tr>
<td>3.1 Computerized Tomographic X-ray Images</td>
<td>45</td>
</tr>
<tr>
<td>3.2 Compressional and Shear Wave Velocity Measurements</td>
<td>45</td>
</tr>
<tr>
<td>3.3 Unconfined Compression Tests</td>
<td>49</td>
</tr>
<tr>
<td>3.4 Confined Compression Tests</td>
<td>49</td>
</tr>
<tr>
<td>3.5 Indirect Tensile Strength Tests</td>
<td>52</td>
</tr>
<tr>
<td>4.0 REFERENCES</td>
<td>53</td>
</tr>
<tr>
<td>APPENDICES</td>
<td></td>
</tr>
<tr>
<td>I: Stress vs Axial Strain and Radial Strain vs Axial Strain Plots for Unconfined Compression Experiments</td>
<td>55</td>
</tr>
<tr>
<td>II: Stress vs Axial Strain and Radial Strain vs Axial Strain Plots for Confined Compression Experiments</td>
<td>80</td>
</tr>
<tr>
<td>III: System Checks using an Aluminum Standard Specimen</td>
<td>85</td>
</tr>
<tr>
<td>IV: Information From the Reference Information Base</td>
<td>89</td>
</tr>
</tbody>
</table>
List of Figures

Figure 1a: Stratigraphic and thermal/mechanical units; UE25 NRG-2 ................. 2
Figure 1b: Stratigraphic and thermal/mechanical units; UE25 NRG-2A ............... 3
Figure 1c: Stratigraphic and thermal/mechanical units; UE25 NRG-2B ............... 4
Figure 1d: Stratigraphic and thermal/mechanical units; UE25 NRG-3 ............... 5
Figure 2: Bulk properties worksheet ............................................................. 13
Figure 3: Geometry used to measure ultrasonic velocities ................................. 16
Figure 4: Apparatus used to measure ultrasonic velocities ............................... 17
Figure 5: Axial and radial deformation instrumentation .................................. 21
Figure 6: Geometry used to measure indirect tensile strength ......................... 27
Figure 7: CT scan for a PTn specimen from UE25 NRG-2A at a depth of 96.0 feet 46
Figure 8: CT scan for a TCw specimen from UE25 NRG-3 at a depth of 218.0 feet 47
Figure 9: CT scan for a TCw specimen from UE25 NRG-3 at a depth of 289.2 feet 48
Figure 10: Axial stress and radial strain as a function of axial strain ................ 50
Figure 11: Axial stress and radial strain as a function of axial strain (TCw) ....... 51
Figure A-1: Axial stress and radial strain as a function of axial strain (system check) 87

List of Tables

Table 1: Unconfined compression tests data summaries .................................. 30
Table 2: Confined compression tests data summaries ..................................... 37
Table 3: Indirect tensile strength tests data summaries .................................. 38
Table 4: Porosity only data summaries ......................................................... 42
Table A-1: System checks .......................................................... 88
1.0 INTRODUCTION

An integral part of the licensing procedure for the potential nuclear waste repository at Yucca Mountain, Nevada, involves characterization of the in situ rheology for the design and construction of the facility and the emplacement of canisters containing radioactive waste. The data used to model the thermal and mechanical behavior of the repository and surrounding lithologies include dry and saturated bulk densities, average grain density, porosity, compressional and shear wave velocities, elastic moduli, and compressional and tensional fracture strengths. In this study, a suite of experiments was performed on cores recovered from boreholes UE25 NRG-2, 2A, 2B, and 3 drilled in support of the Exploratory Studies Facility (ESF) at Yucca Mountain. The holes penetrated the Timber Mountain tuff and two thermal/mechanical units of the Paintbrush tuff. The thermal/mechanical stratigraphy was defined by Ortiz et al. (1985) to group rock horizons of similar properties for the purpose of simplifying modeling efforts. The relationship between the geologic stratigraphy and the thermal/mechanical stratigraphy for each borehole is presented in Figures 1a, b, c, and d. The tuff samples in this study have a wide range of welding characteristics (usually reflected in sample porosity), and a smaller range of mineralogy and petrology characteristics. Generally, the samples are silicic, ash-fall tuffs that exhibit large variability in their elastic and strength properties (see Price and Bauer, 1985).

Seventy-nine cores from UE25 NRG-2, 2A, 2B, and 3 were sent to New England Research, Inc. for bulk and baseline mechanical property measurements. A breakdown of the samples according to tests performed is given below:

<table>
<thead>
<tr>
<th>Type of Test</th>
<th>Number of Samples Tested</th>
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<tr>
<td>Unconfined Compression</td>
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<td>Confined Compression</td>
<td>8</td>
</tr>
<tr>
<td>Indirect Tensile Strength</td>
<td>41</td>
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</table>

On 12 cores there was insufficient material or the material was of such poor quality that mechanical tests could not be performed. On these cores, only the average grain density and porosity were determined.
Figure 1a: The correlation between the stratigraphic and thermal/mechanical units for borehole UE25 NRG-2 at Yucca Mountain, Nevada (per preliminary stratigraphy by J.F.T. Agapito and Associates, Inc., 11/1/93).
Stratigraphic and Thermal/Mechanical Units Summary

**Stratigraphy**

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<thead>
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<td>248.5</td>
<td></td>
</tr>
<tr>
<td>265.7 TD</td>
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</table>

**Thermal/Mechanical Units**

<table>
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</tr>
</thead>
<tbody>
<tr>
<td>165.9</td>
</tr>
<tr>
<td>265.7 TD</td>
</tr>
</tbody>
</table>

Note: Tuff unit "X" (Carr, 1992) is an informal name for numerous pyroclastic flow and fallout tuffs and tuffaceous sedimentary rocks that occur between the Tiva Canyon and Rainier Mesa tuffs.

**Figure 1b:** The correlation between the stratigraphic and thermal/mechanical units for borehole UE25 NRG-2A at Yucca Mountain, Nevada (per preliminary stratigraphy by J.F.T. Agapito and Associates, Inc., 11/1/93).
Figure 1c: The correlation between the stratigraphic and thermal/mechanical units for borehole UE25 NRG-2B at Yucca Mountain, Nevada (per preliminary stratigraphy by J.F.T. Agapito and Associates, Inc., 1/26/94).
Figure 1d: The correlation between the stratigraphic and thermal/mechanical units for borehole UE25 NRG-3 at Yucca Mountain, Nevada (per preliminary stratigraphy by J.F.T. Agapito and Associates, Inc., 11/1/93).
2.0 EXPERIMENTAL PROCEDURE

Measurements were performed on 79 specimens prepared from core recovered from boreholes UE25 NRG-2, 2A, 2B, and 3. When the core was received it was examined to determine the best utilization of the rock material to obtain the maximum mechanical data. The size of the specimen for each type of measurement was a major part of the selection criteria. The nominal dimensions of the cylindrical rock specimens for each of the mechanical tests are given below:

<table>
<thead>
<tr>
<th>Test</th>
<th>Length (mm)</th>
<th>Diameter (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Indirect Tensile Strength</td>
<td>38.1</td>
<td>50.8</td>
</tr>
<tr>
<td>Unconfined Compression</td>
<td>101.6</td>
<td>50.8</td>
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<tr>
<td>Confined Compression</td>
<td>50.8</td>
<td>25.4</td>
</tr>
</tbody>
</table>

The length and diameter have a tolerance of ±0.125 mm. The ends of the cylinders were parallel to within 0.025 mm.

Dry and saturated bulk densities were measured on each of the specimens prepared for unconfined compression, confined compression, and indirect tensile strength tests. Average grain density was determined with the water pycnometry technique using pieces remaining from the subcoring attendant to the preparation of the test specimens.

2.1 Sample Preparation

All specimens prepared for unconfined compression, confined compression, and indirect tensile strength testing were ground, right circular cylinders with dimensions listed above. The dimensions of the specimens were checked and verified according to the Sandia National Laboratories (SNL) Technical Procedure (TP) 51 entitled “Preparing Cylindrical Samples, Including Inspection of Dimensional and Shape Tolerances.”

In some cases, more than one specimen was obtained from a core. This was particularly true for the confined compression tests. This approach was adopted, on a limited basis, for several reasons. The tuff is very heterogeneous. Testing a limited number of specimens at confining pressures of 5, or 10 MPa may not reflect the singular effect of pressure on fracture strength and elastic constants. The determination of the influence of pressure is further complicated by limited sample availability. Certain standard
test methods suggest that measurements should be carried out on specimens greater than 47 mm in diameter, but they allow for smaller specimens, if deemed appropriate, as long as the decision is documented. If the latter approach is adopted, the influence of the smaller specimen volume must be incorporated into the engineering analysis of the data. By preparing a statistically significant number of 25.4 mm diameter specimens from raw core, the effect of pressure could be more clearly distinguished without the complicating effects of variable porosity and pore geometry. Cores of the TCw thermal mechanical unit, recovered from depths of 263.3 and 265.7 feet from borehole UE25 NRG-3 were of sufficient quality and homogeneity to machine 8 specimens for confined compression experiments. Subdivisions of these cores were treated as new specimens, and a Chain-of-Custody form was prepared for each piece of material generated in the subdivision process (per SNL QAIP 20-03, “Sample Control”). The prepared specimens were labeled and stored in containers until the measurement sequence was initiated.

The general measurement sequence for each specimen is given below:

- Dimensions measurement
- Specimen description
- CT Scan (unconfined compression specimens only)
- Drying specimen to a constant weight
- Dry bulk density measurement
- Compressional and shear wave velocities for the dry condition (all unconfined and representative confined compression specimens)
- Saturating specimen to a constant weight with water
- Saturated bulk density measurement
- Compressional and shear wave velocities for the saturated state (all unconfined and representative confined compression specimens)
- Mechanical testing (unconfined compression, confined compression, or indirect tensile strength tests)
- Description of post failure condition of the specimen
- Post-test photograph taken

Average grain density measurements were performed concurrently with the other activities.
2.2 CT Scans of Each Specimen Tested in Unconfined Compression

Prior to testing, a computerized tomography (CT) scan was performed on each specimen designated for testing in unconfined compression. Because CT scans are sensitive to variations in density, this visual representation permits an initial qualitative examination of the distribution of pores and low density zones throughout the rock specimen and provides data for more quantitative analyses, if desired. Given the presence of lithophysae and vapor-phase altered zones within the tuff, the CT scan is a particularly useful technique for evaluating the effects these features may have on the mechanical properties of the tuff at the laboratory scale.

2.3 Drying, Saturation, Bulk Density, Average Grain Density, and Porosity

The dry and saturated bulk densities, average grain density, and porosity were determined for each specimen prepared for a mechanical test. The procedures were developed in accordance with ASTM D 854 “Standard Test Method for Specific Gravity of Soils” and ASTM C 135 “Standard Test Method for True Specific Gravity of Refractory Materials by Water Immersion.”

2.3.1 Procedure for Drying a Specimen

The specimen was dried to a constant weight, and its dry bulk density determined. Drying was carried out at 110 ± 5 °C according to SNL TP-65, “Drying Geologic Specimens to a Constant Weight.” Once the mass change for successive drying cycles had stabilized to within ± 0.05 percent, the dry bulk density, \( \rho_{db} \), was computed.

Previous studies have shown that drying at 110 °C produces no noticeable damage (microcracks) in welded tuff. This was demonstrated by measuring compressional and shear wave velocities before and after heating a TSw2 specimen to 110 °C. Thermal cycling produced no reduction in the velocities suggesting no thermally induced microcracking. If microcracks develop due to differential thermal expansion, the velocities will decrease.
Each specimen is dried in an oven controlled to an accuracy of ± 5 °C. The procedure for drying is outlined below.

1) Place the specimen in the oven. Increase the temperature in the oven to 110 ± 5 °C at a rate less than or equal to 2 °C min⁻¹.

2) Maintain the specimen at 110 ± 5 °C for 120 to 128 hours. Reduce the temperature in the oven at a rate of less than or equal to 2 °C min⁻¹ until the oven temperature is between ambient and 40 °C.

3) Remove the specimen from the drying oven and weigh it three times. The specimen should be weighed within 15 seconds of removal from the oven. Calculate the mean dry mass of the specimen for the three measurements.

4. Repeat steps 1 through 3.

5) Calculate the mass change of the specimen for the successive drying cycles. If the change in mass for successive drying cycles is less than or equal to 0.05%, the process has met the specification and oven drying of the specimen is terminated. If the mass change is greater than 0.05%, steps 1 through 3 must be repeated until the specification is met.

6) Compute the dry bulk density by dividing the dry mass by the specimen volume.

2.3.2 Procedure for Water Saturation

The mechanical tests were performed on nominally water saturated specimens. Saturation of the specimens was achieved in a two-stage process. First, the specimen was pressure saturated at 10 MPa for a minimum of 1 hour. Next, a minimum of two vacuum saturation cycles were performed, according to SNL TP-64, "Procedure for Vacuum Saturation of Geologic Core Samples." Once the mass change for successive saturation cycles had stabilized to within ± 0.05 percent, the saturated bulk density, \( \rho_{SB} \), was computed. The specimens were stored in distilled water following saturation and prior to
testing.

A brief synopsis of the procedure for saturation follows:

1) Place the specimen in a pressure vessel filled with distilled water.

2) Pressurize the vessel to 10 MPa and hold constant for at least one hour.

3) Remove the specimen from the pressure vessel and blot it with a damp lint-free paper towel.

4) Weigh the specimen within 15 seconds of blotting. Record the mass of the specimen in the laboratory notebook.

5) Place the specimen in a container filled with distilled water.

6) Place the water filled container with the specimen in a vacuum chamber.

7) Apply a vacuum to the vacuum chamber.

8) Vacuum saturate the specimen for at least 30 hours.

9) Turn off the vacuum pump, open the valve on the vacuum chamber and allow the pressure to equilibrate with atmospheric conditions.

10) Keep the specimen submerged in water at ambient pressure for at least 16 hours.

11) Remove the specimen from the container and blot it with a damp lint-free paper towel.

12) Weigh the specimen within 15 seconds of blotting. Record the mass of the specimen on the vacuum saturation data sheet.

13) Return the specimen to the water filled container and repeat steps 11 and 12.
Calculate the mean saturated mass of the specimen for the two measurements.

14) Return the specimen to the water filled container and repeat the vacuum saturation procedure in steps 6 through 13. Calculate the mass change for each successive vacuum saturation cycle. If the mass change for successive vacuum saturation cycles is less than or equal to 0.05%, the process has met the specification and the saturation procedure is terminated. If the mass change is greater than 0.05%, steps 6 through 13 must be repeated until the specification is met.

15) Store the specimen in distilled water at ambient pressure and temperature until it is ready for mechanical testing.

16) Compute the saturated bulk density by dividing the saturated mass by the specimen volume.

2.3.3 Average Grain Density Measurement Using Water Pycnometry

The average grain density of each core received from boreholes UE25 NRG-2, 2A, 2B, and 3 was measured using the water pycnometry method. The technique employs a two-stage measurement. First, the mass of a dry, powdered specimen is measured. Next, the volume of the powder is determined. These two measurements are combined to compute the average grain density.

Pieces of core with a mass of approximately 20 to 50 grams are ground to a powder with a particle size of 1.5 mm or less. The powder is dried according to SNL TP-65.

The powdered specimen of tuff is added to a dry, calibrated, water pycnometer with a nominal volume of 100 ml. The step-by-step procedure presented below produces measurements of the dry mass of the powdered specimen and the corresponding volume of the specimen. The technique has been verified using quartz powders with a well characterized density.

1. Pulverize approximately 20 to 50 g of the specimen to a particle size of 1.5 mm or less. The powder is dried in an aluminum drying pan according to SNL TP-65, except cooling is not allowed at any time in order to minimize rehydration.
2. Add the dried sample to a calibrated, clean, dry and numbered pycnometer by pouring it through a clean, dry transfer funnel.

3. Weigh the pycnometer with the dry sample immediately (do not allow it to cool).

4. Add 50 to 60 ml of distilled water to the pycnometer and swirl it to moisten all of the sample powder.

5. Place the pycnometer, with the sample, in an active vacuum for a minimum of 16 hours. For the first one or two hours, watch the pycnometer to ensure that the boiling action does not displace any of the sample from the pycnometer. The vacuum should be regulated depending on the observed phenomena.

6. Remove the pycnometer from the vacuum chamber and pour additional deaired water into the pycnometer until the water level is just below the scribe line. Note that pouring water down the neck reduces the likelihood of entrapping air into the water as it is added to the pycnometer.

7. Use a pipette to add water until the bottom of the meniscus is at the height of the scribe line. It may be necessary to raise the water level higher than the scribe line to wet the sides of the pycnometer for a suitable meniscus. In this case, water is removed to obtain the correct reading.

8. Use a cotton swab to dry the inside of the neck of the pycnometer.

9. Use a lint-free wipe to clean and dry the exterior of the pycnometer.

10. Weigh the pycnometer and its contents.

11. Measure the water temperature in the pycnometer to the nearest 0.2 ° C.

12. Calculate the average grain density (ρ₉) of the specimen using the water pycnometer grain density measurement sheet (Figure 2).

13. Pour the sample and water into a clean container to allow the water to evaporate.

14. Store the sample powder in a container to maintain it in its original condition.
Bulk Properties
Yucca Mountain Project
WATER Pycnometer Grain Density Measurement
per TP-229 Rev. 0

SAMPLE ID:_________________________

WATER PYCNOMETER ID:____________

Date of latest calibration:________Nominal Pycnometer Volume:____________

DATA:

Dry Pyc. Wt.:___________ g (A)

Dry Sample Wt.:___________ g (C) = B-A

Wt. Pyc. + Sample + Water:___________ g (D)

Water Temperature:___________ °C (E)

Water Density at (E) (See Below):___________ g/cc (F)

Wt. Water Only:___________ g (G) = D-B

Volume Water:____________ cc (H) = G/F

Volume Pyc. at (E) from Calibration:____________ cc (I)

Volume Sample:____________ cc (J) = I-H

Sample Grain Density:____________ g/cc (K) = C/J

Absolute Density of Water

<table>
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<tr>
<th>Temp °C</th>
<th>Density</th>
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Figure 2: Bulk properties worksheet used to record measurements and reduce data for the computation of average grain density.
2.3.4 **Dry Bulk Density, Saturated Bulk Density, and Porosity**

The dry bulk density, $\rho_{db}$, is obtained by computing the volume of a test specimen from its external dimensions and dividing it into the mass measured in a dry condition. The density corresponding to the measurement in the saturated condition is the saturated bulk density, $\rho_{sb}$. Preferably, porosity, $\phi$, is computed using the following relation:

$$\phi = \frac{\rho_g - \rho_{db}}{\rho_g} \times 100\%$$

Alternatively, the porosity can be calculated from the dry bulk density and saturated bulk density according to:

$$\phi = \frac{\rho_{sb} - \rho_{db}}{\rho_w} \times 100\%$$

where $\rho_w$ is the density of water.

2.4 **Compressional and Shear Wave Velocity Measurements**

Compressional and shear wave velocities were measured on right circular cylinders with a nominal length to diameter ratio of 2:1. The velocities were measured for both dry and water saturated conditions at ambient temperature in a benchtop apparatus. Specimens were dried at 110 °C according to SNL TP-65 (as in Section 2.3.1). Earlier investigations have shown that heating specimens of Topopah Spring Member tuff to 110 °C does not damage them by the formation of microcracks.

The compressional and shear wave velocity measurements are used for two main purposes. First, a measure of the specimen anisotropy can be directly obtained by comparing the compressional and shear wave velocities measured both parallel and normal to the core axis. Second, compressional and shear wave velocity data, combined with the density of the specimen, are used to compute Young’s modulus and Poisson’s ratio. These elastic moduli will be referred to as dynamic moduli in subsequent discussions. The dynamic moduli can then be compared with the moduli computed for the data on the unconfined compression tests conducted at a strain rate of $10^{-5}$ s$^{-1}$. The elastic constants computed from stress and strain measurements in a deformational experiment are often referred to as static. For porous rocks, the dynamic moduli are greater than the static moduli measured during the constant strain rate experiments and therefore serve as an
upper bound on the static moduli (Simmons and Brace, 1965; Cheng and Johnston, 1981; Haupt et al., 1992).

A self-contained ultrasonic measuring system is used to perform the velocity measurements. A tuff specimen is placed between a matched set of ultrasonic transducers. One transducer serves as the source; the second as the receiver (Figure 3). The travel time through the rock is divided by the sample length to compute the velocity.

Each ultrasonic transducer contains one compressional and one or two polarized shear wave elements. For the measurements parallel to the core axis, one compressional and two orthogonally polarized shear waves are propagated. For measurements normal to the core axis, one compressional and one polarized shear wave velocity are measured. The polarization direction of the shear wave propagating normal to the core axis is parallel to the axis of the core.

The transducers are constructed using piezoelectric crystals with a resonant frequency of 1 MHz. The multicomponent piezoelectric crystals are bonded to a titanium substrate. Titanium has been selected because it has a good acoustical impedance match both to the rock and to the piezoelectrical crystals. The source crystal is excited with a fast rise time pulse generator. The crystal produces a broad band ultrasonic pulse propagated through the adjacent titanium substrate, the rock, the titanium at the opposite end of the core, and into the receiver crystal. The received electrical signal is then amplified and filtered through the receiving section of the pulser-receiver and displayed on a digital storage oscilloscope. The signals are amplified and high pass filtered above 0.3 MHz. The time series displayed on the oscilloscope is then digitized and transferred to a computer for subsequent analysis including picking the first arrival of the compressional and shear wave energy to compute the compressional and shear wave velocities. The accuracy of the travel time is ± 0.02 microseconds.

A diagram of the system is shown in Figure 4. Pneumatic actuators couple the transducer assemblies in both the axial and radial directions. The stress across the interface for both the matched transducer pairs is identical; this is accomplished by adjusting the loading areas in the pneumatic actuators. The titanium pieces for the radial transducers are concave to mate with the rock surface. Because of the geometry of the interface, only polarizations parallel to the core axis are propagated for shear waves in the radial direction.
Figure 3: Schematic diagram of the geometry used to measure compressional and shear wave velocities in tuff. The diagram shows the setup for the measurement of velocities parallel to the core axis.
Figure 4: Schematic of the apparatus used to measure ultrasonic velocities parallel and normal to the core axis under ambient conditions prior to testing in unconfined compression.
2.4.1 Detailed Procedure for Compressional and Shear Wave Velocity Measurements

The detailed procedure for measuring compressional and shear wave velocities on dry and water saturated tuff specimens is presented below.

1. The tuff sample is a ground, right circular cylinder with a nominal length-to-diameter ratio of 2:1. The sample is machined to meet or exceed the tolerances specified in SNL TP-51.

2. Coat the ends of the specimen with a shear wave couplant. Shear wave couplant is a viscous substance that facilitates the propagation of shear waves across the specimen-titanium interface. Shear wave couplant is also applied at the midpoint of the specimen where the velocities normal to the specimen axis are measured.

3. Position the specimen in the ultrasonic velocity measuring apparatus. Ensure that the specimen is lined up with the transducers in the axial direction.

4. Increase the pressure in the pneumatic actuators to load the axial and radial transducer assemblies to the specimen. Ensure that the specimen has not shifted during this procedure and that the specimen is well coupled to the transducer assembly.

5. Turn on the data acquisition system. First, set the signal selection switch for a compressional (P) wave along the axis of the specimen. Observe the received signal on the digital oscilloscope. Adjust the pulse excitation signal gain and/or attenuation to obtain a well-defined signal.

6. Initiate the data acquisition software to store the waveform; capture and store the waveform.

7. Capture the two shear (S1 and S2) wave polarizations with a propagation direction parallel to the core axis and the compressional and shear wave signals for the propagation direction normal to the core axis, following the same procedure used for the compressional wave in steps 5 and 6.
8. Compute the compressional and shear wave velocities by determining the travel time through the specimen and dividing it into the sample length. The travel time is determined by picking the time of the first arrival of the compressional or shear wave energy. The measured travel times are reduced by the travel time through the titanium substrates. The corrected travel time is then divided into the sample length to determine the velocity.

9. Print the stored waveforms, along with the computed compressional and shear wave velocities and place the data in the scientific notebook.

10. Compute the dynamic Young’s modulus \(E, \text{ GPa}\) and Poisson’s ratio \(v\) from the velocity data collected parallel to the core axis and the bulk density of the specimen for the measurement condition. The dynamic elastic moduli are computed as follows:

\[
E = \rho V_s^2 (3V_p^2 - 4V_s^2)/(V_p^2 - V_s^2)
\]

\[
v = (V_p^2 - 2V_s^2)/(2(V_p^2 - V_s^2))
\]

where

\(V_p\) = compressional wave velocity, \(\text{km s}^{-1}\)
\(V_s\) = average shear wave velocity, \(\text{km s}^{-1}\)
\(\rho\) = bulk density for the measurement conditions, \(\text{g cm}^{-3}\)

2.5 **Unconfined Compression to Failure**

The unconfined compression experiments were performed on saturated right-circular cylinders of tuff with a nominal length to diameter ratio of 2:1, at a constant axial strain rate of \(10^{-5} \text{ s}^{-1}\) at room temperature. System checks of the entire test system were conducted during this experimental series to establish the performance of the system using an aluminum specimen with the same nominal dimensions as the tuff test specimens.

A description of the equipment and an overview of the test procedures place the step-by-step procedures in the proper context. All the compression tests were carried out in
a servo-controlled hydraulic loading frame with a capacity of $1.1 \times 10^6$ N. The servocontroller is a self-contained digital unit, which operates in either force or displacement feedback. The rate at which the reference signal is updated can be varied from $10^{-5}$ to $10^3$ times per second. The loading rate or displacement rate depends on the range of the feedback transducers and the time between steps. The feedback transducers are conditioned with amplifiers in the servo-control unit and balanced so that the full-scale output of the transducer corresponds to the maximum range of the reference signal generator. The full-scale output (10 V) is divided into $2^{12}$ discrete steps.

Figure 5 is a schematic diagram of an instrumented specimen. The test assembly consists of the specimen positioned between hardened steel end caps. The specimen is jacketed in a flexible membrane to maintain its moisture level.

For this experimental series, five outputs from a variety of transducers were monitored. The output from each device is conditioned, amplified, converted to digital format, and recorded as a function of time. The outputs from the devices were recorded with a microcomputer acquisition system. Each channel is sampled at a frequency of 4 Hz.

For constant strain rate tests, the loading frame is operated in displacement feedback. The displacement can be controlled to within $\pm 10^{-3}$ mm when the system is in the displacement mode. The accuracy and the reproducibility of the strain rate in this system is $\pm 0.5$ percent.

During each test, the axial and radial displacements of the specimen were measured with Linear Variable Differential Transformers (LVDTs). A schematic of their arrangement is shown in Figure 5. Two LVDTs monitor the axial displacement. The LVDT barrels are secured in a ring which is attached one specimen radius from the upper end of the specimen. The cores for the displacement transducers are on extended rods which attach to a second ring separated from the first by one specimen diameter. The second ring is mounted approximately one specimen radius from the lower end of the specimen.

The most direct way to measure radial strain is with the radial displacement gage developed by Holcomb and McNamee (1984). Their gage consists of an LVDT mounted in a ring which is spring loaded against the surface of the specimen (Figure 5). The core of the LVDT is connected to the spring. As the specimen diameter changes, the spring deflects, changing the position of the core within the barrel of the LVDT in direct proportion to the radial displacement.

The force on the test column is measured with a load cell. The accuracy of the load cell is better than 0.5 percent of its full-scale output; the combined linearity and hysteresis are better than 1.0 percent. The position of the hydraulic piston is observed with
Axial and Radial Deformation Instrumentation

Figure 5: Schematic of the transducer configuration used to measure the axial and radial displacement of specimens during unconfined and confined compression tests.
a displacement transducer. This transducer provides feedback control in constant strain rate
tests and is continuously monitored along with all the transducer outputs. The hydraulic
piston advances at a constant rate; this is equivalent to deforming the specimen at a constant
strain rate only in the linear portion of the stress-strain curve.

Checks of the entire test system are made using a sample of 6061-T6511 aluminum
with the same nominal dimensions as the test specimens. One check is performed before
the experiments on a suite of tuff samples and then after each group of ten tuff samples.
Each check was performed at the same conditions as the experiments on the tuff specimens:
a nominal strain rate of $10^{-5}$ s$^{-1}$, at ambient temperature.

2.5.1 Experimental Procedures for Unconfined Compression Tests

Specimens of tuff were tested to failure at a constant strain rate of $10^{-5}$ s$^{-1}$, under
ambient temperature and pressure conditions. The following sections include the step-by-
step procedures for the mechanical experiments based on SNL TP 219 “Unconfined
Compression Experiments at 22 °C and a Strain Rate of $10^{-5}$ s$^{-1}$.” The test procedure relies
Core Specimens in Uniaxial Compression” and ISRM procedure “Suggested Methods for
Determining the Uniaxial Compressive Strength and Deformability of Rock Materials.”

1. Each tuff specimen is machined according to SNL TP-51, dried according to
SNL TP-65, and saturated according to SNL TP-64. Velocity measurements are
performed after the drying and saturation procedures.

2. List all transducers used for each experiment. The information includes the serial
numbers of the device, signal conditioning amplifier number, the computer channel
on which the output is recorded, and the scaling factor for the amplified output.

3. Visually inspect the test specimen and note any surface irregularities and
imperfections.

4. Jacket the saturated specimen in a thin, flexible membrane. The membrane is
extended 12 mm beyond each end of the specimen. Hardened steel end caps
are then positioned at each end of the specimen.

5. Position the ring supporting the two axial LVDT barrels approximately one
specimen radius from the upper end of the specimen. Carefully center the ring so that it is concentric with the specimen.

6. Position the LVDT ring used to measure the radial displacement at the midpoint of the specimen. The supporting ring for the LVDT (with a range of ± 1.25 mm) is positioned in such a way to ensure that the line between the adjusting screw on the ring and the axis of the core barrel of the LVDT passes through the axis of the specimen and is perpendicular to the axis of the specimen.

7. Position the lower support ring for the axial LVDT concentrically about the specimen approximately one specimen diameter from the upper ring. This ring supports the stainless steel extension rods for the LVDT cores. Ensure that the axes of the LVDT core barrels are aligned parallel with the axis of the specimen. The extension rods are supported with adjusting screws secured with locking nuts.

8. Measure the center-to-center separation of the axial LVDT support rings with a caliper-micrometer. Record this value in the scientific notebook.

9. Place the specimen assembly on the base plug of the load frame.

10. Connect the two axial LVDTs and the radial LVDT to the electrical leads in the base plug.

11. Advance the loading piston, in displacement control, until a small load is exerted on the specimen column (just enough to hold the specimen securely in position).

12. Make the final mechanical adjustments on the LVDTs. Each LVDT is adjusted so that its initial amplified output is approximately 0.10 V. Note that all the LVDTs are wired so that increasing the specimen diameter and shortening the specimen assembly results in an increasing positive output voltage.

13. Retract the hydraulic piston until there is no force on the loading column.

14. Initiate data acquisition. The amplified outputs from five transducers are monitored and recorded using a microprocessor-based data acquisition system. The transducers that are monitored include the three LVDTs, the feedback displacement transducer, and the force cell. All the channels are sampled every
0.25 seconds. Data is stored when the output of one channel deviates from the previous value by a preselected threshold. The threshold for each channel is independently set prior to the experiment.

15. Adjust the setting on the displacement rate controller to the displacement rate that corresponds to a nominal strain rate of $10^{-5} \text{s}^{-1}$. After a final check of all the transducer values, start loading the specimen.

16. Load the specimen to failure.

17. Remove the specimen from the press and examine the manner in which it failed. Record the observations in the scientific notebook. Photograph the specimen. Ensure that the field of view of the photograph includes the specimen identification, TP identification, scale, date, type of test performed, and NER identification.

18. Return the specimen to its original container and return it to storage.

19. Reduce the data. The following elastic constants are computed:

(a) Young's modulus, $E$

$$E = \frac{\Delta \text{(axial stress)}}{\Delta \text{(axial strain)}}$$

(b) Poisson's ratio, $\nu$

$$\nu = \frac{\Delta \text{(radial strain)}}{\Delta \text{(axial strain)}}$$

The elastic constants are computed by performing a least-squares linear fit to the data collected between 10 and 50 percent of the fracture strength. Axial stress is computed by dividing the axial force by the initial cross sectional area of the specimen. Stress is reported in MPa. Axial strain is obtained by dividing the average axial displacement of the axial LVDT support rings by the original ring separation distance. Radial strain is computed by dividing the change in radial displacement observed by the radial LVDT by the initial specimen diameter. All strains are reported in millistrain.
2.6 **Confined Compression to Failure**

Confined compression tests were carried out on specimens from cores with sufficient uniform material that numerous specimens could be obtained. The specimens had a length to diameter ratio of 2:1 with a diameter of 25.4 mm and were tested in a saturated condition. The general procedure for testing these specimens was the same as that described for the unconfined compression tests except that the specimens were jacketed in copper and deformed in a pressure vessel at a fixed confining pressure. The procedure was based on TP-219 and conforms to ISRM “Suggested Methods for Determining the Strength of Rock Materials in Triaxial Compression” and ASTM D 2664 “Triaxial Compressive Strength of Undrained Rock Core Specimens Without Pore Pressure Measurements.” The series of tests was designed so that at least three specimens were tested for each confining pressure. Specimens were tested at confining pressures of 5 and 10 MPa.

Each specimen was jacketed in dead soft copper 0.13 mm thick. The jacket was seated to the rock specimen by subjecting it to a hydrostatic pressure of 10 MPa. Once the jacket was seated, it was visually inspected to ensure that there were no holes that would permit leakage of the confining medium into the rock core during the test. Next, the specimen was instrumented using the same arrangement utilized for the unconfined compression tests (Figure 5).

The instrumented specimen was mounted on the base plug of a 50 MPa capacity pressure vessel. The LVDTs were adjusted. Once the LVDTs were on scale and functioning properly, the base plug of the pressure vessel containing the instrumented specimen assembly was inserted into the pressure vessel. Next, the confining pressure was exerted on the specimen using a servo-controlled hydraulic intensifier. The confining medium was argon gas. The confining pressure was allowed to stabilize. The confining pressure was held constant during the remainder of the experiment to within ± 0.1 MPa.

Once the confining pressure reached thermal equilibrium the specimen was monotonically loaded to failure at a strain rate of 10^-5 s^-1. After the specimen failed, the confining pressure was released and the specimen was removed and inspected. The data were reduced to determine Young’s modulus, Poisson’s ratio, and fracture strength. Young’s modulus and Poisson’s ratio are computed in the same manner as for the unconfined compression tests, between 10 and 50 percent of the stress difference at failure.
2.7 Indirect Tensile Strength Tests

Indirect tensile strength tests, commonly referred to as Brazil tests, were carried out using a procedure adhering to ASTM D-3967 “Splitting Tensile Strength of Intact Rock Core Specimens.” The test is quite simple in principle. A load is applied to a cylindrical specimen with its axis normal to the loading direction. A tensile stress develops in the center of the cylinder. The specimen fails by an extension fracture along the diametral loading plane. The force is increased until the specimen fails. The strength is computed from the force at failure.

The tests were performed in the servo-controlled load frame used for the unconfined and confined compression tests. For these measurements the only transducer was a load cell with a capacity of $4.5 \times 10^4$ N. A schematic diagram of the setup is shown in Figure 6.

The specimens were right circular cylinders, ground to the dimensions and tolerances listed in Section 2.0. The dry bulk density and saturated bulk density were measured prior to testing. The specimens were tested in a nominally water saturated condition.

To obtain reproducible and accurate data, it is important that the loading axis of the test frame passes through the axis of the specimen. As an aid in the alignment of the specimen in the loading column, diametral lines are scribed on the bearing surfaces.

2.7.1 Experimental Procedures for Indirect Tensile Strength Tests

The detailed procedure developed according to ASTM D-3967 is presented below.

1. Each tuff specimen was machined according to SNL TP-51, dried according to SNL TP-65, and saturated according to SNL TP-64. All initial conditions were documented.

2. Mark diametral lines on the ends of the specimen.

3. Cut 2.5 cm wide by 4.0 cm long (slightly longer than the specimens) pieces of notepad-backing cardboard approximately 0.89 mm thick for bearing strips.

4. Place one bearing strip onto the bottom bearing platen with its length parallel to the diametral line scribed onto the platen.
Figure 6: Schematic of the loading arrangement and sample geometry used for the indirect tensile strength tests.
5. Place the specimen onto the bearing strip making certain its full length is supported by the bearing strip. Align the axis of the specimen parallel to the line scribed onto the platen (lines on both ends of the specimen line up with the line scribed on the platen).

6. Place the second bearing strip onto the top of the specimen making certain it will support the full length of the specimen.

7. Advance the loading piston, in displacement control, until a small load is applied (just enough to hold the specimen and the bearing strips in place while positioning them).

8. Make final adjustments to the specimen’s position to ensure the diametral plane of the two lines marked onto the specimen line up with the center of thrust (lines scribed onto the top and bottom bearing planes), and its midpoint along its length is also lined up to within ±0.025 mm of the center of the bearing platens. The midpoint is centered by repeatedly measuring the distance from the specimen ends to the edge of the bearing platens and adjusting its position until they are equal.

9. Initiate the data acquisition system.

10. Load the specimen monotonically to failure in displacement control at the same rate used for the unconfined compression tests. Once the specimen fails, data acquisition is terminated and any remaining load is removed. The specimen is removed from the loading column, described, and photographed.

11. Calculate the indirect tensile strength using the formula:

\[ \sigma_t = \frac{2F}{\pi LD} \]

where

- \( F \) = applied force
- \( L \) = specimen length
- \( D \) = specimen diameter.

The indirect tensile strength is reported in MPa.
3.0 RESULTS

The results of the bulk properties measurements and the mechanical tests are presented in Tables 1, 2, 3, and 4. Tables 1a, b, c, and d present the data associated with the unconfined compression tests. The data include dry bulk density, saturated bulk density, average grain density, porosity, compressional and shear wave velocities for the dry and saturated conditions both parallel [axial] and normal [radial] to the core axis, static Young’s modulus, static Poisson’s ratio, axial stress at failure, and axial strain at failure. These data have been grouped according to depth. The thermal/mechanical unit is indicated for each specimen. For some specimens, reliable compressional and shear wave velocities were not obtained due to poor signal quality. The absence of the data is reflected in Tables 1a, b, c, and d.

The porosity reported in Tables 1, 2, 3, and 4 is the total porosity including occluded porosity. The value is computed from the grain density and dry bulk density. The total porosity is most applicable when elastic constants and strength characteristics are measured. The porosity computed from the dry and saturated bulk densities is typically lower than the total porosity and reflects the interconnected porosity. The latter value is applicable when considering fluid transport properties.

Table 2 presents the data for the confined compression tests. The dry bulk density, saturated bulk density, average grain density, porosity, compressional and shear wave velocities for propagation parallel to the specimen axis at one confining pressure, confining pressure at which the sample was tested, static Young’s modulus, static Poisson’s ratio, axial stress difference at failure, and axial strain difference at failure are presented as a function of depth for the two suites of confined compression tests performed on core recovered from depths of 263.3 and 265.7 feet (TCw) from borehole UE25 NRG-3.

Tables 3a, b, c, and d present the results of the indirect tensile strength tests. The data are presented as a function of depth and include dry bulk density, saturated bulk density, average grain density, porosity, and indirect tensile strength. The thermal/mechanical unit for each specimen is also indicated. No compressional or shear wave velocities were measured on these specimens.

Tables 4a, b, and c list the specimens for which only average grain density, dry bulk density, and porosity were measured. The thermal/mechanical unit for each of these specimens is indicated.

All specimens tested in this study were ideally water saturated. However, the saturation is not 100%. Porosities computed using the saturated bulk density and dry bulk
| Depth, ft: | 170.4 | 174.0 | 178.0 | 179.5 | 180.0 | 188.3 | 196.2 | 200.0 |
| T/M Unit: | TCw   | TCw   | TCw   | TCw   | TCw   | TCw   | TCw   | TCw   |
| Date Tested: | 7/1/93 | 7/1/93 | 7/1/93 | 7/1/93 | 7/1/93 | 7/1/93 | 7/1/93 | 7/1/93 |
| Dry Bulk Density (g/cc): | 2.335 | 2.342 | 2.319 | 2.360 | 2.334 | 2.345 | 2.350 | 2.347 |
| Saturated Bulk Density (g/cc): | 2.403 | 2.405 | 2.391 | 2.417 | 2.399 | 2.405 | 2.411 | 2.408 |
| Average Grain Density (g/cc): | 2.525 | 2.524 | 2.524 | 2.530 | 2.528 | 2.541 | 2.529 | 2.527 |
| Depth, ft: | 170.4 | 174.0 | 178.0 | 179.5 | 180.0 | 188.3 | 196.2 | 200.0 |
| Porosity via Grain Density (%): | 7.5 | 7.2 | 8.1 | 6.7 | 7.7 | 7.7 | 7.1 | 7.1 |
| Dry S1 Velocity (km/s): | 2.943 | 2.938 | 2.926 | 2.925 | 2.923 | 2.909 | 2.877 | 2.866 |
| Dry S2 Velocity (km/s): | 2.911 | 2.977 | 2.889 | 2.931 | 2.918 | 2.914 | 2.879 | 2.865 |
| Dry Radial S Velocity (km/s): | 2.967 | 2.987 | 2.974 | 2.974 | 2.940 | 2.940 | 2.874 | 2.898 |
| Sat. S1 Velocity (km/s): | | | | | | | | |
| Sat. S2 Velocity (km/s): | | | | | | | | |
| Saturated Radial S Velocity, km/s: | | | | | | | | |
| Static Young's modulus (GPa): | 40.1 | 38.7 | 39.8 | 40.1 | 37.1 | 39.8 | 36.5 | 38.7 |
| Static Poisson's ratio: | 0.19 | 0.20 | 0.20 | 0.22 | 0.20 | 0.20 | 0.22 | 0.23 |
| Ultimate Axial Stress (MPa): | 117.5 | 141.8 | 142.7 | 215.8 | 149.1 | 209.2 | 185.6 | 145.3 |
| Ax. Strn at Ult. Ax. Sts. (milstr): | 4.54 | 3.82 | 3.74 | 5.57 | 4.42 | 5.54 | 5.28 | 5.77 |

P is the compressional wave; S1 and S2 are the two orthogonally polarized shear waves. Elastic properties are calculated between 10 and 50 percent of the ultimate differential axial stress.
### Unconfined Compression Tests

Sample IDs are shortened from the "NRG-2A-Depth-SNL-Subdivision" Format

Test Conditions: Saturated samples, ambient pressure and temperature, and a nominal strain rate of 10E-5 sE-1

Nominal Sample Dimensions: Length = 101.60 mm; Diameter = 50.80 mm

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P is the compressional wave; S1 and S2 are the two orthogonally polarized shear waves.

Elastic properties are calculated between 10 and 50 percent of the ultimate differential axial stress.
### Table 1b (Continued)

**SUMMARY DATA SHEET: NRG-2A BOREHOLE**

**Unconfined Compression Tests**

Sample IDs are shortened from the "NRG-2A-Depth-SNL-Subdivision" Format

Test Conditions: Saturated samples, ambient pressure and temperature, and a nominal strain rate of 10E-5 s^-1

Nominal Sample Dimensions: Length = 101.60 mm; Diameter = 50.80 mm

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<td>0.24</td>
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P is the compressional wave; S1 and S2 are the two orthogonally polarized shear waves.

Elastic properties are calculated between 10 and 50 percent of the ultimate differential axial stress.
Table 1c

SUMMARY DATA SHEET: NRG-2B BOREHOLE

Unconfined Compression Tests

Sample IDs are shortened from the "NRG-2B-Depth-SNL-Subdivision" Format

Test Conditions: Saturated samples, ambient pressure and temperature, and a nominal strain rate of 10E-5 sE-1

Nominal Sample Dimensions: Length = 101.60 mm; Diameter = 50.80 mm

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<td>Saturated Radial S Velocity, km/s:</td>
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P is the compressional wave; S1 and S2 are the two orthogonally polarized shear waves.

Elastic properties are calculated between 10 and 50 percent of the ultimate differential axial stress.
### Table 1d

#### SUMMARY DATA SHEET: NRG-3 BOREHOLE

#### Unconfined Compression Tests

Sample IDs are shortened from the "NRG-3-Depth-SNL-Subdivision" Format

- **Test Conditions:** Saturated samples, ambient pressure and temperature, and a nominal strain rate of 10E-5 sE-1
- **Nominal Sample Dimensions:** Length = 101.60 mm; Diameter = 50.80 mm

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*P is the compressional wave; S1 and S2 are the two orthogonally polarized shear waves.

Elastic properties are calculated between 10 and 50 percent of the ultimate differential axial stress.*
Table 1d (Continued)

**SUMMARY DATA SHEET: NRG-3 BOREHOLE**

**Unconfined Compression Tests**

Sample IDs are shortened from the "NRG-3-Depth-SNL-Subdivision" Format

Test Conditions: Saturated samples, ambient pressure and temperature, and a nominal strain rate of 10E-5 sE-1

Nominal Sample Dimensions: Length = 101.60 mm; Diameter = 50.80 mm

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<tr>
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<td>TCw</td>
<td>TCw</td>
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<td>2.295</td>
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<td>2.905</td>
<td>2.905</td>
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<td>6.71</td>
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</table>

P is the compressional wave; S1 and S2 are the two orthogonally polarized shear waves.

Elastic properties are calculated between 10 and 50 percent of the ultimate differential axial stress.
### Table 1d (Continued)

**SUMMARY DATA SHEET: NRG-3 BOREHOLE**

**Unconfined Compression Tests**

Sample IDs are shortened from the "NRG-3-Depth-SNL-Subdivision" Format

Test Conditions: Saturated samples, ambient pressure and temperature, and a nominal strain rate of 10E-5 sE-1

Nominal Sample Dimensions: Length = 101.60 mm; Diameter = 50.80 mm

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<td>Saturated Bulk Density (g/cc):</td>
<td>2.363</td>
<td>2.347</td>
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<td>Average Grain Density (g/cc):</td>
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<td>Porosity via Grain Density (%):</td>
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<td>4.498</td>
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<tr>
<td>Dry S1 Velocity (km/s):</td>
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<tr>
<td>Dry S2 Velocity (km/s):</td>
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<td>Dry Radial P Velocity (km/s):</td>
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<td>4.380</td>
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<td>Dry Radial S Velocity (km/s):</td>
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<tr>
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<td>Sat. S1 Velocity (km/s):</td>
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</tr>
<tr>
<td>Sat. S2 Velocity (km/s):</td>
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<td></td>
</tr>
<tr>
<td>Saturated Radial P Velocity, km/s:</td>
<td>4.550</td>
<td>4.462</td>
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<td>Saturated Radial S Velocity, km/s:</td>
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</tr>
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<td>Static Young's modulus (GPa):</td>
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<td>Static Poisson's ratio:</td>
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<tr>
<td>Ultimate Axial Stress (MPa):</td>
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<td>75.4</td>
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</table>

P is the compressional wave; S1 and S2 are the two orthogonally polarized shear waves.

Elastic properties are calculated between 10 and 50 percent of the ultimate differential axial stress.
### Table 2

**SUMMARY DATA SHEET: NRG-3 BOREHOLE**

**Confined Compression Tests**

Sample IDs are shortened from the “NRG-3-Depth-SNL-Subdivision” Format.

Test Conditions: Saturated samples, ambient temperature, and a nominal strain rate of 10E-5 sE-1.

Nominal Sample Dimensions: Length = 50.80 mm; Diameter = 25.40 mm.

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<td>9/17/93</td>
<td>9/17/93</td>
<td>9/17/93</td>
</tr>
</tbody>
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- **Dry Bulk Density (g/cc):**
  - 2.243
  - 2.329
  - 2.329
  - 2.324
  - 2.306
  - 2.333
  - 2.332
  - 2.329

- **Saturated Bulk Density (g/cc):**
  - 2.348
  - 2.395
  - 2.395
  - 2.391
  - 2.381
  - 2.397
  - 2.396
  - 2.396

- **Average Grain Density (g/cc):**
  - 2.510
  - 2.510
  - 2.510
  - 2.510
  - 2.510
  - 2.510
  - 2.510
  - 2.510

- **Porosity via Grain Density (%):**
  - 10.6
  - 7.2
  - 7.2
  - 7.4
  - 8.1
  - 7.1
  - 7.1
  - 7.2

- **Sat. P Velocity at 10 MPa (km/s):**
  - 4.716
  - 4.755
  - 4.678
  - 4.685
  - 4.633
  - 4.701
  - 4.742
  - 4.708

- **Sat. S1 Velocity at 10 MPa (km/s):**
  - 2.857
  - 2.843
  - 2.788
  - 2.825
  - 2.759
  - 2.819
  - 2.817
  - 2.820

- **Sat. S2 Velocity at 10 MPa (km/s):**
  - 2.832
  - 2.788
  - 2.798
  - 2.742
  - 2.801
  - 2.827
  - 2.812

- **Confining Pressure (MPa):**
  - 5
  - 10
  - 5
  - 5
  - 5
  - 10
  - 10
  - 10

- **Static Young's modulus (GPa):**
  - 27.0
  - 39.3
  - 37.4
  - 36.5
  - 39.4
  - 37.1
  - 37.5
  - 36.8

- **Static Poisson's ratio:**
  - 0.29
  - 0.19
  - 0.20
  - 0.20
  - 0.25
  - 0.21
  - 0.20
  - 0.21

- **Ultimate Diff. Axial Stress (MPa):**
  - 153.2
  - 281.4
  - 290.9
  - 270.1
  - 175.3
  - 359.0
  - 329.1
  - 299.9

- **Ax. Strn at Ult. Ax. Sts. (milstr):**
  - 7.32
  - 7.64
  - 8.42
  - 7.94
  - 6.00
  - 11.25
  - 9.75
  - 9.10

*P* is the compressional wave; *S1* and *S2* are the two orthogonally polarized shear waves.

Elastic properties are calculated between 10 and 50 percent of the ultimate differential axial stress.
**Table 3a**

**SUMMARY DATA SHEET: NRG-2 BOREHOLE**

Indirect Tensile Strength (Brazil) Tests

Sample IDs are shortened from the "NRG-2-Depth-SNL-Subdivision" Format

Test Conditions: Saturated samples, ambient pressure and temperature.
Nominal Sample Dimensions: Length = 38.10; Diameter = 50.80 mm.

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<td>7/2/93</td>
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<td>7.2</td>
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<td>2.529</td>
<td>2.539</td>
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<td>7.2</td>
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### Table 3b

#### SUMMARY DATA SHEET: NRG-2A BOREHOLE

**Indirect Tensile Strength (Brazil) Tests**

Sample IDs are shortened from the "NRG-2A-Depth-SNL-Subdivision" Format

Test Conditions: Saturated samples, ambient pressure and temperature.
Nominal Sample Dimensions: Length = 38.10; Diameter = 50.80 mm.

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<td>5.6</td>
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Table 3c
SUMMARY DATA SHEET: NRG-2B BOREHOLE

Indirect Tensile Strength (Brazil) Tests

Sample IDs are shortened from the "NRG-2B-Depth-SNL-Subdivision" Format
Test Conditions: Saturated samples, ambient pressure and temperature.
Nominal Sample Dimensions: Length = 38.10 mm; Diameter = 50.80 mm.

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<tr>
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<td>Saturated Bulk Density (g/cc)</td>
<td>1.837</td>
<td>1.698</td>
<td>1.684</td>
</tr>
<tr>
<td>Average Grain Density (g/cc)</td>
<td>2.356</td>
<td>2.315</td>
<td>2.312</td>
</tr>
<tr>
<td>Porosity via Grain Density (%)</td>
<td>35.8</td>
<td>43.5</td>
<td>44.3</td>
</tr>
<tr>
<td>Ultimate Tensile Stress (MPa)</td>
<td>2.7</td>
<td>0.4</td>
<td>0.6</td>
</tr>
</tbody>
</table>
### Table 3d

**SUMMARY DATA SHEET: NRG-3 BOREHOLE**

**Indirect Tensile Strength (Brazil) Tests**

Sample IDs are shortened from the "NRG-3-Depth-SNL-Subdivision" Format.

Test Conditions: Saturated samples, ambient pressure and temperature.

Nominal Sample Dimensions: Length = 38.10; Diameter = 50.80 mm.

<table>
<thead>
<tr>
<th>Depth, ft.</th>
<th>15.4</th>
<th>32.1</th>
<th>42.6</th>
<th>48.0</th>
<th>55.7</th>
<th>87.3</th>
<th>93.8</th>
<th>119.6</th>
<th>136.1</th>
<th>136.1</th>
</tr>
</thead>
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<tr>
<td>T/M Unit:</td>
<td>TCw</td>
<td>TCw</td>
<td>TCw</td>
<td>TCw</td>
<td>TCw</td>
<td>TCw</td>
<td>TCw</td>
<td>TCw</td>
<td>TCw</td>
<td>TCw</td>
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<tr>
<td>Date Tested:</td>
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<td>8/18/93</td>
<td>8/19/93</td>
<td>8/19/93</td>
<td>8/19/93</td>
<td>8/18/93</td>
<td>8/18/93</td>
<td>8/18/93</td>
<td>8/18/93</td>
<td>8/18/93</td>
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<tr>
<td>Dry Bulk Density (g/cc):</td>
<td>1.587</td>
<td>1.725</td>
<td>1.932</td>
<td>1.940</td>
<td>1.925</td>
<td>2.178</td>
<td>2.179</td>
<td>2.166</td>
<td>2.166</td>
<td>2.130</td>
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<tr>
<td>Saturated Bulk Density (g/cc):</td>
<td>1.948</td>
<td>2.023</td>
<td>2.149</td>
<td>2.161</td>
<td>2.149</td>
<td>2.291</td>
<td>2.294</td>
<td>2.279</td>
<td>2.279</td>
<td>2.246</td>
</tr>
<tr>
<td>Average Grain Density (g/cc):</td>
<td>2.605</td>
<td>2.582</td>
<td>2.582</td>
<td>2.556</td>
<td>2.557</td>
<td>2.519</td>
<td>2.514</td>
<td>2.510</td>
<td>2.509</td>
<td>2.509</td>
</tr>
<tr>
<td>Porosity via Grain Density (%):</td>
<td>39.1</td>
<td>33.2</td>
<td>25.2</td>
<td>24.1</td>
<td>24.7</td>
<td>13.5</td>
<td>13.3</td>
<td>13.7</td>
<td>16.4</td>
<td>15.1</td>
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<tr>
<td>Ultimate Tensile Stress (MPa):</td>
<td>3.0</td>
<td>2.6</td>
<td>3.3</td>
<td>3.9</td>
<td>4.0</td>
<td>10.1</td>
<td>11.7</td>
<td>9.1</td>
<td>8.9</td>
<td>10.3</td>
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<th>Depth, ft.</th>
<th>195.7</th>
<th>218.0</th>
<th>226.7</th>
<th>256.0</th>
<th>292.4</th>
</tr>
</thead>
<tbody>
<tr>
<td>T/M Unit:</td>
<td>TCw</td>
<td>TCw</td>
<td>TCw</td>
<td>TCw</td>
<td>TCw</td>
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<tr>
<td>Date Tested:</td>
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<td>8/18/93</td>
<td>8/19/93</td>
<td>8/19/93</td>
<td>8/19/93</td>
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<td>Dry Bulk Density (g/cc):</td>
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<td>2.205</td>
<td>2.166</td>
<td>2.297</td>
<td>2.090</td>
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<tr>
<td>Saturated Bulk Density (g/cc):</td>
<td>2.259</td>
<td>2.302</td>
<td>2.290</td>
<td>2.372</td>
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<td>2.512</td>
<td>2.520</td>
<td>2.511</td>
<td>2.526</td>
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<td>Porosity via Grain Density (%):</td>
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<td>12.2</td>
<td>14.1</td>
<td>8.5</td>
<td>17.3</td>
</tr>
<tr>
<td>Ultimate Tensile Stress (MPa):</td>
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<td>10.2</td>
<td>5.3</td>
<td>14.8</td>
<td>7.4</td>
</tr>
<tr>
<td>Depth, ft</td>
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<td>151.3</td>
<td>151.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>----------</td>
<td>-------</td>
<td>-------</td>
<td>-------</td>
<td></td>
<td></td>
</tr>
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<td>T/M Unit:</td>
<td>Pre-Rainier</td>
<td>Pre-Rainier</td>
<td>Pre-Rainier</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Date Tested:</td>
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<td>8/10/93</td>
<td>8/10/93</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dry Bulk Density (g/cc):</td>
<td>1.743</td>
<td>1.663</td>
<td>1.634</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Average Grain Density (g/cc):</td>
<td>2.436</td>
<td>2.437</td>
<td>2.457</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Porosity via Grain Density (%):</td>
<td>28.5</td>
<td>31.8</td>
<td>33.5</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 4b

SUMMARY DATA SHEET: NRG-2B BOREHOLE

Porosity Values for Untestable Sample Intervals

Sample IDs are shortened from the "NRG-2B-Depth-SNL-Subdivision" Format

| Depth, ft: | 48.4 | 79.1 |
| T/M Unit: | UO   | UO   |
| Date Tested: | 11/1/93 | 11/1/93 |
| Dry Bulk Density (g/cc): | 1.198 | 1.242 |
| Average Grain Density (g/cc): | 2.340 | 2.348 |
| Porosity via Grain Density (%): | 48.8 | 47.1 |
### Table 4c

#### SUMMARY DATA SHEET: NRG-2A BOREHOLE

**Porosity Values for Untestable Sample Intervals**

Sample IDs are shortened from the "NRG-2A-Depth-SNL-Subdivision" Format

<table>
<thead>
<tr>
<th>Depth, ft</th>
<th>135.3</th>
<th>166.5</th>
<th>231.1</th>
<th>162.2</th>
<th>162.2</th>
<th>166.5</th>
<th>166.5</th>
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</thead>
<tbody>
<tr>
<td>T/M Unit:</td>
<td>&quot;X&quot;</td>
<td>TCw</td>
<td>TCw</td>
<td>TCw</td>
<td>TCw</td>
<td>TCw</td>
<td>TCw</td>
</tr>
<tr>
<td>Porosity via Grain Density (%)</td>
<td>47.0</td>
<td>33.9</td>
<td>27.0</td>
<td>46.2</td>
<td>44.9</td>
<td>30.7</td>
<td>27.9</td>
</tr>
</tbody>
</table>
density yield consistently lower values than those computed using the average grain density and dry bulk density. In part, the differences can be attributed to an under-estimation of the saturated bulk density due to water loss on the surface of the specimen. However, these errors are small and cannot account for the total discrepancy. The major contribution to the differences is occluded porosity (isolated pores that are not filled during saturation). As a result, the specimens are not 100% saturated prior to mechanical testing; the saturations range between 80 and 95%.

3.1 Computerized Tomographic X-ray Images

Prior to testing, CT imaging was performed on each of the specimens tested in unconfined compression. A single image was obtained through the center of the specimen parallel to the core axis. These data serve as a qualitative measure of the shape and distribution of the pores and density heterogeneity within the specimen. Typical CT images are shown in Figures 7, 8, and 9. Figure 7 is a tomographic image of a nonwelded tuff recovered from a depth of 96.0 feet in borehole UE25 NRG-2A. The specimen shows the characteristic structure of most nonwelded tuffs. Low density lithic fragments have been pressed together. The lighter the color, the lower the density of the material. Some open pore space (shown as white in the scan) is visible in the section. The specimen has a high porosity, ($\phi = 51\%$) and exhibited an extremely low Young’s modulus and unconfined compressive strength.

Two CT scans of specimens of thermal/mechanical unit TCw recovered from borehole UE25 NRG-3 are shown in Figures 8 and 9. These specimens show elliptical low density zones within the specimens. However, in contrast with the CT scan shown in Figure 7, the background surrounding the low density inclusions is much darker and of higher density in these scans. The porosities are 12.4 and 8.7 percent for 218.0 and 289.2 feet respectively.

3.2 Compressional and Shear Wave Velocity Measurements

Compressional and shear wave velocity measurements were performed parallel and normal to the core axis for both dry and saturated conditions on all specimens tested in unconfined compression. Compressional and shear wave velocities were measured parallel to the core axis only for the specimens tested in confined compression. In some cases, data were not collected due to the inability to transmit the waves through the specimen; the
Figure 7: A tomographic image (CT scan) for a specimen of nonwelded tuff recovered from a depth of 96.0 feet from borehole UE25 NRG-2A at Yucca Mountain, NV.
Mountain, NV. 

TCW (covered from a depth of 21.8 feet from borehole U25 NRG-3 at Yucca 

Figure 6: A tomographic image (CT scan) of a specimen of the Yucca Canyon fulk unit.
Figure 9: A tomographic image (CT scan) for a specimen of the Tiva Canyon tuff unit (TCw) recovered from a depth of 289.2 feet from borehole UE25 NRG-3 at Yucca Mountain, NV.
greatest difficulty was encountered measuring shear wave velocities under saturated conditions. Significant difficulty was encountered measuring shear wave velocities under both dry and saturated conditions in many specimens. This problem is reflected in the large number of blank spaces in Tables 1a, b, c, and d. In most cases, compressional and shear wave velocity data were obtained only for specimens of TCw. An examination of the data shows that both compressional and shear wave velocities increased with decreasing specimen porosity.

3.3 Unconfined Compression Tests

The plots of representative data sets for two specimens tested in unconfined compression are presented in Figures 10 and 11. Figure 10 shows the data for a nonwelded high porosity saturated specimen of tuff recovered from borehole UE25 NRG-2A at a depth of 96.0 feet. These data were collected on the same specimen shown in Figure 7. The Young's modulus and fracture strength computed from these data are 4.9 GPa and 6.2 MPa, respectively. In the lower graph of Figure 10, the radial strain is plotted as a function of axial strain. Poisson's ratio is computed from these data over the same stress interval as Young’s modulus. Poisson’s ratio is 0.17.

Figure 11 shows stress-strain data collected on a specimen recovered from UE25 NRG-3 from a depth of 289.2 feet. The CT scan for this specimen is shown in Figure 8. Stress is plotted as a function of axial strain. The Young’s modulus and fracture strength computed from these data are 33.5 GPa and 121.1 MPa, respectively. The lower graph presents radial strain as a function of axial strain. Poisson’s ratio computed between 10 and 50% of the fracture strength is 0.23.

3.4 Confined Compression Tests

The results of confined compression tests on small diameter specimens obtained from cores recovered from depths of 263.3 and 265.7 feet in borehole UE25 NRG-3 are presented in Table 2. These depth intervals were chosen because the specimens have very similar characteristics, and using both intervals gave a more statistically significant test population. Specimens were tested at confining pressures of 5 and 10 MPa. The specimens from each depth were relatively low porosity and are from the TCw thermal/mechanical unit. The differential stress at failure increased with increasing confining pressure. This trend was consistent at both depths.
Figure 10: Axial stress and radial strain are plotted as a function of axial strain for a specimen tested in unconfined compression. The specimen was recovered from a depth of 96.0 feet from borehole UE25 NRG-2A, Yucca Mountain, NV.
Figure 11. Axial stress and radial strain are plotted as a function of axial strain for an unconfined compression experiment performed on a specimen of TCw. The specimen was recovered from a depth of 289.2 feet from borehole UE25 NRG-3, Yucca Mountain, NV.
3.5 Indirect Tensile Strength Tests

Forty-one indirect tensile strength tests were performed on the cores recovered from boreholes UE25 NRG-2, 2A, 2B, and 3. The tensile strengths ranged between 0.4 and 14.8 MPa. The weakest specimens were the nonwelded, high porosity units; whereas the strongest were the low porosity, welded TCw.
4.0 REFERENCES

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Displacement Gage for the Rock Mechanics Laboratory, SAND84-0651, Sandia National Laboratories, Albuquerque, NM.

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Ortiz, T.S., R.L. Williams, F.B. Nimick, B.C. Whittet, and D.L. South, 1985
A Three-Dimensional Model of Reference Thermal-Mechanical and Hydrological Stratigraphy at Yucca Mountain, Southern Nevada, SAND84-1076, Sandia National Laboratories, Albuquerque, NM.

Price, R.H. and S.J. Bauer, 1985

Simmons, G. and W.F. Brace, 1965
APPENDIX I

Stress vs Axial Strain and Radial Strain vs Axial Strain Plots for Unconfined Compression Experiments
NRG-2 Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw

AXIAL STRAIN (millistrain)

AXIAL STRESS (MPa)

(..): 170.4-A
(0): 174.0-A
(X): 178.0-A
(+): 179.5-A
NRG-2 Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw

(.-): 170.4-A
(o): 174.0-A
(X): 178.0-A
(+-): 179.5-A
NRG-2 Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw

AXIAL STRESS (MPa)

AXIAL STRAIN (millistrain)

(.: 180.0-A
(0): 188.3-A
(X): 196.2-A
(+): 200.0-A
Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw
NRG-2A Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: "X"
NRG-2A Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: "X"

(.-): 90.0-A
(o): 96.0-A
(X): 98.4-A
(+): 105.6-A
NRG-2A Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: "X"
NRG-2A Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: "X"

(\ldots): 127.0-A
(o): 136.1-A
(X): 145.3-A
NRG-2A Unconfined Compression

Saturated Sample
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw

AXIAL STRAIN (millistrain)

AXIAL STRESS (MPa)

(\ldots): 172.1-A
NRG-2A Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw

(.): 172.1-A
NRG-2A Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw

AXIAL STRESS (MPa)

AXIAL STRAIN (millistrain)
NRG-2A Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw

(-): 199.4-A
(0): 203.9-A
(X): 209.3-A
(+): 213.0-A
(*): 218.8-A
NRG-2A Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw

AXIAL STRESS (MPa)

AXIAL STRAIN (millistrain)

(.): 223.1-A
(0): 226.4-A
(\(\times\)): 234.9-A
(+): 238.4-A
(*): 254.5-A
NRG-2A Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw
Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: "UO"

NRG-2B Unconfined Compression

AXIAL STRAIN (millistrain)

AXIAL STRESS (MPa)

(•): 2.7-A
(0): 42.3-A
(X): 46.1-A
(+): 86.9-A
(^): 87.6-A
NRG-2B Unconfined Compression

-..: 2.7-A
(0): 42.3-A
(X): 46.1-A
(+): 86.9-A
(*): 87.6-A

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: "UO"
NRG-3 Unconfined Compression

AXIAL STRESS (MPa)

AXIAL STRAIN (millistrain)

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw
NRG-3 Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw

Radial Strain (millistrain)
Axial Strain (millistrain)

(.): 32.1-A
(0): 38.9-A
(X): 42.6-A
(+): 48.0-A
(*): 55.7-A
NRG-3 Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw

AXIAL STRESS (MPa)

AXIAL STRAIN (millistrain)

(•): 93.8-A
(0): 123.2-A
(X): 142.3-A
(+): 154.4-A
(*): 187.1-A
NRG-3 Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw

AXIAL STRAIN (millistrain)

RADIAL STRAIN (millistrain)

(•): 93.8-A
(0): 123.2-A
(χ): 142.3-A
(+): 154.4-A
(*): 187.1-A
NRG-3 Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw

AXIAL STRESS (MPa)

AXIAL STRAIN (millistrain)

(0): 195.7-A
(0): 208.9-A
(X): 218.0-A
(+): 226.7-A
(*): 256.0-A

76
NRG-3 Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw

( . . ) : 195.7-A
( 0 ) : 208.9-A
( X ) : 218.0-A
( + ) : 226.7-A
( * ) : 256.0-A
NRG-3 Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw

AXIAL STRESS (MPa)

AXIAL STRAIN (millistrain)

(.,): 257.6-A
(0): 257.6-B
(X): 263.3-A
(+): 289.2-A
(*): 292.4-A
(-): 297.1-A
NRG-3 Unconfined Compression

Saturated Samples
Ambient Pressure and Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw

AXIAL STRAIN (millistrain)

RADIAL STRAIN (millistrain)

(.): 257.6-A
(0): 257.6-B
(X): 263.3-A
(+): 289.2-A
(*): 292.4-A
(-): 297.1-A
APPENDIX II

Stress vs Axial Strain and Radial Strain vs Axial Strain Plots for Confined Compression Experiments
NRG-3 Confined Compression

(..): 263.3-E
(O): 265.7-B
(X): 265.7-D
(*): 265.7-F

CP = 5 MPa
Saturated Samples
Ambient Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw
NRG-3 Confined Compression

(. .): 263.3-E
(0): 265.7-B
(X): 265.7-D
(*) : 265.7-F

CP = 5 MPa
Saturated Samples
Ambient Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw
NRG-3 Confined Compression

CP = 10 MPa
Saturated Samples
Ambient Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw

AXIAL STRAIN (millistrain)

Differential Axial Stress (MPa)

(0): 263.3-D
(0): 265.7-C
(X): 265.7-E
(*): 265.7-G
NRG-3 Confined Compression

CP = 10 MPa
Saturated Samples
Ambient Temperature
Nominal Strain Rate of 10E-5 sE-1
T/M Unit: TCw
APPENDIX III

System Checks Using an Aluminum Standard Specimen
System Checks Using an Aluminum Standard Specimen

Unconfined compression experiments were performed on a specimen of 6061-T6511 aluminum. The specimen was monotonically loaded at a constant strain rate of $10^{-5}$ s$^{-1}$ to 138 MPa, (i.e., approximately one-half of its yield stress). Young’s modulus and Poisson’s ratio were computed from the stress and strain data. The purpose of the system checks is to ensure that the entire system is performing correctly and that the reported data are accurate.

A system check involves performing the uniaxial compression experiment on aluminum and comparing the observed Young’s modulus and Poisson’s ratio with the standard values reported for the material. If the measured values deviate by more than ±5 percent from the published reference values, corrective measures are taken and no further experiments are performed on tuff until the aluminum calibration experiment yields acceptable elastic constants.

Typical results from a calibration experiment are shown in Figure A-1. Axial stress and radial strain are plotted as a function of axial strain for an aluminum specimen with the same nominal dimensions as the tuff. These data were collected using the procedure specified for the unconfined compression experiments. The specimen is cyclically loaded to one-half its yield stress at a strain rate of $10^{-5}$ s$^{-1}$; Young’s modulus and Poisson’s ratio are computed from the data.

A summary of the system checks performed during the course of the study of the specimens from the UE25 NRG-2, 2A, 2B, and 3 boreholes are presented in Table A-1. The results indicate that the system performed within the specified accuracy during the course of the study.

Compressional and shear wave velocities were measured on the aluminum standard specimen. These data were used to compute the dynamic Young’s modulus and Poisson’s ratio. These values are also shown in Table A-1. The fact that these values for both Young’s modulus and Poisson’s ratio are larger than those given in the literature, suggests that there are minor variations in the properties of aluminum supplied by the manufacturer. The elastic moduli for nonporous materials are frequently computed from the compressional and shear wave velocities. These dynamic values should be used in conjunction with the static measurements. In many cases, the manufacturer’s data are obtained for tension experiments and empirically corrected for compression, which can also lead to a discrepancy.
Figure A-1: Axial stress and radial strain are plotted as a function of axial strain for a specimen of 6061-T6511 aluminum cyclically loaded in unconfined compression.
Table A-1

UE25 NRG 2, 2A, 2B, and 3
System Checks
Aluminum Standard - 6061 T6511

<table>
<thead>
<tr>
<th>Date</th>
<th>Young's Modulus (GPa)</th>
<th>Deviation (%)</th>
<th>Poisson's Ratio</th>
<th>Deviation (%)</th>
</tr>
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APPENDIX IV

Information from the Reference Information Base
Used in this Report

This report contains no information from the Reference Information Base.

Candidate Information for the
Reference Information Base

This report contains no information for the Reference Information Base.

Candidate Information for the
Geographic Nodal Information Study and Evaluation System

This report contains candidate information for the Geographic Nodal Information Study and Evaluation System (GeNESIS) in Tables 1, 2, 3, and 4. The data have been submitted to the SNL Participant Data Archive (PDA) and are indexed in the Automated Technical Data Tracking system (ATDT). The data packages have the following Data Tracking Numbers (DTN): SNL02030193001.003, SNL02030193001.005, SNL02030193001.006, SNL02030193001.007, SNL02030193001.010, SNL02030193001.011, and SNL02030193001.013.
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<td>1</td>
<td>D. A. Dreyfus (RW-1)</td>
<td>Director&lt;br&gt;OCRWM&lt;br&gt;US Department of Energy&lt;br&gt;1000 Independence Avenue SW&lt;br&gt;Washington, DC 20585</td>
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<td>L. H. Barrett (RW-2)</td>
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<td>D. R. Elle, Director</td>
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<td>Senior Project Manager for Yucca Mountain</td>
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</table>
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